

# Transcriptome sequencing reveals *maelstrom* as a novel target gene of the terminal-system in the red flour beetle *Tribolium castaneum*

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## Abstract

Terminal regions of the *Drosophila* embryo are patterned by the localized activation of the Torso-RTK pathway, which promotes the down-regulation of Capicua. In the short-germ beetle *Tribolium*, the function of the terminal system appears to be rather different, as the pathway promotes axis elongation and in addition, is required for patterning the extraembryonic serosa at the anterior. Here we show that Torso signalling induces gene expression by relieving CAPICUA-mediated repression also in *Tribolium*. Given that the majority of Torso target genes remain to be identified, we established a differential gene-expression screen. A subset of 50 putative terminal target genes was screened for functions in early embryonic patterning. Of those, 13 genes show early terminal expression domains and also phenotypes were related to terminal patterning. Among others, we found the PIWI-interacting RNA factor Maelstrom to be crucial for early embryonic polarization. *Tc-mael* is required for proper serosal size regulation and head morphogenesis. Moreover, *Tc-mael* promotes growth-zone formation and axis elongation. Our results suggest that posterior patterning by Torso may be realized through Maelstrom depended activation of posterior wnt-domains.

**KEY WORDS:** *Tribolium*, Short-germ insect, terminal system, anterior-posterior patterning, transcriptome sequencing, *torso*, *torso-like*, *capicua*, *maelstrom*

## Introduction

Anterior-posterior patterning of the *Drosophila* embryo depends on spatial polarity cues provided by maternal coordinate systems (St Johnston and Nusslein-Volhard, 1992). While anterior patterning is mediated by the Bicoid morphogen (Driever and Nusslein-Volhard, 1988), posterior patterning largely depends on Nanos (Nos) and Pumilio (Pum) proteins (Barker et al., 1992; Macdonald, 1992; Nüsslein-Volhard et al., 1987; Wang and Lehmann, 1991). The terminal system acts to pattern the anterior and posterior non-segmented regions of the embryo through the *torso*-mediated receptor tyrosine kinase (Ras/MAPK) pathway (Furriols and Casanova, 2003; Li, 2005). Both Torso receptor and its putative ligand, Trunk, are ubiquitously expressed throughout the oocyte, and it appears that localized Torso activation results from activation of Trunk by Torso-like, whose expression is restricted to ovarian follicle cells (Johnson et al., 2015; Mineo et al., 2015; Savant-Bhonsale and Montell, 1993; Stevens et al., 2003). Stimulation of Torso-signalling results in de-repression of the terminal target genes *tailless* (*tll*) and *huckebein* (*hkb*). This activation is indirect and involves de-repression mechanisms. Terminal target genes are usually repressed and the activation of Torso relieves this repression, which allows *tll* and *hkb* expression at the poles of the embryo. Repression of terminal target genes requires the high-mobility group (HMG) protein Capicua (Cic) and the Groucho (Gro), which acts as co-repressor. In absence of maternal Cic, *tll* and *hkb* are de-repressed and expression expands towards central regions of the embryo. This results in the loss of thoracic and abdominal anlagen (Ajuria et al., 2011; Astigarraga et al., 2007; de las Heras and Casanova, 2006; Jimenez et al., 2000; Liaw and Lengyel, 1993; Paroush et al., 1997).

The red flour beetle *Tribolium* develops as a short-germ embryo. The majority of segments get patterned in a secondary growth process from a so-called growth zone (Klingler, 2004; Richards et al., 2008). At the blastoderm stage, the anterior region of the *Tribolium* egg gives rise to extraembryonic serosa and amnion, while embryonic anlagen are restricted toward the ventral-posterior region of the egg (Handel et al., 2000).

Despite these differences, Torso-signalling was shown to be active at both poles of the *Tribolium* egg (Schoppmeier and Schroder, 2005; Schroder et al., 2000). At the posterior, Torso is required for setting up or maintaining a functional growth-zone. In absence of Torso signalling, invagination fails and embryos are depleted of all growth-zone derived segments (Grillo et al., 2012; Schoppmeier and Schroder, 2005). At the anterior, terminal signalling is crucial

for the formation of extraembryonic membranes. In *Torso* or *tsl* RNAi, the serosa is reduced in size (Schoppmeier and Schroder, 2005; van der Zee et al., 2005).

While putative target genes of the *Tribolium* Torso pathway in the anterior region remain to be identified, posterior expression of *Tc-wingless* (*Tc-wg*), *Tc-tailless* (*Tc-tll*), *Tc-caudal* (*Tc-cad*), and *Tc-forkhead* (*Tc-fkh*) depends on Tor activation (Schoppmeier and Schroder, 2005). Still, Torso RNAi phenotypes cannot be entirely explained by the loss of the known target genes, indicating that crucial components for terminal patterning remain to be identified (Casanova, 2005). By comparing terminal-system loss-of function (*Tc-tsl* RNAi) vs. gain-of-function (*Tc-cic* RNAi) transcriptomes to the wildtype situation, we identified the first comprehensive set of Tor-target genes in a short-germ insect.

## Results

### *Capicua acts as a repressor of Torso target-genes in Tribolium*

To elucidate, whether Capicua fulfils a conserved function in repressing Torso target-genes, we characterized the *Tribolium capicua* ortholog (*Tc-cic*, TC004697). *Tc-cic* mRNA is expressed ubiquitously in unfertilized eggs and also present in early blastoderm embryos (not shown) and thus, resembling the situation in *Drosophila* (Jimenez et al., 2000). Depletion of *Tc-cic* by parental RNAi results in severe patterning defects (Figure 1; supplementary Figure S1). Only about 7% of eggs collected in the first week after eclosion of injected animals were capable of secreting a larval cuticle, whereas 92.4% of the embryos die prior to completion of embryonic development (empty-egg) (Schmitt-Engel et al., 2015). To determine, whether the strong *Tc-cic* RNAi no-cuticle phenotype does not reflect a late function of maintaining embryonic anlagen, we performed time-lapse recordings of live embryos using a transgenic line expressing nuclear-localized GFP driven by the *Tribolium* EF1- $\alpha$  promoter (EFA-nGFP) (Sarrazin et al., 2012) (Figure 2; movies S1 and S2).

In wildtype embryos, at the differentiated blastoderm stage, a clear distinction arises between the wider-spaced serosal cells and the more densely spaced cells of the germ-rudiment (Figure 2 A; movie S1) (Handel et al., 2000). During gastrulation, the embryonic anlagen condense and give rise to the germ-rudiment that progressively becomes covered by the extending extraembryonic membranes (i.e. serosa and amnion) (Figure 2 E, G; supplementary movie S1) (Benton

et al., 2013; Handel et al., 2000). Eventually, the head anlagen become morphologically visible, serosa window closure proceeds, and germ-band elongation can be observed (Benton et al., 2013; Handel et al., 2000) (supplementary movie S1).

Upon *Tc-cic* RNAi, extraembryonic membranes are enlarged and expanded toward the posterior pole of the embryo (Figure 2 B, D; supplementary movie S2). During gastrulation, the serosal cells spread in posterior direction, while the residual embryonic tissue invaginates (Figure 2 F). Eventually, extraembryonic membranes cover the entire egg, while embryonic tissue is restricted to the posterior pole of the egg and becomes internalized completely. This presumably embryonic tissue does not show any signs of morphological differentiation or axis elongation (Figure 2 H; supplementary movie S2). At this point embryonic development stops. Thus, the majority of *Tc-cic* depleted embryos die during or soon after gastrulation.

To analyse early patterning of the serosa anlage, we stained *Tc-cic* RNAi embryos for *zerknüllt-1* (*Tc-zen-1*), which in wildtype is already expressed in the presumptive serosa (Figure 3 G) (Falciani et al., 1996). As expected, in *Tc-cic* depleted embryos the *Tc-zen-1* expression expands toward the posterior, indicating that already the serosa primordium is enlarged (Figure 3 I). Interestingly, the serosa-embryo boundary in the differentiated blastoderm is still oblique in *Tc-cic* RNAi (Figure 3), suggesting that *Tribolium* Cic may have no impact on embryonic DV patterning.

In order to reveal the function of *Tc-cic* in patterning gnathal and thoracic anlagen, we analysed the formation of segment primordia by *Tc-even-skipped* (*eve*) expression (Figure 3 A-F). In the blastoderm, the pair-rule gene *Tc-eve* is expressed in double-segmental stripes. Eventually, these primary domains will resolve into a segmental pattern (Patel et al., 1994; Schoppmeier and Schroder, 2005). *Tc-eve* stripe 1 corresponds to the maxillary primordia and *Tc-eve* stripe 2 covers the anlagen of the first thoracic segment. The third primary domain, however, resembles the anlagen of the posterior growth-zone (Figure 3 B). In *Tc-cic* RNAi embryos, primary *Tc-eve* domains are still present, but shifted towards the posterior pole of the embryo (Figure 3 C). In addition, the distance between *Tc-eve* stripe 1 and the serosa boundary is severely decreased in size, indicating the depletion of the pre-gnathal anlagen (Figure 3 F, bar). This is supported by the loss of the ocular *Tc-wg* domain (Figure 3 L). In wildtype blastoderm, *Tc-wg* is expressed in a posterior domain as well as in the ocular anlagen of the anterior head (Figure 3 J) (Nagy and Carroll, 1994). Upon *Tc-cic* knock-down this anterior domain is no longer

present (Figure 3 L). Thus, gnathal and anterior thoracic segments are initially patterned, while pre-gnathal segments apparently are lost.

While *Tc-cic* RNAi causes the expansion of extraembryonic membranes at the expense of the anterior head anlagen, the depletion of Torso-signalling results in opposing effects (Figure 3). In *Torso* or *tsl* RNAi, the serosa is reduced in size, while the presumptive head region appears enlarged and extended towards the anterior (Figure 3 E). This finding is corroborated by the expansion of *even-skipped* and the reduced expression of the serosal marker *zerknüllt-1* in *Tc-tsl* RNAi embryos (Schoppmeier and Schroder, 2005) (Figure 3 H). These phenotypes suggest that anterior *Tc-cic* RNAi reflects a de-repression phenotype of the terminal-system.

Next we tested whether also posterior downstream gene activity is affected in *Tc-cic* RNAi. To this end we analysed the expression of *Tc-tll*, *Tc-wg*, and *Tc-wntD/8* in early *Tc-cic* and *Tc-tsl* RNAi embryos (Figure 3). In early wildtype blastoderm, *Tc-tll* is expressed in a small posterior domain (Figure 3 P) (Schroder et al., 2000). While the posterior *tll* domain is abolished in *Tc-tsl* RNAi (Figure 3 Q), we observed a massive expansion of *Tc-tll* expression in *Tc-cic* depleted embryos (Figure 3 R).

A de-repression phenotype was also observed for *Tribolium wnt* genes (Figure 3 J-O). At the blastoderm stage, *wg/wnt1* and *wntD/8* are expressed at the posterior pole (Bolognesi et al., 2008a; Bolognesi et al., 2008b). In *Tc-tsl*, posterior *Tc-wg* and *Tc-wntD/8* domains are lost (Figure 3 K, N) indicating that posterior patterning by Torso may be realized, at least in parts, through wnt-signalling. Again, the knockdown of *Tribolium cic* results in the expansion of posterior *Tc-wg* and *Tc-wntD/8* domains toward central regions of the embryo (Figure 3 L, O). Thus, like in *Drosophila* (Jimenez et al., 2000; Paroush et al., 1997), Torso signalling in *Tribolium* induces gene expression by relieving CIC-mediated repression.

#### *Transcriptome sequencing and analysis of candidate genes*

To uncover a comprehensive set of Torso target genes, we performed a differential gene-expression-screen, using Next Generation Sequencing (NGS). To this end, we compared terminal-system loss-of-function (i.e. *Tc-tsl* RNAi) versus gain-of-function (i.e. *Tc-cic* RNAi) transcriptomes to the wildtype situation. Genes being differentially expressed in *Tc-tsl* vs. *Tc-cic* RNAi (i.e. up-regulated in *Tc-tsl* RNAi and down-regulated in *Tc-cic* RNAi or vice versa) likely depend on the activity of the terminal system.

Wildtype (T1), *ts/* RNAi (T2), and *cic* RNAi (T3) embryos were fixed at the differentiated blastoderm stage (7-11h) and total RNA was isolated. Sequencing was done on a SOLiD 4 system (Life Technologies) and reads were mapped to the *Tcas3* genome (<http://bioinf.uni-greifswald.de/gb2/gbrowse/tribolium/>). In total, 12.795 genes showed differences in mRNA expression levels (Figure 4 and supplementary Table S1). We found 1801 genes to be up-regulated in *Tc-ts/* RNAi and down-regulated in *Tc-cic* RNAi and 2790 genes that were up-regulated in *Tc-cic* RNAi but down-regulated in *Tc-ts/* RNAi (see supplementary Table S1 for details).

Candidate genes for further inspection were selected by different criteria. To account for biological background variations, genes were only considered as differentially expressed if the fold change (T1 vs. T2 or T1 vs. T3) was at least 50% compared to the wildtype (n= 2241). Still, since the data was taken from a single technical replicate comparisons of fold change numbers have statistically no significance value.

Expression of all differentially expressed genes was monitored by assessment of maternal and different early zygotic wildtype transcriptomes (our unpublished data, see: <http://bioinf.uni-greifswald.de/gb2/gbrowse/tribolium/>). Genes were only taken into account if they were expressed during early stages of development (i.e. maternal until differentiated blastoderm), reducing the number of putative candidates to 1142. iBeetle phenotypes of candidates were monitored by comparison to the iB-database (<http://ibeetle-base.uni-goettingen.de>). The iBeetle screen was a large-scale RNAi screen *Tribolium*, which identified functions in embryonic and postembryonic development for more than half of the *Tribolium* genes (Schmitt-Engel et al., 2015). Of the differentially expressed genes, more than 300 were represented in the iB-database. This comparison allowed us to exclude genes without any obvious embryonic / larval phenotype, as well as adult lethality, defects in metamorphosis, and egg laying / oogenesis upon pupal RNAi. On the other hand, genes with larval phenotypes related to anterior-posterior patterning were preferentially selected for closer inspection, reducing the number of putative candidates to 122. In addition, *Drosophila* orthologs of all candidates were determined and candidates were classified according to their molecular and biological function (supplementary Table S2). Genes with no *Drosophila* orthologs or with unknown function were preferentially selected.

Based on these criteria, we selected 50 candidate genes and proceeded to screen those for functions in early patterning (supplementary Table S2). Larvae were scored for RNAi phe-

notypes. Phalloidin (f-actin) and Hoechst (DNA) staining was used to monitor embryonic phenotypes. In addition, expression was monitored by whole-mount in-situ hybridization (supplementary Table S2).

We found two genes not to result in any detectable phenotype; five genes affected oogenesis, as we observed the cessation of egg-laying upon RNAi, and in 13 cases all eggs displayed no-cuticle phenotypes. As deduced from Hoechst and  $\beta$ -Tubulin staining, these so-called empty egg phenotypes are due to lethality prior to cellularization (supplementary Table S2). Depletion of 30 genes resulted in larval phenotypes, which in most cases were associated with various quantities of empty-egg phenotypes (supplementary Table S2 and Figure 4).

A single gene did not show any obvious expression at all and six genes were only expressed during later stages of development (i.e. germ-rudiment and subsequently). 30 candidates were expressed ubiquitously during early stages (supplementary Table S2). Still, we identified 13 genes with distinct early terminal expression domains (Figure 4, Figure 5, see Table 1 for details).

Of these 13 candidate genes, five genes were exclusively expressed at the posterior pole or in the segment addition zone (Figure 4, Table 1). Eight of the 13 candidates were expressed at both poles at some point during embryonic development (Figure 4, Figure 5, Table 1). Interestingly, six of these eight genes displayed an exclusive early anterior expression domain, either in freshly laid eggs (i.e. maternal expression - TC004989, TC006235, TC009922, and *Tc-mael*), or undifferentiated blastoderm stages (i.e. early zygotic expression - TC004241 and TC006282), suggesting functions in patterning the anterior anlagen (Figure 4, Figure 5, Table 1).

To control for potential off-target effects, we tested these 13 genes by injecting dsRNA fragments not overlapping with the original dsRNA fragments ("non-overlapping fragments", NOFs). All phenotypes were identical to those observed for the original dsRNA fragments (not shown).

Larval phenotypes are correlated with the expression of the candidates: genes being expressed at the posterior pole predominantly display posterior truncations or aberrations (Figure 4, Table 1). For instance, larvae depleted for *TC004989*, *TC005697*, or *TC003946* show the loss of abdominal segments, as seen after loss of Torso-signalling (Schoppmeier and Schroder, 2005) and RNAi with *TC000868* and *TC013474* results in the depletion of terminal



structures (i.e. Urogomphi and Pygopods). Candidates that are also expressed anteriorly frequently show additional head defects. The knockdown of *TC006181* or *TC006282* not only affects axis elongation, but also results in deformation of the larval head (Figure 4). For two genes (*TC004241* and *TC008122*) we observed the loss of the entire head without obvious axis elongation phenotypes (Figure 4). These head phenotypes resemble that of mild *Tc-cic* RNAi (Figures 1 and 4) and thus may represent gain-of-function phenotypes of the terminal system.

While it remains to be elucidated to which degree these candidate genes are indeed direct target genes of Torso-signalling, our study establishes that transcriptome sequencing facilitates the identification of novel candidate genes for early anterior and posterior pattern formation in *Tribolium*.

#### *Tribolium Maelstrom RNAi results in the loss of growth-zone derived structures*

We selected the *Tribolium* ortholog of the *Drosophila maelstrom* gene for closer inspection. Maelstrom (Mael) is a conserved PIWI-interacting RNA factor (piRNA), consisting of two domains, a HMG box and a MAEL domain (Sato and Siomi, 2015). In *Drosophila*, piRNA factors are required for transcriptional transposon silencing in both, germ and ovarian somatic cells (Sato and Siomi, 2015). In addition, Mael coordinates axis specification through nucleating microtubule-organizing center (MTOC) components (Sato et al., 2011).

The *Tribolium* genome contains a single *maelstrom* ortholog (*Tc-mael*, *TC008172*). *Tc-mael* is expressed maternally and transcripts accumulate at the anterior pole of unfertilized eggs and the polar body (Figure 5 A). During blastoderm stages, *Tc-mael* becomes expressed at the posterior pole. This domain persists until gastrulation (Figure 5 C, E). Later, *Tc-mael* expression is not longer detectable (not shown).

Knocking down *Tc-mael* by RNAi results in truncation phenotypes (Figure 6 and supplementary Figure S2). About 60% of the larvae lack growth-zone derived structures: abdominal segments and terminal structures are lost, resembling the loss of Torso-signalling (Schoppmeier and Schroder, 2005). These posterior truncation phenotypes were sometimes (about 20%) accompanied by aberrations of the larval head, likely representing stronger phenotypes. Head phenotypes, however, are highly variable and may include deformations and / or deletions of pre-gnathal and gnathal segments. While phenotypes likely reveal *Tribolium mael* functions in early embryonic patterning (see below), we observed additional effects of

*Tc-mael* depletion. RNAi with *Tc-mael* results in incorrect dorsal closure, generating completely or partially everted (“inside-out”) larvae (12,5%). These larvae may have escaped the primary truncation effect.

To analyse the impact of *Tc-mael* RNAi, we performed time-lapse recordings of EFA-nGFP embryos (Figure 7 and Movie S3) (Sarrazin et al., 2012). Upon *Tc-mael* RNAi, anterior germ-rudiment formation is impaired: Head anlagen of *Tc-mael* RNAi embryos are less condensed as compared to wildtype (Figure 7 E). During wildtype germ-band elongation the head region extends to an anterior-dorsal position and the germ-band extends posteriorly (Figure 7 and supplementary Movie S1). Upon *Tc-mael* RNAi, the head anlagen still moves in anterior ventral direction, posterior invagination, however, is delayed to some degree (Figure 7 D and Movie S3) and elongation fails (Figure 7 J, K, L; and Movie S3), supporting a function for *Tc-mael* in growth-zone formation and / or axis elongation (see below).

In *Tc-mael* RNAi embryos, the serosal window eventually closes and amniotic cavity formation seems normal (Figure 7 E, F; Movie S3). Unexpectedly, we observed the premature rupturing of the serosa at a posterior-dorsal position (Figure 7 K, L; Movie S3). In wildtype, serosal withdrawal does not occur until the onset of dorsal closure at an anterior-ventral position (Hilbrant et al., 2016; Panfilio et al., 2013).

Rupture of extraembryonic membranes is a coordinated process that involves opening both the serosa and the amnion in an anterior-ventral region, after germ-band retraction (Hilbrant et al., 2016; Panfilio et al., 2013). If the amnion fails to withdraw prior to embryonic dorsal closure, the embryo becomes everted („inside-out”) (Hilbrant et al., 2016; Panfilio et al., 2013). Proper serosal patterning (and subsequent withdrawal) involves the attachment of the serosa to the vitelline membrane at the posterior pole of the egg (Koelzer et al., 2014). In *Tc-mael* RNAi, however, the posterior serosa never seems to come into contact with the vitelline membrane at the posterior egg pole, but rather contracts anteriorly, loses tissue integrity, and ultimately disintegrates (Figure 7 J-L, Movie S3). Consequently, the embryo is stuck inside of amniotic tissue and the embryo closes ventrally rather than dorsally, resulting in inside-out phenotypes.

### *Maelstrom affects anterior patterning and morphogenesis*

To uncover the influence of *Tc-mael* on the formation of extraembryonic membranes and patterning of anterior embryonic anlagen in more detail, we visualized the emergence of these primordia by analysing the *Tc-eve*, *Tc-giant* (*Tc-gt*), and *Tc-zen-1* expression (Falciani et al., 1996; Patel et al., 1994) (Figure 8). Upon *Tc-mael* knockdown the *Tc-zen-1* domain is smaller, indicating a reduction of the serosa primordia (Figure 9 B). The serosa-germ-rudiment border, however, is still oblique.

In wildtype differentiated blastoderm stage, *Tc-gt* is expressed in a wedge-shaped anterior domain comprising of pre-gnathal and gnathal segments (Bucher and Klingler, 2004). At that stage also a secondary domain arises at the posterior pole (Bucher and Klingler, 2004). In *Tc-mael* RNAi, the anterior domain is unaffected, while the posterior *Tc-gt* domain is lost (see below). Also the first and second primary *Tc-eve* domains (*Tc-eve* stripe 1 and stripe 2) are present in blastoderm stages, but reduced and less defined to some degree (Figure 8 B, D).

During gastrulation extraembryonic membranes fold over anterior and posterior embryonic margins, thereby completing the rim of a serosal window (Handel et al., 2000). *Tc-zen-1* is expressed throughout serosal window formation and closure, with high expression levels at the margin of the serosa window (Figure 8 C). Upon *Tc-mael* knockdown *Tc-zen-1* expression is lost from the anterior rim of serosa window (Figure 8 D, G) and head primordia appear to be less condensed and shifted towards the anterior pole of the egg (Figure 8 D). At that stage there are no obvious effects on anterior *Tc-eve* expression: The first and second primary *Tc-eve* stripes have split, while the 3<sup>rd</sup> double-segmental domain is present (Figure 9 G), reflecting that gnathal and thoracic anlagen are present. Thus, *Tc-mael* is required for proper serosal size regulation and head morphogenesis.

### *Maelstrom is required for axis elongation and growth-zone formation*

*Tc-mael* RNAi embryos are depleted of abdominal segments (Figure 7, Figure 8). The fourth primary *Tc-eve* domain, which corresponds to the first and second abdominal segments, does not form properly and in addition, posterior morphology becomes highly irregular (Figure 8 G, H). Already during late differentiated blastoderm stages, the posterior *Tc-gt* domain is lost (Figure 8 J) indicating that posterior elongation is already affected at that stage.

To reveal whether *Tc-mael* is already involved in setting up a functional growth-zone, we monitored the expression of *Tc-wg* and *Tc-wntD/8* (Bolognesi et al., 2008a; Bolognesi

et al., 2008b) (Figure 9). In wildtype, wnt-signalling is required for growth-zone formation and axis elongation (Beermann et al., 2011; Bolognesi et al., 2008a; Bolognesi et al., 2008b). While in *Tc-mael* RNAi the terminal *Tc-wg* domain was basically present, the *wntD/8* is notably reduced or even absent (Figure 9 C), resembling the situation in *Tc-tor* or *Tc-tsl* knockdown. Our results indicate that posterior patterning by Torso may be realized through *Tc-maelstrom* dependent activation of the posterior *Tc-wntD/8* domain.

Next, we monitored *Tc-mael* expression in both, *Tc-tsl* and *Tc-cic* depleted embryos. As expected, we observed an expansion of *Tc-mael* expression in *Tc-cic* RNAi (Figure 10 E), while in *Tc-tsl* RNAi expression was lost (Figure 10 C). Thus, *Tribolium maelstrom* is indeed a target gene of the terminal system in this beetle.

#### *Posterior retraction of short-gastrulation depends on Maelstrom activity*

Given that *Drosophila maelstrom* is also required for achieving proper DV polarity (Clegg et al., 2001; Clegg et al., 1997; Findley et al., 2003; Sato et al., 2011), we monitored DV axis formation upon *Tc-mael* RNAi.

It has been shown before that the dorso-ventral tilt of the serosa / embryo boundary depends on BMP signalling in *Tribolium* (Nunes da Fonseca et al., 2008; van der Zee et al., 2006). In *Tc-dpp* RNAi, this border becomes DV symmetric and is shifted towards the anterior. Depletion *Tc-sog* has opposing effects. The serosa increases at the expense of the head anlagen and the serosa-embryo boundary develops vertical to the egg axis (van der Zee et al., 2006). Thus, the loss of DV asymmetry reveals impaired dorso-ventral (DV) patterning in *Tribolium* (van der Zee et al., 2006).

While also in *Tc-mael* RNAi the serosal anlagen are reduced (Figure 8 B), the serosa-germ-rudiment border is still oblique (Figure 8 B), indicating that DV axis formation is unaffected. This is corroborated by anterior *Tc-gt* expression (Figure 8 J). In *Tc-mael* RNAi, the *gt* domain still shows a pronounced DV asymmetry. Thus, *Tc-mael* has no impact of on early anterior DV axis polarization.

Interestingly, however, we found *Tc-mael* to be required for posterior *Tc-sog* retraction (Figure 11). In wild type, *Tc-sog* is initially expressed in a broad ventral domain, which subsequently clears from the posterior pole (van der Zee et al., 2006) (Figure 11 A and C). While there is no pronounced difference in early stages, *Tc-sog* fails to retract from the pos-

terior pole of *Tc-mael* RNAi embryos (Figure 11 D) and posterior dorsal Dpp activity (as monitored by pMAD staining) (Figure 11 F) is reduced to some degree. Consequently, posterior dorsal tissue (i.e. the dorsal amnion) lacks peak levels of Dpp activity. In wildtype, *Tc-iro* is expressed in a domain marking the anterior border of the germ-rudiment, which will give rise to the anterior amnion. In addition, *Tc-iro* is expressed in the dorsal amnion (Nunes da Fonseca et al., 2008). In *Tc-mael* RNAi embryos, the anterior of *Tc-iro* expression was maintained, while the dorsal most expression domain was absent (Figure 11 H). Thus, in *Tc-mael* RNAi embryos posterior dorsal tissue is lateralized.

## Discussion

### *The function of Tribolium Capicua is restricted to AP patterning*

The depletion of *Tribolium capicua* results in the expansion of terminal anlagen and de-repression of terminal target genes, reflecting de-repression (i.e. gain of function) phenotypes of Torso-signalling. This resembles the situation in *Drosophila* (Jimenez et al., 2000; Paroush et al., 1997), indicating that Torso signalling induces gene expression by relieving Cic-mediated repression also in *Tribolium*.

While the functions of *Tribolium capicua* in terminal patterning may very well be conserved, we did not observe any obvious impact of *Tc-cic* on dorso-ventral pattern formation. In *Drosophila*, the function of Cic is not restricted to Torso-RTK signalling. During *Drosophila* oogenesis, Cic is required to translate EGFR signalling into asymmetric *pipe* expression, which culminates in the establishment of a nuclear Dorsal gradient (Moussian and Roth, 2005).

In *Tribolium*, EGF signalling has broadly conserved roles in setting up the DV axis of the embryo (Lynch et al., 2010). The depletion of the *Tribolium gurken* ortholog (*Tc-Tgf-alpha*) results in the ventralization of embryos, resembling *Tc-dpp* RNAi phenotypes to some degree (van der Zee et al., 2006). In both, *Tc-Tgf-alpha* and *Tc-dpp* RNAi the first morphologically visible sign of DV polarity, the obliqueness of the border between serosa and germ-rudiment, is lost (Lynch et al., 2010; van der Zee et al., 2006). However, as indicated by morphology and *Tc-zen-1* expression, *Tc-cic* RNAi does not affect the obliqueness of the serosa / germ-rudiment border (Figure 2 and 3). Thus, in *Tribolium* there is no obvious antagonistic interaction of EGFR-signalling with Cic in embryonic DV axis formation.

The expansion of the serosa in *Tc-cic* RNAi embryos goes along with de-repression of *Tc-zen-1* (Figure 3 G, I). At first glance this situation resembles that of *Drosophila*, as knock-down of *Dm-cic* results in the lateral expansion of *Dm-zen* expression (Astigarraga et al., 2007; Jimenez et al., 2000; Kim et al., 2011). In *Drosophila*, *zen* is broadly activated by maternal factors (Rushlow et al., 1987). During early blastoderm stages, *cic* participates in the ventral and lateral repression of *zen* by Dorsal (dl) (Astigarraga et al., 2007; Jimenez et al., 2000; Kim et al., 2011). At the poles, dl-mediated repression of *zen* is antagonized by Torso signalling (Rusch and Levine, 1994), which likely depends on MAPK-dependent down-regulation of Cic (Kim et al., 2011). However, the situation in *Tribolium* differs from *Drosophila*. The distinction between serosa and germ-rudiment is primarily set up by inputs from the AP system and secondarily modulated through DV signalling. In particular, *Tc-zen-1* expression and thus, serosa size regulation depends on the combined activity of terminal signalling and *Tc-orthodenticle* (Kotkamp et al., 2010), while the obliqueness of the border between serosa and germ-rudiment is subsequently established by EFGR and BMP signalling (Lynch et al., 2010; Nunes da Fonseca et al., 2010; Nunes da Fonseca et al., 2008; Van der Zee et al., 2008; van der Zee et al., 2006). Given that serosal size regulation, but not the obliqueness of the border between serosa and germ-rudiment is affected in *Tc-cic* RNAi, it appears that eggs produced after *Tc-cic* depletion maintain normal embryonic DV polarity.

#### *Maelstrom is required for serosa morphogenesis and anterior germ-band condensation*

The depletion of *Tribolium maelstrom* by RNAi affects axis elongation, germ-band condensation, and the formation of the extra-embryonic serosa (Figure 6, 7, and 8). In *Drosophila*, Mael coordinates oocyte polarity through interaction with components of the MTOC (Sato et al., 2011). Achieving proper cell polarity is essential for diverse processes, including cell shape changes and cell migration, asymmetric cell division, and morphogenesis (St Johnston and Ahringer, 2010). Also early *Tribolium* development depends on extensive cell-polarization events (Benton et al., 2013; Handel et al., 2000). RNAi with *Tc-mael* disturbs the anterior condensation of the embryonic anlagen (Figure 7 and 8). Hence, in absence of *Tc-mael* cell polarity may not be established appropriately, which in turn may affect early morphogenesis. This would be equivalent to the situation in *Drosophila* (Clegg et al., 2001; Clegg et al., 1997; Findley et al., 2003; Sato et al., 2011). Although we currently cannot exclude that *Tc-mael* RNAi phenotypes are due to impaired transposon silencing, we posit that *Tc-Mael* coordinates cell

polarity (possibly by acting on MTOC organization) at least in serosa morphogenesis and anterior germ-band condensation in a Torso-signalling dependent manner.

While serosa phenotypes indeed resemble aspects of *Tc-tsl* or *Tc-tor* RNAi phenotypes, we observed additional defects in head development upon *Tc-mael* knock-down. At first glance *Tc-mael* RNAi head phenotypes are not related to terminal patterning. As mentioned before, anterior Torso-signalling is involved in serosal size regulation (Schoppmeier and Schroder, 2005). The loss or reduction of this extraembryonic membrane by *Tc-tsl* or *Tc-zen-1* RNAi results in the expansion of the head anlagen, which is largely compensated later in development (Schoppmeier and Schroder, 2005; van der Zee et al., 2005). Nevertheless, a small fraction of larvae depleted for *Tc-zen-1* or *Tc-tsl* do also display head defects (van der Zee et al., 2005) (our unpublished observation). These phenotypes are highly variable and thus, may rather reflect defects during size compensation and head morphogenesis instead of specific functions in head patterning.

#### *Tribolium maelstrom* in posterior patterning

At the posterior pole, *Tc-mael* function is necessary for growth-zone formation. Upon *Tc-mael* knock-down, expression of *Tc-wntD/8* is lost and posterior elongation is terminated prematurely (Figures 6 and 9). Given that *Tc-wntD/8* is under control of Torso-signalling (Figure 3) (Bolognesi et al., 2008b; Schoppmeier and Schroder, 2005), our results suggest that posterior patterning by Torso may be realized, at least in part, through Maelstrom dependent activation or positioning of posterior wnt-domains.

On the other hand, the knock-down of *Tc-wntD/8* only results in a small percentage of embryos lacking abdominal segments and additional removal of *Tc-wg* only enhanced the penetrance of this phenotype to some degree (Bolognesi et al., 2008a). As effects of *Tc-mael* RNAi on growth-zone formation and axis elongation are considerably more severe, *Tc-mael* likely fulfils additional functions in posterior patterning, which may involve DV patterning.

In *Tc-mael* RNAi embryos, posterior *Tc-sog* expression fails to retract and as consequence, posterior dorsal tissue, i.e. the dorsal amnion lacks peak levels of Dpp activity. Changes in Dpp activity also affect morphogenetic movements during gastrulation (Nunes da Fonseca et al., 2010; van der Zee et al., 2006). Upon *Tc-dpp* RNAi, for instance, the germ-band is symmetrically folded inward at the posterior pit and extends toward the anterior. In *Tc-mael* RNAi embryos, however, we did not observe any strong effects on posterior invagination, and

posterior amniotic fold formation is normal, suggesting that posterior morphogenetic movements might be largely unaffected. Thus, even though *Tc-mael* also contributes - in a Torso-independent manner- to proper DV axis formation, we deem it unlikely that this function has impact on early posterior axis polarization or growth-zone formation.

During oogenesis, *Drosophila mael* mutants fail to establish cytoplasmic polarity, and to normally accumulate Gurken. Also other polarity markers (e.g. *osk*, *BicD*) fail to accumulate in the posterior ooplasm, and *bcd* mRNA becomes localized at both poles (Clegg et al., 2001; Clegg et al., 1997; Findley et al., 2003; Sato et al., 2011). The *Tribolium grk* ortholog has no major role in AP axis formation, but rather acts in DV patterning (Lynch et al., 2010). Moreover, *Tc-wntD/8* is not maternally expressed, indicating that additional potentially localized factors might be involved in wntD/8 regulation.

An important role for localized maternal determinants has long been postulated for several insect taxa including crickets, beetles, and flies. However, thus far the existence of two morphogenetic centers (anterior and posterior) has only been proven for the long-germ wasp *Nasonia vitripennis* (Rosenberg et al., 2009). *Tribolium* utilizes anterior mRNA localization to some degree (Fu et al., 2012). Posterior localized maternal determinants, however, have not been identified yet (Schmitt-Engel et al., 2012). While posterior patterning in *Tribolium* may depend on spatially restricted polarity cues (i.e. the terminal system) that regulate short-range acting target genes required for axis elongation, like wnt-signaling (Beermann et al., 2011; Bolognesi et al., 2008a; Bolognesi et al., 2008b) rather than on long-range acting gradients (i.e. Nanos) (Schmitt-Engel et al., 2012), it's tempting to speculate that *Tc-mael* promotes growth-zone formation by localizing an as yet unknown posterior maternal determinant.



## Methods

### *Histology*

Fixation and single in situ and immunofluorescence staining were performed using standard protocols (Patel et al., 1989; Tautz and Pfeifle, 1989). Embryos were subsequently counterstained with Hoechst 33342.

### *RNAi*

Parental RNAi and dsRNA synthesis was performed as described (Bucher and Klingler, 2004; Bucher et al., 2002). dsRNAs were injected into female pupae at a concentration of 1 µg/µl. RNAi phenotypes were confirmed by injection of non-overlapping-dsRNA fragments. RNAi for all fragments resulted in identical phenotypes, excluding the possibility that an unrelated gene was affected (off-target effects).

### *Cuticle preparation*

First instar larvae were cleared in lactic acid/10% ethanol overnight at 60 °C. Cuticles were transferred to a drop of lactic acid on a slide. The cuticular autofluorescence was detected on a Zeiss Axiophot microscope, and maximum projection images were created from z stacks using the Analysis D software (Olympus).

### *Transcriptome Sequencing*

Sequencing was done on a SOLiD 4 system (Life Technologies) using 50bp fragment reads. Mapping to the Tcas3 reference genome (<http://bioinf.uni-greifswald.de/gb2/gbrowse/tribolium/>) and counting were performed using LifeScope version 2.5, also by Life Technologies. In order to filter out adapters, ribosomal and tRNAs, a filter reference file was created using the adapter sequences provided from Life Technologies, large and small subunit rRNA sequences from the reference and the results of a tRNAscan-SE (v1.1) run over the reference sequence.

Samples T1 (wildtype), T2 (*tsl* RNAi) and T3 (*cic* RNAi), mapped read counts were 15,517,184; 6,172,710 and 31,441,775; respectively. Counting yielded read counts for 143,603 exons in 16,543 transcripts as provided by the Tcas3 reference. Normalization and comparison

of expression levels between all 3 samples was performed using the *differential expression analysis for sequence count data algorithm* (DESeq) (Anders and Huber, 2010).

#### *Data access*

The complete dataset, including all relevant parameters, was deposited at NCBI under accession number PRJNA311251.

#### *Live imaging*

For time-lapse imaging, the EFA-nGFP-strain of *Tribolium castaneum* was used (Sarrazin et al., 2012). Embryos (4 to 6 h) were treated with 12.5% bleach for two times 30 seconds and then rinsed with tap water. Embryos were then mounted in a hanging drop of Sigma Halocarbon oil 700.

For time-lapse imaging, a Leica TCS SP5 II Confocal System was used. Stacks were recorded about every 12 minutes for an interval of 12 hours at 10x magnification and 20°C. Processing of the image stacks was done in LAS AF Version 2.4.1 build 638 and Fiji (Schindelin et al., 2012).

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## **Author contributions**

Conceived and designed the experiments: FP, MS. Acquisition of data: FP, MW, NK, AS, AE. Analysis and interpretation of data: FP, MW, SU, MS. Writing the manuscript: MS

## **Competing financial interests:**

All authors declare no competing financial interests.

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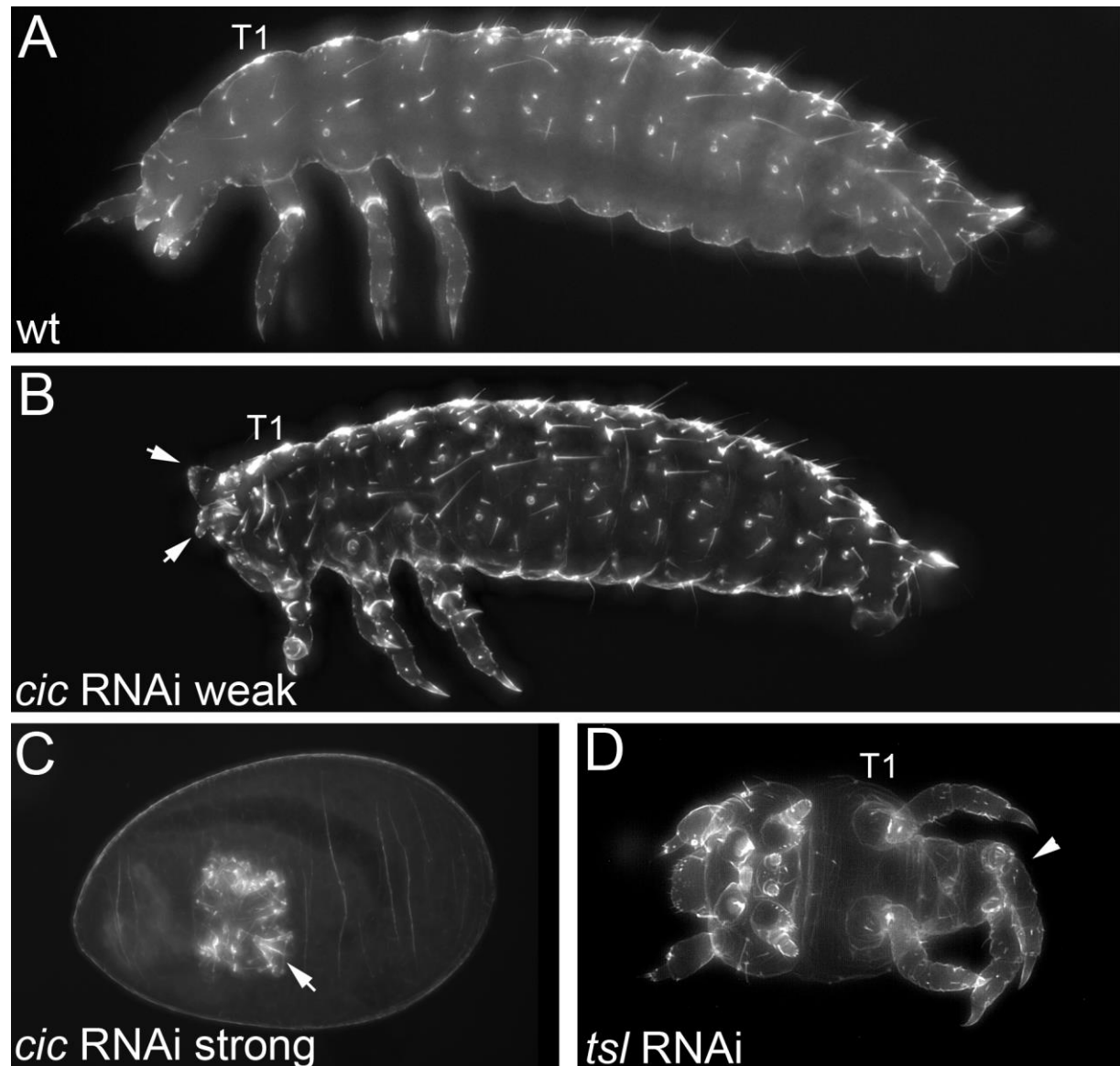
**Wang, C. and Lehmann, R.** (1991). Nanos is the localized posterior determinant in *Drosophila*. *Cell* **66**, 637-647.



Tc-#	iB-#	<i>Dm</i> ortholog	Molecular function	Relative expression	
				<i>tsl</i> RNAi	<i>cic</i> RNAi
TC004989	iB_00793	cathD	aspartic-type endopeptidase activity	↓	↑
TC006235	iB_04097	Dsp1	DNA binding, bending; transcription factor binding	↓	↑
TC009922	-	ap2	protein transporter activity	↓	↑
TC004241	iB_03713	-	hydroxylase function	↑	↑
TC006282	-	geminin	DNA binding	↓	↑
TC006181	iB_01008	CG43129	unknown	↓	↑
TC008122	iB_04429	CG8408	unknown	↓	↑
TC005697	iB_00909	CG9323	RNA helicase activity, G-quadruplex RNA binding	↓	↑
TC003946	iB_00642	crol	metal ion binding; nucleic acid binding	↓	↑
TC000868	iB_03094	eIB	metal ion binding; nucleic acid binding; protein binding	↓	↑
TC007740	iB_04375	-	unknown	↑	-
TC013474	-	sens	transcription factor activity, sequence-specific DNA binding	↓	↑

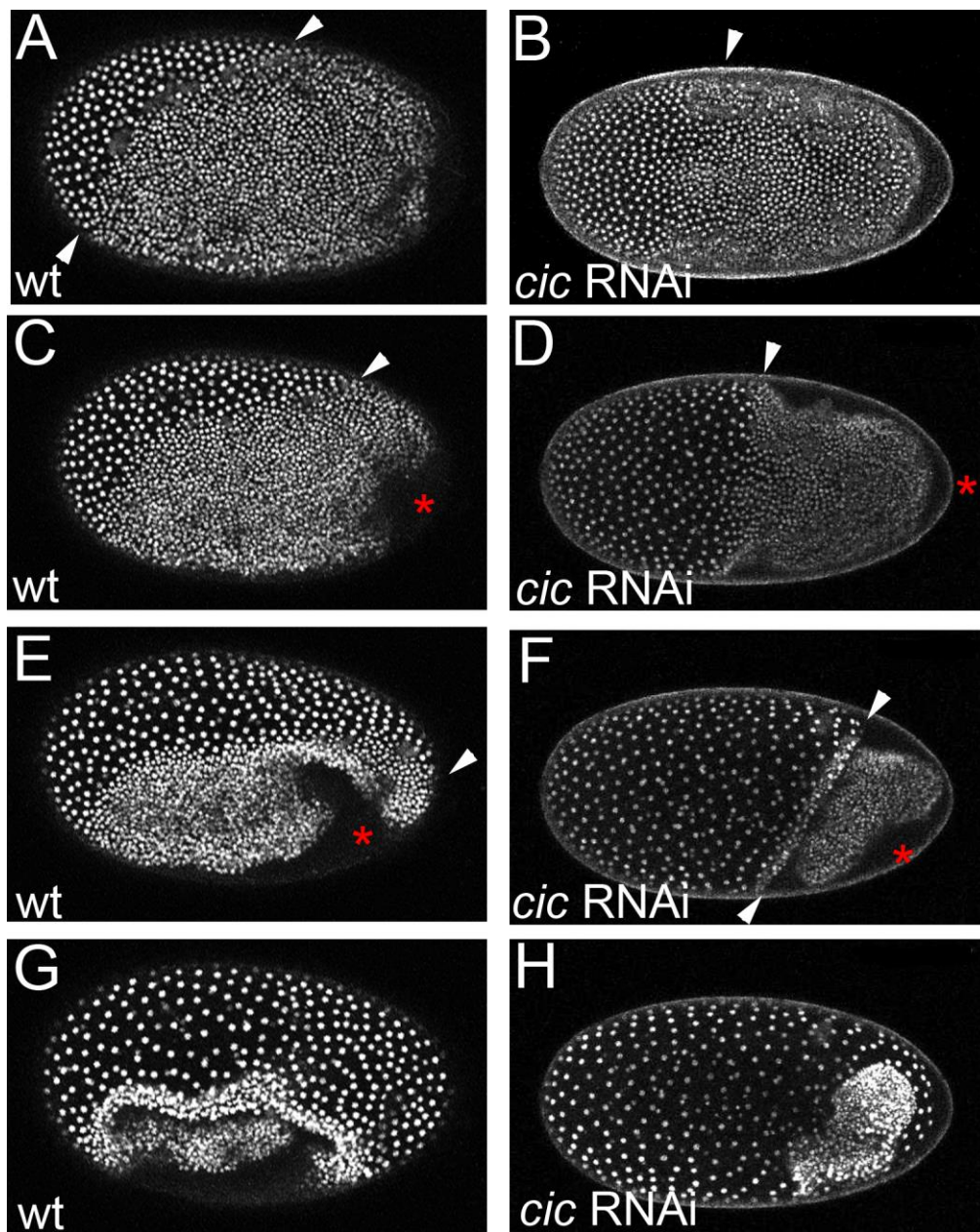
Table 1: Official gene set number (Tc), iBeetle (iB) number (<http://ibeetle-base.uni-goettingen.de>), *Drosophila* ortholog, molecular function and changes in relative expression of candidate genes. Candidate genes were sorted according to figure 4.

## Figures



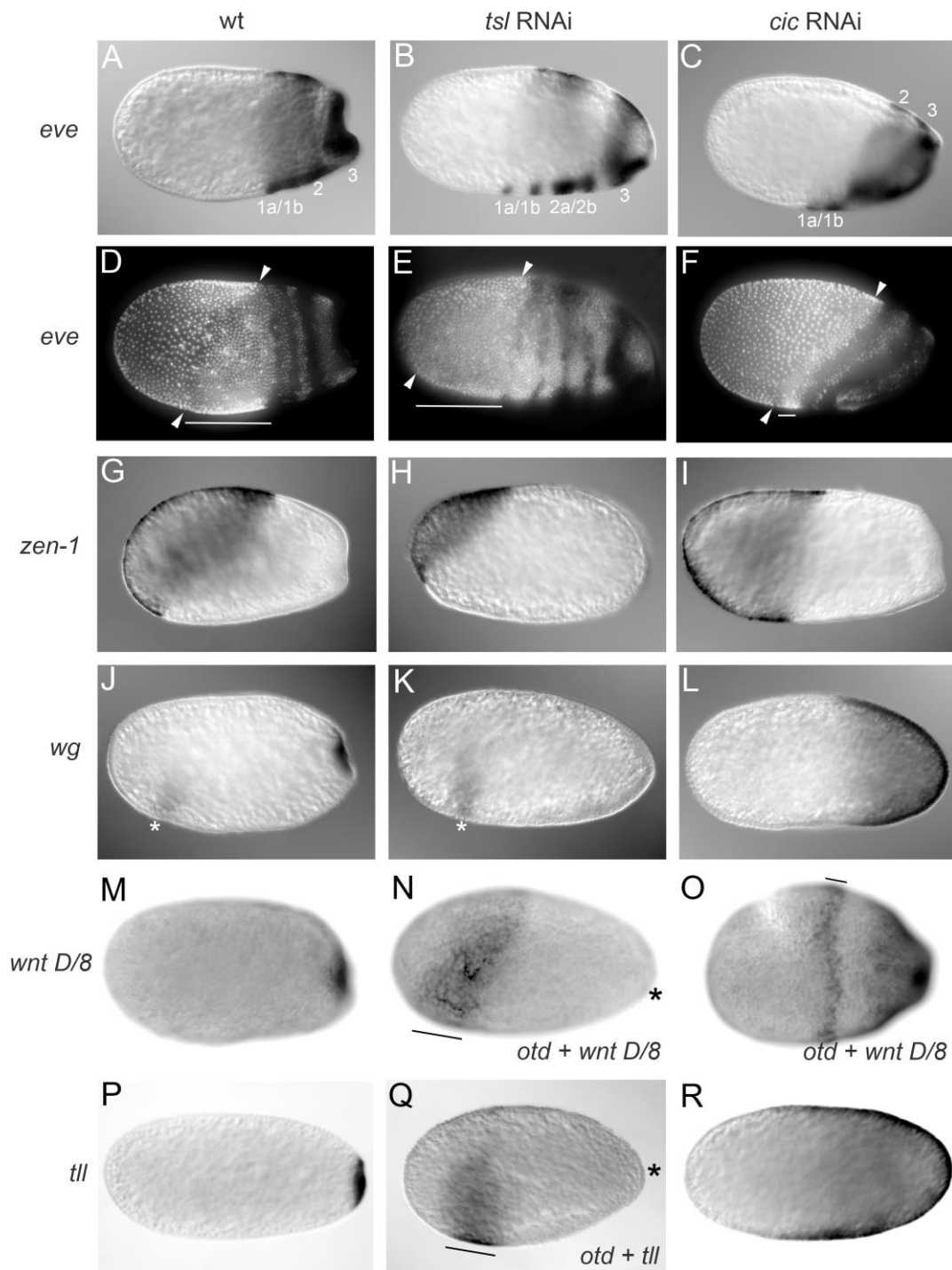
**Figure 1: Larval *Tc-capicua* and *Tc-torso-like* RNAi phenotypes**

Cuticle of wildtype (A), *Tc-cic* (B-C), and *Tc-tsl* (D) RNAi larvae. (B) Mildly affected *Tc-cic* RNAi larva display head defects (arrows). (C) In strong phenotypes, cuticle remnants with Urogomphile-like structures were obvious (arrow). (D) *Tc-tsl* RNAi larvae are depleted of growth-zone derived segments. The first thoracic segment (T1) is unaffected, T2 is reduced (arrowhead). All panels in all pictures: anterior to the left. A-B: lateral views and C, D: ventral view.



**Figure 2: Live imaging of *Tc-cic* RNAi**

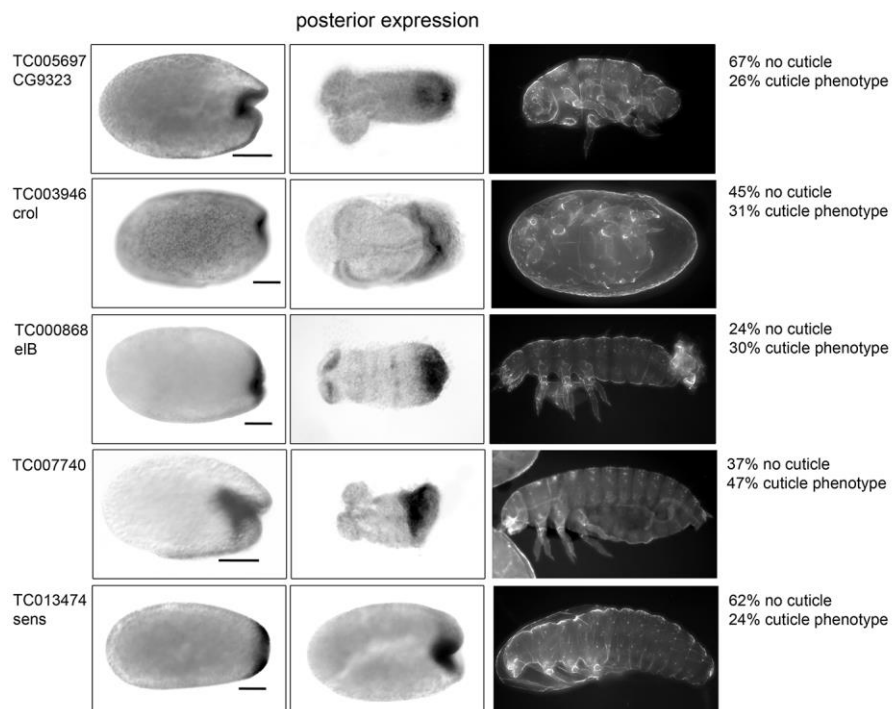
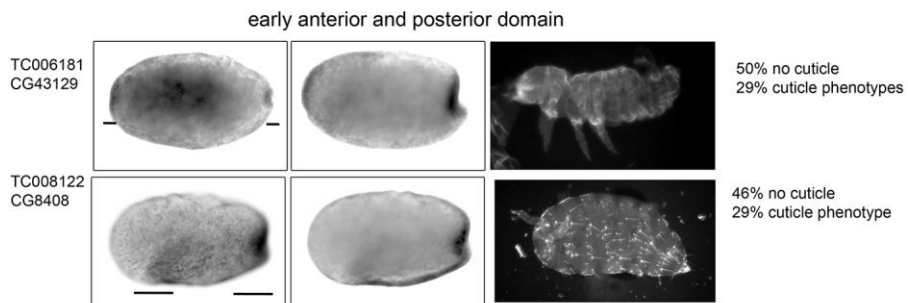
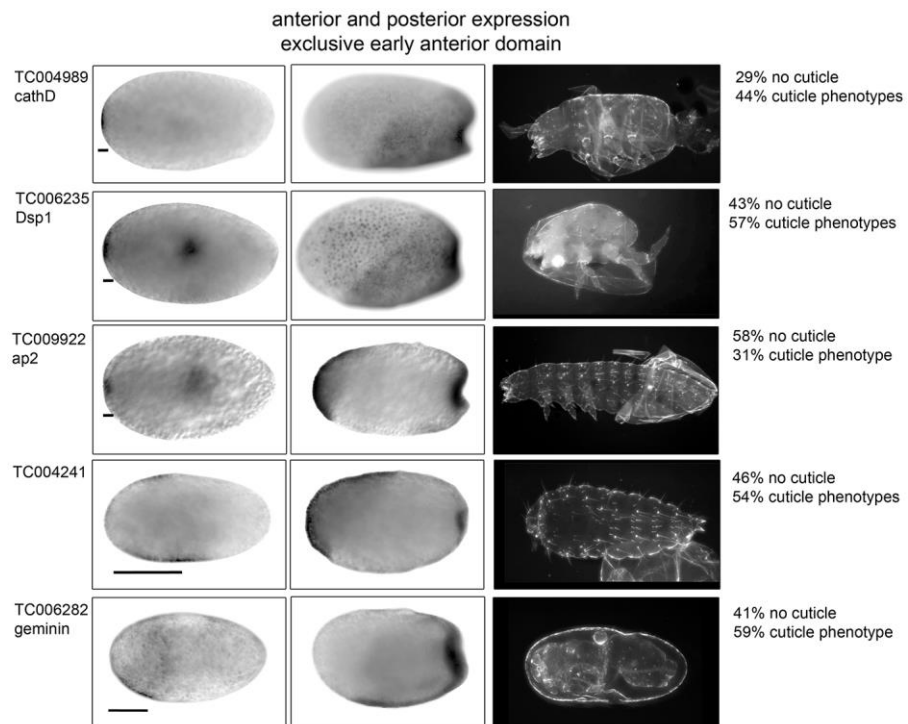
Live imaging of wildtype and *Tc-cic* RNAi in the transgenic *Tribolium* EFA-nGFP line. Columns show nGFP labelled stage matched embryo at representative time-points. (A, C, E, G) lateral views of wt embryos at differentiated blastoderm (A), posterior-pit (C), invagination (E), and serosa closure stages (G). (B, D, F, H) corresponding stages of *Tc-cic* RNAi. Arrows mark the serosa / embryo boundary, asterisks the site of gastrulation. See text for details.



### Figure 3: Expression of terminal target genes in *Tc-cic* and *Tc-tsl* RNAi

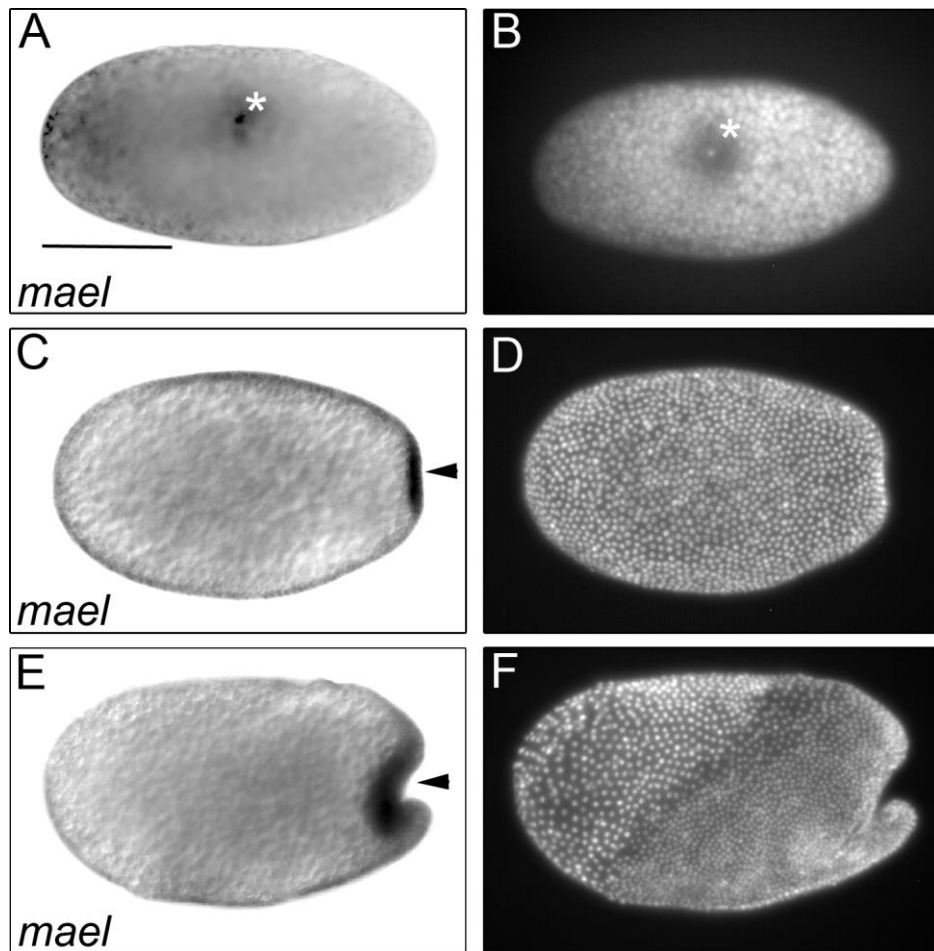
Expression of *Tc-eve* (A-C), *Tc-zen-1* (D-F), *Tc-wg* (J-L), *Tc-wnt D/8* (M-O), and *Tc-tll* (P-R) in wildtype (A, D, G, J, M, P), *Tc-tsl* (B, E, H, K, N, Q), and *Tc-cic* RNAi (C, F, I, L, O, R). Embryos in (A-C) were subsequently stained for the nuclear marker Hoechst 33342 (D-F). Embryos in (N, O, Q) were double stained for *Tc-otd* to visualize the head anlagen (bars). In wildtype (A, B) *Tc-eve* is expressed in three primary stripes (1-3) that eventually will split and give rise to segmental (1a, 1b) domains. In *Tc-tsl* RNAi (B, E), *Tc-eve* stripes are shifted toward the anterior, reflecting the expansion of the head anlagen (bar) at the expense of the serosa (arrowheads). In *Tc-cic* depleted embryos, (C, F) the serosa is expands (arrowheads), while the head anlagen (bar) are reduced. (G) *Tc-zen-1* is expressed in the anlagen of the serosa. Upon *Tc-tsl* RNAi (H) *Tc-zen-1* expression reduced, while *Tc-zen-1* is expanded in *Tc-cic* RNAi (F). At the blastoderm stage, *Tc-wg* (J) is expressed in the ocular domain (asterisk) and at the posterior pole. In *Tc-tsl* RNAi (K) the posterior domain is absent. *Tc-cic* RNAi (L), results in the expansion of the posterior *Tc-wg* domain, while the head domain is lost. In wildtype blastoderm stages, *Tc-wnt D/8* and *Tc-tll* (M, P) are expressed at the posterior pole. After the knockdown of *Tc-tsl* (N, Q) these domains are absent (asterisk). *Tc-cic* RNAi (O, R) results in the expansion of both domains.





#### **Figure 4: Results of the RNAi and expression screen**

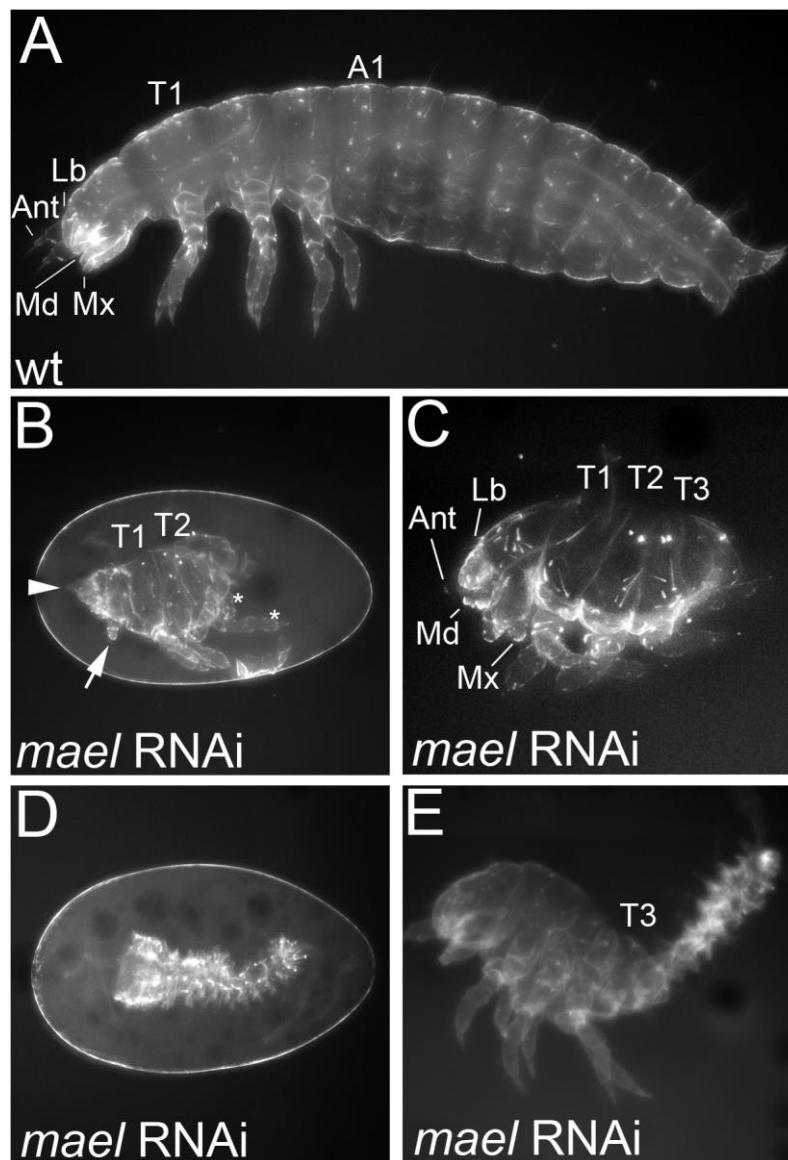
Candidate genes were monitored for expression and function in AP patterning by RNAi. Depletion of genes resulted in larval phenotypes, which in most occasions were associated with various quantities of no-egg phenotypes.



**Figure 5: Expression of *Tc-mael***

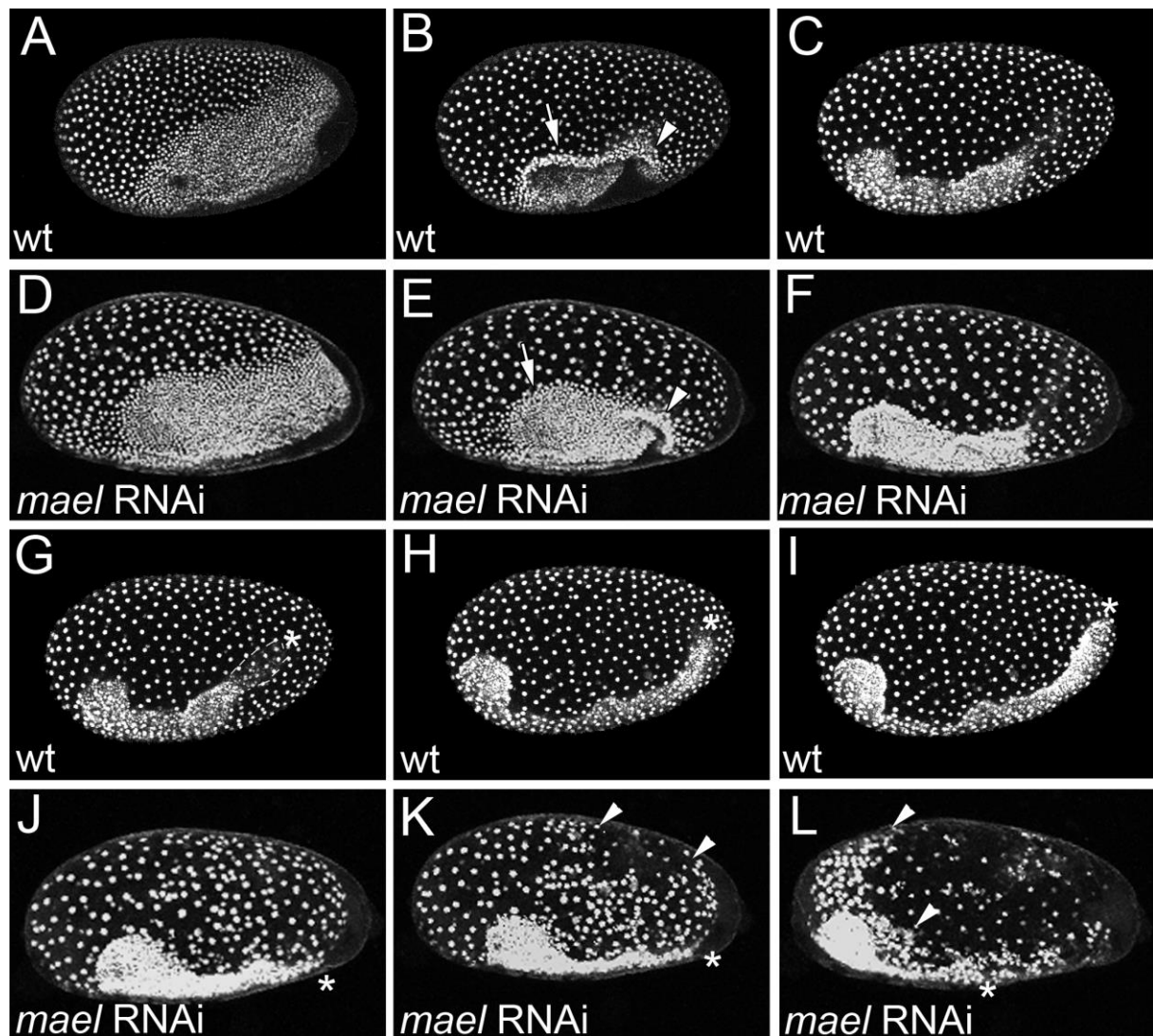
Expression of *Tc-mael*. Embryos in (A,C,E) were subsequently stained for Hoechst 33342 (B,D,F). (A) In unfertilized eggs, *Tc-mael* is expressed around the polar body (asterisks in A, B) and in an anterior domain of the embryo (bar in A). The embryo shown in A and B is the same as that in Figure 10A and B. (C) During undifferentiated blastodermal stages, *Tc-mael* is expressed in a distinct posterior expression domain (arrow) that is maintained in differentiated blastodermal stages and invagination (D).





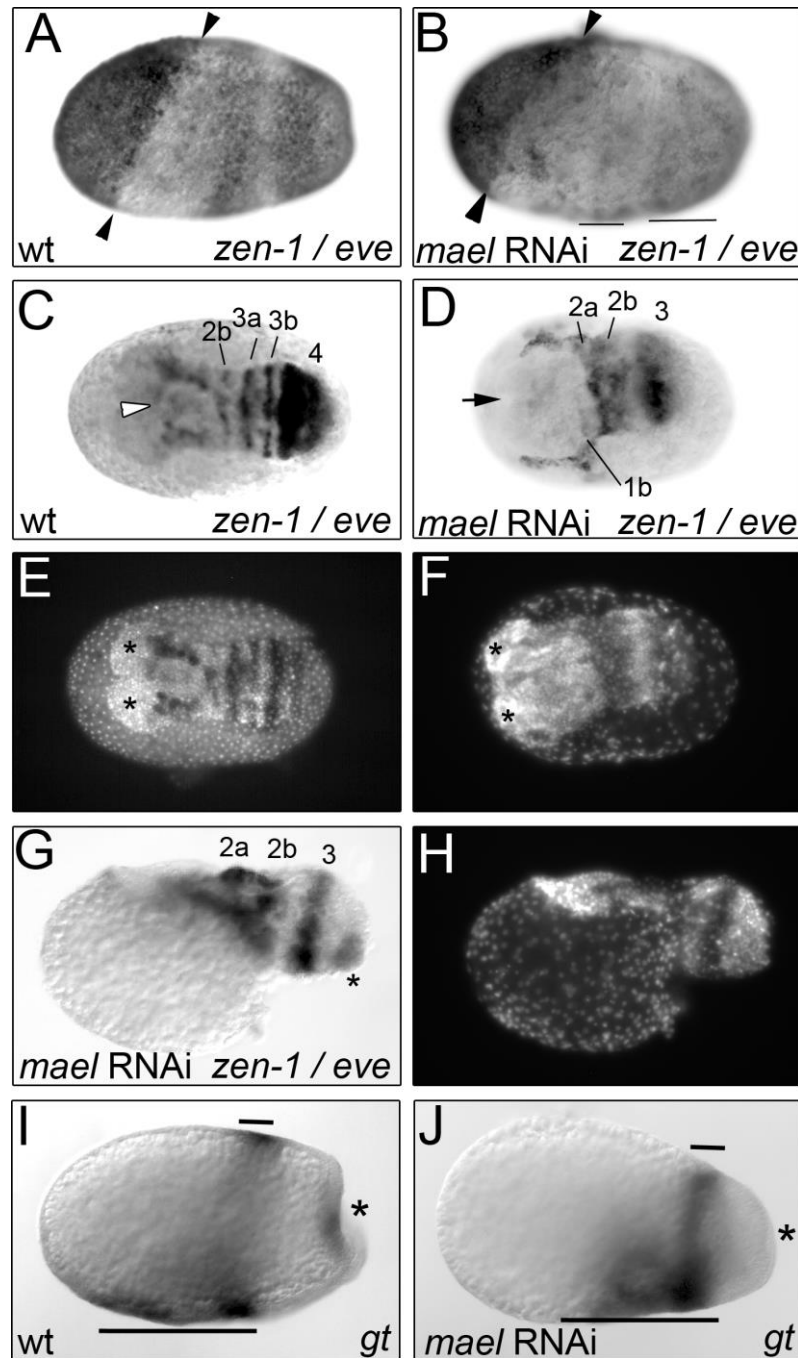
**Figure 6: *Tc-mael* RNAi**

Cuticle of Wildtype (A) and *Tc-mael* RNAi larvae (B-D). (B) Only remnants of pre-gnathal and gnathal segments are left (arrows). The third thoracic segment is still formed (asterisks), but all abdominal segments are deleted. (C) Weaker phenotypes show the reduction of the head capsule. All abdominal segments are deleted. (D) *Tc-mael*-RNAi also results in inside-out phenotypes affecting the whole larva or (E) only posterior segments.



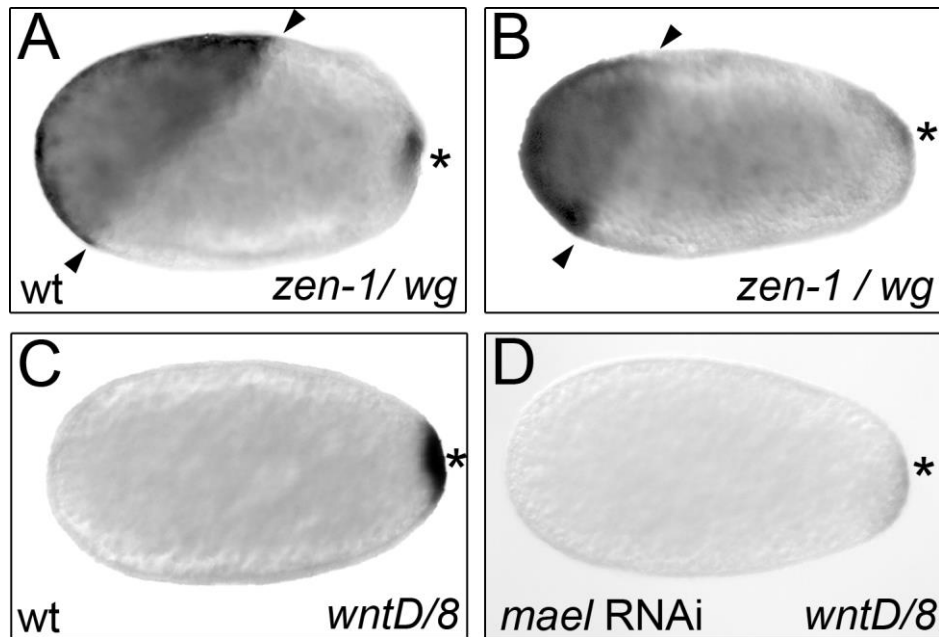
**Figure 7: Live imaging of *Tc-mael* RNAi**

Live imaging detection of wildtype (A-B, G-H) and *Tc-cic* RNAi (D-F, J-L) in the transgenic *Tribo-*lium** EFA-nGFP line. Columns show nGFP labelled stage matched embryo at representative time-points.(A-C) In wildtype, the embryonic anlagen gives rise to germ-rudiment that becomes covered by the serosa. (D-F) Upon *Tc-mael* RNAi, head anlagen are expanded and less condensed (E, Arrow). Also the rim of the serosa window is initially less condensed (E, Arrow and arrowhead). In wildtype, the germ-band extends posteriorly (G-I, asterisks). In *Tc-mael* RNAi posterior elongation fails (J-L, asterisks ). In addition, the serosa ruptures prematurely (K, L).



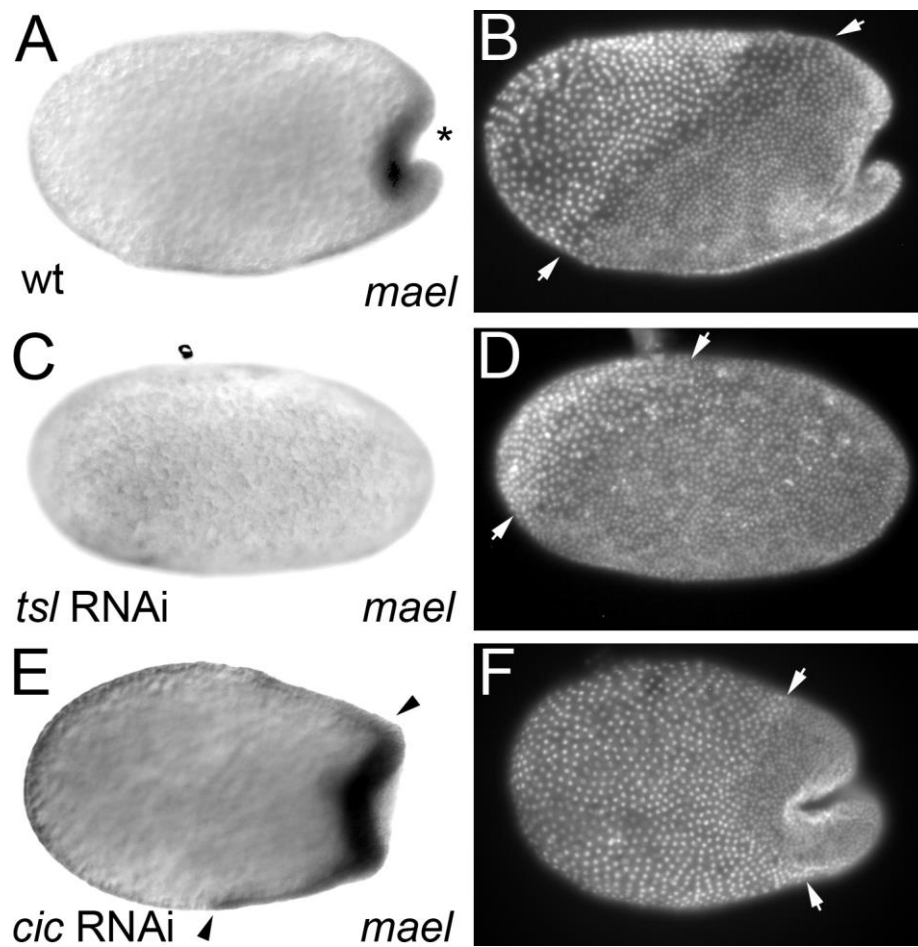
**Figure 8: Expression of *Tc-zen-1*, *Tc-eve*, and *Tc-gt* in *Tc-mael* RNAi**

Embryos double-stained for *Tc-eve* and *Tc-zen-1*(A-H) and *Tc-gt* (I,J). Embryos in (C, D, G) were subsequently stained for Hoechst 33342 (E, F, H). In *Tc-mael* RNAi, the serosa anlagen is reduced (B, arrowheads). The first and second primary *Tc-eve* domains (B, bars) are present (C, E). In wildtype, *Tc-zen-1* is expressed at the margin of the serosa window. (D, F) In *Tc-mael* knockdown, *Tc-zen-1* expression lost from the anterior rim of serosa window (D, arrow). The head primordia is less condensed (D arrow, F asterisks). G) The fourth primary *Tc-eve* domain does not form properly and in addition, posterior morphology eventually becomes highly irregular. I) In wildtype blastoderm embryos, *Tc-gt* is expressed in a head domain (bars) and at the posterior pit (asterisk). J) The posterior *Tc-gt* domain is lost in *Tc-mael* RNAi (asterisk).



**Figure 9: Expression of wnt-genes in *Tc-mael* RNAi**

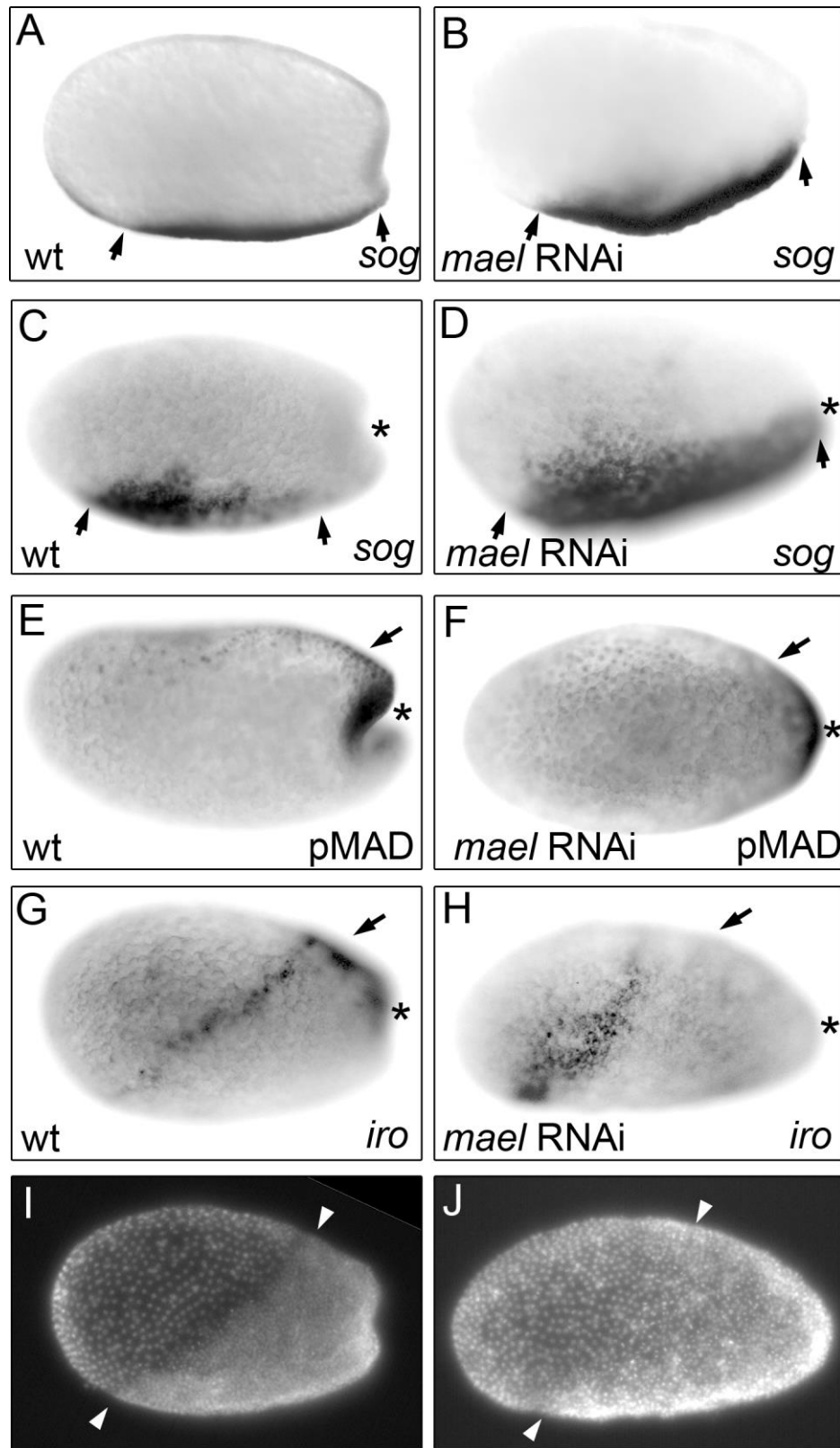
Expression of *Tc-wg* (A,B) and *Tc-wntD/8* (B, D) in wildtype (A, C) and *Tc-mael* RNAi (C, D). Embryos in (B, D) were subsequently stained for Hoechst 33342. Embryos in (A and B) were double staining with *Tc-zen1* (A) In wildtype undifferentiated blastoderm, *Tc-zen-1* covers the anlagen of the serosa and *Tc-wg* is expressed at the posterior pole. (B) In *Tc-mael* RNAi, the *Tc-zen-1* expression domain is reduced, while posterior *Tc-wg* expression is still present. (C) *Tc-wntD/8* is expressed at the posterior pole (asterisk). (D) Upon *Tc-mael* RNAi this domain is notably reduced (asterisk) or even absent.



**Figure 10: *Tc-mael* is a target gene of the terminal system**

Expression of *Tc-mael* in wildtype (A), *Tc-tsl* RNAi (C), and *Tc-cic* RNAi (E). Embryos in (A, C, E) were subsequently stained for Hoechst 33342 (B, D, F). The embryo shown A and B is the same as that in Figure 5A and B. (C) In *Tc-tsl* RNAi, *Tc-mael* expression is lost, while the *Tc-mael* domain expands upon *Tc-cic* RNAi (E). Arrows point to the serosa - embryo boundary.





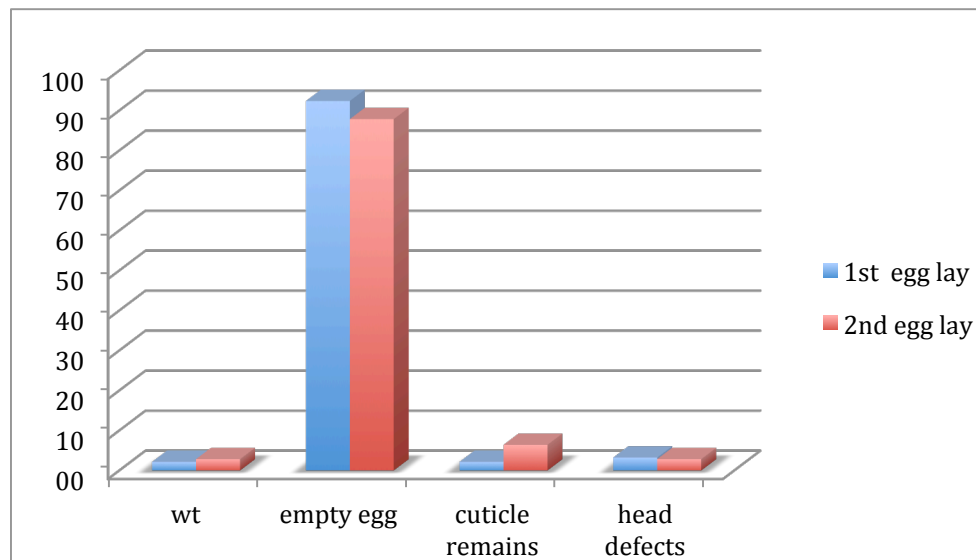
### Figure 11: *Tc-mael* RNAi in dorsoventral patterning

Wildtype (A, C, E, G, I) and *Tc-mael* RNAi (B, D, F, H, J) embryos stained for *Tc-sog* (A-D), pMAD (E, F), and *Tc-iro* (G-H). Embryos in (G, H) were subsequently stained for Hoechst 33342 (I, J). In wildtype blastoderm (A) *Tc-sog* is expressed in a broad ventral domain (arrows), which (C) eventually retracts from the posterior pole (asterisk). Upon *Tc-mael* RNAi the (B) early *Tc-sog* domain might be somewhat stronger (arrows), otherwise there is no pronounced difference. (D) Later, *Tc-sog* fails to retract from the posterior pole (asterisk). (E) Wildtype, Dpp activity (as monitored by pMAD staining) peaks in the posterior-dorsal region of the embryo (arrow, asterisk). In *Tc-mael* RNAi posterior pMAD signal is still present (asterisk), while dorsal dpp activity seems to be reduced (arrow). In wildtype (G, I), *Tc-iro* is expressed in the anterior and dorsal (arrow) amnion that expands towards the posterior pole (asterisk). In *Tc-mael* RNAi embryos, (H, J), anterior *Tc-iro* expression is maintained and still oblique. The posterior-dorsal expression domain is absent (arrow, asterisk).

## Supplementary Figures

**Figure S1: phenotypic analysis of *Tc-cic* RNAi experiments**

A)



RNAi phenotypes (%)	1st egg lay	2nd egg lay
wt	2,2	2,9
empty egg	92,4	87,9
cuticle remains	2,2	6,4
head defects	3,3	2,9

B)

RNAi (n)	1st egg lay	2nd egg lay
wt	2	4
empty egg	85	123
cuticle remains	2	9
head defects	3	4
total	92	140

Summary of RNAi experiments. Percentages (A) and numbers (B) of the observed phenotypic defects obtained with RNAi treatments using *Tc-cic* dsRNA (1µg /µl).

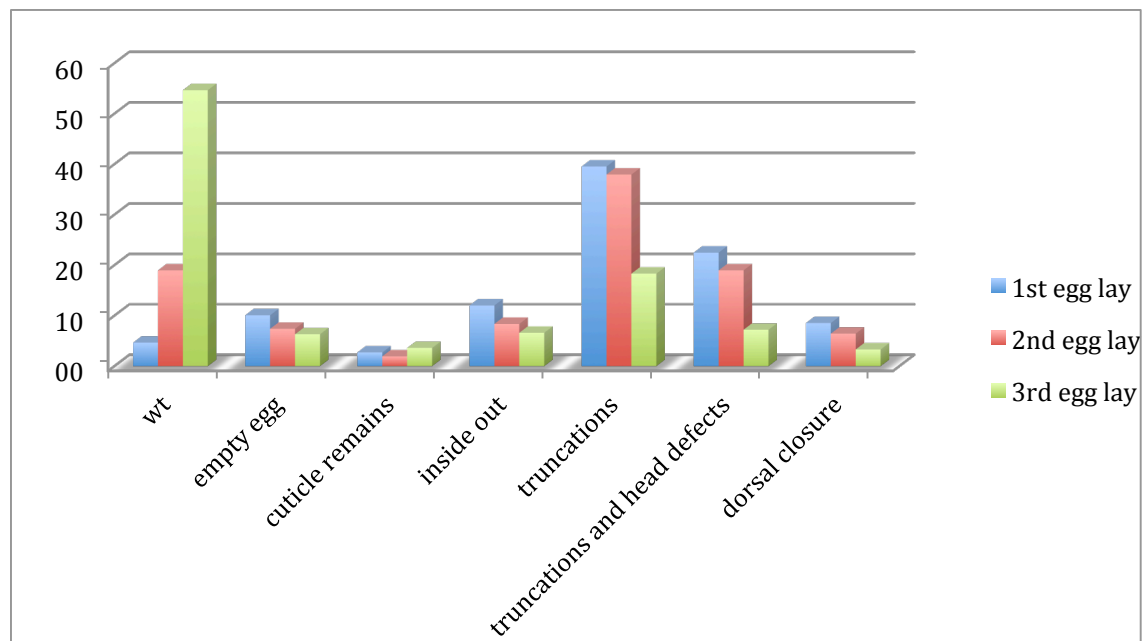
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**Figure S2: phenotypic analysis of *Tc-mael* RNAi experiments**

A)



RNAi phenotypes (%)	1st egg lay	2nd egg lay	3rd egg lay
wt	4,7	19,0	54,7
empty egg	10,1	7,4	6,3
cuticle remains	2,7	1,9	3,6
inside out	12,0	8,4	6,6
truncations	39,5	37,9	18,3
truncations and head defects	22,5	19,0	7,2
dorsal closure	8,5	6,4	3,3

B)

RNAi Phenotypes (n)	1st egg lay	2nd egg lay	3rd egg lay
wt	12	59	182
empty egg	26	23	21
cuticle remains	7	6	12
inside out	31	26	22
truncations	102	118	61
truncations and head defects	58	59	24
dorsal closure	22	20	11
total	258	311	333

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Relative expression values for wildtype (T1), tsl RNAi (T2), and cic RNAi (T3) as revealed by SOLiD Next Generation Sequencing. In total, 12,795 genes showed differences in mRNA expression levels. All genes were categorized according to their expression values. We found 1801 genes to be up-regulated in Tc-tsl RNAi and down-regulated in Tc-cic RNAi (class: +/-) and 2790 genes that were up-regulated in Tc-cic RNAi but down-regulated in Tc-tsl RNAi (class: -/+). In addition, 2157 genes were up-regulated (class: ++) in both, Tc-cic and Tc-tsl RNAi and 4589 genes down-regulated (class: --) in both, Tc-cic and Tc-tsl RNAi. Some genes that were not expressed in wildtype became expressed after Tc-tsl (class: +0) or Tc-cic RNAi (class: 0+).

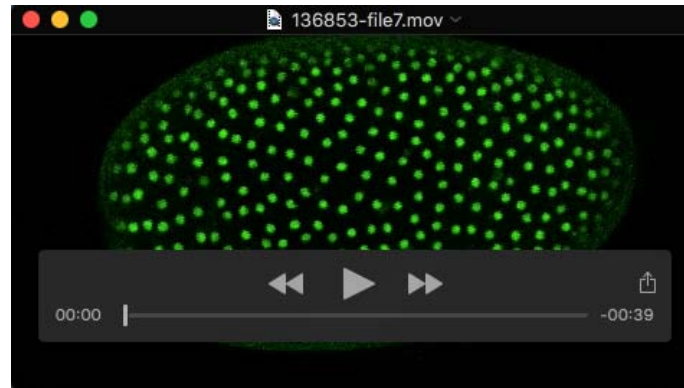
[Click here to Download Table S1](#)

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50 genes were selected for closer analysis. Relative fold change (T2 or T3 divided by T1) in tsl RNAi (T2) or cic RNAi (T3) is shown in comparison to wildtype (T1) (tsl RNAi vs wt: T2:T1, cic RNAi vs wt: T3:T1) Larvae were scored for RNAi phenotypes. Hoechst and  $\beta$ -Tubulin staining were used to visualize early embryonic phenotypes ("early defects"). Expression of candidate genes was monitored by whole-mount in-situ hybridisation.

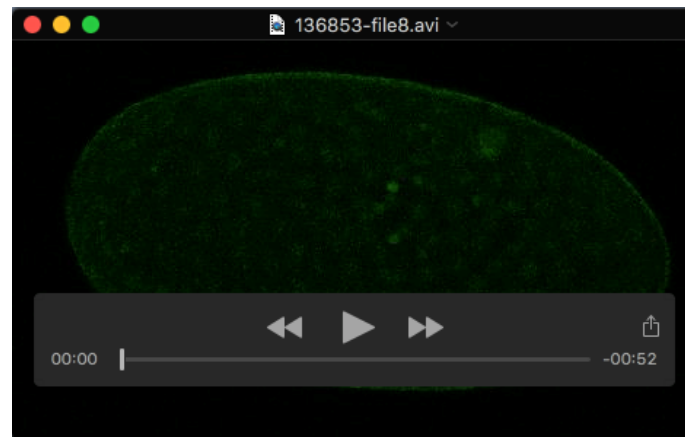
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## Supplementary Movies



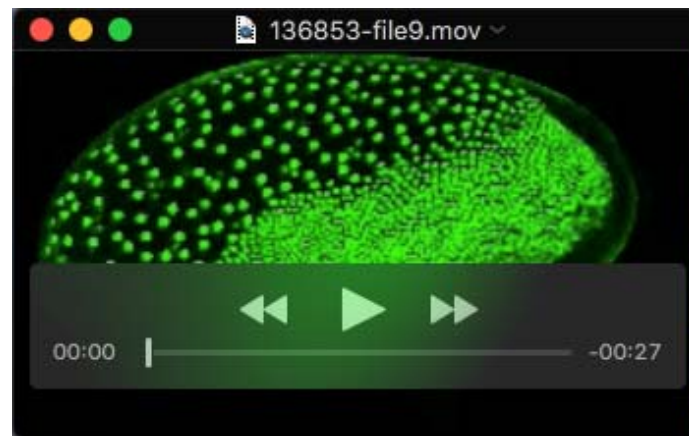
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12 hours Time-lapse fluorescence recording of a *Tribolium* embryo expressing nuclear-localized GFP under a ubiquitous promoter (transgenic line EFA-nGFP). Stacks were recorded every 12 minutes at 10x magnification and 20°C. After synchronous cell divisions, the embryonic anlagen condense and give rise to germ-rudiment that progressively becomes covered by the extending serosa. The germ-band extends posteriorly, bending around the posterior pole. Anterior to the left.



### Movie S2: Live imaging of early *Tc-capicua* RNAi embryo

12 hours Time-lapse fluorescence recording of a *Tribolium* embryo expressing nuclear-localized GFP under a ubiquitous promoter (transgenic line EFA-nGFP). Stacks were recorded every 12 minutes at 10x magnification and 20°C. Upon *Tc-cic* depletion, the serosa anlagen expands at the expense of the anterior head anlagen. Eventually extraembryonic membranes cover the entire egg. Embryonic tissue is restricted to the posterior pole of the egg and becomes internalized completely. Anterior to the left.



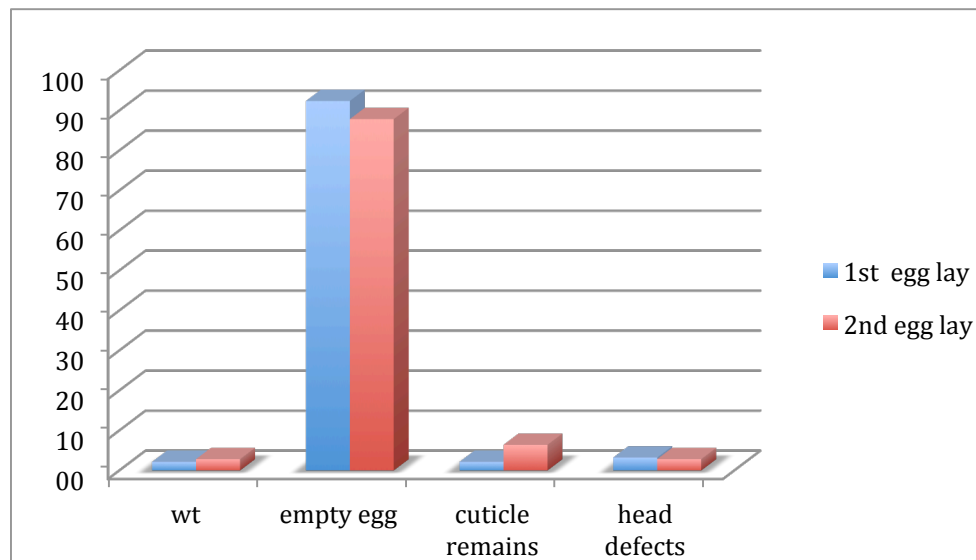
### Movie S3: Live imaging of early *Tc-maelstrom* RNAi embryo

12 hours Time-lapse fluorescence recording of a *Tribolium* embryo expressing nuclear-localized GFP under a ubiquitous promoter (transgenic line EFA-nGFP). Stacks were recorded every 12 minutes at 10x magnification and 20°C. Upon *Tc-mael* RNAi, head anlagen are expanded and less condensed as compared to wildtype. Serosa window formation remains incomplete to some degree and RNAi posterior elongation fails. Eventually, the serosa ruptures premature. Anterior to the left.

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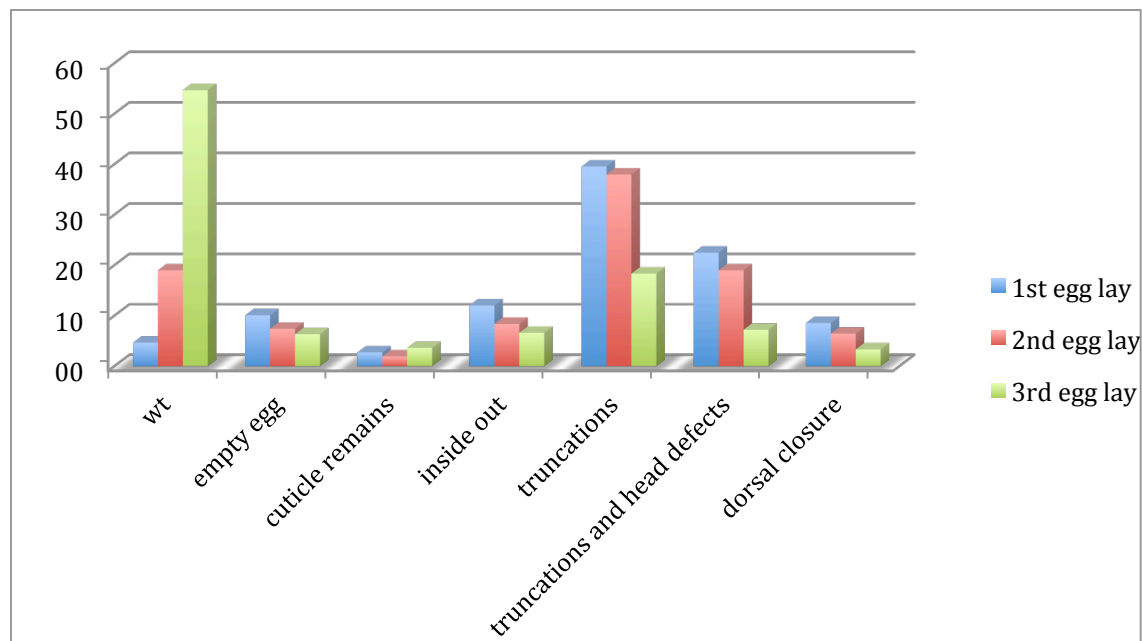
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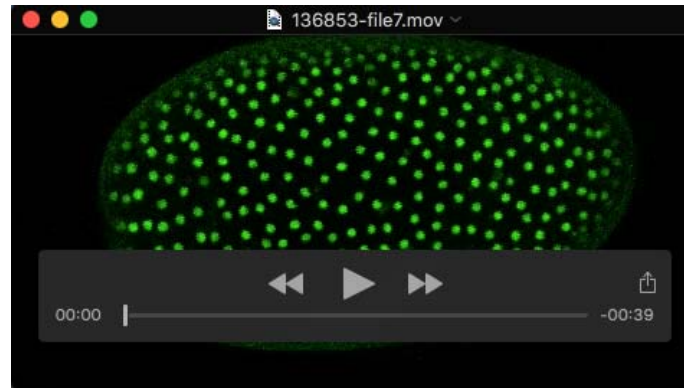
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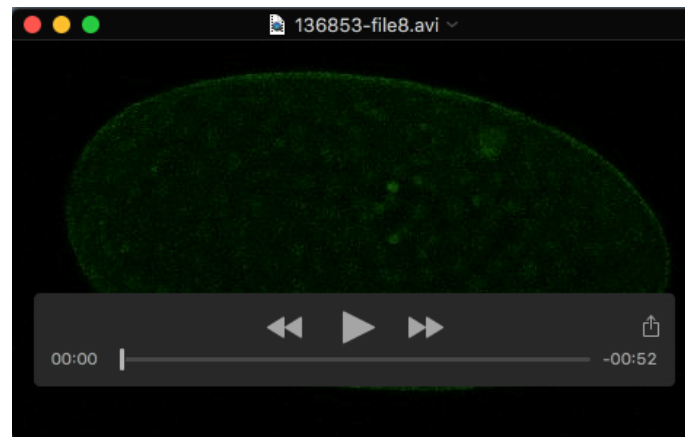
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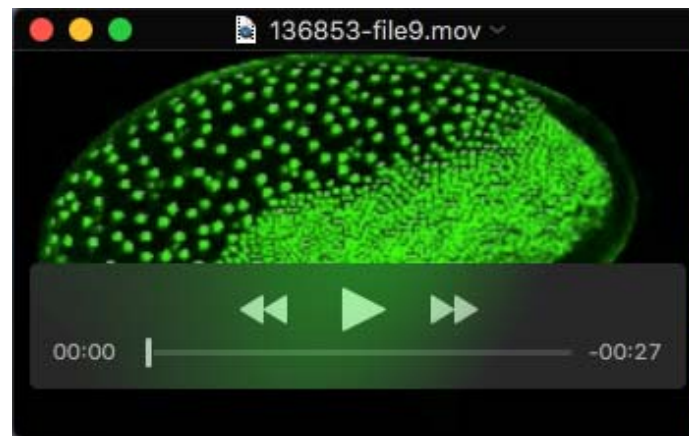
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