Signalling crosstalk at the leading edge controls tissue closure dynamics in the *Drosophila* embryo

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Abstract

Tissue morphogenesis relies on proper differentiation of morphogenetic domains, adopting specific cell behaviours. Yet, how signalling pathways interact to determine and coordinate these domains remains poorly understood. Dorsal closure (DC) of the *Drosophila* embryo represents a powerful model to study epithelial cell sheet sealing. In this process, JNK (JUN N-terminal Kinase) signalling controls leading edge (LE) differentiation generating local forces and cell shape changes essential for DC. The LE represents a key morphogenetic domain in which, in addition to JNK, a number of signalling pathways converges and interacts (anterior/posterior -AP- determination; segmentation genes, such as Wnt/Wingless; TGFβ/Decapentaplegic). To better characterize properties of the LE morphogenetic domain, we sought out new JNK target genes through a genomic approach: 25 were identified of which 8 are specifically expressed in the LE, similarly to *decapentaplegic* or *puckered*. Quantitative *in situ* gene profiling of this new set of LE genes reveals complex patterning of the LE along the AP axis, involving a three-way interplay between the JNK pathway, segmentation and HOX genes. Patterning of the LE into discrete domains appears essential for coordination of tissue sealing dynamics. Loss of anterior or posterior HOX gene function leads to strongly delayed and asymmetric DC, due to incorrect zipping in their respective functional domain. Therefore, in addition to significantly increasing the number of JNK target genes identified so far, our results reveal that the LE is a highly heterogeneous morphogenetic organizer, sculpted through crosstalk between JNK, segmental and AP signalling. This fine-tuning regulatory mechanism is essential to coordinate morphogenesis and dynamics of tissue sealing.

Author summary

Dorsal closure of the *Drosophila* embryo is used as a paradigm to study epithelial sealing and is related to wound healing. This vital process relies on the dorsal migration of the two lateral ectodermal sheets and is necessary for the protective epidermis to completely envelop the embryo. The row of cells located at the front of migration, called the leading
edge, is the organizing center of the process, where key signaling pathways turn on specific gene expression. Here we used a genomic approach to identify new genes whose expression is restricted to the leading edge. A quantitative analysis revealed differential expression along the anterior-posterior axis of the leading edge, which was considered for a long time as homogeneous or amorphous. We demonstrate that anterior-posterior cues provide an orthogonal coordinate system specifying cell identity along the whole leading edge, making it a highly patterned morphogenetic center. We further show that these anterior-posterior cues are functionally important, controlling the dynamics of dorsal closure and participating to the robustness of the process. Our work sheds new light on the role of anterior-posterior cues in epithelial tissue sealing related to wound-healing.

Introduction

Epithelial morphogenesis is orchestrated at the cellular level through local shape changes and tension-based dynamics. In this process, cell-cell signalling plays an essential role in coordinating gene expression programs with tissue behaviour. One of the best-studied morphogenetic movements is embryonic dorsal closure (DC) in *Drosophila*. Following germ band retraction, the embryo is only partly enveloped by the epidermis, leaving a large dorsal area covered by the transient squamous amnioserosa. DC uses a combination of signalling pathways to generate mechanical forces, cell shape changes and cell adhesion rearrangements causing the two lateral epidermal sheets to spread dorsalwards and fuse at the dorsal midline through a zipping mechanism.

At the onset of DC, the dorsal-most epidermal cells, a.k.a leading edge cells or LE cells, polarise under the influence of the canonical Wingless (Wg) pathway and elongate along the dorso-ventral axis [1–3]. LE polarization leads to a particular organization of the dorsal cell membrane (that in contact with the amnioserosa), which in particular loses adherens junction markers (such as E-Cadherin, ECad) and septate junction markers (such as Discs-large, Dlg) [2, 4, 5]. This redistribution results in the formation of actin-nucleating centers (ANC) from either side of the dorsal membrane that will transmit, along with the adherens junctions, mechanical forces [2, 5, 6]. LE cells start synthesising an ANC-linked acto-myosin cable that generates a contractile force allowing their alignment during dorsal migration [7, 8]. The acto-myosin cable also participates in a ratchet-like process induced by the asynchronous pulsation of the amnioserosa cells [9]. By progressively contracting and disappearing by cell engulfment and apoptosis, the amnioserosa supplies an additional positive force, whereas the stretching lateral ectoderm opposes a resistance force to tissue progression [8, 10]. The two contralateral ectodermal sheets then adhere to each other starting from the two canthi of the hole in a process called zipping, providing a fourth force to DC [6, 8, 11]. Non-muscle myosin II is involved in the generation of the four forces thanks to its motor and contractile activities [12].

The LE is the organizing centre of DC and is specified by JNK signalling. Mutations in components of the JNK pathway such as DJNKK/hemipterous (hep) or DJNK/basket (bsk) strongly affect DC, leaving embryos with dorsal holes in their cuticle [1, 13, 14]. JNK signalling is specifically activated in the LE cells, controlling the expression of the target genes puckered (puc, as revealed by the lacZ-containing enhancer-trap puc-Z), decapentaplegic (dpp) and scarface (scaf) [1, 15–17]. Recently, a feed-forward loop between the JNK and Dpp activities was shown to regulate the expression in the LE of three proteins, the myosin VI homologue Jaguar, the microtubule-binding protein Jupiter and the integrin-linked Zasp52 [18]. Other JNK target genes, whose expression is not limited to the LE during DC, have also been described, such as...
the profilin-coding gene *chickadee*, the transcription factor *cabut*, the integrin-coding genes *scab* and *myospheroid*, and the trafficking gene *Rab30* [19–22].

A specific feature of DC is that it occurs in a field of cells that is not uniform along the anterior-posterior (AP) axis, encompassing the thoracic and abdominal regions. In addition, the ectodermal cells (LE and lateral ectoderm) are divided in repeated, segmental units (T1–T3 and A1–A8 segments). Whereas the HOX genes define the identity of the segments along the AP axis [23], the segment polarity genes are responsible for the elaboration of cellular patterning within each segment of the *Drosophila* embryo leading to the formation of anterior and posterior compartments [24]. Therefore, it is important to characterize the interaction between these orthogonal signalling events (JNK vs. AP/segmentation signalling) and determine how this interaction controls tissue morphogenesis.

Previous studies have shown that the Wg pathway collaborates with JNK to induce *dpp* expression in the LE [3, 25]. It was also demonstrated that the dynamics of closure presents robust asymmetric properties along the AP axis [10, 26]; for instance, the anterior speed of closure is faster than the posterior one, which could be due to localized apoptotic forces present in the anterior amnioserosa [10]. Yet, whether and how segmentation and AP cues impact on DC is currently unknown. In this study, we characterized the genomic response of the JNK pathway during DC, revealing a whole set of new target genes, several of them being specifically transcribed in the LE. Quantification of these new intra-LE expression profiles uncovers a complex organization of the LE that depends on crosstalk between JNK, HOX and segmentation pathways. In this network, HOX genes can have positive or negative activities, regulating segmental features during closure. For instance, loss of the posterior HOX gene *abdominal-A* (*abd-A*) or *Abdominal-B* (*Abd-B*) leads to closure delays of the most posterior segments, indicating that they control the timing of closure. Thus, crosstalk between the AP and JNK systems shapes the LE organizing centre for proper tissue sealing dynamics.

**Results**

**JNK transcriptional response during DC**

To better characterize the LE and its role during DC, we used microarrays to identify genes expressed under the control of the JNK pathway in the *Drosophila* embryo (Figs 1A and S1A). Three different conditions for JNK activity were tested: wild-type (WT; used as reference), loss-of-function (LOF) and gain-of-function (GOF). GOF embryos were obtained by ectopically expressing the active form of the DJNKK/Hep (Hep<sup>act</sup>) protein in the ectoderm using the 69B-GAL4 driver, while LOF embryos were generated using a hept<sup>75</sup>/hept<sup>1</sup> hetero-allelic combination [1]. The extent of JNK activity and sample homogeneity in the three groups of embryos were assessed through *in situ* hybridization using a *dpp* probe (S1A Fig).

Total RNAs were prepared from carefully-staged embryos and analysed by microarray. The statistical comparison of WT and GOF conditions identified 1648 independent genes (corresponding to 1679 probes) which are regulated by a factor equal to or higher than 1.5, of which 1001 (1023 probes) are activated by JNK and 647 (656 probes) are inhibited (S1B Fig). For the WT-LOF comparison, 113 distinct genes (117 probes) are uncovered with a fold change limit of 1.5, with 35 (37 probes) activated in the *hep* mutant (i.e. inhibited by JNK) and 78 (80 probes) inhibited (S1B Fig).

These results revealed a smaller number of genes controlled in LOF compared to GOF. This difference is not surprising when comparing the extent of JNK activity (evaluated through the expression of *dpp*) in GOF/LOF vs. wild-type embryos. Indeed, JNK activity is strongly increased in GOF embryos, while only LE cells (representing approximately a mere 200 cells of the whole embryo) lose *dpp* mRNA in LOF embryos (S1C Fig). This difference in amplitude...
A 3 genetic conditions control: WT
GOF: $69B>$hep^act
LOF: hep'/hep^75

2hr embryo collections

stage 13 + stage 14 + stage 15

RNA samples

GOF (FC+/-1.5): 1001 JNK-activated genes
647 JNK-repressed genes

LOF (FC+/-1.5): 78 JNK-activated genes
35 JNK-repressed genes

microarray

GOF: filtering

194 genes

in situ hybridization

31 JNK target genes
(10 LE genes)

B WT enGAL4>$hep^act

CG1146
GstE3
CG1890
FHOS
CG4199
GlCAT-S
Pak
Rab30
CG2811
CG10916
ALX

WT enGAL4>$hep^act

CG1572
CG34417
CAH2
Jra
scb
al
sick
PvT2
Zasp52
igpr

C

relative FC

WT
GOF
LOF
between GOF and LOF likely explains the fact that only a few genes are found in common between the two lists (S1 Table). For this reason, we focused on the WT-GOF comparison, and in particular on the genes that are activated by the JNK pathway.

We initially performed a functional classification of 1648 GOF genes based on Gene Ontology (GO) terms, using the David bioinformatics tool which has the advantage of making family groups from GO terms and ranking them according to enrichment scores [27]. Functional classification of the 1001 up-regulated genes (S2 Table) and 647 down-regulated genes (S3 Table) revealed that during DC, JNK signalling controls a wide range of biological functions including cell cycle, cytoskeleton, proteolysis, mRNA transport, and regulation of transcription.

Identification of 31 JNK target genes

To further identify bona fide JNK target genes, we performed in situ hybridization on en-GAL4/UAS-hep<sup>act</sup> embryos, in which the JNK pathway is activated in a striped pattern, as seen with the puc control (Fig 1B). Because of the great number of identified genes, we first made a selection of genes based on their potential localization and function (membrane, secreted, motors, etc.) and on available expression data from public databases (such as FlyExpress). In addition, comparison of our embryonic data set with Drosophila Schneider-2 cells induced by the lipopolysaccharide JNK activator, allowed the extraction of a list of genes having a common transcriptional response to JNK activity [28, 29]. In total, we selected and tested 194 genes by in situ hybridization and identified 31 bona fide JNK target genes (Fig 1B; S4 Table). The response of these 31 genes to JNK signalling was further confirmed by quantitative RT-PCR (Fig 1C). Of the 31 genes identified, some have already been described as transcriptional targets of the JNK pathway (jra/jun, scab, Zasp52, scaf, Rab30 and reaper) [17, 18, 21, 22, 30, 31], while others have been linked to the JNK pathway and/or to DC without being described as transcriptional targets of JNK (Puk, icarus and ALiX) [32–34], thus validating our approach and genomic data. Interestingly, 10 out of 31 genes show specific expression in the LE, which is dependent on JNK activity as their expression is lost in the hep mutant, similarly to puc or dpp (Fig 2). Therefore, transcriptional profiling of embryos undergoing DC allowed the identification of several new JNK target genes (representing 25 genes plus the previously known ones: jra/jun, scab, Zasp52, reaper, scaf and Rab30; i.e. 31 genes in total), including 10 (8 new in addition to scaf and Zasp52) with a specific expression in the LE, like the well-known LE JNK target genes puc and dpp (hereafter referred to as “LE genes”).

Spatio-temporal heterogeneity of gene expression at the LE

Surprisingly, we noticed that the expression of most of the LE JNK target genes was not spatially and temporally homogeneous along the LE. First, the onset and arrest of gene expression show temporal regulation during DC. Genes can start to be expressed from either stage 11 (corresponding to full germ-band extension; puc-Z, CG34417, dpp, al, Zasp52), stage 12 (during germ-band retraction; scaf, sick) or at the onset of DC at stage 13 (CG5835, hui, yb) (Fig 3A). Termination of expression also varies from gene to gene. The expression of dpp, al and
Zasp52 decreases very rapidly after the start of DC and is no longer visible at stage 14 while DC is still in progress. In contrast, other genes lose their LE expression after the complete fusion of the lateral sheets of the epidermis. To our knowledge, this is the first time that early, intermediate and late JNK-target gene expression is described in a developmental process (Fig 3A), whereas it is known that the JNK pathway has distinct phases of strength and duration [35–38]. Second, our initial in situ hybridization experiments suggested heterogeneous expression along the LE (Fig 2). In order to precisely define the expression pattern of each LE gene, we developed CurvedPeriodicity, a program to quantify the mRNA levels specifically in the LE from fluorescent in situ hybridizations (FISH). The FISH experiments were coupled to immuno-fluorescence (FISH-IF) revealing the En protein to delineate the LE borders and position the segment boundaries. We extracted from CurvePeriodicity the fluorescence intensity per segment, showing the existence of 4 intra-LE profiles with either uniform expression along the entire length of the LE (dpp and CG34417), or expression domains biased towards the anterior (puc-Z, al, sick, Zasp52 and Pvf2), posterior (hui and yb) or central (scaf, CG5835, CG4199) portions of the LE (Fig 3B). Pvf2 is a remarkable example of the ‘anterior’ group, being only expressed in the three thoracic segments (posterior expression in Fig 2 corresponds to cardiac cells). In the central domain, expression can be either lowered (scaf) or increased (CG5835, CG4199) compared to more terminal parts.

Fig 2. New genes showing JNK-dependent expression in the LE. In situ hybridizations showing expression in the LE of dpp (control) and of the 10 LE genes (1st and 3rd columns; arrows). LE-specific expression is lost in hep mutant embryos (2nd and 4th columns; arrows), indicating that the JNK pathway drives LE expression of these genes. Of note, expression is not uniform along the whole LE.

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Fig 3. Spatio-temporal regulation of JNK target gene expression at the LE. A) Temporal expression (black lines) of the twelve LE genes from stage 11 (full extension of the germ band) to stages 14/15 (end of DC), including stage 12 (germ band retraction) and stage 13 (onset of signalling and tissue closure dynamics).
In addition to this global LE heterogeneity along the AP axis, we also found striking differences in the segmental expression of JNK target genes (Fig 3C). Quantitative analysis revealed five different categories of segmental profiles: i) a linear profile, with no significant change along the segment (puc, CG34417, CG4199), ii) anterior and iii) posterior profiles, showing stronger expression in the anterior (scaf) or posterior (CG5835, Zasp52) compartment, respectively, iv) a parasegmental boundary profile in which highest expression is observed between the anterior and posterior compartments of the same segment (dpp, hui) and v) a segmental boundary profile in which highest expression is found at the segment boundary (sick, al, yb, Pvf2). Altogether, these results identify a highly complex, spatio-temporal regulation of the JNK transcriptional response during DC both along the anterior-posterior axis and within segments, raising the question of the mechanisms setting these different transcriptional programs at the LE.

The segmentation gene en is a repressor of LE gene expression

To address this question, we used the JNK target gene scaf as readout because it exhibits a strong expression in the LE that undergoes both AP and segmental regulations (Figs 3B, 3C and 4A). Quantification in the A and P compartments from each segment revealed that scaf is down-regulated in the P compartments of the central abdominal segments, from A1 to A6 (Fig 4C). The posterior factor En has been shown to have both activating and repressive transcriptional activities [39], making it a potential scaf regulator in this process. In en null mutant embryos, scaf expression is no longer segmented (Figs 4A and 5A). Moreover, the overexpression of en with the pannier (pnr)-GAL4 driver in the whole dorsal ectoderm leads to down-regulation of scaf expression, but strikingly, this inhibition only occurs in the abdominal segments. Of note, en had no influence on the non-segmented expression of the puc-Z JNK reporter gene (Fig 4A). Altogether, these results indicate that en is a LE negative regulator, repressing the posterior expression of a subset of JNK target genes, including scaf. To assess if En acts directly or through a relay mechanism, we expressed a mutant form of En in which the repressor domain has been replaced by the VP16 transactivation domain (VP16En) [39]. In contrast to the wild-type form, the overexpression of VP16En has no inhibitory effect on scaf expression, supporting a direct repressive activity of En (Fig 4B and 4C). This result also indicates that the effect of en occurs independently of the auto-regulatory loop involving wg and hh. Additionally, we observed that in VP16En-overexpressing embryos, the scaf expression level in the posterior compartments of central segments is higher than in WT embryos, indicating that VP16En can compete with endogenous En (Fig 4C). Altogether these results indicate a direct role of the segment polarity gene en in patterning the LE during DC, shown by the control of the JNK target gene scaf (Fig 4D).

The HOX gene abd-A collaborates with en to repress scaf expression

Despite being expressed in all segments, the repressor activity of En is restricted to central abdominal segments, indicating the existence of additional regulatory mechanisms to
circumscribe En action and refine LE patterning. We thus tested the role of the HOX genes, which control segment identity along the AP axis. The HOX gene *abd-A* is expressed and acts in the abdominal segments. In *abd-A* null mutant embryos, the expression of *scaf* is increased in segments A1-A7 (i.e. in the *abd-A* domain; Figs 5A and S4B), indicating that *abd-A* negatively regulates *scaf*. Interestingly, quantification of *scaf* expression at the segmental compartment level shows that it is no longer repressed in the P compartments of central segments (A1-A6). This *abd-A* mutant phenotype is therefore similar to the *en* mutant phenotype, suggesting that both genes work together to control *scaf* in the LE. When overexpressed, *abd-A* leads to decreased *scaf* expression in the posterior segments (Figs 5B and S4C). In the thoracic segments where *abd-A* is normally not expressed, we observed *scaf* down-regulation in the P compartments.
compartments where en is also present, indicating that the inhibition of scaf expression in the P compartment requires both En and Abd-A. This phenotype explains why the overexpression of En has a restricted effect in the abdominal segments where abd-A is expressed, with no effect in the thoracic segments where abd-A is absent (Fig 4A). Therefore, En and Abd-A collaborate to down-regulate scaf in the central segments, where their pattern of expression overlaps. In conclusion, En and Abd-A are two repressors regulating LE gene expression (Fig 5C).

The HOX genes Scr, Antp, Ubx and Abd-B are activators of scaf expression

We extended our study to the other HOX genes that are expressed in the dorsal ectoderm during DC (Fig 5F) by testing the role of Sex combs reduced (Scr; expressed in the T1 segment), Antennapedia (Antp; expressed in the T1-T3), Ultrabithorax (Ubx; strongest expression in A1, declining until A7) and Abd-B (strongest expression in A8, declining towards A6) [23]. Null mutants for either Hox gene showed reduced scaf mRNA levels in their expression domain, suggesting they all have a positive role on scaf transcription (Figs 5D and S4D). Consistently, upon overexpression, these HOX genes were all able to ectopically up-regulate scaf expression (Fig 5E). Therefore, in contrast to abd-A which acts as a repressor, Scr, Antp, Ubx and Abd-B activate the expression of scaf in their respective functional domain.

All these results reveal that key transcription factors controlling AP axis and segmentation work together generating a complex gene regulatory network to control gene expression along the LE. This LE network, which involves both activating (JNK, Scr, Antp, Ubx and Abd-B) and repressive (abd-A and en) activities, leads to a remarkable transcription profile of scaf in the LE, with high expression at the poles of the embryo and low expression in the P compartments of A1-A6 segments (Fig 5F).

HOX cofactor requirement for LE gene expression

HOX proteins cooperate with cofactors to activate transcription [40]. Homothorax (Hth) and Extradenticle (Exd) are known to interact with each other and with HOX proteins to form a transcriptional complex binding specific cis-regulatory sequences [41]. Hth contains a homeodomain (HD) in its C-terminal part that is responsible for DNA binding. In addition, Hth promotes Exd nuclear translocation and its binding to DNA [40, 42–44]. In order to inquire the role of HOX cofactors on scaf expression, we thus analysed embryos mutants for hthP2, a strong hypomorphic allele resulting in an absence of Hth protein expression [45]. As a result, Exd cannot be translocated into the nucleus. In absence of cofactor function, scaf expression decreases (Fig 5G). This is especially the case in the more extreme segments, whereas the effect is only moderate in the central segments. Indeed, we still observe in the region A1-A6 of the hthP2 mutant the segmented pattern of scaf expression, with the same level of transcripts in the posterior compartments than wild-type embryos (Fig 5G, right panel and S4E Fig). These results indicate that the HOX cofactors Hth and Exd are required for the positive regulation of scaf expression in the LE by Scr, Antp, Ubx and Abd-B, but do not participate in the negative action of En and Abd-A. Therefore, in addition to the core LE network involving the transcription factors A1, HOX and En, the HOX cofactors have a distinctive contribution along the AP axis according to the identity of the segments, further refining LE patterning.

HOX genes are important regulators of JNK-dependent LE patterning

To determine whether HOX genes have a more general role on JNK target genes, we investigated the profiles of hui, yb and Pf2 genes, which like scaf display a complex pattern of expression along the LE. hui and yb exhibit a stronger expression in the posterior part of the embryo
Fig 5. HOX regulation of scaf expression. A) Quantification of scaf expression in the abd-A mutant (n = 15). First bar diagram: expression along the AP axis of the abd-A mutant (black) is higher than in WT embryos (blue), but only in the abdominal segments where abd-A is active (yellow). Second bar diagram: compartmental expression of scaf (anterior in blue and posterior in green) showing that posterior compartment expression is highly increased compared to WT embryos (shown in Fig 4C, first panel). Third bar diagram: average expression in the central A1 to
A6 segments of WT (see Fig 4C) and abd-A embryos, which shows the same level of expression in the A and P compartments in the abd-A mutant. B) Quantification of scaf expression in embryos overexpressing abd-A (pnr > abd-A; n = 20). First bar diagram: compared to WT embryos (blue curve), overexpression of abd-A (black curve) leads to scaf mRNA down-regulation in the posterior region. Middle bar diagram: abd-A overexpression also induces scaf mRNA diminution in the posterior compartments (i.e. en-expressing cells) of thoracic segments, where Abd-A is normally absent, indicating that Abd-A requires En for its repressive function. Right bar diagram: average expression in the thoracic segments (shown in Fig 4C): *: p < 0.05; **: p < 0.01; ***: p < 0.001. In the right panel, only the segments from A1 to A6 were quantified, averaged and compared to WT embryos (see Fig 4C).

('posterior' group), whereas Pvf2 expression is limited to the thoracic segments ('anterior' group; see Fig 3B). In abd-A and Abd-B mutants, the expression of hui and yb is diminished in the respective HOX domains, indicating that both genes activate the expression of hui and yb, which is responsible for their stronger expression at the posterior pole of the embryo (Fig 6A and 6B). For Pvf2, we observed a strong expression increase from T3 to A6 segments in the abd-A mutant and a weak up-regulation in the A6/A7 segments in the Abd-B mutant (Fig 6C). Thus, the restricted expression of Pvf2 in the thoracic segments is due to the inhibitory action of the two abdominal genes, especially abd-A. These results indicate that abd-A and Abd-B can either act as repressors or activators to shape the asymmetric expression profiles of JNK target genes in the LE.

**abd-A and Abd-B control the posterior closure of the embryo by regulating the zipping process**

In order to assess the functional relevance of Hox gene regulation during DC, we analysed the phenotype of HOX gain- and loss-of-function embryos. Overexpression of Abd-B in the dorsal ectoderm results in a strong DC phenotype, as revealed by holes present in the embryonic dorsal cuticles (Fig 7A). Therefore, 'posteriorisation' of the LE through Abd-B overexpression, which leads to ectopic activation of JNK target genes (Fig 5E and 5F), is detrimental to DC, indicating the importance of proper patterning of the LE along the AP axis. In contrast, abd-A overexpression does not create cuticle holes (Fig 7A), which might be due to the regulation of a different set of target genes compared to Abd-B. To characterize the role of the HOX genes in their normal domain of expression, we analysed the effect of their loss-of-function. First, we observed the cuticles of the abd-A and Abd-B mutants, which resemble that of WT embryos (Fig 7A). The hth mutant exhibits a strong DC phenotype, with holes located in the anterior part. This regionalisation of the phenotype is likely reflecting the higher expression of hth in the thoracic part of the embryo [45]. In addition, this strong phenotype is probably due to the effect on several HOX genes, in particular the thoracic ones (Scr, Antp and Ubx), as previously shown (Fig 5G). We then decided to analyse in more details the DC phenotype of the HOX genes by analysing segment closure during early DC. In the WT embryo, closure of the posterior segment A8 and closure of the anterior segment T1 take place approximately at the same time (Fig 7B). In the hth mutant, when A8 has just closed, the segment T1 is far from being closed, revealing a strong delay in the closure of the anterior part of the embryo (Fig 7B and 7C). We also analysed the abd-A and Abd-B mutants, and both show a delay in the closure of the posterior segment A8 (Fig 7B and 7C). Therefore loss of HOX gene function triggers DC...
defects in their respective field of action, and the severity of the phenotype depends on the number of HOX genes affected.

To get better insight into the role of individual HOX genes on the dynamics of DC, we performed live imaging on Abd-B mutant embryos, which display a clear delay in DC (Fig 7D; S1 Movie). Despite this delay, they eventually close and show no hole in their dorsal cuticles (Fig 7A), reflecting the high degree of adaptability and robustness of DC in challenged conditions [8, 46, 47]. Analysis of the point of closure at the LE indicated that closure of the Abd-B mutant was shifted towards the posterior, compared to WT embryos whose final point of closure is located at the middle of the LE (Fig 7E). Early delay observed in fixed embryos (Fig 7B and 7C) is thus maintained throughout DC to trigger an asymmetric closure, indicating that no compensatory mechanism takes place. The posterior shift and closure delay seen in Abd-B embryos can be explained by a progressive reduction of the posterior speed of closure. Indeed, this speed shows a clear reduction of 59% (mid DC) and 78% (late DC) (S5A Fig). The anterior speed also decreases, but to a less extent (58% in late DC; see Discussion).

The DC defect of the hth mutant seems to be generated by an inefficient zipping in the anterior part, but the strong phenotype makes it difficult to interpret. We therefore analysed the zipping zone of the abd-A and Abd-B mutant embryos. Antibody stainings indicated that the establishment and positioning of LE markers and of the zipping zone are correct, such as the ANC-localised protein Enabled (Ena), the adherens junction protein ECad and the septate
Fig 7. DC phenotypes of HOX mutant embryos due to zipping defects. A) Cuticles of a WT embryo (top left), abd-A and Abd-B-overexpressing embryos with *pnr-GAL4* (top middle and top right, respectively), *hth*, abd-A and Abd-B mutants (bottom panels, as indicated). The overexpression of Abd-B induces holes in the...
cuticles (arrow; 48% of the dead embryos, n = 200), revealing strong closure defects. The hth mutant also produces a DC phenotype, with holes located in the anterior part (arrow; 76% of the dead embryos, n = 117). In contrast, the cuticles of abd-A-overexpressing embryos, abd-A and Abd-B mutants resemble that of WT embryos. B) Anti-En immunostaining of the hth (top right), abd-A (bottom left) and Abd-B (bottom right) mutants compared to WT embryos (top left). For hth, the distance in the anterior (D-A) between the two opposing En-expressing compartments of T1 is determined when the A8 segment just closed. For abd-A and Abd-B, the distance in the posterior (D-P) between the two opposing En-expressing compartments of A7 is determined when the T1 segment just closed. The red lines delimitate the posterior boundaries of the T1 and A7 segments. C) Quantification of D-A (μm +/- s.e.m., top panel) for the hth mutant and D-P (μm +/- s.e.m., bottom panel) for abd-A and Abd-B mutants. In WT embryos, T1 is almost closed (D-A = 8.3 μm +/- 1.6, n = 20) when A8 just closed, but the hth mutant shows a strong anterior opening (D-A = 52.3 μm +/- 4.5, n = 17). In the posterior part of WT embryos, A7 is usually closed (D-P = 1.6 μm +/- 0.5, n = 18) when T1 just closed. In contrast, the abd-A and Abd-B mutants exhibit a posterior delay of closure (D-P = 16.3 μm +/- 3.0, n = 19, and D-P = 11.0 μm +/- 2.0, n = 16, respectively). D) Still images of arm-GFP and arm-GFP;Abd-B live embryos from S1 Movie. Compared to the dynamics of the arm-GFP (WT) embryo (left images), the closure of the Abd-B mutant (right images) is slowed down: at 160 minutes, the abd-B mutant embryo is still undergoing closure while the WT embryo is closed. Height (H) between the two opposing LE and length (L) of the projected LE at the midline between the two zipping zones that are used in C) are depicted in the top right panel. H is quantified at fixed positions along L, as indicated. 

Discussion

Our identification of several new JNK target genes during DC and analysis of their quantitative expression pattern uncover the complex transcriptional response taking place in the LE morphogenetic domain. Results reveal an intricate regulatory network integrating multiple signalling layers. In this process, AP positional information and JNK signalling cooperate to
generate a highly patterned, yet apparently smooth and regular LE. Mutant analysis shows that LE partitioning into discrete domains is important to control the coordination, hence dynamics of the whole closure process.

The LE is a major component of DC, being the site of JNK activity and actin cable assembly; it also provides an active boundary with the amnioserosa, driving epidermal spreading and seamless tissue sealing. Therefore, it is important to determine its morphogenetic and signalling features and how these are dynamically controlled. To this end, first we identified a new set of target genes whose expression in the dorsal ectoderm is dependent on JNK activity during DC. Transcriptome analysis allowed us to identify 1648 independent genes which are up- or down-regulated in JNK activated embryos. Filtering of this large set led us to focus on a group of 194 genes whose expression was analysed by quantitative in situ hybridization in different genetic conditions. Transcriptional profiling unveiled 31 Drosophila JNK target genes of which only a fraction was already known, including jra/jun, reaper, Zasp52 and scab [21, 30, 31]. Amongst novel targets were also Scaf and Rab30 the role of which we have previously described during DC [17, 22]. Two categories of JNK target genes can be distinguished: genes that are specifically expressed in the LE and genes whose expression is more ubiquitous in the dorsal ectoderm. Genes belonging to the latter category may play a general role in the ectoderm under the control of different pathways, for example in the case of Rab30. In contrast, LE-specific genes likely play a specific role during DC, as is the case for puc, dpp and scaf. However, it is also possible that some of the new genes, despite being expressed in the embryo in a JNK-dependent manner, are not involved in DC. These target genes thus remain under the control of JNK, but are functionally ‘silent’ during DC. This behaviour is best illustrated by the reaper gene, whose expression is JNK-dependent in the embryo (Fig 1B and 1C), but which does not seem to have any function in the LE, only later during development or at the adult stage.

Surprisingly, quantitative analysis of LE-specific gene expression profiles showed a variety of previously uncharacterized expression patterns along the LE, with two levels of regulation, AP and segmental. These observations reveal a new property of the LE which appears highly patterned along the AP axis, contrasting with the homogenous and linear structure previously envisioned. In addition, the higher order regulation that emerges from these results provides every LE cell with its own identity through the cross-talk between JNK, AP and segmental information. Such cell-level patterning through signalling crosstalk [48] is likely essential for coordination and robustness of closure as well as segment matching. In this view, recent work showed that Wg and JNK interact at the LE to control the formation of specific mixer cells at segment boundaries [49].

Previous work showed that, instead of acting independently, HOX and segmentation genes can be coupled to regulate target genes in the embryo [50]. Here, we reveal an additional layer of regulation involving the ‘morphogenetic’ JNK signalling pathway. During DC, JNK acts as a tissue-specific switch whose activity can be regulated by HOX and segmentation pathways, providing positional information and segmental organization to a moving tissue boundary. Thus, a multi-layered or ‘onion-like’ regulatory model allows for several levels of regulation/information to pile up in order to regulate individual cellular behaviours important for tissue morphogenesis. Each layer can act positively or negatively on LE target gene expression, generating a complex repertoire of regulatory pathways. Distinct categories of expression profiles have been identified in this study through the analysis of individual target genes, with likely more gene-specific patterns to be anticipated. For example, the same HOX gene (abd-A or Abd-B) can have activating or repressive activity according to the target gene, as is the case for the transcription factor En [39]. Molecular functional characterization of cis-regulatory elements controlling LE gene expression will bring a more detailed view of how transcription
factor complexes are formed, how specificity of DNA recognition is achieved and how activating or repressive activities are regulated to generate LE patterning.

scaf proves to be a remarkable case among the JNK target genes, summarizing the different levels of regulation that can be integrated into a single promoter. Not only is it strongly expressed in the LE in a JNK-dependent manner, but it is also regulated by both the segmentation gene en and the HOX genes. In particular scaf displays a transcriptional response induced by all the trunk HOX genes tested, being positively controlled by Scr, Antp, Ubx and Abd-B and negatively by abd-A. It can therefore be considered as a general HOX target gene, i.e. regulated by most Hox paralogs, as previously defined [40]. Another example of a general target is the Drosophila gene optix, which is activated by the head HOX genes labial and Deformed (Dfd) and inhibited by the trunk HOX genes [51]. Nonetheless the general HOX target genes do not represent the majority. A genomic analysis in the Drosophila embryo identified more than 1500 genes regulated by at least one of the six HOX paralogs tested (Dfd, Scr, Antp, Ubx, abd-A, Abd-B) [52]. Only 1.3% of these genes are regulated by the six paralogs and 1.5% by the five paralogs that we used in our study. Interestingly more than 40% of the ~1500 HOX target genes are also present in the JNK genomic data set that we obtained. This strong overlap well reflects the fact that the LE runs along most of the body AP axis encompassing the thorax and abdomen. More importantly, it also indicates that AP patterning plays a crucial role in the regulation of DC, as shown in this study.

Live imaging and mathematical modelling revealed asymmetries in the geometry and zipping process along the AP axis [8, 53], which can be attributed to local constraints induced by head involution and apoptosis [10, 26]. Head involution is concomitant with DC and induces tension in the anterior part of the embryo, explaining why the DC phenotypes are almost exclusively observed in the anterior part, leading to the so-called ‘anterior-open phenotype’. The exception to this rule is the experimental manipulation of the posterior zipping rate through localized laser ablation of the amnioserosa close to the canthus, which induces a strong delay of posterior closure [26]. Our results with the abd-A and Abd-B mutants show that posterior delay can also be obtained in genetically-perturbed embryos. However, while anterior zipping is slightly up-regulated when posterior zipping is laser-targeted [26], we showed that the anterior speed of closure is diminished in the Abd-B embryo. Thus, compensatory mechanisms may only appear when tissue integrity is severely impaired. Apoptosis was also proposed to participate in the asymmetric properties of DC [10]. Delamination of apoptotic cells in the anterior amnioserosa produces forces that are responsible for a higher rate of anterior zipping. However, the phenotype that we observed with the abd-A or Abd-B mutation cannot be attributed to defects in this mechanism, as the rate of apoptosis is already very low in the posterior amnioserosa. Therefore, our data reveal a genetic control of zipping through precise transcriptional regulation in the LE. Overall, our work provides a framework for apprehending how the HOX selector genes and their cofactors collaborate with other signalling pathways to generate specific transcriptional responses allowing morphogenetic patterning and proper coordinated development.

Materials and methods

Fly stocks

The following fly stocks were used in this study: w1118 (BDSC#3601) as WT flies, y w hep’ and y w hep77 [1], UAS-hepact (BDSC#9306), pucE69 (puc-Z) [54], 69B-GAL4 (BDSC#1774), pnr-GAL4 (BDSC#3039), 69B-GAL4 (gift from A. Brand), AS-GAL4 (c381-GAL4; BDSC#3734), enX31 [55], UAS-en [56], UAS-HA::VP16::en [39], Scr17 (BDSC#3400), AntpNsrC41 (BDSC#1830), Ubx-1 (BDSC#626), abdA M1 and AbdB D18 [57], UAS-Scr::HA, UAS-Ubx::HA and UAS-abdA::HA [58],
UAS-Antp [59], UAS-AbdBm (BDSC#913), UAS-AbdBi (VDRC#12024)[60], UAS-Dicer2 (BDSC#24650), hthP2 [61], TM3,dfd-lacZ [62]. For live imaging, we used arm::GFP (BDSC#8556 on chromosome II; BDSC#8555 on chromosome III) and created the recombinant line arm::GFP,Abd-BD18 that we balanced with TM3,twi-GAL4;UAS-GFP (BDSC#6663) to select the homozygote mutant embryos. We also created the following line for the Abd-B RNAi: UAS-Dicer2;UAS-CD8::RFP,UAS-Abd-Bi, allowing the selection of AbdBi embryos thanks to the expression of CD8::RFP.

Microarray screen

Three genetic conditions were compared: wild-type (WT, w1118), gain of function (GOF, 69B-GAL4 > hep625) and loss of function (LOF, hep1/hep75). The 69B-GAL4 driver allows a uniform expression in the ectoderm. For the LOF embryos, we crossed y w hep1/Y females to y w hep1/Y males to obtain y w hep625/y w hep1 virgin females that were then crossed to y w hep1/Y males. 100% of the embryos coming from this last cross are mutant for DJNKK because hep1 behaves as a total loss of maternal function while being zygotically viable [1].

Two hour egg collections were incubated at 25˚C during the time necessary to obtain 85 to 90% embryos undergoing DC (~9 hrs). Because of a certain variability of development that we were not able to control, a fraction of each embryo was systematically tested by dpp in situ hybridization, and the collections that did not correspond to the definite criterion (above 85% of stage 13–14 embryos) were eliminated. Embryos from three biological replicates for each genetic condition were dechorionated with 50% bleach, frozen in nitrogen liquid and stored at -80C. Embryos were homogenized in RLT buffer (QIAGEN)/beta-mercaptoethanol using a conventional rotor–stator homogenizer. Total RNA was prepared with the Qiagen RNeasy mini kit for animal tissues according to the manufacturer’s instructions. Biotinylated cRNA were prepared according to the standard Affymetrix protocol from 2 μg total RNA. The biotinylation, the hybridization and the scan were done at the Affymetrix Platform located at the Institut Curie, Paris. Bioinformatics analysis was performed using RMA ("Robust Multi-array Average") and SAM ("Significance Analysis Microarray") under R/Bioconductor. The false discovery rate (FDR) of the GOF analysis is excellent (0.85%), evaluating the number of false positive to 14 out of the 1648 genes identified. The FDR associated with the LOF comparison is 3.19%, i.e. corresponding to 3 or 4 false positive out of the 113 isolated genes. The enrichment scores were obtained using the DAVID Bioinformatics Resources [27]. Comparison of microarray data sets was performed with THEA [29]. Microarray data have been deposited in the Gene Expression Omnibus (accession number GSE21805).

In situ hybridizations

The embryos were prepared using a standard protocol. Briefly, embryos resulting from 16hr collections at 25˚C, or 29˚C for the overexpression experiments, were dechorionated in 50% bleach, fixed in 4% formaldehyde and devitellinized in a heptane:methanol (1:1) mix. Embryos were then freshly used or kept at -20˚C for a few weeks. Digoxigenin (DIG)-labelled RNA probe synthesis was performed with T3, T7 or Sp6 RNA polymerases as recommended (Promega, New England Biolabs) from cDNA collections (Drosophila Genomics Resource Center). Classical in situ hybridization (ISH) experiments were done using a standard protocol with the anti-DIG antibody conjugated to alkaline phosphatase (1/2000; Roche Diagnostics). Staining was performed with NBT/BCIP reagent (Sigma). For fluorescent ISH coupled to immunofluorescence (FISH-IF), the second fixation step was accomplished using freshly prepared paraformaldehyde and the proteinase K treatment was omitted. Primary antibodies were: rabbit anti-En (1/100; Santa Cruz), mouse anti-Scr (1/50; DSHB), mouse anti-Antp (1/50; DSHB),
mouse anti-Ubx (1/50; DSHB), mouse anti-abd-A (1/100; DSHB), mouse anti-Abd-B (1/50; DSHB), mouse anti-HA (1/500; Covance), chicken anti-β Galactosidase (1/500; GeneTex).

Sheep anti-DIG antibody coupled to horseradish peroxidase (1/500; Roche Diagnostics) was joined to the secondary antibodies: anti-rabbit Al488 (1/400; Molecular Probe), anti-mouse Cy5 (1/100; Jackson ImmunoResearch), anti-chicken DyLight649 (1/200; Life Technologies). Revelation was done twice for 5–10 minutes with Tyramide Signal Amplification (PerkinElmer). Embryos were mounted in Mowiol 4–88 Reagent (Calbiochem). Images were acquired with a LSM 710 Zeiss confocal microscope using a 40X Objective.

mRNA signal quantification at the LE

Confocal Z sections that entirely encompass the LE of 15 to 20 embryos per genotype (except for the en mutant: n = 10) were acquired as 2048X2048–12 bit images and maximum intensity projections were created. To quantify the fluorescence signal in the LE, we developed CurvedPeriodicity, a standalone user-friendly program coded in Matlab (the Mathworks). The steps of the image pre-treatment are 1) to correct the background measured in the amnioserosa to linearize the intensity signal to fluorescence, 2) to extract the ribbon containing the LE and to make it linear, 3) to split the ribbon by semi-automatic demarcation of the segment boundaries, and 4) to normalize each segment in length. From these data, the software quantifies the mRNA signal as the mean intensity projections along the LE. It yields both the signal distribution per segment along the LE ($d_{LE}$) and the mean distribution along a Normalized Segment ($d_{NS}$). Segments are composed, on average, of 10 cells with 7 in the anterior compartment and 3 in the posterior compartment. Therefore $d_{NS}$ was divided in 10 regions and the parasegmental boundary was set between the 7th and 8th cells. CurvedPeriodicity calculates the average signal from each cell of each segment, and also the AP ratio per segment. Taking advantage of the normalization, the software is able to pool the confocal images per condition, synthesizing the data and extracting the mean and s.d. for $d_{LE}$ and $d_{NS}$ and to export it directly into an Excel file. In case of negative numbers (i.e. expression below background), the expression level was set to 0. For the AP analysis, the mean of the signals from each segment from T1 to A7 (n embryos; n = 15 to 20) was calculated. For the compartmental analysis, the signals for a given cell of all the segments (n segments = 150 to 200, with 10 segments per embryo) were averaged.

Quantitative PCR

We used the same RNA samples (control: w^{1118}, GOF: 69B-GAL4 > hep^{act}, LOF: hep/hec^{75}) than the ones prepared for the microarray analysis. Reverse-transcription was performed with SuperScript III Reverse Transcriptase (Invitrogen Life Technology) after DNase I digestion with a mix of oligo-dT and random primers. Q-PCR was performed with the Mastermix Plus for SYBR Green containing Rox (Eurogentec) with the endogenous rp49 gene for normalization. The list of primers that were used is available upon request. Standard curves of all the couples of primers presented an efficacy of amplification comprised between 95% and 110% with a coefficient of determination R² of at least 0.995. For each condition we did three technical replicates from one biological sample. Results were analysed with the StepOne software v2.1 (Applied Biosystems).

Cuticle preparations

Embryos were collected for 24 hours and incubated at 25°C or 29°C to let the wild-type larvae crawl away. The cuticles of unhatched (dead) embryos were prepared as following: embryos were dechorionated, mounted in 100 μl of a mix (1:1) made of lactic acid and Hoyer’s
mounting medium (30 g of gum Arabic, 50 ml of distilled water, 200 g of chloral hydrate, 20 ml of glycerol), and incubated overnight at 65˚C. Images were taken with a Nikon Coolpix 990 camera under a Leica DMR microscope.

**Immunostaining and quantification**

Embryos resulting from 16hr collections at 25˚C were dechorionated in 50% bleach, fixed in 4% formaldehyde and devitellinized in a heptane:methanol (1:1) mix. Embryos were then quickly rinsed with PBS-triton X-100 0.1% and the following primary antibodies were then used: rabbit anti-En (1/200; Santa Cruz), mouse anti-Ena (1/100, DSHB), rat anti-ECad (1/100, DSHB) and/or mouse anti-Dlg (1/200, DSHB). After washes, secondary antibodies were added: anti-rabbit Al488 (1/400; Molecular Probe), anti-mouse Al488 (1/400; Molecular Probe), anti-mouse Al546 (1/400; Molecular Probe), anti-rat Al546 (1/400; Molecular Probe) and/or anti-rat Al647 (1/400; Jackson ImmunoResearch). After washes, embryos were mounted in Mowiol 4–88 Reagent (Calbiochem). Images were acquired with a LSM 780 Zeiss confocal microscope using a 25X or 63X Objective.

Closure delay was quantified by measuring with ImageJ the distance between the matching posterior compartments (stained with En) of the T1 (D-A) or A7 (D-P) segment when the opposing segment (A8 or T1, respectively) just closed. For the zipping zone, quantification was done with ImageJ by tracing a line from the point where the two opposing LE are in close contact to the point where a stable junction (adherens junction with ECad or septate junction with Dlg) is formed.

**Live imaging and quantification**

Embryos collected from overnight egg-laying were dechorionated (50% bleach) and stage 13 embryos (at the onset of DC) were selected under the binocular. They were mounted in Halocarbon 95 oil between two cover slips separated by spacers, glued on their ventral part. Two hydrating chambers (watered cotton) were positioned on the sides. 1024X1024 Z-stacks (1 to 2 μm/image) were acquired over 4 hours with a Zeiss 780 confocal microscope. Distances necessary for calculation of the speed of closure, H and L (Fig 7) were measured from maximum intensity projections of optimized Z sections with FIJI. Stages of DC (expressed as % of DC) were estimated by calculating the ratio of the seam (fused LE) and L (projected LE at the mid-line). L was divided in 8 sub-domains (corresponding approximately to 8 segments), as previously described [63].

**Supporting information**

S1 Fig. *Microarray analysis of the JNK response during DC.* A) *In situ* hybridizations of stage 13 embryos showing *dpp* expression in the three conditions used to prepare the RNAs for microarray analysis: WT (*w^{118};* top), GOF (69B-GAL4 > *hep*{	extsuperscript{ok}}; middle) and LOF (hep{	extsuperscript{1}/hep{	extsuperscript{75}}}; bottom). Activation of the JNK pathway in the ectoderm with 69B-GAL4 leads to a lateral expansion of *dpp* expression (dotted area), whereas in the hep{	extsuperscript{1}/hep{	extsuperscript{75}}} mutant, *dpp* expression is lost specifically in the LE, but not in the other tissues. B) Distribution of the number of genes that are up-regulated (green) and down-regulated (red) by the JNK pathway (with a fold change (FC) superior to 1.5 or inferior to -1.5) in the GOF (top) and LOF conditions (bottom).

(TIF)

S2 Fig. *Anterior-posterior expression of the twelve LE genes.* FISH-IF (first and third columns) and mRNA signal quantifications at the LE (second and fourth columns) are shown for
each LE gene. mRNA staining is in red and anti-En immuno-fluorescence is in green. Bar diagram: average mRNA signal intensities at the LE in each segment, expressed as fluorescent intensity (a.u. +/- s.e.m.). This figure is a supplement of Fig 3B.

(TIF)

S3 Fig. Segmental expression of the twelve LE genes. FISH-IF (columns 1, 2, 4 and 5) and mRNA signal quantifications at the LE (columns 3 and 6) are shown for each LE gene. mRNA staining is shown in red alone (columns 1 and 4) or merged with anti-En immuno-staining (in green)(columns 2 and 5). Bar diagram: average mRNA signal intensities at the LE in the ten cells of the segments, expressed as fluorescent intensity (a.u. +/- s.e.m.). This figure is a supplement of Fig 3C.

(TIF)

S4 Fig. Regulation of scaf expression by en and the HOX genes. A) Quantification of scaf expression (expressed as fluorescent intensity; a.u. +/- s.e.m.) in the LE of en mutant embryos (n = 10, FISH-IF shown in Fig 4A). The anterior limit for quantification corresponds to the dorsal ridge abutting the head segments, whereas the posterior limit was set just anteriorly to the place where scaf expression starts to expand in the more lateral epidermis (see Fig 4A). The LE was then divided in 10 equal segments for quantification with CurvedPeriodicity. Top panel: quantification in the A (blue) and P (green) compartments along the AP axis. Bottom panel: quantification in the A and P compartments of segments A1 to A6 of the en mutant, compared to WT embryos, showing the absence of negative regulation on scaf expression in the P compartments. B) FISH-IF showing scaf expression (scaf mRNA is in red either merged or not with anti-En staining in green) in the abd-A mutant (quantification shown in Fig 5A). C) FISH-IF in embryos overexpressing abd-A with pnr-GAL4 (quantification shown in Fig 5B). D) scaf expression in Scr (first column), Antp (second column), Ubx (third column) and Abd-B (fourth column) mutants. First row: quantification of scaf expression (expressed as fluorescent intensity; a.u. +/- s.e.m., n = 15 for each mutant) in each compartment (anterior: blue; posterior: green) of each segment along the AP axis. Each main expression domain of the corresponding HOX gene is indicated by a color: grey for Scr, pink for Antp, blue for Ubx and orange for Abd-B. Second row: FISH-IF showing scaf mRNA (red). Third row: FISH-IF showing scaf mRNA (red) with anti-En staining (green). This figure is a supplement of Fig 5D.

(TIF)

S5 Fig. abd-A and Abd-B posterior closure defects. A) Speed of closure (nm/s +/- s.e.m.) in the anterior and posterior regions of control (pink; n = 9) and Abd-B mutant (purple; n = 6) embryos during early DC (seam < 33%), mid DC (33% < seam > 66%) and late DC (seam > 66%). A strong reduction of the posterior closure speed was observed from mid DC (59%) to late DC (78%) in Abd-B embryos. The anterior speed also decreased, especially during late DC (58%). This panel is a supplement of Fig 7D and 7E. B) and C) Dlg-defined zipping zones of WT, abd-A and Abd-B embryos. B) Anti-Dlg (red) immunostainings showing the posterior zipping zones (Zip-P) in the A6 segment of a WT embryo (left) and of abd-A (middle) and Abd-B (right) mutants. The zipping zone is defined by the area where the two opposing LE are in close contact till the formation of a stable septate junction (marked with Dlg). C) Quantification of the anterior (A, blue) and posterior (P, green) zipping zones of the WT embryo and the two HOX mutants. Whereas no variation is observed in the anterior part (no statistical significance), the Zip-P of abd-A (7.7 μm +/- 0.5) and Abd-B (7.3 μm +/- 0.7) are reduced compared to WT embryos (11.2 μm +/- 1.0). As there is no homogeneity of variance for Zip-A, the
Kruskal & Wallis test was performed with R, whereas multiple comparisons of Zip-P between WT, \textit{abhd-A} and \textit{Abd-B} were performed using the Dunnett test (\(^*: p < 0.05; **: p < 0.01; ***: p < 0.001\)). For WT embryos, \(n(\text{Zip-A}) = 7\) and \(n(\text{Zip-P}) = 13\); for \textit{abhd-A}, \(n(\text{Zip-A}) = 9\) and \(n(\text{Zip-P}) = 14\); for \textit{Abd-B}, \(n(\text{Zip-A}) = 11\) and \(n(\text{Zip-P}) = 15\). This figure is a supplement of Fig 7F and 7G. D) and E) Inhibition of \textit{Abd-B} expression in the dorsal ectodermal cells induces posterior DC phenotype. 

D) Anti-En (green) and anti-RFP (red) immunostainings of embryos expressing \textit{Abd-B} RNAi \((\textit{Abd-B}^{i})\) and CD8::RFP either with \textit{pnrt-GAL4} (left panels) or with \textit{AS-GAL4} (right panels). Whereas expression in the AS alone has no effect (when A8 closes, T1 is just about to close), expression in the dorsal ectoderm and transiently in the AS with \textit{pnrt-GAL4} triggers a delay in the posterior closure: when A8 closes, the anterior segments T1, T2 and T3 are all closed (blue arrow). E) Quantification of the penetrance of the posterior DC defects (\%) of \textit{pnr} > \textit{Abd-B}^{i}, CD8::RFP \((n = 12)\) and \textit{AS} > \textit{Abd-B}^{i}, CD8::RFP \((n = 16)\). The Fisher’s exact test (nonparametric test for two independent samples and a binomial distribution) was used (***: \(p < 0.001\)).

S1 Movie. \textit{arm-GFP} (left) and \textit{AbdB}^{D18,arm-GFP} (right) live embryos. Frames were taken every 20 minutes.

S1 Table. Common genes of the GOF and LOF microarray screen with their corresponding fold change (FC).

S2 Table. Over-represented Gene Ontology (GO) terms in the list of JNK up-regulated genes in the GOF screen (using DAVID).

S3 Table. Over-represented Gene Ontology (GO) terms in the list of JNK down-regulated genes of the GOF screen (using DAVID).

S4 Table. Description of the 31 JNK target genes.

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