



Double abdomen in a short-germ insect: Zygotic control of axis formation revealed in the beetle *Tribolium castaneum*

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The distinction of anterior versus posterior is a crucial first step in animal embryogenesis. In the fly *Drosophila*, this axis is established by morphogenetic gradients contributed by the mother that regulate zygotic target genes. This principle has been considered to hold true for insects in general but is fundamentally different from vertebrates, where zygotic genes and Wnt signaling are required. We investigated symmetry breaking in the beetle *Tribolium castaneum*, which among insects represents the more ancestral short-germ embryogenesis. We found that maternal *Tc-germ cell-less* is required for anterior localization of maternal *Tc-axin*, which represses Wnt signaling and promotes expression of anterior zygotic genes. Both RNAi targeting *Tc-germ cell-less* or double RNAi knocking down the zygotic genes *Tc-homeobrain* and *Tc-zent1* led to the formation of a second growth zone at the anterior, which resulted in double-abdomen phenotypes. Conversely, interfering with two posterior factors, *Tc-caudal* and Wnt, caused double-anterior phenotypes. These findings reveal that maternal and zygotic mechanisms, including Wnt signaling, are required for establishing embryo polarity and induce the segmentation clock in a short-germ insect.

axis formation | germ cell-less | homeobrain | Torso signaling | short-germ segmentation

One of the most important first steps in animal embryogenesis is the establishment of anterior-posterior polarity (AP polarity), distinguishing head from tail. If this process fails, mirror image duplications of embryonic structures may develop like the “double-abdomen” embryos induced by classical manipulations of insect embryos by ligation or UV-irradiation experiments (1–3). Based on these types of experiments, autonomous patterning mechanisms were proposed for the insect egg (4).

In the fly *Drosophila melanogaster*, mRNA of the *bicoid* gene is localized to the anterior pole from where the Bicoid protein diffuses to form an anterior-to-posterior gradient. An opposing gradient of Nanos protein forms from posterior to regulate Hunchback translation, which thus forms another anterior gradient. In addition, the graded activity of the Torso pathway directs patterning near both poles of the *Drosophila* embryo. Together, these maternal morphogenetic gradients provide positional information and initiate zygotic segmentation gene expression in a concentration-dependent manner (5, 6). Mutations in *bicoid* result in duplication of some posterior structures at the anterior, while symmetric larvae with “double abdomen” form after double inactivation of both Bicoid and Hunchback (7) and a number of other genotypes (8, 9) where *bicoid* and *nanos* localization is impeded. In this system, zygotic genes interpret the maternal information but are not involved in axis formation themselves. However, axis formation in *Drosophila* is derived and *bicoid* is not present outside higher dipterans (10). In the lower dipteran *Chironomus*, another long-germ fly, axis formation relies on maternal anterior localization of *panish*, a putative repressor related to the Wnt component *pangolin*. Knockdown of *panish* leads to double abdomina, while knockdown of the zygotic gene *tailless* leads to double heads, revealing an alternative genetic mechanism for axis

formation (11). In the wasp *Nasonia*, a more distantly related long-germ insect, *orthodenticle (otd)* provides maternal morphogenetic gradients at both poles. *giant*, which is maternally localized at the anterior pole, substitutes for the permissive role of *bicoid* at the anterior (12, 13), but axis duplication phenotypes have not been described.

However, all these long-germ embryos lack the predicted autonomous posterior patterning mechanism. In contrast, typical short-germ insect embryos form posterior segments sequentially from a “growth zone” or segment addition zone (SAZ), which is specified at the posterior pole of the blastoderm. Further, in short-germ insects, extraembryonic tissue is formed at the anterior blastoderm, rather than head structures as in *Drosophila*. The red flour beetle *Tribolium* is a model for short-germ embryogenesis, where the formation of abdominal segments requires *Tc-caudal* expression as well as Wnt and Torso signaling at the posterior pole. Torso signaling appears to be the only conserved maternal signal. It had been suggested that *Tc-otd* and *Tc-hunchback* substitute for the anterior morphogen in the beetle *Tribolium* (12, 14). However, more recent data do not support a morphogenetic role of these genes in anterior patterning (15, 16). Instead, anterior localization of maternal *Tc-axin* mRNA represses Wnt signaling, providing a permissive environment for anterior development, as in vertebrate embryos (17–20). Ubiquitously distributed *Tc-caudal* mRNA is translationally repressed by anterior *Tc-Mex-3* and *Tc-Zen2*, but their knockdown does

Significance

One of the first crucial steps of animal development is to distinguish the anterior versus the posterior pole of the embryo, i.e., the AP axis. If this process fails, embryos may develop two mirror image tails or heads. In the fly *Drosophila*, the mother provides the signals required for AP axis formation, while in vertebrates, gene activity of the embryo is required as well. We identified two genes whose knockdown leads to double-tail phenotypes in the beetle *Tribolium*, representing the insect-typical short-germ embryogenesis. Intriguingly, embryo polarity depends on zygotic gene activities and Wnt signaling. Hence, short-germ insect axis formation is more similar to vertebrates than the mechanism employed by *Drosophila*.

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not lead to double-abdomen phenotypes (21). In summary, the mechanism of symmetry breaking and subsequent AP-patterning in short-germ embryos is predicted to be different but has remained obscure. Notably, genetically induced mirror image phenotypes have not been observed so far. Likewise, the predicted autonomy of SAZ patterning has not been demonstrated.

Results

Unexpected Genes in AP Axis Formation. In an ongoing genome-wide RNAi screen we searched for cuticle phenotypes affecting the anterior-posterior body axis (22, 23). We found two genes whose knockdown led to double-abdomen phenotypes. One of these dsRNAs, iB_02693, targeted the ortholog of the *Drosophila* *germ cell-less* gene (*gcl*), which in the fly is required at the posterior pole for germ cell development. It encodes a BTB domain protein localized at the nuclear lamina in both *Drosophila* and vertebrates, where it appears to be involved in transcriptional regulation (24–26) or cytoskeleton organization (27, 28). Knockdown of *Tribolium* *germ cell-less* (*Tc-gcl*) results in double-abdomen phenotypes with high penetrance (>95%), where head and thorax are replaced by a mirror image abdomen (Fig. 1 *B* and *C* and *SI Appendix*, Fig. S1). Interestingly, in some specimens the total number of segments of these two abdomina (2×10 abdominal segments) exceeds the normal number of segmentation-clock-derived segments (16 segments), demonstrating the autonomy of the duplicated SAZ. Some segmentation and dorsal closure defects occur as well.

We found maternal *Tc-gcl* transcripts to be localized to the anterior half of the developing oocyte and the freshly laid egg (Fig. 1 *G–J*). We generated an antibody detecting Tc-Gcl. As in *Drosophila*, we found the protein at the nuclear envelope, which was marked with a nuclear pore protein (Fig. 1 *M* and *N*). Later,

a posterior domain with similarity to *Tc-vasa* expression developed de novo (Fig. 1 *I* and *SI Appendix*, Fig. S2 *C, G, and H*). However, neither dsRNA injected into embryos, nor into L5 larvae, resulted in strongly reduced egg production, suggesting that in *Tribolium*, *gcl* may be dispensable for germ cell development. dsRNA injection into early embryos did not lead to lethality or cuticle phenotypes, indicating a predominantly maternal function of *Tc-gcl*.

The second dsRNA (iB_04564) causing double abdomina targeted the ortholog of *homeobrain*, which is a homeodomain transcription factor expressed in the head and brain primordium of *Drosophila*, but for which a function has not been described (29). Knockdown of *Tribolium homeobrain* (*Tc-hbn*) resulted in anterior deletions or double-abdomen phenotypes (Fig. 1 *D–F*). Both penetrance and expressivity of double abdomina were lower compared with *Tc-gcl* knockdown (*SI Appendix*, Fig. S1 *O and P*), and thoracic segments remained present even in the strongest phenotypes (*SI Appendix*, Fig. S1). *Tc-hbn* is expressed as one of the earliest zygotic genes in an anterior cap of the blastoderm, which later retracts from the pole and eventually forms bilateral domains in the head anlagen (Fig. 1 *J–L* and *SI Appendix*, Fig. S2). Both phenotypes were reproduced by RNAi using nonoverlapping dsRNA fragments in the same genetic background. It was unexpected to find axis duplication upon knockdown of a zygotic gene since the current insect paradigm holds that maternal morphogens establish the axes, while zygotic genes then interpret them.

To characterize the mechanism of axis duplication we scrutinized the expression of SAZ markers in RNAi embryos (Fig. 2). *Tc-gcl* RNAi induced a mirror image duplication of *Tc-even-skipped* (*Tc-eve*) and *Tc-vasa* expression from early blastoderm stages onward, while *Tc-caudal* (*Tc-cad*) was expressed ubiquitously (Fig. 2 *B, E*,

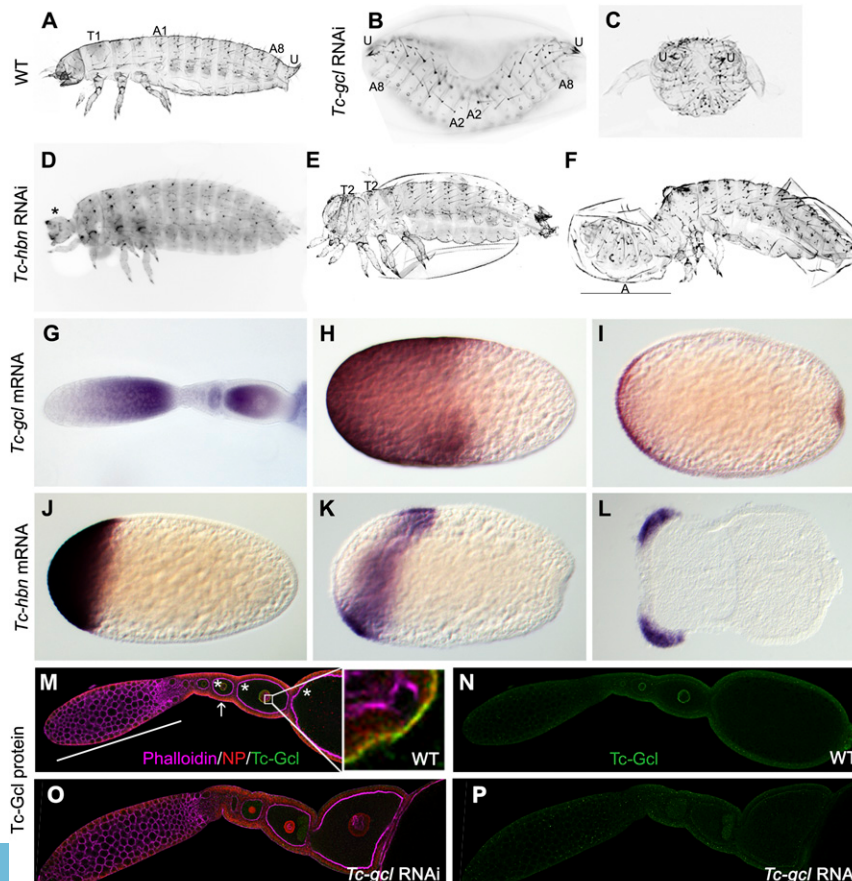


Fig. 1. *Tc-gcl* and *Tc-hbn* RNAi phenotypes and expression. (A) Wild-type first-instar larval cuticle with thoracic (T) and abdominal (A) segments and urogonphii (U). (B and C) *Tc-gcl* RNAi larvae lacking the thorax but displaying mirror image abdomina. (D–F) *Tc-hbn* RNAi phenotypes range from loss of the labrum and antennae (asterisk in D) to specimen with partially duplicated thorax (E) and incomplete double abdomen (F). (G–I) *Tc-gcl* mRNA is found in nurse cells and in the anterior of developing oocytes (G) and in the anterior half of freshly laid eggs (H). In differentiated blastoderm embryos, expression is found at the anterior and posterior poles (I). (J–L) Zygotic *Tc-hbn* mRNA is expressed in an anterior cap (J) but later retracts from the pole (K) to form domains in the head lobes of germ rudiments (L). (M and N) *Tc-Gcl* protein distribution in ovaries. *Tc-Gcl* (green signal; single channel shown in N) is found at the nuclear envelope, which is marked by a nuclear pore protein (red signal; see *Inset* for blow-up). White bar, nurse cells; stars, oocytes; white arrow, follicle cells. (O and P) After *Tc-gcl* RNAi, the nuclear staining is reduced (single channel shown in P). Anterior is to the Left in A–L; A–F are not in the same scale.

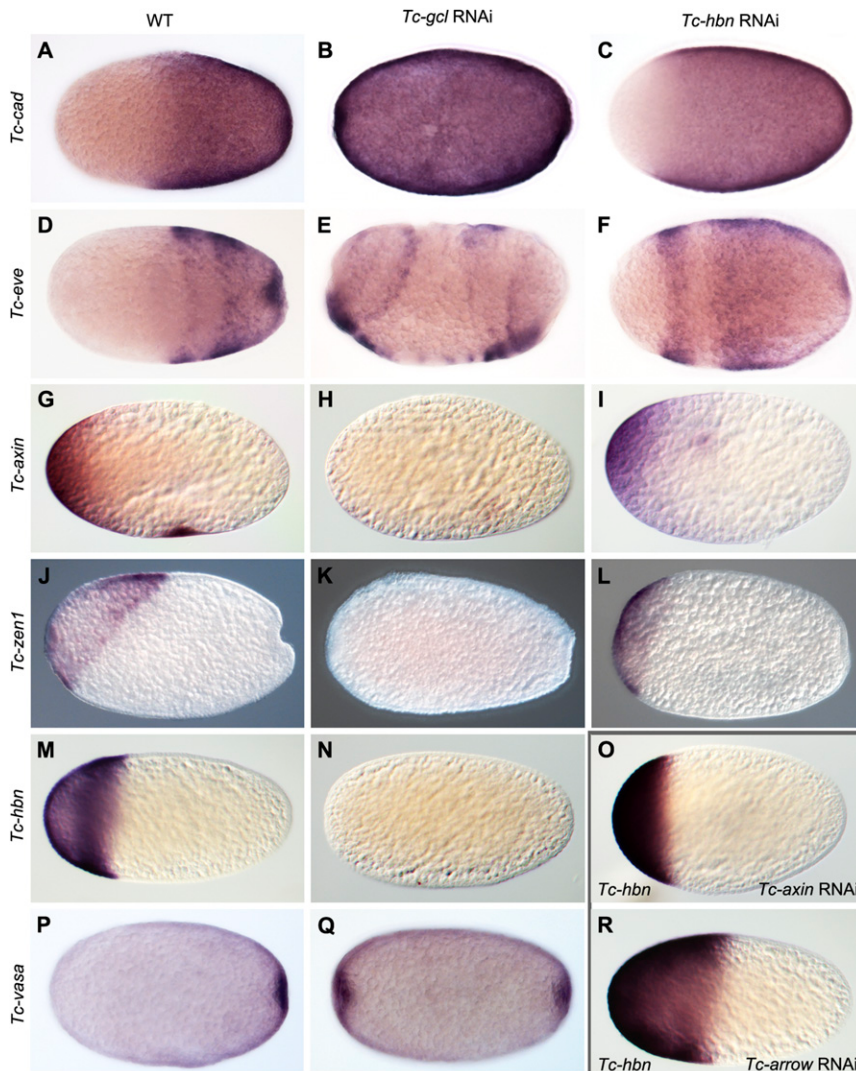


Fig. 2. Change of embryo fate map after *Tc-gcl* and *Tc-hbn* RNAi. (A–C) *Tc-cad* mRNA is expressed in the posterior half of WT blastoderm embryos (A) but is ubiquitously distributed after *Tc-gcl* RNAi (B). It is shifted anteriorly after *Tc-hbn* RNAi (C). (D–F) Striped *Tc-eve* mRNA expression at late WT blastoderm embryos (D) is duplicated after *Tc-gcl* RNAi (E) but shifted anteriorly in *Tc-hbn* RNAi (F). (G–I) Maternal *Tc-axin* mRNA at the anterior pole (G) is lost after *Tc-gcl* RNAi (H) but not affected after *Tc-hbn* RNAi (I). (J–L) *Tc-zn1* expression in the serosa (J) is lost after *Tc-gcl* RNAi (K) and reduced upon *Tc-hbn* RNAi (L). (M–O and R) *Tc-hbn* expression at the anterior pole (M) is absent after *Tc-gcl* RNAi (N). In *Tc-axin* RNAi (Wnt derepression) the boundary is shifted toward anterior (O), while it is shifted posteriorly after *Tc-arrow* RNAi (Wnt repression; R). (P and Q) *Tc-vasa* mRNA expression marking the posterior pit in WT (P) is duplicated after *Tc-gcl* RNAi (Q).

and Q). Early *Tc-wingless* (*Tc-wg*) expression in the SAZ was duplicated as well (SI Appendix, Fig. S3 A and B), while its segmental pattern prefigured the duplication seen in the cuticles (SI Appendix, Fig. S3). In *Tc-hbn* knockdown embryos, in contrast, the anterior expression boundary of these genes was only shifted anteriorly during blastoderm stages (Fig. 2 C and F), while *Tc-vasa* expression was not altered. Later, in germ band embryos, duplicated expression domains of *Tc-cad* and *Tc-wg* arose in the head, which developed into a mirror image SAZ (SI Appendix, Fig. S4 A–I).

Time and location of ectopic SAZ formation in *Tc-gcl* knockdown embryos corresponds to a region with active Torso signaling, in line with the notion that Torso signaling is required for SAZ formation (20, 30). The fact that a functional SAZ can form at both poles of the blastoderm as well as later in the head of postblastodermal embryos, strongly supports the idea of autonomy of SAZ patterning; i.e., the SAZ has self-organizing capabilities.

Tc-Gcl Acts Upstream in Anterior Patterning. Next, we sought to establish the genetic interactions involved in ectopic SAZ formation. The maternal expression of the anteriorly localized morphogen *Tc-axin* (18), a negative regulator of Wnt signaling, was strongly reduced or absent upon *Tc-gcl* knockdown (Fig. 2 G and H). This was not a general effect on maternally localized mRNAs because *Tc-pangolin* (*Tc-pan*) remained properly localized at the anterior pole. Both the expressions of *Tc-zn1*, a

marker for extraembryonic serosa (Fig. 2 J and K), and of *Tc-hbn* were abolished (Fig. 2 M and N). Conversely, *Tc-gcl* expression was not altered in *Tc-axin*, *Tc-pan*, or *Tc-hbn* knockdown. These results place *Tc-gcl* upstream of both maternally localized *Tc-axin* and of the earliest known zygotic anterior markers, *Tc-zn1* and *Tc-hbn*.

The lack of anteriorly localized *Tc-axin* could be due to reduction of transcription in the nurse cells, reduced transport of *Tc-axin* mRNA to the oocyte, or due to lack of anterior localization within the oocyte. To distinguish between these possibilities, we performed *in situ* hybridization in wild-type and *Tc-gcl* RNAi ovaries. We found that in RNAi ovaries, the signal in oocytes was reduced compared with the one in nurse cells (Fig. 3 J–M). qPCR confirmed that the *Tc-axin* message was significantly reduced 3.5-fold in both freshly laid eggs and ovaries (which included late oocytes; SI Appendix, Fig. S6). Hence, Tc-Gcl appears to act in transport either by a nontranscriptional role in cytoskeletal regulation or by transcriptional regulation of another factor required for mRNA transport (27, 28).

We wondered if the *gcl* phenotype can be phenocopied through direct interference with early Wnt signaling. Knocking down the Wnt pathway components *Tc-axin*, *Tc-arrow*, or *Tc-pan* did not produce double abdomina; using molecular markers to probe early embryos, we found that the expression boundaries of *Tc-hbn* and *Tc-zn1* were shifted posteriorly in Wnt-low and anteriorly

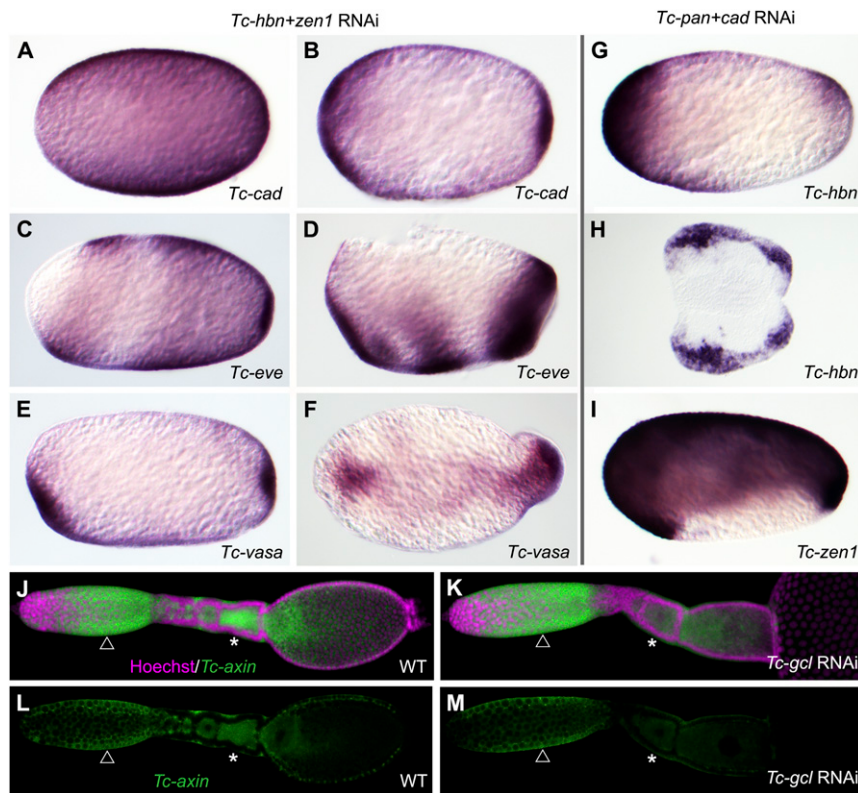


Fig. 3. Zygotic genes required for anterior and posterior structures. (A–F) *Tc-hbn*+*Tc-zen1* double RNAi leads to mirror image expression of SAZ markers like *Tc-cad* (A and B), *Tc-eve* (C and D), and *Tc-vasa* (E and F). (G–I) Upon *Tc-pan*+*Tc-cad* double RNAi, anterior markers become expressed at the posterior pole. *Tc-hbn* mRNA develops an additional posterior domain (G) and marks duplicated posterior head lobes in germ rudiments (H). *Tc-zen1* expression covers both poles and dorsal tissue (I). (J–M) In *Tc-gcl* RNAi ovaries, *Tc-axin* mRNA remains expressed in nurse cells (compare arrow in J and L with K and M) but is reduced in oocytes (compare stars in J and L with K and M). This indicates that *Tc-axin* transport is affected in *Tc-gcl* RNAi. Respective WT embryos are shown in Fig. 1.

in Wnt-high situations (*SI Appendix*, Fig. S5). However, none of the treatments completely abolished their expression. Further, the shift of *Tc-hbn* was small in early blastoderm embryos but became prominent subsequently (*SI Appendix*, Fig. S5 B–D and T–V). Hence, interference with early Wnt signaling alone did not phenocopy the *Tc-gcl* phenotype in our hands, while recently, double-abdomen phenotypes have been described upon *Tc-axin* RNAi (31).

We next tested whether *Tc-axin* might cooperate with some other *gcl*-dependent factor to repress anterior SAZ formation. However, double-RNAi treatments of *Tc-axin* combined with either *Tc-zen1*, *Tc-zen2*, *Tc-hbn*, *Tc-tll*, or *Tc-pan* also did not produce double abdomen. Only in very few *Tc-axin*/*Tc-zen1* double-RNAi embryos did we observe a small ectopic *Tc-cad* domain in the head anlagen of posterior pit-stage embryos (*SI Appendix*, Fig. S4P). Likewise, double knockdown of *Tc-pan* with either *Tc-zen1* or *Tc-hbn* did not lead to double abdomen on the cuticle level.

Drosophila gcl dampens Torso signaling by targeting the receptor for degradation (32). Hence, we tested for a similar role in anterior patterning in *Tribolium*. In wild-type embryos, activated MAPK emerges first at the anterior pole and remains more active than at the posterior pole throughout the blastoderm, suggesting that anterior Torso signaling is not negatively affected by anteriorly localized *Tc-Gcl* in the embryo. Subsequently, the anterior domain expands and retracts into a sub-terminal ring, while the activity directly at the anterior pole becomes weaker compared with the posterior domain (*SI Appendix*, Fig. S7). This late effect might reflect dampening of anterior Torso signaling by accumulating *Tc-Gcl* protein translated from maternally contributed *Tc-gcl* mRNA. In *Tc-gcl* RNAi embryos, the early pattern of activated MAPK is comparable to wild type, while from blastoderm stages onward, symmetric MAPK

activity reflects the mirror image symmetry of the embryo (*SI Appendix*, Fig. S7).

Zygotic Genes Contribute to Axis Formation. It was unexpected that RNAi targeting zygotic *Tc-hbn* resulted in axis duplication. However, this phenotype was weaker and less penetrant than the one of maternal *Tc-gcl*, suggesting *Tc-hbn* might be only one of several zygotic effectors. Indeed, combining *Tc-hbn* with *Tc-zen1* (but not *Tc-zen2*) resulted in a substantially more penetrant and stronger double-abdomen phenotype (*SI Appendix*, Fig. S1Q), including some additional segmentation and dorsal closure defects similar to *Tc-gcl* RNAi (*SI Appendix*, Fig. S4 Q and R). Moreover, we observed near-homogeneous expression of *Tc-cad* and an increased anterior shift of *Tc-eve* expression in blastoderm embryos (Fig. 3 A–D), while *Tc-vasa* displayed mirror image expression at both poles (Fig. 3 E and F). Further, we found mutual activation of *Tc-hbn* and *Tc-zen1* RNAi (Fig. 2L and *SI Appendix*, Fig. S4 S–U). In summary, the zygotic genes *Tc-hbn* and *Tc-zen1* act synergistically in suppressing an anterior SAZ, thereby preventing axis duplication.

We wondered whether zygotic genes might likewise be involved in specifying the posterior pole. In *Tribolium*, both posterior *Tc-cad* expression and Wnt signaling are required for SAZ formation (17, 21, 33). Hence, we hypothesized that *Tc-cad* activity at the posterior pole cooperates with Wnt signaling to repress anterior patterning. Indeed, in the double knockdown of *Tc-cad* and *Tc-pan*, we found *Tc-hbn* and *Tc-zen1* to become ectopically expressed at the posterior pole of the blastoderm, indicating the ectopic specification of anterior tissues (Fig. 3 G and I). Such ectopic expression was not found in *Tc-pan* single knockdown (*SI Appendix*, Fig. S5), while *Tc-cad* single RNAi did not lead to the

formation of extraembryonic tissue at the posterior pole (34). In germ-rudiment embryos, both *Tc-hbn* expression and embryo morphology indicated that a second head anlage had developed at the posterior (Fig. 3H and *SI Appendix*, Fig. S4 J–O; these embryos did not produce cuticles).

Discussion

A Model for Axis Formation in Short-Germ Embryos. In the freshly laid egg (Fig. 4, Top), maternally contributed *Tc-cad* mRNA and Tc- β -Catenin protein are initially evenly distributed (35). Torso signaling is active at both poles, being required for the formation of the serosa at the anterior and the SAZ at the posterior pole (20, 30, 36). Symmetry breaking is initiated by maternal Tc-Gcl, which is required to establish *Tc-axin* mRNA at the anterior pole. There, Tc-Axin protein leads to degradation of Tc- β -Catenin, resulting in repression of canonical Wnt signaling at the anterior pole (Fig. 4, Middle and *SI Appendix*, Fig. S8). In the absence of Wnt signaling, Torso signaling contributes to *Tc-zn1* expression (36) and allows for expression of zygotic *Tc-hbn* at the anterior pole (Fig. 4, Bottom). Once activated, *Tc-hbn* and *Tc-zn1* form a zygotic feedback loop, which further promotes anterior fates by repressing *Tc-cad*. Two factors repress translation of maternal *Tc-cad* mRNA: *Tc-zn2*, which is activated by *Tc-zn1*, and *Tc-mex3* (21, 37). Later, *Tc-zn1* is required for specifying the extraembryonic serosa. Hence, our data indicate that *Tc-gcl* acts predominantly via the

Wnt pathway, although we cannot exclude additional functions. At the posterior pole, Wnt and Torso signaling together with maternal *Tc-cad* initiate a feedback loop between Wnt signaling and *Tc-cad* to form a signaling center that specifies the SAZ (Fig. 4, Bottom and *SI Appendix*, Fig. S8). Indeed, Wnt activates *Tc-cad* (38) and *Tc-wg* (38) expression in *Tribolium*, but it remains unclear whether *Tc-cad* activates the expression of Wnt ligands like it does in spiders (39) but not in *Gryllus* (40).

This model explains the phenotypes we observe: Removing localized *Tc-axin* by *Tc-gcl* RNAi results in derepression of Wnt signaling and *Tc-cad* translation at the anterior, which together with Torso signaling initiate a SAZ at the anterior pole like they do at the posterior pole (33, 36, 38, 41). In this model, all three components are required for specifying the SAZ, while Wnt signaling and *Tc-cad* are sufficient to suppress the expression of *Tc-hbn* and *Tc-zn1* in the posterior Torso domain.

Zygotic Regulation Instead of Maternal Gradients. In the long-germ embryos of both *Drosophila* and *Nasonia*, maternal morphogenetic gradients of transcription factors regulate zygotic genes in a concentration-dependent manner (6, 12, 42). Zygotic genes are not involved in axis specification, and respective mutants lead to loss of body regions but not to axis duplications. Our data suggest that this paradigm does not hold true for short-germ insects. Here, initial asymmetry provided by maternal signals needs to be reinforced by two zygotic signaling centers located at both poles, respectively. Hence, in contrast to *Drosophila*, knockdown of zygotic genes (*zen1* and *hbn*) does lead to posterior axis duplications. Wnt and *cad* probably form another zygotic feedback loop required for maintaining the growth zone. A more central role of zygotic control in AP axis formation is corroborated by results in the dipteran *Chironomus*, where knockdown of the zygotic gene *tailless* is sufficient to produce double heads (11). Likewise, dorsal-ventral axis formation is based on zygotic self-regulatory circuits in *Tribolium* and on a self-organizing gene network in the hemimetabolous insect *Oncopeltus*, contrasting the hierarchical maternal control in *Drosophila* (43).

Taken together, these insights indicate that insect and vertebrate axis formation have more in common than had been assumed based on the *Drosophila* system. First, initial symmetry breaking based on maternal molecules leads to zygotic interactions, which are required to establish the axes (44–46). Second, the repression of anterior Wnt signaling is required in the vertebrate anterior neural plate (47, 48), the anterior of ancestral animal embryos (49) and *Tribolium* (18), but not in *Drosophila*. It is tempting to speculate that the same is true for *Chironomus*, where the anterior determinant has similarity to a Wnt pathway component (11). Intriguingly, *Tribolium* uses a *Drosophila*-like mechanism (anterior localization of maternal mRNAs) to realize a vertebrate-like situation (i.e., anterior repression of Wnt signaling).

Materials and Methods

RNAi. Initial validation was performed in the same genetic background as in the iBeetle screen (23). All further experiments were done in the pig19 strain (50). dsRNA was synthesized with the Ambion T7 MEGAscript kit (Life Technologies), and pupae, larvae and embryos were injected as described previously (51, 52) using FemtoJet (Eppendorf). Off-target effects for *Tc-gcl* and *Tc-hbn* were tested by nonoverlapping fragments (*SI Appendix*, Fig. S1). dsRNA concentrations are listed in *SI Appendix*, Table S2.

Generation of a Tc-Gcl Polyclonal Antibody. The N terminus (amino acids 1–57) was cloned into pET SUMO vector, generating a fusion with a His-SUMO tag and expressed in BL21-DE3 Rosetta cells. Antibodies were produced and affinity purified in guinea pigs by Eurogentec.

Ovary WMISH and Antibody Staining. Embryonic and ovary WMISH were performed as described previously (38, 53, 54). Phospho-p44/42 MAPK Rabbit mAb (Cell Signaling Technology, Inc.) with 1:1,000 dilution was used to analyze the Torso signaling during the early embryogenesis. Images were acquired with a Zeiss AxioPlan 2 LSM510.

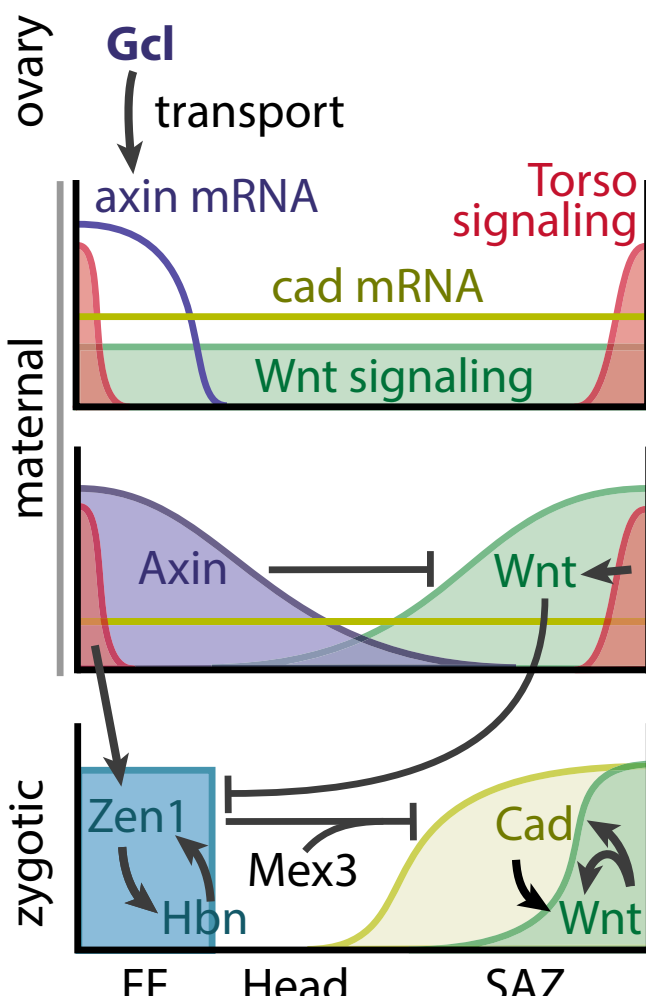


Fig. 4. Model for axis formation *Tribolium castaneum*. See text for details. EE, extraembryonic tissue; SAZ, segment addition zone.

Real-Time qPCR. Total RNAi was extracted from 0- to 4-h-old RNAi and WT embryos and RNAi and WT ovaries using TRIzol (Ambion) (55). One microgram for embryos and 500 ng for ovaries was used for the first strand cDNA synthesis (SuperScript III; Thermo Scientific). qRT PCR was performed in CFX96 Real-Time PCR System (Bio-Rad Laboratories) with HOT FIREpol Eva-Green qPCR Mix Plus (ROX) (Solis BioDyne). Three biological and three technical replicates and no-RT and no template control were performed. Relative fold-change expression between WT and RNAi tissues was calculated by double-delta Ct method with RPS18.3 as ref. 56. Significance was tested with the Welch Two Sample t test.

- See *SI Appendix* for more details on the methods used.
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