



# An evolutionary perspective on germ cell specification genes in insects

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# An evolutionary perspective on germ cell specification genes in insects

A dissertation presented

by

Benjamin Ewen-Campen

to

The Department of Organismic and Evolutionary Biology

In partial fulfillment of the requirements
for the degree of
Doctor of Philosophy
in the subject of
Biology

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### An evolutionary perspective on germ cell specification genes in insects

#### **ABSTRACT**

This dissertation investigates the embryonic specification of a specific group of cells: the germ cells. Germ cells, which give rise to sperm and egg, are the only cells in sexuallyreproducing animals that directly contribute hereditary information to the next generation. Germ cells are therefore a universal cell type across animals, and represent a profound novelty that likely arose near the base of the animal phylogeny. Yet despite their conserved, essential function in all animals, there is surprising diversity in the mechanisms that specify these cells during embryonic development. In this dissertation, I address the diversity of germ cell specification mechanisms in insects. I focus on two species, the milkweed bug Oncopeltus fasciatus (Hemiptera) and the cricket Gryllus bimaculatus (Orthoptera), which both branch basally to the Holometabola (those insects which undergo metamorphosis, including the well-studied fruit fly Drosophila melanogaster), and thus provide important phylogenetic breadth to our understanding of germ cell specification across insects. Using functional genetic approaches, I show that germ cell specification in both Oncopeltus and Gryllus differs fundamentally from germ cell specification in *Drosophila*. Specifically, I provide evidence that germ cells arise via inductive cell signaling during mid-embryogenesis, rather than via maternally-supplied cytoplasmic determinants localized in the oocyte, as is the case for *Drosophila*. These data suggest that Drosophila employs an evolutionarily derived mode of germ cell specification. In further support of this hypothesis, I show that several of the genes required for *Drosophila* germ cell specification perform other functions in both Oncopeltus and Gryllus. I demonstrate that one of these genes, oskar, which is the only gene both necessary and sufficient for germ cell specification in *Drosophila*, instead functions in nervous system of the cricket, both during embryonic development and in the adult brain. I suggest that the evolution of the derived mode of

iii

germ cell specification seen in *Drosophila* may have involved co-opting *oskar* into the germ cell specification pathway from an ancestral role in the nervous system.

# TABLE OF CONTENTS

Title Page	i
Copyright	ii
Abstract	iii
Table of Contents	v
Acknowledgements	vi
Dedication	viii
Introduction	1
Chapter 1: The maternal and embryonic transcriptome of the milkweed bug Oncopeltus fasciatus	26
Chapter 2: oskar predates the evolution of germ plasm in insects	54
Chapter 3: Germ cell specification requires zygotic mechanisms rather than germ plasm in a basally branching insect	
Chapter 4: Evidence against a germ plasm in the milkweed bug <i>Oncopeltus</i> fasciatus, a hemimetabolous insect	107
Chapter 5:  oskar functions in adult neural stem cells to influence long-term memory formation in the cricket Gryllus bimaculatus	126
Discussion and Outlook	144
Appendix A: The molecular machinery of germ line specification	154
Appendix B: Preliminary experiments on potential roles for <i>oskar</i> in the  Drosophila nervous system	170

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Announcement of Liz Ewen's dissertation defense (1979), "Immigrant Working Women in New York 1870-1920," on display in the Ewen Library (aka my uncle's bathroom).

INTRODUCTION

... Life is a continuous stream ...

The individual body dies, it is true, but the germ cells live on.

- E.B. Wilson, 1900, p 8-10

Germ cells: bridging the generation gap

Every sexually reproducing animal begins as a single cell: a fertilized egg. This egg cell,

prior to being fertilized, originates within the mother's body as the result of a dividing germ cell

in her ovary. Of course, the germ cells within the mother's ovary, like all the other cells in her

body, trace back to a single fertilized egg cell yet again, which itself arose from a dividing germ

cell within the ovary of the previous generation. This direct cellular link continues back through

the generations, every organism tracing to a single cell division event in its mother. Germ cells

thus directly connect the generations, serving as the physical and genetic link between parents

and offspring. As cell biologist E.B. Wilson wrote in 1900, "the body is, as it were, an offshoot

from the germ cell," (Wilson, 1900).

Given the seeming immortality of the germ cells (or "PGCs", for Primordial Germ Cells

as they are called when they first form in the embryo), the mechanisms that make these cells

different from the mortal somatic cells have intrigued biologists for centuries. The restriction of

reproductive potential to a small, dedicated group of germ cells appears to be a universal process

across sexually reproducing animals, one that probably arose at the dawn of animal

multicellularity (Buss, 1987). The indispensable function of germ cells might lead one to assume

that germ cell specification during development is a highly conserved process. However, despite

the homology of germ cells in all metazoans, the mechanisms that specify germ cells during

development have turned out to be remarkably diverse in different taxa (Extavour and Akam,

2003), raising questions about how this process has evolved across the animal phylogeny. This

1

dissertation is concerned with how PGC specification mechanisms have evolved in a particular group of animals: the insects.

#### Germ cell specification in Drosophila

Arguably our most detailed understanding of PGC specification in any animal comes from the fruit fly, *Drosophila melanogaster*. Over a century of embryological research has demonstrated that in this species, as in many other holometabolous insects (a monophyletic group of insects that undergo complete metamorphosis) PGCs are specified via cytoplasmic components that localize to the oocyte posterior while the egg cell is developing within the mother's ovary (reviewed in Mahowald, 2001). This specialized cytoplasm, termed "germ plasm," was first observed using histological techniques in a variety of holometabolous insects around the turn of the 20<sup>th</sup> century, including *Drosophila* in 1923 (**Figure 1.1A**) (Huettner, 1923). The germ plasm was shown to be asymmetrically inherited during the earliest cell divisions into a small group of so-called "pole cells" at the embryonic posterior (Hegner, 1914; Mahowald, 2001).

Beginning in the 1930s, several lines of experimental evidence in *Drosophila* demonstrated that if germ plasm was destroyed, for example by UV irradiation, it would lead to adults that lack germ cells, indicating that germ plasm is necessary for germ cell formation (Figure 1.1B) (reviewed in Mahowald, 2001). The sufficiency of germ plasm to autonomously induce PGC fate was decisively established in the 1970s, in a series of technically virtuosic germ plasm transplantation experiments conducted in Anthony Mahowald's laboratory. In these experiments, germ plasm from a donor embryo was transplanted to ectopic locations in a host embryo, where it was found to induce ectopic pole cells (Figure 1.1C) (Illmensee and Mahowald, 1976a; 1974). To demonstrate conclusively that these ectopic pole cells were functional, Illmensee and Mahowald took the additional step of transplanting the resulting ectopic pole cells

into the posterior of a second host embryo, where they formed functional germ cells (Illmensee and Mahowald, 1976b; 1974).

Together, these findings established that germ plasm was necessary and sufficient for PGC formation. However, the molecular components of germ plasm remained a mystery until the first genetic screens for maternal effect genes in *Drosophila*. These screens uncovered a group of "grandchildless" mutants, representing genes that function in the mother's ovary to correctly specify the PGCs in her offspring (Ephrussi et al., 1991; Lehmann and Nüsslein-Volhard, 1991; Schüpbach and Wieschaus, 1986). In subsequent years, detailed analysis of these and additional genes has provided an exquisitely detailed understanding of the molecular mechanisms that orchestrate PGC specification in *Drosophila*.

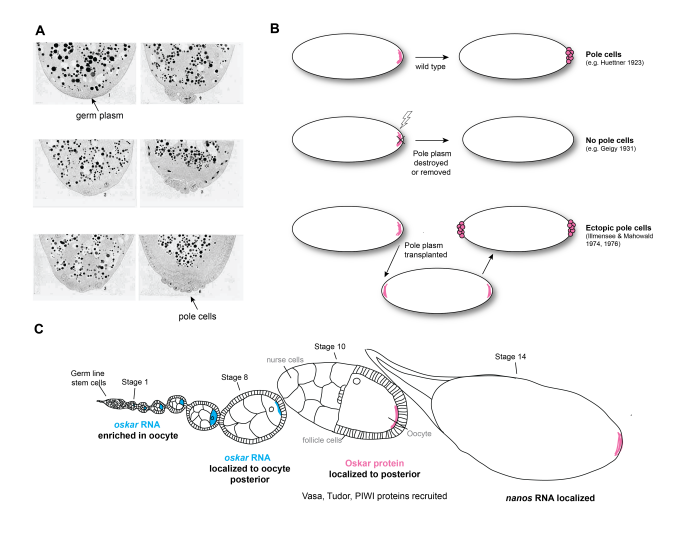


Figure 1.1 (previous page). Germ cell formation in Drosophila melanogaster. (A) Huettner's (1923) original description of germ plasm and pole cell formation in Drosophila. Germ plasm is visible as dark, punctate staining at the posterior cortex, which is subsequently taken up by the pole cells during cellularization. Posterior is down. (B) Summary of the experiments demonstrating that germ plasm is both necessary and sufficient for PGC formation. Posterior is to the right. (Reviewed in Mahowald, 2001). (C) Simplified summary of the molecular basis for germ plasm assembly during Drosophila oogenesis. Posterior is to the right. Ovariole traced from (Horne-Badovinac and Bilder, 2005). Figure adapted from (Lehmann and Ephrussi, 1994), with additional data from (Bardsley et al., 1993; Harris and Macdonald, 2001; Lasko and Ashburner, 1990).

During oogenesis, each *Drosophila* oocyte develops in conjunction with 15 clonallyrelated "nurse cells," which remain connected to the oocyte via cytoplasmic bridges (Figure 1.1C). These nurse cells synthesize mRNAs and proteins that are transferred into the oocyte, many of which are subcellularly localized within the oocyte and thereby establish the axes of the embryo. Among these nurse cell-derived factors is oskar RNA, which is transcribed by the nurse cell nuclei and transported via microtubules as an untranslated RNA to the oocyte posterior (Figure 1.1C) (Kim-Ha et al., 1995; 1991; Zimyanin et al., 2008). Upon posterior localization, oskar is translated into two protein isoforms, dubbed "Long Osk" and "Short Osk," which are thought to anchor Oskar to the posterior and to recruit downstream germ plasm components, respectively (Figure 1.1C) (Markussen et al., 1995). Via unknown biochemical mechanisms – Oskar is a novel protein lacking any functionally characterized domains – Short Osk recruits additional germ plasm components including Vasa, Tudor, PIWI-family proteins, and nanos, the latter of which has additional roles in abdominal patterning (Harris and Macdonald, 2001; reviewed in Mahowald, 2001). Following cellularization, the so-called "pole cells" at the posterior of the embryo inherit these cytoplasmic factors, which induce germ cell fate and repress somatic differentiation programs (reviewed in Seydoux and Braun, 2006).

Interestingly, while many of the germ plasm factors are necessary for germ cell formation, only Oskar has been shown to be sufficient for inducing germ cell formation in ectopic locations (Ephrussi and Lehmann, 1992; Smith et al., 1992). Given the sufficiency of Oskar to assemble

functional germ plasm, this gene is considered to represent the most upstream member of the germ cell specification pathway in *Drosophila*.

#### Patterns of germ cell specification mechanisms across Metazoa

The mode of germ cell specification in *Drosophila*, generically termed "cytoplasmic inheritance," represents one of the two modes of germ cell specification that have been described in animals. This mode, whereby maternally supplied cytoplasmic factors specify PGC fate, occurs in nearly all widely studied model organisms for developmental biology. The nematode *Caenorhabditis elegans*, the zebrafish *Danio rerio*, the frog *Xenopus laevis*, and likely also the chicken *Gallus gallus*, all form germ cells via cytoplasmic inheritance of maternally provided germ plasm.

In contrast, the mouse *Mus musculus* specifies PGCs via "zygotic induction," whereby cell-cell signaling induces PGC specification in the absence of a maternally supplied germ plasm. Accordingly, cell transplantation experiments in mouse embryos demonstrate that, as late as the initiation of gastrulation, distal embryonic cells that are experimentally grafted proximally into the location of normal PGC formation are capable of responding to local signals and differentiating into PGCs (Tam and Zhou, 1996). Thus, lineage restriction of the PGCs involves inductive signals and does not occur until well after the activation of the zygotic genome (reviewed in Saitou and Yamaji, 2012). Interestingly, once germ cells are specified in the mouse, they express orthologs of many of the genes found in *Drosophila* germ cells, including Vasa (Fujiwara et al., 1994; Toyooka et al., 2000), Nanos (Tsuda, 2003), Tudor-domain proteins (Chuma et al., 2006; Hosokawa et al., 2007), and PIWI proteins (Carmell et al., 2007; Deng and Lin, 2002; Kuramochi-Miyagawa et al., 2004), although several of these genes have sex-specific functions in the mouse.

The signaling pathways involved in mouse germ cell specification have been characterized in great detail via genetic analyses and cell culture experiments. During the second half of embryonic day 5, several BMP ligands are secreted from the extraembryonic ectoderm and visceral endoderm and synergistically signal via SMAD1/5 to upregulate the expression of a trio of transcription factors, *Blimp1*, *Prdm14*, and *Tcfap2c*, in a small subset of embryonic cells (reviewed in Saitou and Yamaji, 2012). These three transcription factors are necessary for PGC development, and are sufficient to induce functional PGCs *in vitro* from cultured epiblast-like cells (Magnúsdóttir et al., 2013; Nakaki et al., 2013). Together, these transcription factors upregulate hundreds of PGC-specific genes and down-regulate hundreds of soma-specific genes (reviewed in Saitou and Yamaji, 2012). Of the BMP ligands required for PGC specification, BMP4 is uniquely capable of causing epiblast cells to adopt functional PGC fate *in vitro*, a process that also requires Wnt3 signaling from nearby cells (Ohinata et al., 2009) and acts at least partially through the transcription factor *brachyury* to regulate the transcription of *Blimp1*, *Prdm14*, and *Tcfap2c*, which together upregulate hundreds of additional genes involved in PGC development (Aramaki et al., 2013).

Although the majority of widely studied model organisms specify PGCs via cytoplasmic inheritance, two lines of evidence suggest that zygotic induction may be the ancestral mode of PGC specification in Metazoa, and that the cytoplasmic inheritance mode has evolved independently multiple times in various lineages. First, an analysis of embryological literature covering nearly all animal phyla suggests that zygotic induction is in fact the more common mode of specification (Extavour and Akam, 2003). Further, this analysis revealed that cytoplasmic inheritance tends to be nested within clades whose basally branching members are thought to employ zygotic induction (Extavour and Akam, 2003; Extavour, 2007). Thus, given the phylogenetic distribution of PGC specification modes, parsimony suggests that cytoplasmic inheritance is a derived character that has arisen multiple times.

The second line of evidence that cytoplasmic inheritance has evolved independently several times is that although there is conservation of several of the molecular components of germ plasm across animals (reviewed in Ewen-Campen et al., 2010), the molecular mechanisms that assemble these components into functional germ plasm in different taxa are highly disparate, driven by genes that show no signs of homology. For example, although Oskar protein is necessary and sufficient to recruit all of the cytoplasmic factors necessary to generate functional germ cells in *Drosophila* (Ephrussi and Lehmann, 1992), this gene is a novel, insect-specific gene, and its role in germ cell specification is unique to holometabolous insects (Lynch et al., 2011).

In the zebrafish *Danio rerio*, which lacks *oskar*, formation of germ plasm in the oocyte and its asymmetric distribution during early cell cleavages requires *bucky-ball*, a novel vertebrate-specific protein with unknown molecular function (Bontems et al., 2009; Marlow et al., 2008). In *C. elegans*, the *pgl-1* and *pgl-3* genes are not *oskar* orthologs, but have a function analogous to *oskar*, as they can promote the assembly of germ plasm granules (P granules), including Vasa orthologs, in ectopic cellular contexts (Hanazawa et al., 2011; Updike et al., 2011; Wang and Seydoux, 2013). However, it is noteworthy that when P granules are formed ectopically, they are insufficient to confer germ cell fate on the cells containing them (Gallo et al., 2010), unlike analogous experiments with *Drosophila* or *Xenopus* (Ephrussi and Lehmann, 1992; Illmensee and Mahowald, 1976b; 1974; Tada et al., 2012). In the context of wild type *C. elegans* embryogenesis, the localization of germ plasm involves dynamic P granule dissasembly in the anterior and reassembly in the posterior, mediated via MEX5/6 and PAR-1, respectively, in a process with no clear molecular homology to germ plasm assembly in any other model system (Gallo et al., 2010).

In sum, the patchy phylogenetic distribution of germ plasm, together with the fact that distinct developmental and molecular mechanisms are used to form germ plasm in those species where it is observed, suggests that the cytoplasmic inheritance mode of PGC specification has evolved independently multiple times.

There are profound mechanistic differences between the cytoplasmic inheritance and zygotic induction modes of PGC specification. For those species that specify PGCs via cytoplasmic inheritance, the operative molecular mechanisms are those which maintain early embryonic cells in an essentially undifferentiated, totipotent state (reviewed in Seydoux and Braun, 2006). In *Drosophila*, for example, transcription is globally repressed in PGCs, a process mediated by the *pgc* protein, a small peptide which inhibits the transcriptional activity of RNA polymerase II by interfering with a critical phosphorylation event (Hanyu-Nakamura et al., 2008). A remarkably similar mechanism has independently evolved in *C. elegans*, where the germ cell-specific protein PIE-1 interferes with RNA polymerase II phosphorylation, effectively repressing transcription in the germ cells (Wang and Seydoux, 2013). There is also evidence that transcription is globally repressed at the level of chromatin modifications in both species (reviewed in Seydoux and Braun, 2006).

In contrast, for those species that specify PGCs inductively, mechanisms exist for reestablishing totipotency *de novo* in the PGCs. In the mouse, mechanisms exist to de-differentiate the PGCs towards the totipotent state that they held several days prior. Namely, PGCs must "undo" several aspects of somatic differentiation, including a DNA demethylation across the entire genome to remove imprinting and re-activate a recently de-activated X-chromosome, the upregulation of "pluripotency" factors including Nanog, Oct4 and Sox2, the active repression of somatic differentiation genes, and a variety of global changes in histone modifications (reviewed in Saitou and Yamaji, 2012).

Given the fundamental differences in the molecular mechanisms at play in these two modes of PGC specification, it is interesting to ask how this process has evolved across the animal tree. Namely, how could the cytoplasmic inheritance mode have arisen (likely multiple

times) from an ancestral mode of zygotic induction<sup>1</sup>? In order to address this issue, comparative studies of related species that differ in their PGC specification mechanisms are needed.

#### Insects: a case study in the evolution of PGC specification mechanisms

Insects provide an attractive opportunity to study the evolution of PGC specification mechanisms because basally branching taxa are thought to specify PGCs using zygotic induction, whereas derived lineages (including *Drosophila melanogaster*) form PGCs via cytoplasmic inheritance (Extavour and Akam, 2003). Thus, comparative developmental studies of basally branching taxa are poised to shed light on how cytoplasmic inheritance can evolve, a process which appears to have occurred repeatedly in a variety of phyla (Extavour and Akam, 2003; Extavour, 2007). Furthermore, an understanding of the specific signaling pathways that underlie zygotic induction in basally branching insects may provide new insight into how PGC identity can be conferred on embryonic cells, and whether this process has any conservation with other species that utilize the zygotic induction mode of PGC specification.

In the insects, the hypothesis that germ plasm is a derived character confined principally to Holometabola and their close relatives is based on surveys of embryological literature from the past 150 years (Extavour and Akam, 2003; Matsuda, 1976; Nieuwkoop and Sutasurya, 1981), coupled with experimental data from a small number of species (Lynch et al., 2011; Mahowald, 2001). Beginning in 1908, when Hegner experimentally removed the posterior cytoplasm from just-laid beetle embryos and observed the loss of germ cells (Hegner, 1908), cytoplasmic

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9

<sup>&</sup>lt;sup>1</sup> Some have also asked *why* these processes would evolve, i.e. what are the selective advantages of one specification mechanism versus the other. For example, Johnson and colleagues (2011) have suggested that in vertebrates, the uncoupling of PGC specification from somatic patterning in the cytoplasmic inheritance mode could allow for greater body plan evolution and thus increased speciation. In contrast, Buss (1988) found evidence for the opposite association across Metazoa (with many stated exceptions). In addition, Buss (1983) invokes the need to "police" potential cell-cell conflicts in any multicellular context as the operative selective pressure on the timing of PGC specification. From the standpoint of an experimental embryologist, these hypotheses seem difficult to test, and these ideas are not pursued further in this dissertation.

inheritance has been experimentally demonstrated in several additional species of Coleoptera (beetles), Diptera (flies and mosquitoes), and Hymenoptera (wasps, bees, and ants) (Ewen-Campen et al., 2013a; Hegner, 1914; Lynch et al., 2011; Mahowald, 2001). Moreover, pole cells have been cytologically identified in many additional holometabolous insects, including Lepidoptera (butterflies and moths), and several less well-studied orders such as Neuroptera (lacewings), Strepsiptera (twisted-winged flies), and Siphonaptera (fleas) (reviewed in Ewen-Campen et al., 2013a; Matsuda, 1976). Taken together, these data suggest that germ plasm and pole cells were present in the common ancestor of Holometabola (**Figure 1.2**).

Further, there is evidence that the germ plasm and pole cells in holometabolous insects are formed via homologous molecular mechanisms. Although functional molecular studies of *oskar*, *vasa*, and *nanos* have only been undertaken in *Drosophila melanogaster* and the wasp *Nasonia vitripennis*, these studies have revealed a largely conserved molecular program for germ plasm assembly in these species (Lynch et al., 2011; Lynch and Desplan, 2010). Given that *D. melanogaster* (Diptera) and *N. vitripennis* (Hymenoptera) phylogenetically bracket the holometabolous insects (Wiegmann et al., 2009), these studies support the hypothesis that germ plasm was present at the base of Holometabola, and suggest that its assembly involved *oskar* (**Figure 1.2**).

Interestingly, however, there are species within nearly all of the Holometabolous orders that lack cytologically discernable germ plasm and/or pole cells, including the moth-midge *Clogmia albipunctata* (Diptera) [but see (Jiménez-Guri et al., 2014)], the flour beetle *Tribolium castaneum* (Coleoptera), the honey-bee *Apis mellifera* (Hymenoptera), and the silkworm *Bombyx mori* (Lepidoptera) (reviewed in Ewen-Campen et al., 2013b; Lynch et al., 2011). Furthermore, in those species for which genomic data is available, the *oskar* gene is absent from the genome of those holometabolous species that lack germ plasm (Jiménez-Guri et al., 2013; Lynch et al., 2011). These observations have led to the hypothesis that the presence of *oskar* in the genome correlates with the cytoplasmic mode of PGC specification in insects, such that both the *oskar* 

gene and the cytoplasmic inheritance mode of PGC specification have been repeatedly lost in several holometabolous lineages (Lynch et al., 2011).

In stark contrast to the Holometabola, there is no experimental evidence of a germ plasm in any of the species that branch basally to Holometabola (the hemimetabolous insects, which do not undergo metamorphosis), and PGC specification in these taxa remains essentially a mystery. Histological observations of these taxa have typically failed to identify a discernable germ plasm or pole cells (Anderson, 1972a; Ewen-Campen et al., 2013a; reviewed in Matsuda, 1976; Nieuwkoop and Sutasurya, 1981). Instead, the embryological literature on hemimetabolous insects suggests that PGCs are not identifiable until later stages of development, in some cases not until the embryo is fully segmented, implying that zygotic induction must specify PGCs (See Table S1 in Ewen-Campen et al., 2013a).

The reported timing and location of PGC origin in hemimetabolous insects varies widely between species. In many Hemiptera ("true bugs"), cytologically-defined PGCs are first visible as a group of cells facing the yolk at the posterior of the blastoderm just prior to gastrulation (**Figure 3A**) (Butt, 1949; Heming and Huebner, 1994). At this stage, cellularization has occurred (Butt, 1949) and anterior segments have been specified, as revealed by *engrailed* expression (PZ Liu and T. Kaufman, 2004). These species are examples of relatively early PGC specification amongst the hemimetabola, but it is important to note this stage is still markedly later than PGC formation in *Drosophila*, and that germ plasm is not reported in either of these species, making it unclear if there could be a maternal contribution to PGC specification.

Reports from Orthoptera (crickets, grasshoppers, and locusts) differ from those in Hemiptera, and also vary widely between species. In the cricket *Achaeta domestica*, putative PGCs are reported at the egg posterior during the earliest stages of mesoderm formation [Heymons 1895, cited in (Matsuda, 1976)] whereas in two separate studies of the locusts *Concocephalus* (previous called *Xiphidium*) and *Locusta*, PGCs are reported to arise from mesodermal structures after the completion of segmentation (**Figure 1.3B**)

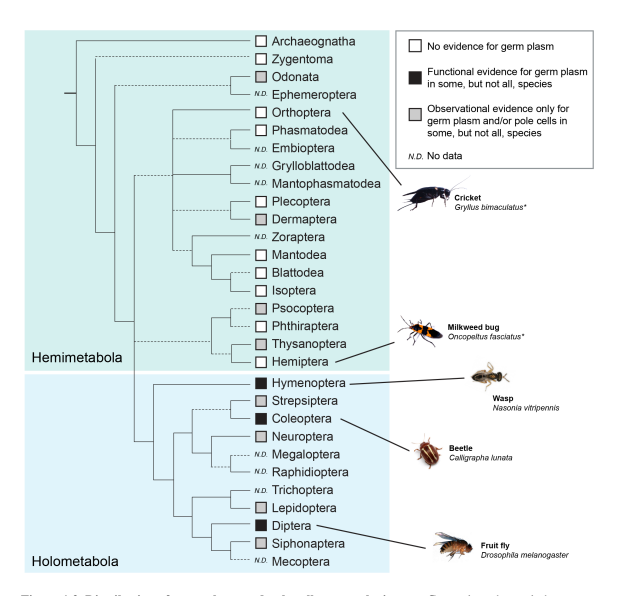
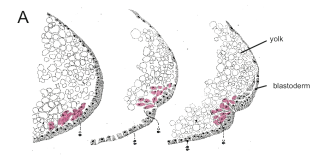


Figure 1.2. Distribution of germ plasm and pole cells across the insects. Germ plasm has only been demonstrated experimentally in holometabolous insects. Cytological evidence suggests that germ plasm is rare in hemimetabolous insects, and is absent from the closest hexapod outgroups, suggesting that zygotic induction is the ancestral mode of PGC specification in insects. Images are shown of species studied in this dissertation (*Gryllus* and *Oncopeltus*, indicated with astrices), or holometabolous species for which experimental evidence for germ plasm has been given, including *C. lunata*, one of the original beetle species studied by Hegner (1908). References are given in Supplemental Table 1 of (Ewen-Campen et al. 2013), with additional data for Odonata (Ando, 1962), Dermaptera (Singh, 1967), Neuroptera (Matsuda, 1976), and Strepsiptera (Matsuda, 1976). *N.D.* refers to orders for which, to my knowledge, germ cell origin has not been reported. Phylogeny from (Trautwein et al., 2012); dashed lines indicate uncertain relationships and/or possible polyphyly.

(Photo credits: *Gryllus* and *Oncopeltus* by David Behl, *Nasonia* from www.bioecologysrl.it, *Calligrapha* from bugguide.net, and *Drosophila* from http://www.lifesciencesfoundation.org/events-The\_Drosophila\_melanogaster\_genome.html)

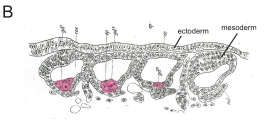
(Roonwal, 1937; Wheeler, 1893). A variety of PGC origin sites are reported for other hemimetabolous species, ranging from the "early" specification exemplified by Hempitera to the "late" specification exemplified by some Orthoptera. The distribution of these observations across species do not appear to follow a discernable phylogenetic pattern within the Hemimetabola (Extavour and Akam, 2003; Matsuda, 1976; Nieuwkoop and Sutasurya, 1981). In addition, PGC specification has not been reported, to my knowledge, for several of the smaller hemimetabolous orders, including Zoraptera, Mantophasmatodea, Grylloblattodea, Mantodea, and Ephemeroptera (Matsuda, 1976).

In spite of the notable variation observed between hemimetabolous species, there is general agreement amongst those who have reviewed this literature that PGC specification in holometabolous insects largely occurs via germ plasm and pole cells, whereas in hemimetabolous insects there is no evidence of germ plasm, implying that PGC specification must occur via other, unknown mechanisms (Anderson, 1972a; 1972b; Extavour and Akam, 2003; Matsuda, 1976; Nieuwkoop and Sutasurya, 1981). It should be noted, however, that a small number of hemimetabolous species are reported, based solely on histological observations, to specify PGCs very early, in a manner reminiscent of pole cell formation in *Drosophila* (Figure 1.3C-F). In one species of Thysanoptera (thrips), a cytologically identified germ plasm is localized to the oocyte posterior during late stages of oogenesis, and the cleavage nuclei that migrate into this germ plasm appear to become PGCs (**Figure 1.3C**) (Heming, 1979). In a species of Psocoptera (book lice), although germ plasm is not reported in the oocytes or just-laid eggs, PGCs form at the embryonic posterior just as the blastoderm forms, similar to pole cells (**Figure 1.3D**) (Goss, 1952). These two orders fall within the sister assemblage to Holometabola (Figure 1.2), and thus suggest that germ plasm and pole cells may have originated before the holometabolous radiation. In addition, there are also reports of germ plasm and pole cells in one species of Dermaptera (earwigs, Figure 1.3E) (Singh, 1967) and of pole cells (but not germ plasm) in one species of Odonata (dragonflies and damselflies, Figure 1.3F) (Ando, 1962), both of which branch much



"Early" PGC specification at the blastoderm posterior The milkweed bug *Oncopeltus fasciatus* (Hemiptera)

"The events that lead up to the formation of the embryo begin to occur early on the second day. At this time a clump of cells (gc [germ cells]) appears at the posterior pole of the egg situated within the yolk in close proximity to the inner surface of the blastoderm." (Butt 1947)

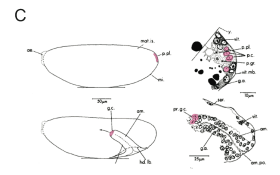


"Late" PGC specification in the segmental mesoderm of the abdomen

The locust Xiphidium (Orthoptera)

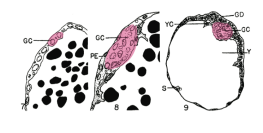
D

"In earlier stages, careful scrutiny failed to reveal any differentation of the mesoderm cells...it is not, therefore until the somites are established as distinct sacs that unmistakable primitive germ-cells make their appearance..."(Wheeler 1893)



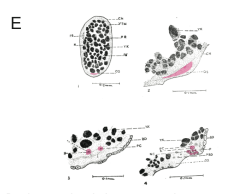
#### Putative germ plasm in the oocyte posterior The thrip *Haplothrips verbasci* (Thysanoptera)

"In fully developed eggs of *H. verbasci*, the pole plasm (p. pl) is situated at the posterior end...Pole (germ) cells (p.c.) first become recognizeable late in stage B when a few cleavage energids enter the pole plasm...Although I do not have experimental evidence to prove the existence of pole plasm and germ cell determinants in eggs of *H. verbasci*, I believe that both occur." (Heming 1979)



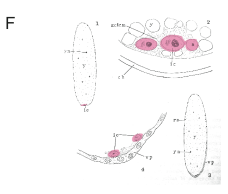
#### Putative pole cells during blastoderm formation The book louse *Liposcelis divergens* (Psocoptera)

"The germ cells become differentiated when the primary epithelium is completed. As far as could be determined, no structures that could possibly be identified as oosomes or germ cell determinants were found in these eggs..." (Goss 1952)



#### Putative germ plasm in the oocyte posterior The earwig *Labidura riparia* (Dermaptera)

"In a section of freshly laid egg of *Labidura riparia* an area of distinct cytoplasmic differentiation has been observed near the posterior end...some of them [cleavage nuclei] along with their trailing cytoplasm move into the Oosome to give rise to the primordial germ cells..." (Singh 1967)



#### Putative germ plasm in the oocyte posterior The dragonfly Epiophlebia superstes (Odonata)

"They are possible the primordial germ cells or "pole cells" which are found at the posterior of the eggs at the blastoderm formation...but final decision needs experimental prove, i.e. by cauterize method" (Ando 1962)

**Figure 1.3** (*previous page*). Examples of histological descriptions of PGC formation in various hemimetabolous insects. Images and quotations are excerpted from primary descriptions of PGC formation, with germ plasm and/or PGCs false-colored in pink. (A) An example of "early" PGC formation seen in many hemimetabolous insects, at the posterior of the blastoderm, in the milkweed bug *Oncopeltus fasciatus*. (B) An example of "late" PGC formation in the segmented abdominal mesoderm, from the locust *Concocephalus* (formerly *Xiphidium*). (C-F) Four atypical examples of hemimetabolous species described as possessing germ plasm and/or pole cells, although note that none of these examples have been reinvestigated using molecular techniques.

more deeply in the insect tree (**Figure 1.2**). It will be very interesting to re-examine these exceptional species using experimental techniques and molecular markers in order to test whether this mode is truly homologous to PGC specification in holometabolous insects such as *Drosophila*.

There is a major limitation of relying solely on histological and cytological data to determine the presence or absence of germ plasm: in some cases, the use of molecular markers can reveal a cryptic germ plasm that eluded prior histological studies. Indeed, in chicken, zebrafish, and *Amphioxus*, historical observations had suggested an inductive mode of PGC specification, but analysis of *vasa* gene expression revealed the presence of a maternally-supplied germ plasm (Gomez et al., 2011; Tsunekawa et al., 2000; Yoon et al., 1997). Thus, the use of molecular markers and functional genetic tests for PGCs and/or germ plasm is essential in order to test for the presence of germ plasm in any organism.

In the case of hemimetabolous insects, PGC gene expression studies have previously been limited to just three species, and have been far from conclusive. Studies of *vasa* expression in two orthopterans, the grasshopper *Schistocerca americana* and the cricket *Gryllus bimaculatus*, did not reveal a germ plasm in oocytes or early embryos, and left PGC specification an open question in these species (Chang et al., 2009; 2007; Mito et al., 2008). The parthenogenic embryos of the asexual-phase pea aphid *Acyrthosiphon pisum*, a hemipteran with a highly derived mode of viviparous development, appears to asymmetrically localize Nanos, but not Vasa, to the oocyte posterior and subsequently to the PGCs (Chang et al., 2009; 2007). It is not clear if this is the case for embryos developing in the sexual phase of *A. pisum*, as PGCs have not been

identified prior to segmentation stages (Miura et al., 2003). In addition, functional studies of "germ cell genes" have not previously been conducted in any hemimetabolous insects.

In sum, the mechanisms of PGC specification in basally branching insects remain unknown, and there is reason to believe it occurs in the absence of germ plasm. Thus, a molecular and functional understanding of PGC specification in hemimetabolous insects is necessary in order to understand how such a system may have evolved into the well-characterized mode observed in *Drosophila* and other holometabolous insects.

#### **Outline of this dissertation**

In this dissertation, I aim to add to our understanding of germ cell specification in basally branching insects, and thus to our knowledge of how germ cell specification mechanisms evolve. I focus on two hemimetabolous insects, the milkweed bug *Oncopeltus fasciatus* (Hemiptera) and the cricket *Gryllus bimaculatus* (Orthoptera), which occupy important phylogenetic positions in the insect phylogeny. *Oncopeltus*, a hemipteran, falls within the sister assemblage to Holometabola (along with Thysanoptera [thrips], booklice Psocoptera [booklice], and Phthiraptera [body lice]) (**Figure 1.2**) and therefore offer a relatively close comparison to such species as *D. melanogaster* and *N. vitripennis*. *Gryllus*, an orthopteran, branches far closer to the base of Insecta, offering a deeper phylogenetic comparison with Holometabola and with Hemiptera (**Figure 1.2**).

In addition to their informative phylogenetic positions, *Oncopeltus* and *Gryllus* are experimentally tractable for embryological studies, a non-trivial requirement for many evolutionarily interesting animals (Abzhanov et al., 2008). Both species breed in the laboratory at high density and low cost, laying hundreds to thousands of eggs each day, year-round. More importantly, protocols have been developed in both species for such essential embryological methods as gene expression analysis and functional knockdown via RNA interference (RNAi)

(Paul Liu and T. C. Kaufman, 2009a; 2009b; 2009c; Mito and Noji, 2008), which are essential for investigating PGC specification on a cellular and molecular level.

In Chapter 1 (Ewen-Campen et al., 2011), I describe a *de novo* embryonic and ovarian transcriptome database for the milkweed bug, *Oncopeltus*. For "non-model organisms" such as *Oncopeltus* and *Gryllus*, the rate-limiting step in developmental studies often continues to be the identification and cloning of individual genes-of-interest. Most commonly, individual genes are cloned using the time-consuming and failure-prone technique of degenerate PCR, limiting many studies to focusing on small numbers of candidate genes. With the advent of new sequencing technologies, however, it has become possible to assemble transcriptomic or genomic data *de novo* in the absence of a sequenced genome, and simultaneously identify thousands of genes simultaneously. In this paper, I present a simple conceptual framework for sequence assembly and annotation, and generate an easy-to-use database of over 10,000 genes expressed during oogenesis and embryogenesis of *Oncopeltus*. This work also established a cDNA synthesis protocol and analysis workflow which was later extended to the cricket *Gryllus bimaculatus* (Zeng et al., 2013) and the crustacean *Parhyale hawaiiensis* (Zeng et al., 2011), and laid the technical foundation for all subsequent work in this dissertation.

In Chapter 2 (Ewen-Campen et al., 2012), I focus on a particularly fascinating gene involved in the evolution of PGC specification: *oskar*. The only gene known to be both necessary and sufficient for germ cell specification in *Drosophila* (or any animal, for that matter), *oskar* sits atop the PGC specification pathway in *Drosophila* (Ephrussi and Lehmann, 1992). However, previous studies have suggested that this gene is absent from all hemimetabolous and non-insect genomes, and *oskar* was therefore believed to be a novel gene that evolved at the base of Holometabola (Lynch et al., 2011). In the course of sequencing the *Gryllus* transcriptome, I made the surprising discovery that *oskar* is present in this species, the first example of an *oskar* ortholog outside of Holometabola. I then showed that *Gryllus oskar* is involved in neural development rather than germ cell development. These results demonstrate that *oskar* evolved far

earlier in insect evolution than previously thought, and that its well-studied role in *Drosophila* germ cells is a relatively recent evolutionary innovation, possibly the result of co-option from an ancestral role in the nervous system.

In Chapter 3 (Ewen-Campen et al., 2013a), I describe in detail how PGCs arise in the cricket *Gryllus bimaculatus*. Using a variety of molecular markers, I provide evidence that PGCs first arise in close association with abdominal mesoderm, during abdominal elongation, long after cellularization has occurred, in the absence of a maternally-supplied germ plasm. I show that mesoderm is required for PGC specification using RNAi against *twist*, a transcription factor necessary for mesoderm formation. *twist* RNAi dramatically reduces PGCs, consistent with the hypothesis that PGCs arise from a pool of cells that also generate mesoderm. Lastly, I show that neither *vasa* nor *piwi* is required for PGC specification in *Gryllus*, suggesting that the roles of these genes in PGC specification in *Drosophila* may also be a derived character. In *Gryllus*, both *vasa* and *piwi* are instead involved in spermatogenesis in the adult male, reminiscent of their role in the mouse.

In Chapter 4 (Ewen-Campen et al., 2013b), I extend my studies of hemimetabolous PGC specification to the milkweed bug, *Oncopeltus fasciatus*. I perform an *in situ* hybridization screen of 19 genes with germ plasm expression in *Drosophila*, and show that none of these genes localizes posteriorly in oocytes or early embryos. I identify three *bona fide* molecular markers of PGCs (*vasa*, *tudor*, and *boule*), and show that transcripts all three of these markers are expressed ubiquitously in earliest stages of development, and only localize to germ cells after the cellularized blastoderm stage, just prior to gastrulation. These gene expression data argue against the presence of a germ plasm in *Oncopeltus*, and are consistent with previous histological observations of related species that suggested a post-cellularization origin of PGCs at the embryonic posterior (Butt, 1949; Heming and Huebner, 1994). Lastly, I show that, as in *Gryllus*, knockdown of these PGC markers does not disrupt PGC formation, and instead that *vasa* RNAi disrupts spermatogenesis.

In Chapter 5, I present unpublished results demonstrating the surprising finding that *Gb-oskar* functions in the adult brain during long-term olfactory learning and memory in the cricket *Gryllus*. In collaboration with the Mizunami lab (Hokkaido University), I show that RNAi against *Gryllus oskar* in adult crickets interferes with long-term (1 day) olfactory learning in an odor association assay, although short term (1 hour) associative learning is not disrupted. I show that *Gb-oskar* is expressed in a population of proliferative neuroblasts in the adult mushroom body, the anatomical substrate implicated in olfactory learning in insects. These data, together with recently published work on *oskar* in the *Drosophila* nervous system (Xu et al., 2013), demonstrate that *oskar* plays a conserved role in the nervous system in holometabolous and hemimetabolous insects, and support the hypothesis that the neural function may therefore be ancestral.

The research in this dissertation provides the first functional genetic investigation of PGC specification in basally branching hemimetabolous insects, and supports the hypothesis that the cytoplasmic inheritance mode of PGC specification known from *Drosophila* is a derived character, having evolved from a zygotic induction mode in ancestral insects. Furthermore, the evolution of germ plasm in insects may have involved the co-option of the *oskar* gene from an ancestral role in the nervous system to a novel function atop the PGC specification pathway.

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# **CHAPTER 1**



#### **RESEARCH ARTICLE**

Open Access

# The maternal and early embryonic transcriptome of the milkweed bug *Oncopeltus fasciatus*

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#### **Abstract**

**Background:** Most evolutionary developmental biology ("evo-devo") studies of emerging model organisms focus on small numbers of candidate genes cloned individually using degenerate PCR. However, newly available sequencing technologies such as 454 pyrosequencing have recently begun to allow for massive gene discovery in animals without sequenced genomes. Within insects, although large volumes of sequence data are available for holometabolous insects, developmental studies of basally branching hemimetabolous insects typically suffer from low rates of gene discovery.

Results: We used 454 pyrosequencing to sequence over 500 million bases of cDNA from the ovaries and embryos of the milkweed bug *Oncopeltus fasciatus*, which lacks a sequenced genome. This indirectly developing insect occupies an important phylogenetic position, branching basal to Diptera (including fruit flies) and Hymenoptera (including honeybees), and is an experimentally tractable model for short-germ development. 2,087,410 reads from both normalized and non-normalized cDNA assembled into 21,097 sequences (isotigs) and 112,531 singletons. The assembled sequences fell into 16,617 unique gene models, and included predictions of splicing isoforms, which we examined experimentally. Discovery of new genes plateaued after assembly of ~1.5 million reads, suggesting that we have sequenced nearly all transcripts present in the cDNA sampled. Many transcripts have been assembled at close to full length, and there is a net gain of sequence data for over half of the pre-existing *O. fasciatus* accessions for developmental genes in GenBank. We identified 10,775 unique genes, including members of all major conserved metazoan signaling pathways and genes involved in several major categories of early developmental processes. We also specifically address the effects of cDNA normalization on gene discovery in *de novo* transcriptome analyses.

**Conclusions:** Our sequencing, assembly and annotation framework provide a simple and effective way to achieve high-throughput gene discovery for organisms lacking a sequenced genome. These data will have applications to the study of the evolution of arthropod genes and genetic pathways, and to the wider evolution, development and genomics communities working with emerging model organisms.

[The sequence data from this study have been submitted to GenBank under study accession number SRP002610 (http://www.ncbi.nlm.nih.gov/sra?term=SRP002610). Custom scripts generated are available at http://www.extavourlab.com/protocols/index.html. Seven Additional files are available.]

#### **Background**

New and emerging model organisms occupy an increasingly important part of the developmental biology and developmental genetics research landscape. While studying a huge diversity of animals has long been the norm in the classical fields of experimental embryology and

functional morphology [see for example [1-3]], the molecular biology revolution and the advent of the "model system" concept [4] created demand for a small number of highly genetically manipulable organisms that could be intensively studied [5]. Research on these "big six" [sensu 6] genetic model organisms has led to enormous advances in our understanding of general principles of embryogenesis. However, placing these general principles in an evolutionary context requires broader taxonomic sampling. Many researchers have

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highlighted the need for developing new model organisms for specific comparative, evolutionary and ecological questions [6-8]. It has also been suggested, however, that the single gene expression approach of the last several decades of evolutionary developmental biology ("evo-devo") has outlived its usefulness, and that what are needed are not more model organisms, but rather a smaller number of groups chosen for the ability to functionally manipulate genes [9,10]. Sophisticated gene expression techniques and even stable germline transgenesis have been developed in a large array of models outside of the "big six" [see for example [11,12]]. The ancient history of the small RNA processing machinery [13,14] means that gene knockdown is a feasible goal for most organisms, as long as the sequences of genes of interest are available.

While whole genome sequencing is an increasingly viable option for some organisms, many new models, particularly within the arthropods, lack the large community resources necessary to finance and maintain annotation of a genome. For these reasons, many researchers studying nontraditional model organisms have turned to Sangersequenced EST libraries [see for example [15,16]]. In principle this method of gene discovery can lead to highthroughput expression and functional genetic analyses of multiple genes [see for example [17]]. In practice, however, most non-traditional organism studies are still subject to a gene discovery bottleneck. This is largely because at the scale needed to uncover rare developmental transcripts, Sanger-based EST sequencing quickly becomes technically and financially prohibitive for many labs working on organisms with smaller research communities. In addition, those smaller-scale EST projects that have been carried out are often not publically available in easily searchable formats, and their potential contribution to the developmental and evolutionary biology fields is thus limited.

Next-generation sequencing (NGS) offers comparative and evolutionary developmental biologists a way to obtain orders of magnitude more developmental gene data than ever before, at a fraction of its former cost. Several studies have demonstrated the feasibility of NGS for identifying SNPs for population studies and gene sequences for use as phylogenetic markers [18-35]. Unfortunately, the lack of suitable protocols for cDNA preparation, and of established pipelines for analysis have left this tool under-utilized by many evo-devo researchers. Furthermore, according to some estimates [35], few of these studies have been carried out at a scale large enough to provide significant recovery of rare transcripts, and therefore of developmental genes. Here we present an optimized protocol for synthesizing cDNA for 454 Titanium pyrosequencing, as well as a simple workflow for de novo assembly of the data without a reference genome, annotation and analysis of the

dataset, and a demonstration of its utility for comparative developmental genetics.

A large body of literature is dedicated to the development and genomics of holometabolous insects (insects undergoing complete metamorphosis between embryonic and adult stages). Tens of holometabolous insect genomes are now available, thanks largely to work on Drosophila melanogaster, other drosophilids, and dipteran disease vectors [36,37]. In contrast, relatively little is known about the development of hemimetabolous insects, which undergo incomplete metamorphosis. Although several of these insects are amenable to laboratory culture and a variety of experimental manipulations, molecular developmental studies are scarce, and gene discovery rates remain low. Notable exceptions among the Hemiptera are the aphid Acyrthosiphon pisum and the Chagas' disease vector Rhodnius prolixus, whose genomes are completed and in progress respectively [38,39]. However, the aphid genome has undergone extensive duplications and gene loss, possibly due to its unusual reproductive and ecological characteristics [38]. The mammalian blood feeding needs of R. prolixus make it a sub-optimal organism for developmental studies.

The milkweed bug *Oncopeltus fasciatus* (Figure 1A-D) has emerged as a promising hemipteran system for studying the molecular development of hemimetabolous insects [40-42]. It can be reared easily and cheaply in the laboratory, and has a long history as a laboratory animal for classical embryology and pattern formation studies [43-45]. More recently, robust protocols for *in situ* hybridization, live imaging of embryogenesis, and RNAi-mediated gene knockdown have been developed and successfully applied to the study of the evolution of development [see for example [46,47]].

Here we present the results of the sequencing and *de novo* assembly of the *Oncopeltus* ovarian and early embryonic transcriptome. We outline an assembly and analysis framework using a combination of existing tools and freely available custom-made command line computational tools, which we hope will make this approach to gene discovery accessible to comparative developmental biologists. We identify homologues of genes involved in all major signaling pathways and developmental processes, including biologically verified splicing isoforms for some genes. We also address the need for library normalization in these studies, and show that at large enough scales of NGS, large numbers of developmental genes can be discovered even with omission of a normalization step.

#### **Results and Discussion**

# Assembling the ovarian and embryonic transcriptome of O. fasciatus

We prepared cDNA from ovaries and early to mid-staged embryos of *O. fasciatus*, covering oogenesis and all major

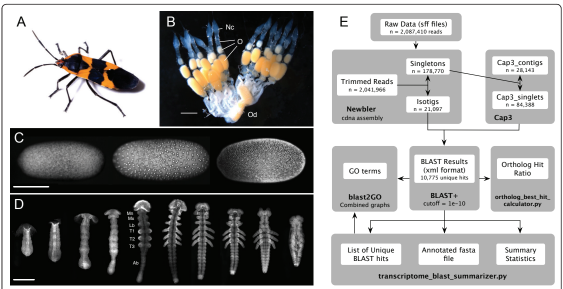


Figure 1 Introduction to *Oncopeltus fasciatus* and the workflow for producing a *de novo* transcriptome assembly. (A) An adult milkweed bug, *Oncopeltus fasciatus*. (B) Ovaries of adult female. Anterior is up. Oocytes (O) are visible in progressive stages of growth before reaching a common oviduct (Od). Oocytes are cytoplasmically connected to nurse cells (NC) in the anterior of each ovariole. Scale bar = 1.0 mm. (C-D) The stages of *O. fasciatus* embryogenesis represented in this transcriptome. Embryos are stained with Sytox Green (Invitrogen) to visualize nuclei. Scale bars = 0.5 mm. (C) Development proceeds from left to right. Anterior is to the left. The cellularized blastoderm forms during the first ~20% of development (~0-24 hours at 28°C), as nuclei reach the surface of the yolk and repeatedly divide. (*D*) Germ band extension and segmentation occur from ~20-60% of development (~24-72 hours at 28°C). Development proceeds from left to right. Anterior is up. Mn = mandibular segment; Mx = maxillary segment; Lb = labial segment; T1-T3 = leg-bearing thoracic segments 1-3; Ab = abdomen. (*E*) The flow of information during this *de novo* transcriptome assembly project. Data files are represented as white boxes within grey boxes that indicate the computer programs used to generate these files. All of the computer programs used are freely available. Ortholog\_best\_hit\_calculator.py and transcriptome\_blast\_summarizer.py are custom python scripts available at http://www.extavourlab.com/protocols/index.html (see text for details). Photograph in (A) courtesy of David Behl.

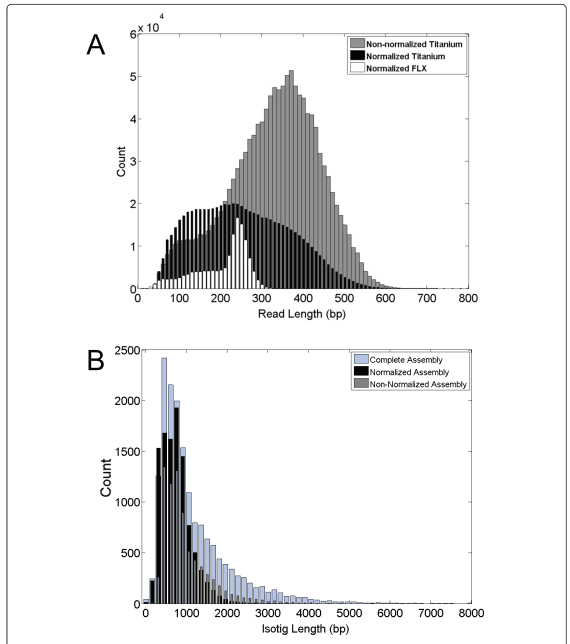
stages of embryonic patterning (Figure 1B-D). These cDNA samples were prepared using a protocol optimized for preparation of small or limiting samples for 454 pyrosequencing (see Materials and Methods). From these libraries, we generated a total of 2,087,410 sequence reads (Table 1). This includes reads generated using GS-FLX technology as well as both normalized (N) and nonnormalized (NN) cDNA sequenced using the GS-FLX Titanium platform. As expected, the reads generated using GS-FLX Titanium technology were substantially longer than those generated using GS-FLX technology (Table 1, Figure 2A). However, the N sample gave an

unexpectedly low number of reads, which were on average shorter than those generated by the NN sample (Table 1; Figure 2A). Given that a pilot run of one lane (1/8 plate) of this same normalized cDNA sample generated roughly equal number and size-distribution as a NN pilot study (Additional file 1), we suspect that a technical error reduced the sequencing efficiency of this plate. Despite the comparatively low yield of this normalized cDNA, it still generated more than 600,000 high quality reads that we therefore included in subsequent analyses.

We used the cDNA assembly algorithm of Newbler v2.3 (Roche) to screen the reads for adaptor sequence

Table 1 Sources of O. fasciatus sequence reads

		•					
Tissue	Normalized?	cDNA prep	454 Platform	No. Plates	No. Reads	Median Read Length	Accession #
Ovary	Υ	SMART	GS-FLX	1/4	65,394	225	SRR057570.2
Embryonic	Υ	SMART	GS-FLX	1/4	71,911	230	SRR057571.1
Ovarian and Embryonic	Υ	Modified SMART	GS-FLX Titanium	1 + 1/4	656,782	244	SRR057572.1
Ovarian and Embryonic	N	Modified SMART	GS-FLX Titanium	1 + 1/8	1,293,323	313	SRR057573.1
Total				2 + 7/8	2,087,410	301	SRP002610.1



**Figure 2 Effects of normalization and 454 sequencing chemistry on read length and isotig length.** (A) Titanium sequencing chemistry (grey, black) generally results in longer read lengths when compared with FLX chemistry (white). However, the normalized sample run with Titanium chemistry (black) had shorter read lengths than the non-normalized sample (grey). This result is likely due to a technical error in that particular sequencing run, since a 1/8 plate run of the same sample showed a read length distribution comparable to that of the non-normalized sample (Additional file 1). (β) Isotig length distributions from assemblies of Titanium-sequenced data. The longest isotig per isogroup is shown. The number of bases in the non-normalized (grey) and normalized (black) samples has been equalized to eliminate possible bias due to the greater number and length of reads obtained from the run of the normalized sample (see (A)). The isotigs generated from the normalized cDNA tended to be shorter than those produced by the non-normalized cDNA (see also Table 2). Pooling all FLX and Titanium reads generates an assembly with more, longer isotigs (blue).

and then assemble the cleaned reads (see Note Added in Proof for a comparison with Newbler v2.5). After quality trimming and adapter screening, 2,041,966 reads (97.8%) were used in the assembly. Of these, 1,773,450 (86.9%) assembled either wholly or partially into contigs, and 178,770 (8.8%) remained as singletons. The remaining reads were excluded as either originating from repeat regions (9,875 reads; 0.05%), outliers (26,943 reads; 1.3%), or too short (<50 base pairs: 52,928 reads; 2.6%).

To our knowledge, Newbler v2.3 and higher are the only assembly programs that address alternative splicing and can output multiple isoforms per gene. Newbler v2.3 explicitly accounts for alternative splicing by creating a hierarchical assembly composed of three elements: contigs, isotigs, and isogroups. For consistency, we follow their terminology. Contigs are stretches of assembled reads that are free of branching conflicts. In other words, contigs can be thought of as exons or sets of exons that are always co-transcribed. Isotigs represent a particular continuous path through a set of contigs, i.e. a transcript. An isogroup is the set of isotigs arising from the same set of contigs, i.e. a gene. Different isotigs within an isogroup are thought to represent alternative isoforms of the same gene. Note that it is possible for an isogroup to contain only one isotig, and it is also possible for an isotig to be composed of only one contig.

After the initial Newbler assembly, we noticed substantial redundancy among the singletons. We therefore subjected the 178,770 unassembled singletons to a secondary assembly with CAP3 [48]. This secondary assembly reduced the number of singletons from 178,770 to 112,531 (28,143 cap3\_contigs and 84,388 cap3\_singlets). Thus, in total, our assembly generated a total of 133,628 sequences, including isotigs, cap3\_contigs and cap3\_singlets (Table 2).

Our data assembled into 22,235 contigs, organized among 21,097 isotigs (Figure 2B). The isotig N50 length was 1,735 bp (in other words, 50% of the bases are incorporated into isotigs  $\geq$  1,735 bp), and 14,460 (68.5%) of the isotigs contained only one contig. The 21,097

isotigs fell into 16,617 isogroups, of which 14,562 (87.6%) contain only one isotig (average number of isotigs per isogroup = 1.3).

The average coverage among contigs was 23.2 reads/bp (median coverage = 6.9 reads/bp) (Additional file 2). This coverage value is more than twice as high as the highest reported value from a *de novo* transcriptome assembly to date [summarized in [20]]. Such deep coverage should be helpful for overcoming the presence of insertion/deletion errors in the individual raw reads [49].

To test whether our assembly would have been aided by the inclusion of nucleotide sequence from *Rhodnius* prolixus, the most closely related hemipteran to O. fasciatus whose genome is currently being sequenced [39], we used the BLASTN algorithm to compare our isotigs (the longest isotig per isogroup) with the published ESTs of R. prolixus with an e-value cut-off of 1e-6. Consistent with previous observations of extremely low levels of conservation between insect genomes [50] we found that only 53 out of 16,617 isotigs had hits to R. prolixus ESTs on the nucleotide level. These results suggest that de novo sequencing and assembling efforts will be necessary for most insect species, even when sequence data are available for other members of the same order. We note, however, that a recent study [51] has shown that it may be possible to incorporate EST data from different species into a de novo assembly by using amino acid sequence rather than nucleotide sequence.

#### Validation of predicted alternate isoforms

To examine whether the alternative isoforms predicted by Newbler v2.3 are in fact present in developing embryos of *O. fasciatus*, we first focused on a gene of particular interest to developmental biologists, *nanos*. This conserved metazoan gene was first described as a loss of function mutation in *Drosophila melanogaster* [52], and is necessary for germ cell and posterior somatic development [reviewed in [53]]. Newbler v2.3 predicted this gene to encode two alternative isotigs within a single isogroup (Figure 3B). The two isotigs differ in that the

Table 2 O. fasciatus transcriptome assembly statistics

	Full Assembly	Normalized Assembly	Non-Normalized Assembly
Assembled reads (base pairs)	1,773,450 (508,738,047)	389,605 (84,353,140)	336,568 (108,372,883)
Isogroups ("genes")	16,617	10,581	7,591
Isotigs ("transcripts")	21,097	11,353	8,346
Isotig N50	1,735	846	1,162
Mean # isotigs per isogroup	1.3	1.1	1.1
Contigs ("exons")	22,235	11,839	8,731
Mean # contigs per isotig	1.9	1.2	1.3
Singletons (singletons after secondary CAP3 assembly)	178,770 (112,531)	110,265 (N/A)	52,585 (N/A)

To enable comparison, we equalized individual assemblies of Normalized and Non-Normalized samples to contain the same number of base pairs before assembly.

longer contains an additional 100-bp exon that is absent from the shorter (Figure 3B). We designed PCR primers against sequences present in both isotigs (Figure 3B arrows), which amplified two bands differing by  $\sim 100$  bp from a pool of embryonic cDNA (Figure 3C). Sequencing of these two bands confirmed that they differ exactly as predicted by Newbler v2.3 (Figure 3D).

Importantly, a previous version of Newbler (v2.0), which does not account for alternative splicing, failed to join together the three fragments which were linked by Newbler v2.3 (Figure 3A). Because of this, Newbler v2.0 (and presumably other assemblers which do not address branching within contigs) predicted three separate contigs, only one of which could be identified as *nanos* with BLASTX, as the others fall in poorly conserved regions of the gene. Thus, the ability of Newbler2.3 to handle branching conflicts between reads allows this program to assemble longer continuous sequences, which are therefore in turn more easily annotated using BLAST.

To further characterize the accuracy of Newbler's predictions of alternative transcript isoforms, we randomly selected 10 isogroups that contained exactly two alternative isotigs differing by the presence/absence of a single contig (Additional file 3). As we did for nanos, we designed primers to flank the region differing between the two predicted isoforms (Additional file 3A), and performed RT-PCR on O. fasciatus embryonic cDNA. In eight of ten instances, we observed bands of the predicted sizes following agarose gel electrophoresis (Additional file 3B,C). However, in four of the eight positive cases, additional, unpredicted bands were present (Additional file 3). In one of the ten cases, we observed two RT-PCR products, but only one of them was of the predicted size (Additional file 3C, lane 6). Taken together, these results suggest that Newbler v2.3 has a low rate of false positives in the prediction of multiple splicing isoforms. Including our investigation of nanos, only one of 11 test cases (9.1%) produced a single RT-PCR product where Newbler v2.3 had predicted multiple products.

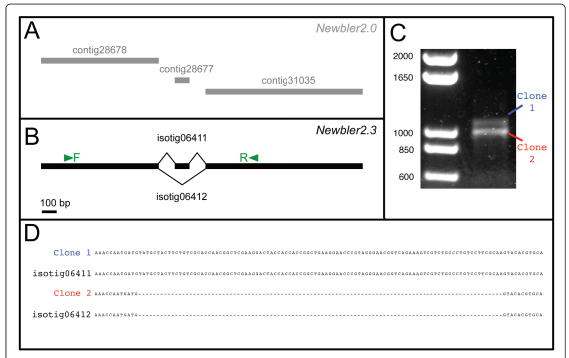


Figure 3 Newbler 2.3 correctly identifies splicing isoforms of *nanos*. (A) Newbler v2.0 identified three separate contigs that map to an O. fasciatus nanos homologue that we had previously identified by degenerate PCR (Ewen-Campen & Extavour, unpublished). Newbler v2.0 failed to identify these contigs as belonging to the same transcript because of branching conflicts amongst the reads joining these contigs. BLASTX against the RefSeq protein database identified only contig 31035 as being a putative *nanos* homologue; the other two contigs lie outside the conserved Nanos domain and obtain no BLAST hits. (B) Newbler v2.3 predicted that the same three contigs identified by Newbler v2.0 belonged to two isotigs, or splicing isoforms. (C) RT-PCR with specific primers F and R shown in (B) resulted in two bands of the predicted sizes of the isotigs predicted by Newbler v2.3. (D) Sequencing the bands from (C) revealed that they were identical to the sequences of the predicted isotigs from (B).

However, we observed that roughly half of the time, Newbler v2.3 failed to predict all of the isoforms identified via RT-PCR.

#### Transcriptome annotation

A BLASTN search of our dataset for the 93 existing GenBank accessions for *O. fasciatus* sequences yielded a hit result for 56% of the accessions, with an e-value cutoff of 1e-10. This result may be due in part to the short length of some of the GenBank sequences. Accordingly, we found that accessions with hits in the database were significantly longer (mean length 729 bp) than accessions without hits (mean length 397 bp) (unpaired Student's t-Test: t = 2.89, DF = 91, p = 0.0048). Of greater relevance to developmental applications of this dataset, however, was our finding that 85% of *O. fasciatus* developmental genes with existing GenBank accessions (n = 32) are represented in our transcriptome.

We then used BLASTX to map the 133,628 O. fasciatus sequences (isotigs, cap3\_contigs and cap3\_singletons) against the entire RefSeq Protein database with an e-value cut-off of 1e-10. To simplify these statistics, we report only the BLAST results for the longest isotig per isogroup, under the assumption that all isotigs within an isogroup share nearly identical BLAST results. Of 16,617 isotigs, 7,219 (43.4%) had at least one hit. Of the 28,143 cap3\_contigs, 2,594 (9.2%) had hits, and of the 84,388 cap3\_singlets, 2,367 (2.8%) had hits. These values are higher than comparable BLAST statistics of most other published studies of 454-generated de novo transcriptomes [24-26,30,32,33], likely because deeper sequencing increases the length of assembled sequences and thereby makes these sequences more likely to be identified via BLAST. The unidentifiable sequences likely originate from UTRs or non-conserved portions of protein-coding sequences. Of the top BLAST hits, 89.3% were genes from arthropod sequences (Additional file 4). Of the 12,180 O. fasciatus sequences with BLAST hits, 1,455 hit non-overlapping segments of the same top BLAST hit (i.e. potentially unassembled portions of the same transcript), and 825 hit overlapping segments of the same top BLAST hit (i.e. potential paralogs). Excluding those 1,455 potentially double-counted BLAST hits, our transcriptome identified a total of 10,775 genes. The assembled sequences generated in this study, as well as pre-computed BLAST results, are available as flat files from the authors upon request.

To explore and summarize the functional categories of the genes sequenced in this study, we obtained the Gene Ontology (GO) terms associated with the top 20 BLAST hits of each sequence using Blast2GO [54]. Among the 7,059 genes for which we obtained GO terms, we observed a wide diversity of functional categories represented on all levels of the Gene Ontology database

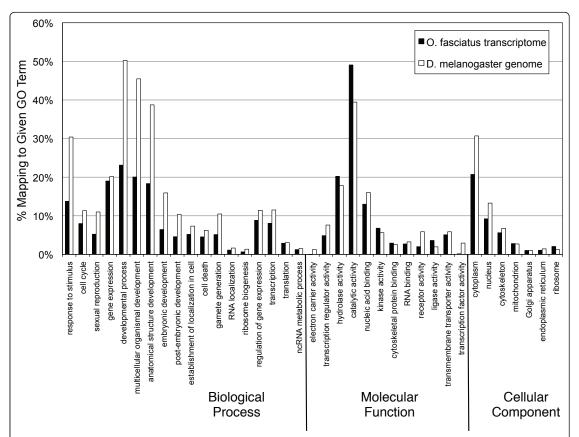
(Figure 4). The *O. fasciatus* sequences fall into GO categories with a roughly similar distribution to that of the well-annotated *Drosophila melanogaster* genome, suggesting that our sequence data contain a large diversity of genes involved in a variety of biological processes, and do not contain any notable biases towards particular categories of genes.

#### Assessing coverage of the O. fasciatus transcriptome

We wished to know how thoroughly our sequencing efforts sampled the true diversity of transcripts present in our cDNA samples. This is a two-part question: first, of the genes truly expressed during *O. fasciatus* oogenesis and embryogenesis, how many did we identify? And second, of these identified genes, how thoroughly had we assembled their full-length transcripts?

To address the first question, we created eight separate assemblies of progressively larger sub-samples of our total reads and tallied the total number of genes identified via BLASTX. The number of newly discovered genes began to plateau after ~1.5M reads (1 7/8 plates in our case) (Figure 5 black line). However, the N50 isotig length continued to increase roughly linearly over this range of reads (Figure 5 grey line). These results suggest that additional sequencing of this sample is unlikely to identify substantially more genes, but may continue to lengthen the existing sequences. Although in the absence of a sequenced genome it is not possible to accurately estimate how many genes are in fact present in the O. fasciatus transcriptome, we note that while several developmental genes of interest were identified in this study, others were not. (Tables 3, 4 and see below). Because these data suggest that we have sequenced these specific cDNA samples quite deeply, some form of specific target enrichment may be necessary for future attempts to discover additional genes not identified in this dataset.

To address the second question, we employed a method proposed by O'Neil and colleagues [20] for addressing the question of how closely our sequences approached full-length transcripts. Their metric, the "ortholog hit ratio," compares the length of the newly discovered sequence that obtains a BLAST hit versus the full length of its top hit [20]. Thus, an ortholog hit ratio of one implies that a transcript has been assembled to its true full length, while values over one suggest insertions in the query sequence relative to its top BLAST hit. We note the caveat that many genes contain relatively poorly conserved regions that may fail to obtain a BLAST hit at all, causing the ortholog hit ratio to be an underestimate in these cases (Additional file 5). In our dataset, many of the O. fasciatus isotigs appear to be nearly fully assembled, while the singletons predictably tend to represent small portions of their top



**Figure 4 GO term distribution of BLAST hits from the** *O. fasciatus* **transcriptome compared with those from the** *D. melanogaster* **genome**. Several GO categories are shown within the top-level divisions of Biological Process, Molecular Function, and Cellular Component. Column heights reflect the percentage of annotated sequences in each assembly that mapped to a given Biological Process GO term. The relative percentages of genes falling into GO categories are comparable between our *O. fasciatus* transcriptome (black) and the *D. melanogaster* transcriptome (white).

BLAST hit in RefSeq (Figure 6). In total, of the 7,219 isotigs with BLAST hits, 3,953 (54.8%) had ratios > 0.5 and 2,689 (37.2%) had ratios > 0.8.

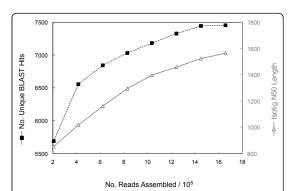
We also asked, for those *O. fasciatus* sequences of developmental genes already present in GenBank that overlapped with transcriptome hits (n = 23), whether our transcriptome data provided any net gain in transcript sequence compared to the GenBank accession sequence. In 15/23 cases (68%), the transcriptome data extended the known sequence beyond that reported in GenBank by an average of 349 bp (range: 82-1,366 bp). In most cases, additional 3' sequence was obtained (Figure 7).

#### Assessing the value of cDNA normalization

Reducing the representation of highly abundant transcripts (i.e. normalizing the cDNA) is often considered

essential to capture sequence from genes expressed at lower levels, including many important developmental genes [see for example [55-57]]. However, we hypothesized that current next-generation sequencing technologies could provide sufficiently deep sequence to render normalization largely unnecessary for construction of *de novo* transcriptomes for comparative developmental biologists. To address this question, we assessed the relative contribution of the N and NN cDNA to our final assembly using several strategies.

First, to test whether our normalization protocol successfully reduced the presence of highly abundant transcripts, we created separate assemblies from the N and NN cDNA samples (equalizing the total number of bases to reduce the contribution of additional sequence found in the NN sample). The N assembly contained a greater number of isotigs that were shorter on average



**Figure 5** Assessing coverage of the *O. fasciatus* transcriptome. Randomly chosen subsets of increasing numbers of Titanium reads were used to generate progressively larger sub-assemblies. The number of reads in each sub-assembly (X axis) is plotted against the number of unique BLAST hits in each sub-assembly (left Y axis: black), and against the N50 isotig length (right Y axis: grey). For this analysis BLAST was performed against the SwissProt database. The number of unique BLAST hits plateaus when the assembly is composed of approximately 1.5 million reads. However, the N50 isotig length maintains an approximately constant rate of increase.

than those in the NN assembly (Figure 2B). Additionally, more singletons were generated in the N assembly relative to the NN assembly (Table 2). Further, similar to the results obtained by Bellin and colleagues [27], we observed the predicted decrease in the maximum number of reads per contig in the N assembly compared to the NN assembly (Figure 8A, B), demonstrating that the normalization procedure successfully reduced the sequencing of highly abundant transcripts. These statistics, which could be interpreted to suggest that the N reads generated an inferior assembly, may result from the shorter average length of reads in the N sample (Figure 2A). Indeed, Newbler rejected 7.9% (30,780) of the N reads as too short, compared to only 1% (3,935) of the NN reads. However, these assembly statistics could also indicate greater heterogeneity in the N sample, which would suggest that normalization might increase the number of new genes identified.

To discriminate between these possibilities, we explored the contribution of the N and NN reads to the genes discovered in our full assembly. We used BLASTN to map one plate's worth of raw reads from the N sample and from the NN sample (equalized to contain the same number of base pairs) against the complete assembled transcriptome, with an e-value cutoff of 1e-4. We then explored the GO annotation of those genes hit exclusively by only one of these two samples. We observed similar overall GO term distributions between the N and NN samples (Figure 8C). We found that a small number of GO terms (n = 20) were

significantly differentially represented in the two samples, albeit generally with very few sequences in each GO term (Additional file 6). For example, we were surprised to see that three of the four terms statistically over-represented in the N sample were related to ribosome function (14/750 (1.9%) of the N hits were annotated with 'ribosomal subunit', compared to 1/1124 (0.09%) NN hits; FDR-corrected p-value = 0.006). In contrast, several terms related to active transmembrane transport were over-represented in the NN sample (Additional file 6) possibly indicating that normalization may have reduced the representation of genes involved in certain basic metabolic processes.

As an additional way to investigate the contribution of the N and NN samples to identifying specific genes of interest for our studies, we manually examined the results of mapping the N and NN samples to the fully assembled transcriptome. Of the 79 genes of interest that we investigated, four (5.1%) were uniquely present in the N sample, whereas nine (11.4%) were uniquely present in the NN sample, and the remaining 66 (83.6%) were present in reads of both the N and NN samples (Tables 3, 4). Although this may be an artifact of sequencing depth (i.e. low-abundance genes of interest may be present in only one of the two cDNA samples simply due to sampling effects rather than the normalization protocol *per se*), our data suggest that the normalized cDNA sample did not contribute disproportionately to gene discovery.

#### Gene discovery for developmental studies

The ultimate goal of this sequencing project was to identify a wide diversity of candidate genes involved in developmental processes. Traditionally, such gene discovery in "non-model" organisms has required degenerate PCR, which is labor-intensive, expensive, and prone to failure. The annotated transcriptome assembly we present here allows researchers to identify genes of interest via simple text searches, or via BLAST searches. To demonstrate the usefulness of these data for largescale gene discovery, we report here the identification of several components from each of the seven widely studied metazoan signaling pathways (Table 3) as well as many genes involved in specific developmental processes (Table 4). We note that the majority of these gene fragments are of suitable length for immediate application of such widely used techniques as in situ hybridization and RNAi-based functional knockdown. In cases of functional experiments where full-length proteins are desirable, such as protein overexpression, RACE PCR will likely be required. Importantly, we note that many genes of interest were present among the singletons, many of which are long enough for immediate use as sequences for in situ hybridization probes or RNAi

Table 3 Selected signaling pathway genes identified in the O. fasciatus transcriptome

			•	Pro	esent in:
Pathway	# Hits	Hit ID (I/C/S)	Length (range)	Normalized	Non-Normalized
HEDGEHOG					
cubitus interruptus	3	I,S	225-906	Υ	Υ
fused	2	1	516-1582	Υ	Υ
patched	2	C, S	225-418	N	Υ
smoothened	2	I	1270-1604	Υ	Υ
JAK/STAT					
domeless	1	1	4028	Υ	Υ
hopscotch (janus kinase)	3	I, C	473-2644	Υ	Υ
Signal transducer and activator of transcription	4	I	444-3270	Υ	Υ
NFKB/TOLL					
cactus	7	I, C	629-1748	Υ	Υ
dorsal (Nuclear factor NF-kappa-B)	2	1	1308-3926	Υ	Υ
relish	1	1	2650	Υ	Υ
Toll	11	I, C, S	215-4323	Υ	Υ
NOTCH					
fringe	1	1	877	Υ	Υ
Hairless	1	1	1053	Υ	Υ
hairy (Enhancer of split/HES-1)	1	1	2530	Υ	Υ
mind bomb	7 (6 <sup>†</sup> )	I,C,S	335-1185	Υ	Υ
Notch	1	S	235	Y*	N
Notchless	1	1	2035	Υ	Υ
Presenilin	1	1	1661	Υ	Υ
Serrate/Jagged	2	S	246-300	Y*	Υ
strawberry notch	7	I,S	191-3519	Υ	Υ
Suppressor of Hairless	3	I,C	375-697	Υ	Υ
WNT					
armadillo	5	I,S	348-3001	Υ	Υ
dishevelled	2	I	954-1321	Υ	Υ
frizzled	3	C,S	194-500	N	Υ
Wnt family (wingless, WNTs)	6	C,S	207-508	Υ	Υ
TGF-BETA					
decapentaplegic (BMP2/4)	1	C	547	Υ	Υ
glass bottom boat (BMP5/7)	2	1	510-737	Υ	Υ
SMADs (Mad, Smad2/3, Smad4/Medea)	7	I,C	276-2276	Υ	Υ
Type I Receptor (saxophone/thickveins/activin receptor type I)	5	I,C	236-2466	Υ	Υ
Type II Receptor (punt, wishful thinking)	3	I	259-5038	Υ	Υ
RECEPTOR TYROSINE KINASES					
Epidermal growth factor receptor	7 (5 <sup>†</sup> )	I,C,S	229-715	N	Υ
rhomboid	2	С	229-602	N	Υ
HORMONE SIGNALING (ECDYSONE, NUCLEAR HORMONE)					
disembodied (ecdysteroidogenic P450)	1	I	1835	Υ	Υ
Ecdysone receptor	2	I,C	231-1393	Υ	Υ

Table 3 Selected signaling pathway genes identified in the O. fasciatus transcriptome (Continued)

Ecdysone-induced protein 63E	1	I	1479	Υ	Υ
ecdysoneless	1	1	4158	Υ	Υ
Nuclear hormone receptor E78	1	1	3150	Υ	Υ
Nuclear hormone receptor HR3	2	1	529-737	Υ	Υ
phantom (cytochrome P450 306a1)	2	C	344-575	N	Υ
shade (cytochrome 450 314A1)	1	1	2125	Υ	Υ
shadow (cytochrome 450 315A1)	1	1	1650	Υ	Υ
ultraspiracle nuclear receptor	1	C	245	γ*	N
without children	2	1	1155-1357	Υ	Υ

Hit ID indicates if gene hits were found among isotigs (I), Cap3-assembled contigs (C), or unassembled singletons (S). Sequence length (range) indicates the shortest and longest S, C or I hit sequences for each gene. These results were generated by BLASTing the raw reads from the N and NN samples against the full assembly. When multiple sequences were obtained via name search, they were tested to see whether they could be made to form a contig with Sequencher or CLC Combined Workbench (see Methods). Asterisk indicates hits only present in normalized GS-FLX reads. X(Y¹) indicates that the X sequences with hits could be assembled into Y contigs.

templates, emphasizing the importance of including these in NGS gene discovery studies.

Although we identified a diverse array of genes, some well-studied genes known to be expressed during embryogenesis were not easily identified in this study. For example, our BLAST results only contained three genes from the Hox cluster (fushi tarazu, Antennapedia, and Abdominal-B), although orthologs of all the canonical arthropod Hox genes are known to be present in O. fasciatus [58]. However, using the O. fasciatus Hox gene sequence fragments available from NCBI as a BLAST query against our transcriptome did reveal sequences for all Hox genes except Sex combs reduced. It is

possible that these genes are expressed at very low levels during the developmental stages sampled here, suggesting that enrichment techniques may be necessary to more easily identify certain genes of interest. We do note, however, that *fushi tarazu*, the only Hox cluster gene not previously identified in *O. fasciatus*, was identified in both N and NN samples of this transcriptome dataset (Table 4).

## Case study: gene discovery for endocrine regulation of development

In addition to surveying the transcriptome for genes involved in embryonic patterning and other developmental

Table 4 Selected developmental process genes identified in the O. fasciatus transcriptome

Process	# Hits	Hit ID (I/C/S)	Length (range)	Normalized	Non-Normalized
GERM PLASM					
Argonaute 3	2 (1 <sup>†</sup> )	I	2042-2231	Υ	Υ
germ cell-less	2 (1 <sup>†</sup> )	1	630-1817	Υ	Υ
maelstrom	1	1	994	Υ	Υ
nanos	1	1	1961	Υ	Υ
piwi/aubergine	1	1	2888	Υ	Υ
pumilio	2	1	424-2574	Υ	Υ
staufen	3	1	599-2100	Υ	Υ
Tudor	2	1	2719-3299	Υ	Υ
vasa	1	C	330	Υ	Υ
ANTERIOR-POSTERIOR DETERMINATION					
GAP					
hunchback	1	1	1429	Υ	Υ
Kruppel	1	S	250	N	Υ
ocelliless (orthodenticle)	1	S	207	Υ	N
TERMINAL GROUP					
huckebein	1	I	589	Υ	Υ
torso-like	2 (1 <sup>†</sup> )	I,C	430-1868	Υ	Υ
PAIR RULE					

Table 4 Selected developmental process genes identified in the O. fasciatus transcriptome (Continued)

fushi tarazu	1	I	788	Υ	Υ
hairy (Enhancer of split/HES-1)	1	1	2530	Υ	Υ
odd skipped	1	С	346	N	Υ
SEGMENT POLARITY					
armadillo	5	I,S	348-3001	Υ	Υ
cubitus interruptus	3	I,S	225-906	Υ	Υ
engrailed	1	S	227	Y*	N
fused	2	1	516-1582	Υ	Υ
pangolin	2	I,C	492-544	N	Υ
patched	2	C, S	225-418	N	Υ
Nnt family (wingless, Wnts)	6	C,S	207-508	Υ	Υ
OORSO-VENTRAL AXIS					
ractus	7	I, C	629-1748	Υ	Υ
decapentaplegic (BMP2/4)	1	C	547	Υ	Υ
gastrulation-defective	1	1	1773	Υ	Υ
nudel	4	I,S	322-1458	Υ	Υ
pipe	1	C	266	Ν	Υ
hort gastrulation	2	C	254-615	Υ	Υ
nake	1	1	1789	Υ	Υ
pätzle	2	I	993-3170	Υ	Υ
· oll	11	I, C, S	215-4323	Υ	Υ
MOLTING/METAMORPHOSIS					
cuticular proteins (including CP 49Ae and adult cuticle protein)	4	I,C	404-566	Υ	Υ
lisembodied (ecdysteroidogenic P450)	1	1	1835	Υ	Υ
Ecdysone receptor	2	I,C	231-1393	Υ	Υ
75	3	I,S	257-649	Υ	Υ
Ecdysone-induced protein 63E	1	I	1479	Υ	Υ
ecdysoneless	1	I	4158	Υ	Υ
tz transcription factor 1	1	1	807	Υ	Υ
normone receptor 4	2	I	1003-2114	Υ	Υ
uvenile hormone acid methyltransferase	5	I	548-2871	Υ	Υ
uvenile hormone binding protein	1	1	1099	Υ	Υ
uvenile hormone epoxide hydrolase	5	I,S	255-2859	Υ	Υ
uvenile hormone esterase	4	I	850-2382	Υ	Υ
uvenile hormone esterase binding protein	1	1	1057	Υ	Υ
luvenile hormone-inducible protein	7	1	456-2757	Υ	Υ
Methoprene-tolerant	1	1	3415	Υ	Υ
luclear hormone receptor E78	1	1	3150	Υ	Υ
luclear hormone receptor HR3	2	1	529-737	Υ	Υ
hantom (cytochrome P450 306a1)	2	C	344-575	N	Υ
hade (cytochrome 450 314A1)	1	I	2125	Υ	Υ
hadow (cytochrome 450 315A1)	1	1	1650	Υ	Υ
akeout	3	I	591-1011	Υ	Υ
ultraspiracle nuclear receptor	1	C	245	Y*	N
without children	2	I	1155-1357	Υ	Υ

Hit ID indicates if gene hits were found among isotigs (I), CAP3-assembled contigs (C), or unassembled singletons (S). Sequence length (range) indicates the shortest and longest S, C or I hit sequences for each gene. These results were generated by BLASTing the raw reads from the N and NN samples against the full assembly. When multiple sequences were obtained via name search, they were tested to see whether they could be made to form a contig with Sequencher or CLC Combined Workbench (see Methods). Asterisk indicates hits only present in normalized GS-FLX reads. X(Y¹) indicates that the X sequences with hits could be assembled into Y contigs. Boldface indicates genes also present in Table 3.

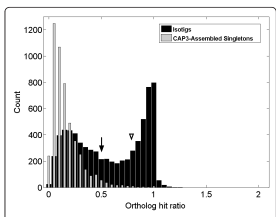


Figure 6 Ortholog hit ratio analysis of isotigs and CAP3-reassembled singletons. An ortholog hit ratio of one implies that a transcript has been assembled to its true full length. For isotigs (black), a majority (54.8%) appear to contain at least 50% of the full length transcript sequence (arrow), while over one-third (37.2%) appear to represent at least 80% of the full length transcript sequence (arrowhead). Most singletons (grey) represent much smaller percentages of full-length transcripts.

processes, we asked whether we could also identify genes known to be employed in biological processes during postembryonic development of holometabolous insects. Recent studies have suggested that many of the genes used during holometabolous insect metamorphosis may also play important roles during embryogenesis in hemimetabolous insects [59,60]. To investigate this, we searched the O. fasciatus transcriptome for expression of key ecdysteroidand juvenile hormone (JH)-related genes. We identified transcripts for many of the known ecdysteroid biosynthesis genes, including cytochrome P450 genes encoded by the Drosophila Halloween family, such as shade (CYP314A1), shadow (CYP315A1), phantom (CYP306A1) and disembodied (CYP302A1) (Table 4). We also detected expression of ecdysone response genes. In particular, we identified many of the ecdysone-regulated genes that play key roles during molting and metamorphosis, including E75, HR3, and HR4 (Table 4). The presence of these genes in the ovaries and early embryos of O. fasciatus corroborates recent studies that implicate ecdysone-response genes in key developmental processes during embryogenesis [59-61]. As might be expected for a situation where ecdysone regulates embryonic development but not molting, transcripts encoding insect peptide hormones implicated in eclosion behavior, such as ecdysis-triggering hormone, eclosion hormone and crustacean cardioactive peptide, were not detected. JH biosynthesis and response genes were also isolated (Table 4). JH has been shown to play a role in promoting embryonic development and tissue maturation [62]. The expression of these genes, together with that of JH esterase and JH binding proteins, is consistent with previous studies implicating tight control of JH during embryogenesis [63].

#### **Conclusions**

We have used 454 pyrosequencing to create an early developmental transcriptome for the milkweed bug O. fasciatus in the absence of a reference genome. Although genomic sequence data will be necessary in the future for linkage or cis-regulatory analyses, at the early stages of establishing new model organisms, one of the most important goals is often gene discovery. In this regard, while no transcriptome generated in this way can realistically be "complete" in the sense of containing full length transcripts for all expressed genes, we propose that for many evolutionary developmental biology studies, the approach described here is a useful one for fast, high-throughput gene discovery. A high priority for comparative developmental biology research is gene expression and function analyses. By sequencing at great depth and testing a variety of cDNA preparation methods (normalized, non-normalized, embryo- and ovaryspecific), we have generated tens of thousands of gene sequences of sufficient lengths for the commonly used developmental techniques of in situ hybridization and RNAi-mediated gene knockdown. These data can also be used for phylogenetic, population genetic, and functional genomic applications, provide a starting point for identification of genomic regulatory sequences, and assist with assembly of hemipteran genomes sequenced in the future.

#### Note added in Proof

While this article was in review, Kumar and Blaxter [64] published a comparison of de novo assemblers for 454 transcriptome data, and reported important shortcomings of Newbler v2.3 compared to other available assemblers. Specifically, the authors reported that Newbler v2.3 produced the smallest assembly (i.e. the smallest number of base pairs incorporated into contigs) of the assemblers tested. The authors argue that this poor performance is likely because Newbler v2.3 inexplicably discards portions of read overlap information. In contrast, a newer, currently unreleased version of Newbler, v2.5, produced the most complete assembly of all those tested. Kumar and Blaxter (2010) therefore strongly advise all de novo 454 transcriptome assembly projects which have used Newbler v2.3 to recompute their assemblies with Newbler v2.5.

To address this concern, we obtained a pre-release version of Newbler v2.5 from Roche and reassembled the *O. fasciatus* data, again using the -nosplit flag. In contrast to Kumar and Blaxter (2010), we observed much less dramatic differences between the assemblies

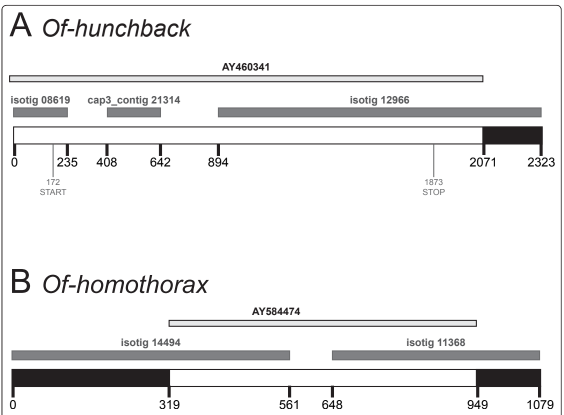


Figure 7 The *O. fasciatus* transcriptome adds sequence data to existing GenBank accessions, which in turn improves annotation of transcriptome sequences. (A) Extended contig for *Of-hunchback* (bottom), comprising the complete mRNA GenBank accession (top, light grey), two isotigs and one CAP3 contig from the transcriptome (middle, dark grey). The largest isotig provides an additional 252 bp of 3' UTR sequence to the GenBank sequence (black). Comparison with the GenBank sequence enabled isotig 08619 and cap3\_contig 21314 to be assembled into the same contig. (B) Extended contig for *Of-homothorax* (bottom), with a partial mRNA GenBank accession (top, light grey) and two transcriptome isotigs (middle, dark grey). Both isotigs extend beyond the known GenBank sequence at the 3' and 5' ends, extending the known region by 449 bp in total (black). Both isotigs had been identified as *homothorax*, and because they did not overlap, they were classified as belonging to the same transcript rather than being paralogs. The GenBank sequence bridges an 87 bp gap between the isotigs, confirming that both sequences are fragments of a single gene.

produced by Newbler v2.3 and Newbler v2.5 (Additional file 7). For example, Kumar and Blaxter (2010) report that Newbler v2.5 increased their total assembly size by 39% compared to Newbler v2.3. For the *O. fasciatus* data analyzed here, Newbler v2.5 increased the total assembly size by less than 1% (Additional file 7). Further, we observed very similar numbers of isogroups, isotigs, and singletons between the two assemblies (Additional file 7). We did observe a 16% increase in the number of contigs reported by Newbler v2.5, but this difference was markedly less than the 80% increase observed in the data analyzed by Kumar and Blaxter (2010). After BLASTing all of the assembled isotigs and cap3-assembled singletons against the RefSeq database, we identified a total of 10,886 unique

BLAST hits, compared to 10,775 genes identified using Newbler v2.3.

These results suggest that, although we did observe a modest increase in assembly size using Newbler v2.5, the analyses presented in the current study are largely robust against differences between currently available versions of Newbler. One possible explanation for the difference between these results and those observed by Kumar and Blaxter (2010), is the greater sequencing depth performed in the current study. If in fact the poor performance of Newbler v2.3 involves discarding information in regions of low coverage, the fact that our dataset includes ~2.4x more reads than that analyzed by Kumar and Blaxter (2010) may explain the reduced improvement that Newbler v2.5 provided our dataset.

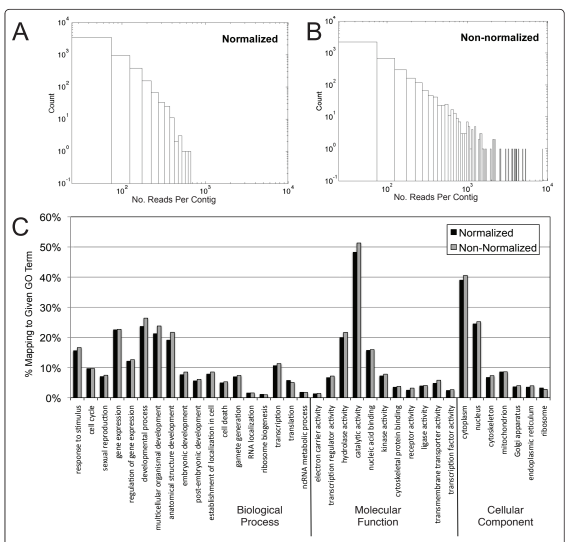


Figure 8 Normalization decreases coverage of highly abundant genes, but does not change the GO term distribution of contigs. In both samples, most contigs are composed of  $<10^2$  reads. However, the non-normalized sample (A) contains contigs with many more reads per contigs than the normalized sample (B). In other words, normalization preferentially decreases the number of reads of those contigs with the most reads. (C) GO term distributions do not differ dramatically between pyrosequenced libraries of N versus NN cDNA. However, see Additional file 6 for exceptions. Column heights reflect the percentage of annotated sequences in each assembly that mapped to a given GO term. Note that the GO terms shown represent the results of mapping the N and NN reads against the complete assembly, rather than those obtained via independent assemblies of N and NN reads.

We also suggest that the reduced number of genes identified via BLAST observed by Kumar and Blaxter (their Table five) may result from the fact that the authors excluded singletons from their analyses. If Newbler v2.3 indeed fails to assemble regions of low coverage and instead retains those reads as singletons, many genes of interest may only be present as singletons. Indeed, we observed many genes of interest exclusively represented

as singletons (Tables 3 and 4). Thus, for the purpose of gene discovery, we emphasize that future *de novo* transcriptome projects should analyze singletons as an important source of useful gene sequence.

Although our results do not appear to be greatly sensitive to which version of Newbler is used, we agree with Kumar and Blaxter (2010) that future transcriptome project should use utilize the most current

available version of Newbler, or whichever assembler algorithm they find most useful for their data.

#### Methods

#### Animal culture

The *O. fasciatus* specimens sequenced in this study were originally purchased from the Carolina Biological Supply Company (Burlington, NC) and were maintained in the laboratory on sunflower seeds under a 12h:12h light/dark cycle at 28°C.

#### cDNA Synthesis

For our pilot study using the GS-FLX platform, total RNA was isolated from mature ovaries (Figure 1B) and from mixed-stage embryos representing the first three days of development (roughly 60% of embryogenesis at  $28^{\circ}\text{C}$ ; Figure 1C, D) using TRIzol (Invitrogen), following the manufacturer's protocols. For each RNA sample, approximately 5  $\mu g$  of cDNA was prepared using the SMART cDNA library construction kit (Clontech, CA, USA). The cDNA was normalized using Evrogen's Trimmer-Direct cDNA Normalization kit (Evrogen, Moscow, Russia), and subsequently digested with SfiI to partially remove the SMART adapters. The size distributions of total RNA and cDNA were assessed on 1.0% agarose gels following each step of the protocol.

To prepare cDNA for sequencing on the GS-FLX Titanium platform, we followed a modified version of the SMART cDNA protocol [65] that has been optimized for cDNA quality and yield from small quantities of total RNA. A helpful guide that formed the initial basis for the optimization of this protocol was once available online from Evrogen, but has since been removed. At the time these libraries were prepared, Roche had not yet provided a specific protocol for cDNA library preparation for 454 pyrosequencing. Subsequently, the company has released a cDNA protocol that requires approximately 500 ng of purified mRNA (typically requiring isolation of 10 to 50 µg of total RNA). While useful for larger tissue samples, the Roche cDNA preparation protocol is difficult to apply to samples in which RNA quantity is limiting, as is the case with many non-model organisms. The protocol we present here does not require the loss-prone step of mRNA purification, and we have found that it produces sufficient quantities of high-quality cDNA when 5 µl of the RNA (18S and 28S bands) can be visualized on a 1% agarose gel stained with ethidium bromide. Compared with the original SMART protocol, we have optimized the primers, PCR conditions, and downstream purification steps to maximize the yield of double-stranded cDNA required for 454 pyrosequencing. We initially optimized this protocol for Roche's original 454 library preparation protocol (not specific to cDNA), which required input of double-stranded DNA amounts of 2.5-10  $\mu g$  (in our experience, typically 10-20  $\mu g$  prepared cDNA as measured by UV absorbance). However, newer protocols from Roche require only 500 ng double-stranded cDNA, limiting the need for a secondary amplification step, as described here, for samples with highly limiting quantities of total RNA.

After separately isolating total RNA from mature ovaries (Figure 1B) and from each of the first three days of embryogenesis (Figure 1C, D) as described above, each RNA sample was treated with DNAse to remove potential genomic contamination. Equal amounts of each sample were then pooled for use as a template for first strand cDNA synthesis. Due to concerns that the poly (T) primer used in the SMART kit could interfere with pyrosequencing, the 3'-primer used was modified in two ways: (1) the poly(T) was interrupted every fourth base by the inclusion of a cytosine [sensu 30]; and (2) the primer contained an MmeI site which allowed most of the poly(T) to be removed during digestion. This 3'-primer (PD243Mme-30TC, 5'-ATT CTA GAG CGC ACC TTG GCC TCC GAC TTT TCT TTT CTT TTT TTT TCT TTT TTT VN-3') was used during first strand synthesis and for all subsequent amplification steps. Because MmeI also cleaves relatively commonly within eukaryotic genes, it may not always be desirable to use this enzyme for library preparation. As an alternative, we have additionally found that a similar 3' primer containing an SfiI cleavage site (PD243-30TC, 5'-ATT CTA GAG GCC ACC TTG GCC GAC ATG TTT TCT TTT CTT TTT TCT TTT TTT TTT VN-3') is also effective in producing cDNA that yields highquality 454 data (data not shown).

For first-strand synthesis, 3 µg of total RNA (in 6 µl) and 2 µl 3' primer (12 µM) were mixed and denatured at 65°C for 5 minutes, then placed on ice. Reverse transcription reactions using SuperScript II (Invitrogen) in the manufacturer's recommended buffer were performed for 50 minutes at 42°C using twice the recommended concentration of enzyme, 1 µl of Protector RNAse inhibitor (Roche) to avoid RNA degradation, 2  $\mu l$  5' primer (12  $\mu$ M), 2  $\mu$ l 10 mM DTT, and 1  $\mu$ l 10 mM dNTPs. Template-switching essential for the SMART technique was achieved using a 5' primer (PD242, 5'-AAG CAG TGG TAT CAA CGC AGA GTG GCC ACG AAG GCC rGrGrG-3') with three RNA nucleotides at its 3' end, which contains an SfiI site. Reactions were then heat-inactivated for 15 minutes at 70°C and diluted 1:5 in milliQ water in preparation for PCR amplification. Contrary to some expectations, SuperScript III reverse transcriptase (Invitrogen) may be substituted in this protocol with equivalent results (data not shown).

To maximize yield during cDNA amplification, the first round of amplification was conducted using a 2:2:1

mix (v:v:v) of Hemo KlenTaq (New England Biolabs), Phusion (New England Biolabs), and PfuTurbo (Stratagene) polymerases. This mixture of enzymes was determined empirically to provide the highest yield of cDNA with a range of input first-strand concentrations. Cesium KlenTaq AC (DNA Polymerase Technologies) and the hot start versions of Phusion and PfuTurbo polymerases in the same ratio may be also substituted at this step without sacrificing yield; this may produce fewer PCR artifacts in the final cDNA preparation. Buffer conditions (MgCl2 and DMSO) were also empirically optimized to maximize yield and minimize PCR artifacts. Reactions were performed in 100 µL total volume in 1X Phusion HF buffer, 1.5 µL polymerase mix, 5 µL first-strand cDNA (previously diluted 1:5 in  $H_2O$ ), 1  $\mu L$ 3' primer (PD243Mme-30TC, 12  $\mu$ M), 1  $\mu$ L 5' primer (PCRIIA, 5'-AAG CAG TGG TAT CAA CGC AGA GT-3', 12 µM), and a final concentration of 1% DMSO, 1.5 mM MgCl<sub>2</sub> (in addition to the MgCl<sub>2</sub> already present in the HF buffer), and 200 µM dNTPs. Reactions were cycled with the following program: 1 minute at 95°C, followed by 16-20 cycles of 30 seconds at 95°C (see below for determining optimal number of cycles), 30 seconds at 66°C, and 3 minutes at 72°C, and a final 10 minutes at 72°C. After cooling to room temperature, 10 μL 3M NaOAc pH 5.5 was added to each 100 μL secondary PCR reaction followed by purification with the QiaQuick PCR purification kit (Qiagen) using the manufacturer's recommended protocol. For all purification steps, samples were eluted with TM buffer (10 mM Tris-HCl pH 8.5, 1 mM MgCl<sub>2</sub>) to prevent strand separation of double-stranded cDNA.

To produce sufficient cDNA for sequencing, Advantage 2 (Clontech) polymerase was used under the manufacturer's recommended conditions during the second round of amplification using the same primer concentrations and 1 µl of undiluted primary PCR product. We recommend testing a range of dilutions of the primary PCR product to obtain the desired quantity of amplified cDNA in 9-10 PCR cycles. In cases of highly limiting RNA concentration, we have also found that a secondary PCR reaction using a 1:1:1 mix of Phusion, Cesium Klen-Taq AC, and Deep Vent (exo-) (New England Biolabs) polymerase in ThermoPol reaction buffer supplemented with 1.5 mM MgSO<sub>4</sub> and 1% DMSO produces the highest yield of secondary PCR product (note that this polymerase mix does not produce optimal results when used for first-round amplification). Secondary PCR reactions were cycled using the same parameters as the primary PCR but running for approximately 10 cycles.

To prevent overcycling during both rounds of PCR amplification, each reaction was prepared in duplicate, and one reaction was spiked with 1  $\mu$ l of 1:750

SybrGreen I (Invitrogen). The spiked reactions were monitored in real time on an Mx3005P QPCR machine (Stratagene Inc.), and the samples were removed when amplification began to plateau. To increase the representation of double-stranded cDNA, two cycles of "chase PCR" were conducted following each round of cDNA amplification after the optimal number of cycles had been reached. Excess primers were added (1.5  $\mu$ L of each, 12  $\mu$ M primer per 100  $\mu$ L reaction), and each reaction was subjected to two additional non-denaturing cycles of 1 minute at 77°C, 1 minute at 65°C, and 3 minutes at 72°C, followed by a 10 minute extension at 72°C.

Following the second round of amplification and PCR purification, the cDNA samples were double-digested with SfiI and MmeI (40 and 26 units per 150 µl reaction, respectively). cDNA species <500 bp were then removed using Chroma Spin 400 columns (Clontech) which had been equilibrated with TM buffer following the manufacturer's protocol. It should be noted that the Chroma Spin column protocol suggested in the Clontech SMART cDNA kit is non-optimal, and that following the protocol provided with the separately purchased columns is less labor-intensive and produces a higher yield of size-selected cDNA. Equilibration of Chroma Spin columns is critical for maximizing the yield of doublestranded cDNA as required by the Roche library preparation protocols. Following size selection, cDNA was blunt-ended with the NEB Quick Blunting kit (New England Biolabs) and purified once more with the Oia-Quick kit. After each step of cDNA synthesis, the size distribution was checked on 1.0% agarose gels, and the cDNA samples were quantified using a Qubit (Invitrogen), after observing that the NanoDrop 1000 (Thermo Scientific) did not reliably quantify ds-cDNA (C. Dunn, personal communication).

To prepare normalized cDNA for GS-FLX Titanium sequencing, 1  $\mu$ l of the twice-amplified, purified cDNA sample described above was subjected to Evrogen's DSN-treatment protocol, followed by a single round of further amplification, SfiI/MmeI digestion, and size selection. Approximately 5  $\mu$ l of normalized and nonnormalized cDNA were synthesized.

#### 454 Titanium Pyrosequencing

For the pilot study using the GS-FLX platform, EnGen-Core (University of South Carolina) conducted the final steps of library preparation, including nebulization, adaptor-ligation, and sequencing of each sample (½ plate each). For sequencing using the Titanium platform, the samples were nebulized, adaptor-ligated, and pyrosequenced by the Institute for Genome Science and Policy DNA Sequencing Facility (Duke University).

#### Sequence Assembly

Raw reads were assembled using the cDNA assembly algorithm of Newbler v2.3 (Roche) with default assembly parameters. An adaptor-trimming step was included in the assembly (the "-v" flag), and the "-nosplit" flag was also used to reduce the generation of extremely short contigs that might otherwise have been created. All of the raw reads generated in this study have been submitted to the NCBI Short Read Archive (Study Accession Number: SRP002610.1).

Because redundancy was observed among the singletons generated by Newbler v2.3, the singletons were reassembled using CAP3 [48], with '-z' option set to 1. Prior to this secondary assembly, the singletons were screened for adaptor sequences using both cross\_match [66-68] and a custom python script (Casey Dunn, personal communication), We note that Newbler can also be used to produce a .fasta and corresponding .qual files of trimmed reads using the '-tr' option. The final assembly thus consists of three types of sequences: Newbler-assembled sequences, cap3\_contigs, and cap3\_singlets, all of which were subjected to subsequent analyses.

#### Sequence Annotation

Sequences were first mapped against the RefSeq Protein database [[69], downloaded from ftp://ftp.ncbi.nih.gov/ blast/db/ on April 27, 2010] using BLASTX. All BLAST searches were conducting using BLAST v2.2.23+ [70] with an e-value cut-off of 1e-10. We then used Blast2GO v1.2.7 [54] to retrieve the Gene Ontology (GO) [71] terms and their parents associated with the top 20 BLAST hits for each sequence. To avoid potentially double-counting sequences that might represent un-assembled portions of the same transcript, a custom python script ("transcriptome\_blast\_summarizer.py", available at http://www.extavourlab.com/protocols/index.html) was used to identify sequences with identical top BLAST hits prior to GO annotation. If multiple sequences hit non-overlapping portions of the same top BLAST hit, we used the conservative assumption that these sequences represented unassembled portions of the same transcript, and therefore only tallied the GO terms of one of these sequences. However, if multiple sequences hit overlapping portions of the same top BLAST hit, we considered these sequences potential paralogs and retained them all. Thus, the counts of sequences in each GO term only include one sequence per top BLAST hit, unless the multiple sequences mapped to overlapping portions of the same BLAST hit. These counts were used to compare the distribution of sequences among specific GO terms between the transcriptomes of O. fasciatus and the Drosophila melanogaster genome. For this comparison, we used a precomputed GO annotation of the D. melanogaster genome [72].

The FASTA formatted transcriptome data set file was examined in TextWrangler (v. 3.1, Bare Bones Software, Inc.). Candidate genes were sought via whole gene names and, where possible, via the gene name abbreviations, while avoiding irrelevant hits. The FASTA header annotation of transcriptome sequences includes the top 20 BLASTx hits to the RefSeq database as described above.

Sequencher (v4.8, Gene Codes Corporation; default settings: minimum 20 bp overlap between sequences, ≥85% sequence identity) and CLC Combined Workbench (v5.6.1, CLC Bio) were used to examine whether transcriptome sequences could be further assembled.

#### Estimating sequencing depth

To estimate how thoroughly our sequencing efforts sampled the O. fasciatus transcriptome, eight progressively larger subsets of the reads were independently assembled. The total number of genes was then identified via BLASTX. For these smaller assemblies, reads from one plate each of normalized and non-normalized reads were combined in random order and sampled without replacement. For each assembly, we BLASTed the longest isotig of each isogroup, and all of the singletons, against the SwissProt database [[73], downloaded from ftp://ftp.ncbi.nih.gov/blast/db/ on April 21, 2010]. We used the relatively small SwissProt database in order to reduce computation time. However, the absolute values of BLAST hits against this database are likely to be underestimates of those values that would have been obtained from a larger database such as RefSeq or nr. If multiple isotigs or contigs hit non-overlapping portions of the same top BLAST hit, only one of these sequences was counted. However, because frequent cases of identical, unassembled singletons were observed, we counted only one singleton per top BLAST hit, regardless of whether these hits overlapped or not.

We used a custom python script to calculate the ortholog hit ratio. This script, "ortholog\_hit\_ratio\_calculator.py" is available at http://www.extavourlab.com/protocols/index.html).

#### Assessing the importance of cDNA normalization

To assess the relative contribution of cDNA normalization to the quality of our assembly, the screened, raw reads from both normalized (N) and non-normalized (NN) samples were mapped against the complete assembly of all reads using the BLASTN algorithm [70] with an e-value cut-off of 1e-4. Based on these results, the Fisher's Exact Test was used to identify over- and under-represented terms in each gene list. This test was performed using Blast2GO (two-tailed, removing double IDs so that only those genes hit uniquely by either N or NN reads were considered). The BLASTN results were also investigated

using text searches to find whether certain genes of interest were present in only one of the two cDNA samples.

#### **Additional material**

Additional file 1: Normalized sample did not perform equally in pilot and full sequencing runs. (A) For the normalized sample, the read lengths of the full plate sequencing runs (white) were shorter than those obtained by the 1/8 plate run (grey). (B) The read length distribution of the non-normalized sample was comparable for both 1/8 plate (grey) and full plate (white) sequencing runs.

Additional file 2: Distribution of average coverage (reads/bp) within contigs in the *O. fasciatus* transcriptome. The coverage within contigs is calculated by dividing the total number of base pairs contained in the reads used to construct a contig by the length of that contig. Note that Newbler v2.3 discards those contigs <100 bp.

Additional file 3: RT-PCR validation of bioinformatically predicted multiple isoforms. (A) Schematic of experimental design. Ten isogroups were randomly selected, each containing exactly two isotigs that differed by the presence/absence of a single contig. PCR primers were designed to flank the differing region. (B) Band sizes predicted by Newbler v2.3 for ten randomly selected isogroups containing exactly two isotigs. (C) Agarose gel following RT-PCR using primers against the sequences described in (B). Ladder sizes are given in base pairs on the left. Blue arrowheads: bands of the sizes predicted by Newbler v2.3; red arrowheads: bands not predicted by Newbler v2.3.

Additional file 4: Identity of taxa with top BLAST hits. "Isotigs" refers only to the longest isotig of each isogroup; "Singletons" refers to the Newbler-generated singletons after secondary CAP3 assembly. The category "other" is the summation of all those species obtaining very low numbers of BLAST hits.

Additional file 5: O. fasciatus assembly isotigs have ortholog hit ratios similar to predictions from fully genome-sequenced databases. When isotigs from the O. fasciatus transcriptome are BLASTed against the RefSeq protein database, ortholog hit ratios show a similar profile to those obtained when the complete Acyrthosiphon pisum gene prediction set (downloaded from http://www.aphidbase.com/aphidbase/downloads/) is BLASTed against the predicted gene set of Drosophila melanogaster (r5.28 downloaded from ftp://ftp.flybase.net/genomes/Drosophila\_melanogaster/) with an e-value cut-off of 1e-10.

Additional file 6: GO terms enriched in Normalized (N) and Non-Normalized (NN) cDNA samples. N (assembly generated from full plate of normalized cDNA) and NN (assembly generated from an equalized number of base pairs of non-normalized cDNA) reads were BLASTed against the full transcriptome assembly, and the results were used to generate "test" and "reference" sets for a Fisher's Exact Test. FDR: false discovery rate.

Additional file 7: Comparison of *de novo* transcriptome assemblies produced by Newbler v2.3 and Newbler v2.5. Number of BLASTx hits reflects a search against RefSeq Protein database with an e-value cut-off value of 1e-10.

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#### Authors' contributions

BEC helped design the research, performed the experiments, collected and analyzed the data, and wrote the manuscript. NS contributed new protocols and helped write the manuscript. KAP helped analyze the data and write the manuscript. YS helped analyze the data and write the manuscript, and obtained funding for the research. SR helped design the research and review the manuscript, and obtained funding for the research. CE proposed the idea for the research, helped design the research and analyze the data, wrote the manuscript and obtained funding for the research. All authors read and approved the final manuscript.

#### Competing interests

The authors declare that they have no competing interests.

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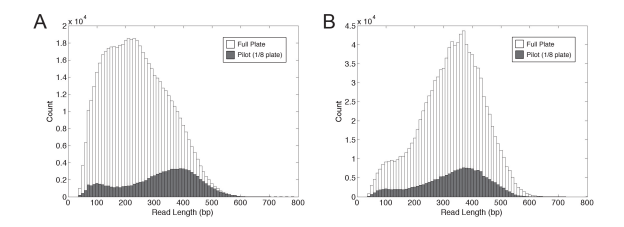
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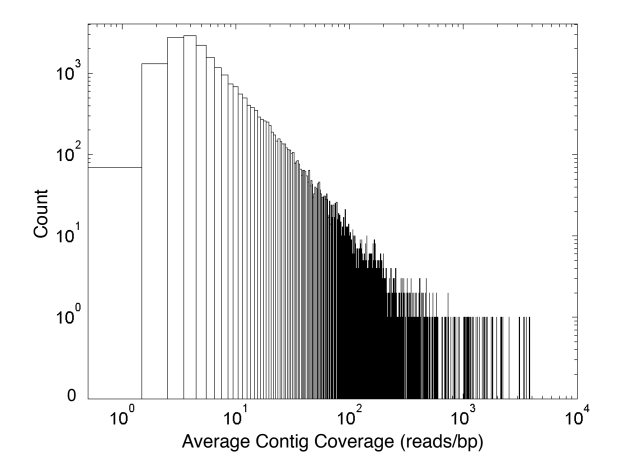
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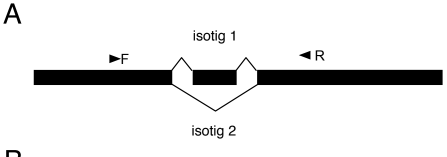




Additional File 1. Normalized sample did not perform equally in pilot and full sequencing runs. (A) For the normalized sample, the read lengths of the full plate sequencing runs (white) were shorter than those obtained by the 1/8 plate run (grey). (B) The read length distribution of the non-normalized sample was comparable for both 1/8 plate (grey) and full plate (white) sequencing runs.

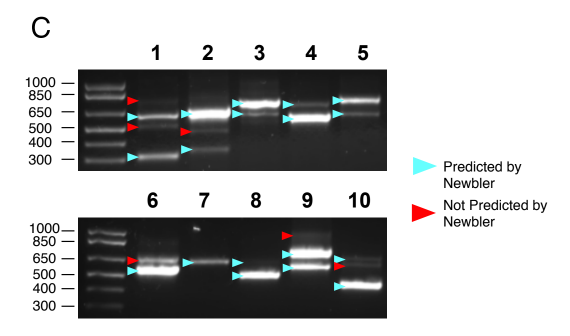


Additional File 2. Distribution of average coverage (reads/bp) within contigs in the *O. fasciatus* transcriptome. The coverage within contigs is calculated by dividing the total number of base pairs contained in the reads used to construct a contig by the length of that contig. Note that Newbler v2.3 discards those contigs <100 bp



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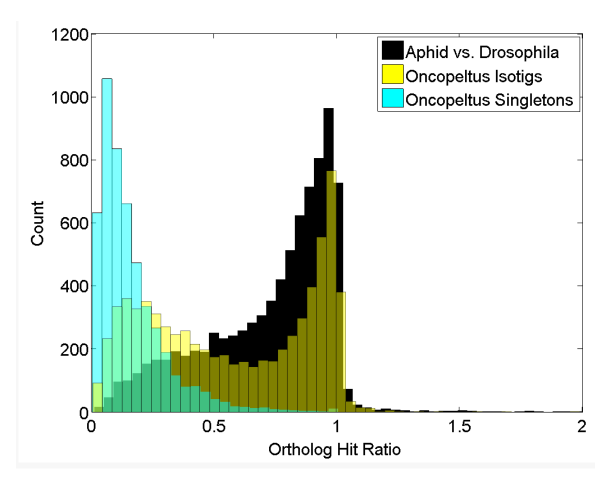
Lane	Isogroup	Predicted Band 1	Band 1 Present?	Predicted Band 2	Band 2 Present?	Additional Bands Present?
1	935	625	Υ	319	Υ	Υ
2	948	657	Υ	347	Υ	Υ
3	984	756	Υ	628	Υ	N
4	1045	759	Υ	609	Υ	N
5	1082	789	Υ	623	Υ	N
6	1133	701	Ν	579	Υ	Υ
7	1134	652	Υ	519	N	N
8	1144	659	Υ	515	Υ	N
9	1162	759	Υ	585	Υ	Υ
10	1179	700	Υ	437	Υ	Υ



Additional File 3. RT-PCR validation of bioinformatically predicted multiple isoforms. (A) Schematic of experimental design. Ten isogroups were randomly selected, each containing exactly two isotigs that differed by the presence/absence of a single contig. PCR primers were designed to flank the differing region. (B) Band sizes predicted by Newbler v2.3 for ten randomly selected isogroups containing exactly two isotigs. (C) Agarose gel following RT-PCR using primers against the sequences described in (B). Ladder sizes are given in base pairs on the left. Blue arrowheads: bands of the sizes predicted by Newbler v2.3; red arrowheads: bands not predicted by Newbler v2.3.

Top BLAST hit taxa	Isotigs	Singletons (CAP3-assembled)	Total
Holometabola	4,311	2,650	6,961
Hemimetabola	2,156	1,640	3,796
Deuterostomes	358	234	592
Non-hexapod arthropods	62	58	120
Non-bilaterian metazoa	47	48	95
Non-metazoa	21	0	21
Others	264	331	595

Additional File 4. Identity of taxa with top BLAST hits. "Isotigs" refers only to the longest isotig of each isogroup; "Singletons" refers to the Newbler-generated singletons after secondary CAP3 assembly. The category "other" is the summation of all those species obtaining very low numbers of BLAST hits.



Additional File 5. *O. fasciatus* assembly isotigs have ortholog hit ratios similar to predictions from fully genome-sequenced databases. When isotigs from the *O. fasciatus* transcriptome are BLASTed against the RefSeq protein database, ortholog hit ratios show a similar profile to those obtained when the complete *Acyrthosiphon pisum* gene prediction set (downloaded from <a href="http://www.aphidbase.com/aphidbase/downloads/">http://www.aphidbase.com/aphidbase/downloads/</a> <a href="webcite">webcite</a>) is BLASTed against the predicted gene set of *Drosophila melanogaster* (r5.28 downloaded from <a href="ftp://ftp.flybase.net/genomes/Drosophila melanogaster/">ftp://ftp.flybase.net/genomes/Drosophila melanogaster/</a> <a href="webcite">webcite</a>) with an e-value cut-off of 1e-10.

GO Term	FDR	# Normalized (total n = 750)	# Non-Normalized (total n = 1124)	Enriched
establishment of localization (GO:0051234)	0.0253	107	227	NN
transport (GO:0006810)	0.0283	106	224	NN
transporter activity (GO:0005215)	0.0475	50	121	NN
ATPase activity (GO:0016887)	0.0174	35	99	NN
establishment of localization in cell (GO:0051649)	0.0411	32	88	NN
vesicle-mediated transport (GO: 0016192)	0.0111	16	61	NN
active transmembrane transporter activity (GO:0022804)	0.0001	11	62	NN
ATPase activity, coupled to movement of substances (GO:0043492)	0.0001	7	51	NN
hydrolase activity, acting on acid anhydrides, catalyzing transmembrane movement of substances (GO:0016820)	0.0001	6	52	NN
primary active transmembrane transporter activity (GO:0015399)	0.0001	6	49	NN
P-P-bond-hydrolysis-driven transmembrane transporter activity (GO:0015405)	0.0001	6	49	NN
ATPase activity, coupled to transmembrane movement of substances (GO:0042626)	0.0001	6	49	NN
GTPase activator activity (GO:0005096)	0.0480	6	31	NN
ATPase activity, coupled to transmembrane movement of ions (GO:0042625)	0.0475	2	20	NN
ATPase activity, coupled to transmembrane movement of ions, phosphorylative mechanism (GO:0015662)	0.0325	1	18	NN
lipid transporter activity (GO:0005319)	0.0475	0	13	NN
cytosolic part (GO:0044445)	0.0031	21	4	N
structural constituent of ribosome (GO:0003735)	0.0282	17	4	N
ribosomal subunit (GO:0033279)	0.0059	14	1	N
large ribosomal subunit (GO:0015934)	0.0381	9	0	N

Additional File 6. GO terms enriched in Normalized (N) and Non-Normalized (NN) cDNA samples. N (assembly generated from full plate of normalized cDNA) and NN (assembly generated from an equalized number of base pairs of non-normalized cDNA) reads were BLASTed against the full transcriptome assembly, and the results were used to generate "test" and "reference" sets for a Fisher's Exact Test. FDR: false discovery rate.

	Newbler v.3	Newbler 2.5
Total bases assembled	19,921,298	20,096,403
Isogroups ("genes")	16,629	16,849
Isotigs ("transcripts")	21,097	20,985
Isotig N50	1,735	1,651
Mean # isotigs per isogroup	1.3	1.2
Contigs ("exons")	22,235	25,955
Mean # contigs per isotig	1.9	1.8
Singletons (singletons after secondary cap3 assembly)	178,770 (112,531)	168,807 (114,487)
Total # unique genes identified BLASTx	10,775	10,886

Additional File 7. Comparison of *de novo* transcriptome assemblies produced by Newbler v2.3 and Newbler v2.5. Number of BLASTx hits reflects a search against RefSeq Protein database with an e-value cut-off value of 1

## **CHAPTER 2**

## Report

# oskar Predates the Evolution of Germ Plasm in Insects

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#### Summary

oskar is the only gene in the animal kingdom necessary and sufficient for specifying functional germ cells [1, 2]. However, oskar has only been identified in holometabolous ("higher") insects that specify their germline using specialized cytoplasm called germ plasm [3]. Here we show that oskar evolved before the divergence of higher insects and provide evidence that its germline role is a recent evolutionary innovation. We identify an oskar ortholog in a basally branching insect, the cricket Gryllus bimaculatus. In contrast to Drosophila oskar, Gb-oskar is not required for germ cell formation or axial patterning. Instead, Gb-oskar is expressed in neuroblasts of the brain and CNS and is required for neural development. Taken together with reports of a neural role for Drosophila oskar [4], our data demonstrate that oskar arose nearly 50 million years earlier in insect evolution than previously thought, where it may have played an ancestral neural role, and was co-opted to its well-known essential germline role in holometabolous insects.

#### **Results and Discussion**

Animal germ cells can be specified either through the cytoplasmic inheritance of maternally deposited germ plasm or through inductive cell signaling [5]. In Drosophila melanogaster, germ cells form by incorporating germ plasm deposited and localized at the oocyte posterior. Germ plasm assembly requires oskar [6], which is necessary and sufficient for germ cell specification [1, 2]. Oskar is localized at the oocyte posterior, where it promotes the accumulation of conserved germline factors including Vasa, PIWI, and Tudor proteins [7] and induces posterior patterning by recruiting nanos mRNA [8]. Despite its essential role in Drosophila germ cell formation and axial patterning, oskar has only been identified in the genomes of a small number of holometabolous insects, all of which specify their germ cells via germ plasm [3]. In contrast, oskar is absent from the genomes of holometabolous insects that lack germ plasm, and neither oskar nor germ plasm have been identified in any insect taxa that branch basally to Holometabola [3]. The prevailing hypothesis is therefore that oskar arose as a novel gene at the base of Holometabola coincidentally with the evolution of insect germ plasm [3].

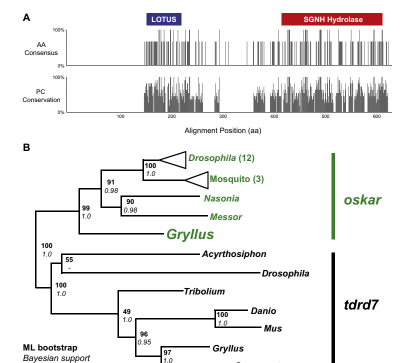
Here we report the first discovery of an oskar ortholog in a basally branching insect that lacks germ plasm, the cricket

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Gryllus bimaculatus (Orthoptera). We unexpectedly detected Gb-oskar in a combined Gryllus ovarian and embryonic de novo transcriptome. Three lines of evidence confirm that Gb-oskar is a bona fide oskar ortholog. First, Gb-Oskar is the reciprocal best protein BLAST hit against the protein products of all known oskar genes from flies, mosquitoes, ants, and the wasp Nasonia, Second, Gb-oskar encodes the LOTUS (aka OST-HTH) and SGNH hydrolase domains that characterize all oskar orthologs [3, 9, 10] (Figures 1A and see Figure S1 available online). Physicochemical conservation is pronounced within both the LOTUS and SGNH hydrolase domains of Gb-Oskar (37.1% and 36.3%, respectively), while amino acid identity is less strongly conserved (11.4% and 21.2%, respectively), consistent with previous observations of oskar orthologs [3, 11]. Third, phylogenetic reconstruction clearly places Gb-oskar with other known oskar genes and not within the tdrd7 genes (Figure 1B), a conserved metazoan gene family that also contains the LOTUS domain [9, 10]. Thus, in contrast to previous hypotheses that oskar first arose in the lineage leading to the Holometabola [3], our analyses demonstrate that oskar was present at least ~50 million years earlier than previously thought [3] in the common ancestor of Orthoptera and Holometabola [12].

The presence of oskar in the genome of an insect branching basal to the Holometabola is surprising because neither germ plasm nor pole cells have been described in any of these taxa. Instead, germ cells are thought to arise from the mesoderm during midembryogenesis in orthopterans [13] and most other early diverging insects [14]. We therefore asked whether Gb-oskar plays a conserved role in germ cell formation in Gryllus or whether this function arose later during insect evolution. We examined the expression of Gb-oskar mRNA and protein (Figure S2) during oogenesis and embryogenesis. In stark contrast to oskar expression in Drosophila [15] and Nasonia [3], Gb-oskar mRNA and protein do not accumulate at the posterior of Gryllus oocytes and instead are distributed ubiquitously throughout all stages of oogenesis (Figures 2A-2D). In just-laid eggs, Gb-oskar mRNA does not localize posteriorly and is barely detectable by in situ hybridization (Figure 2E), although RT-PCR confirms that Gb-oskar is expressed throughout all stages of embryogenesis (Figure 2J). Gryllus primordial germ cells, marked by Gb-piwi and Gbvasa transcript (Figures 2H and 2I) and protein (Figures 2H' and 21'; Figures S2G and S2H) expression, arise during abdominal segmentation stages, but expression of both Gb-oskar mRNA and Gb-Oskar protein remains at low levels in all embryonic cells throughout these stages and does not become enriched in primordial germ cells before or during their formation (Figures 2G and 2G'). The expression pattern of Gb-oskar therefore does not support a role for this gene in germ cell formation.

To directly test whether *Gb-oskar* is functionally required for germ cell formation in *Gryllus*, we knocked down *Gb-oskar* function during oogenesis and embryogenesis using maternal RNAi (mRNAi) and embryonic RNAi (eRNAi), respectively [16] (Figures 2, S3A, and S3B). In contrast to *Drosophila* and *Nasonia*, maternal knockdown of *Gb-oskar* did not reduce egg laying (Figure S3C), affect ovary morphology, or impede



the progress of oogenesis (Figure S3D). Embryonic *Gb-oskar* knockdown did not cause any of the posterior patterning defects seen in *Drosophila osk* mutants [6] (Figure S3E, Table S1), and these embryos were morphologically wild-type and hatched at normal rates (Figure S3F). These data show that in contrast to the known requirement for *oskar* in anterior-posterior (A-P) axial patterning in holometabolous insects [3, 6], *oskar* does not direct *Gryllus* axial patterning. Further, *Gb-oskar* eRNAi embryos produced germ cells that expressed Gb-Piwi and Gb-Vasa (Figures 2K–2N) and ultimately pro-

duced functional ovaries in adulthood (Figure 2P).

In contrast to the essential and conserved role that oskar plays in Holometabolous germ cell formation [3, 8], our analyses of Gb-oskar gene expression and function show that this gene is not required for germ cell formation in Gryllus. We therefore hypothesized that Gb-oskar has a distinct somatic function in Gryllus that may reflect an ancestral function for this gene. Consistent with this hypothesis, we observed that Gb-oskar has a specific somatic expression pattern during midembryogenesis: Gb-oskar mRNA and protein are enriched in neuroblasts along the A-P axis (Figures 3A-3C"). Neuroblasts are neural stem cells that arise from the ventral ectoderm and produce neurons of the CNS in Drosophila and other Pancrustacea (insects and crustaceans) [17]. Gb-oskar expression in the neuroblasts begins during the earliest stages of neurogenesis and persists throughout all stages examined. In addition to this embryonic expression pattern, Gb-oskar is also expressed in the adult brain

We examined embryonic nervous system development in Gb-oskar knockdown conditions and found that Gb-oskar eRNAi embryos exhibit morphological defects of the axonal

Figure 1. Domain Structure and Phylogenetic Analysis of *Gb-oskar* 

(A) ClustalW protein alignment of Oskar orthologs from Gryllus bimaculatus, Nasonia vitripennis, and Drosophila melanogaster (see Figure S1 for alignment showing amino acids). Sequence identity at the amino acid level is not strongly conserved (top graph), but physicochemical conservation is more pronounced (bottom graph), specifically within the conserved LOTUS (blue) and SGNH hydrolase (red) domains in all three proteins, meaning that the chemical properties of these domains are conserved at a local level despite amino acid divergence and suggesting that these regions may represent functional domains of Oskar. (B) Phylogenetic reconstruction of Gb-oskar with known oskar (green) and tdrd7 (black) orthologs. The best-scoring unrooted ML topology is shown: bold, ML bootstrap values; italics, Bayesian posterior probability.

tracts that are consistent with an impairment of neuroblast divisions: lateral connectives are often broken or reduced in width compared to controls (Figures 3D–3D", p < 0.001), anterior commissure of absent (Figures 3E–3E", p < 0.01), and posterior commissure formation is similarly disrupted (Figures 3F–3F", p < 0.025). These axonal defects suggest that *Gb-oskar* may be required for

proper neuronal determination and are reminiscent of the axonal scaffold defects of *Drosophila miranda* and *prospero* mutants, which disrupt neuroblast divisions resulting in neuronal misspecification [18].

Aberrant neuroblast divisions can be assayed by examining the expression of even-skipped, which is expressed in a stereotyped subset of ganglion mother cells and neurons within each segment, including the aCC and pCC neurons that are homologous across insects [19]. We found that 34.5% of Gb-oskar eRNAi embryos displayed significant defects in aCC/pCC specification (Figures 3G–3G", p < 0.01) that were never observed in controls, indicating that Gb-oskar is required for proper neuroblast division.

Our results demonstrate a role for Gb-oskar in the development of the cricket CNS, in contrast to its well-known role in germ plasm formation in holometabolous insects. These divergent functions of oskar suggest at least two possible evolutionary scenarios. First, oskar's ancestral role in insects could be that of germ plasm assembly as seen in Drosophila, and the CNS function we report here could be a derived trait in the branch of the insect tree leading to Gryllus. However, several lines of evidence support a second scenario, whereby oskar's neural function is ancestral to Orthoptera (crickets, locusts, and grasshoppers) and Holometabola, which diverged approximately 380 million years ago (Mya) [12], thus implying that its role in assembling germ plasm is a derived trait in higher insects (Figure 4). In support of this interpretation, we note that insect germ plasm itself is thought to be a derived trait within insects and unique to Holometabola [5], consistent with our observation that Gryllus possesses an oskar ortholog yet lacks germ plasm. Moreover, in Drosophila adults, oskar is expressed in the brain and is required for place learning, as are

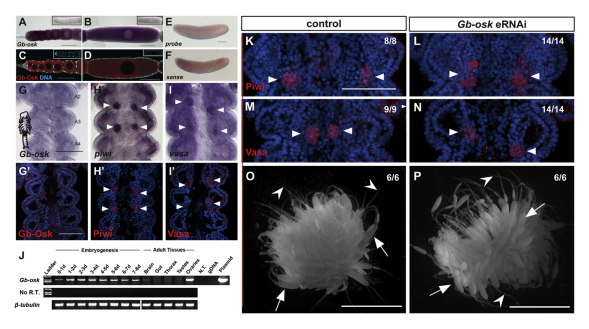


Figure 2. Gb-oskar mRNA and Protein Do Not Accumulate in Germ Plasm or Embryonic Germ Cells and Are Not Required for Germ Cell Formation or Development

(A–D) *Gb-oskar* mRNA (A and B) and protein (C and D) are expressed ubiquitously in oocytes and do not accumulate asymmetrically at the posterior. Insets show sense probe (A and B) and preimmune serum (C and D) controls. (E) *Gb-oskar* transcript levels are low and ubiquitous in blastoderm stage embryos. (F) Gb-oskar sense control. (G–I') In fully segmented embryos, *Gb-oskar* mRNA (G) and protein (G') are not enriched in embryonic germ cells, which express *piwi* and vasa mRNA (H and I) and protein (H' and I'). Arrowheads in (H)–(I') indicate germ cell clusters. A2, A3, and A4 indicate abdominal segments 2, 3, and 4. Black region of embryo schematic in (G) shows the region displayed in (G)–(I'). (J) RT-PCR analysis of *Gb-oskar* throughout embryogenesis and in different adult tissues. β-tubulin was used to ensure equal quantities of RNA template in the cDNA synthesis reaction and as a gel loading control. The amplified *Gb-oskar* band is 2,149 bp and was amplified with 35× PCR cycles. The highest levels of *Gb-oskar* are detected throughout embryogenesis and in adult ovaries. Lower levels are detected in the adult brain, and faint expression is detected in the adult thoracic muscles and testes. No expression is detected in the adult gut. N.T., no template control; gDNA, genomic DNA control; Plasmid, *Gb-oskar* amplified from the full-length plasmid clone; No R.T., no reverse transcriptase control. (K–N) Vasa- and Piwi-positive germ cells form in *Gb-oskar* eRNAi embryos (L and N) as in *DsRed* eRNAi controls (K and M). (P and O) *Gb-oskar* eRNAi embryos raised to adulthood form functional ovaries (P), which contain functional germaria (arrowheads) and late-stage oocytes (arrows) as in uninjected controls (O). Numbers indicate sample sizes. Left shows anterior in (A)–(E) and top shows anterior in (G)–(I') and (K)–(N). Scale bar represents in (A)–(B) and 5 mm in (N) and 5 mm in (N) and (O). Validation of all Gb-specific antibodies is shown in Figure S2; validation of *Gb-oskar* eRNAi is shown in Figure S3 and T

several other *Drosophila* genes with germline functions [4, 20], suggesting that these genes may have an ancient association with the nervous system.

Multiple Drosophila genes originally characterized for their role in germ cell development, including nanos, pumilio, and Staufen, have subsequently been shown to function in the nervous system, where they regulate translation in such processes as dendrite morphogenesis, synaptic growth, asymmetric neuroblast divisions, and neuronal specification [4, 21-23]. The co-occurrence of multiple germ plasm genes in the CNS of D. melanogaster raises the intriguing possibility that these genes may function within an evolutionarily conserved functional module [24], which could facilitate their co-option to a novel context such as holometabolous germ plasm. Consistent with this hypothesis, we find that Gb-vasa mRNA and protein are coexpressed with Gb-oskar in neuroblasts (Figures S2I-S2K), suggesting that a functional link between oskar and other germline genes predates the evolution of germ plasm. Moreover, expression of germline genes in the nervous system has also been observed in other insects belonging to both Hemimetabola (piwi in aphids [25]) and Holometabola (vasa in ants [26]).

If oskar were to acquire expression in germ cells due to its functional linkage with other germ plasm genes, an evolutionary change in its transcriptional, translational, or functional regulation might then have feasibly allowed its co-option to a critical function in the germline specification pathway. Consistent with this possibility, we note the presence of extremely low levels of Gb-oskar in Gryllus germ cells (Figures 2G and 2G'), although it appears to play no essential germ cell function (Figures 2K-2P). Co-option of oskar to assemble germ plasm probably involved molecular evolution of its regulation and function; accordingly, we find that Gb-oskar is not regulated by Drosophila oskar translational machinery in Drosophila oocytes or embryos (Figure S4, Table S2), suggesting that oskar's translational regulation mechanisms have evolved extensively in the lineage leading to Drosophila. However, it is also possible that specific features of the Gb-oskar coding sequence, or its incompatibility with Drosophila UTRs, may have prevented the translation of Gb-Oskar in our transgenic

We have shown that oskar was present nearly 50 million years earlier in insect evolution than previously thought [3, 12] and must therefore have been lost several times in

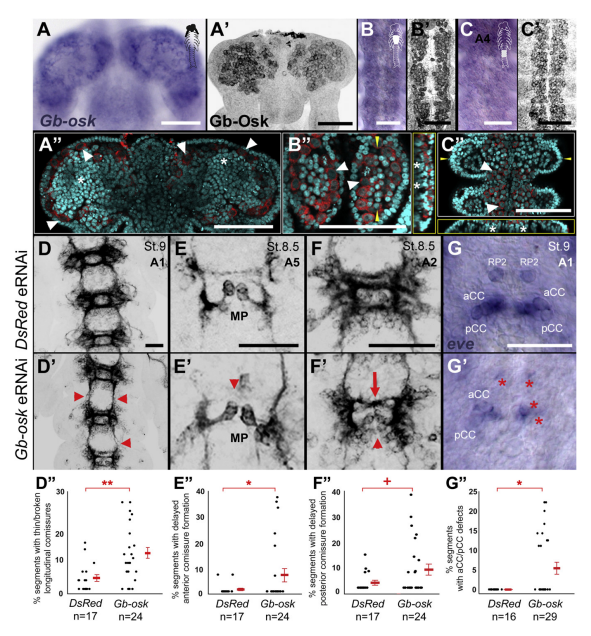


Figure 3. Gb-oskar Is Expressed in Neuroblasts and Is Required for Neural Development

(A- C") Gb-oskar mRNA (A, B, and C) and protein (A', B', and C') accumulate in neuroblasts of the brain (A and A'), thorax (T1-T3; B and B'), and abdomen (A4-A6; C and C'). (A"), (B"), and (C") show single optical sections of Gb-Osk expression (red, Gb-Osk; cyan, nuclei), revealing the highest levels of Gb-Osk expression in neuroblasts (cells with large nuclei and diffuse chromatin; arrowheads) and absent or lower levels in neuroblast daughter cells (cells with smaller, denser nuclei; asterisks). Bright staining near midline of head in (A') and (A") is nonspecific staining of the extraembryonic membranes. Large panels of (B") and (C") show single optical sections through T3 and A5 neuromeres, respectively; anterior is on top in both central panels. Yellow framed boxes to the left and bottom of (B") and (C"), respectively, show orthogonal sections at the plane indicated by yellow arrowheads in the large panels; in these orthogonal sections, ventral is on the right in (B") and on top in (C"). White arrowheads indicate Gb-Osk-expressing neuroblasts immediately dorsal to the ectoderm and ventral to the underlying daughter neurons and neuronal precursors (asterisks), which show little to no expression of Gb-Osk. (D-F') Ventral views of embryonic abdominal segments of DsRed eRNAi controls (D, E, and F) and Gb-oskar eRNAi in embryos of the same developmental stage (D', E', and F') labeled with axonal marker anti-HRP. The percent of Gb-oskar eRNAi embryos that exhibit thin or broken longitudinal connectives (arrowheads) is 83.3% (D and D'). Gb-oskar eRNAi embryos show delayed formation of both anterior (E and E') and posterior (F and F') commissures (arrowheads) relative to the development of the midline precursors (MP). Abnormal or fused anterior commissures also appear in a greater proportion of Gb-oskar eRNAi embryos than in controls (arrow in F'). (G and G') Formation of aCC and pCC neurons is impaired in Gb-oskar eRNAi embryos (asterisks) (G') but never

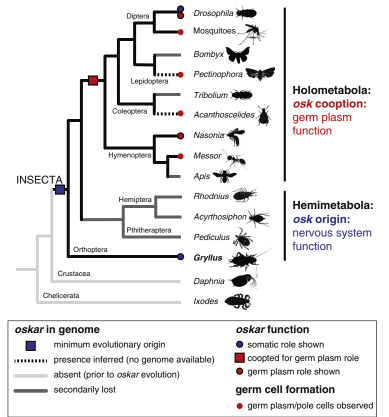


Figure 4. Phylogenetic Hypothesis of oskar Origin and Function across Arthropods

Species shown have sequenced genomes, allowing unambiguous determination of the presence or absence of an oskar ortholog; dotted lines indicate exceptions to this rule and show taxa in which pole cells and/or germ plasm have been observed but that lack available genome sequence. oskar is absent from the sequenced genomes of noninsect arthropods (light gray lines). Boxes indicate proposed origin of oskar in the last common ancestor of Holometabola and Hemimetabola (blue) and putative co-option of oskar to a germline role in the Holometabola (red). Circles indicate that oskar plays known roles in the nervous system (blue) or presumably in the germline (red); circles outlined in black indicate that there is functional evidence for the described role. Evidence for evolution of oskar translational regulation in the lineage leading to D. melanogaster is shown in Figure S4 and Table S2.

some insect lineages (Figure 4). Indeed, completely sequenced genomes of holometabolous insects lacking germ plasm or pole cells confirm that *oskar* has been lost in these lineages [3]. Germ cell specification via germ plasm is thought to have arisen independently in multiple bilaterian taxa [5], but how germ plasm evolved has remained unclear. Our results suggest a novel molecular mechanism for this process in insects: co-option of the *oskar* gene into the top of the germ plasm assembly hierarchy.

#### **Experimental Procedures**

#### Animal Husbandry, Gene Expression, and Functional Analysis

G. bimaculatus husbandry, gene expression analysis, RNAi experiments, egg-laying analysis, and axonal scaffold visualization were carried out as previously described [16].

#### Gene Cloning and Phylogenetic Analysis

Full-length *Gb-oskar* was recovered from a *G. bimaculatus* transcriptome and its identity confirmed by both Bayesian and Maximum Likelihood analysis. Details of sequence analysis are available in Supplemental Experimental Procedures.

#### **Antibody Generation**

Rabbit polyclonal antibodies were raised against an N-terminal and a C-terminal peptide from Gb-Oskar (Figures S1 and S2A) (Open Biosystems), recombinant proteins of full-length Gb-Vasa, and a 774 amino acid fragment of Gb-Piwi (McGill Biology CIAN facility). Details of antibody construction and validation are described in Supplemental Experimental Procedures.

#### **Accession Numbers**

The Genbank accession numbers for the *Gb-oskar*, *Gb-piwi-like*, and *Gb-tdrd7* sequences reported in this paper are JQ434102, JQ434103, and JQ434104, respectively.

#### Supplemental Information

Supplemental Information includes four figures, two tables, and Supplemental Experimental Procedures and can be found with this article online at http://dx.doi.org/10.1016/i.cub.2012.10.019.

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in controls (G). aCC/pCC are located in the corners where the longitudinal connectives meet the posterior commissure; these axonal scaffolds are visible in (G). The out-of-focus darkened spots adjacent to the in-focus aCC/pCC neurons are U/CQ neurons present ventral to the dorsally located aCC/pCC neurons. (D", E", F", and G") Quantification of neural defects illustrated in (D), (E), (F), and (G); thick red bars at the bottom of plots show mean values  $\pm$  SE. Statistical significance of differences between treatments (red brackets) based on chi-square tests: \*\*p < 0.001, \*p < 0.01, \*p < 0.025. Anterior is shown on the top in all panels. Scale bar represents 100  $\mu$ M in (A)-(C") and 50  $\mu$ M in (D)-(G"). Embryonic stage and/or the most anterior segment shown are indicated in top right corner in (C), (D), (E), (F), and (G); stages and segments shown in (D), (E), (F), and (G) apply to (D'), (E'), (F'), and (G'), respectively. Validation of *Gb-oskar* eRNAi is shown in Figures S2 and S3.

this work. We thank Elke Küster-Schöck of the McGill Biology Cell Imaging and Analysis Network (Canada) for assistance with antibody creation, Andrés Leschziner for western blot assistance, Anne Ephrussi, Satoru Kobayashi, Sam Kunes, and Akira Nakamura for reagents, and members of the Extavour laboratory for discussion.

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Current Biology, Volume 22

## **Supplemental Information**

## oskar Predates the Evolution

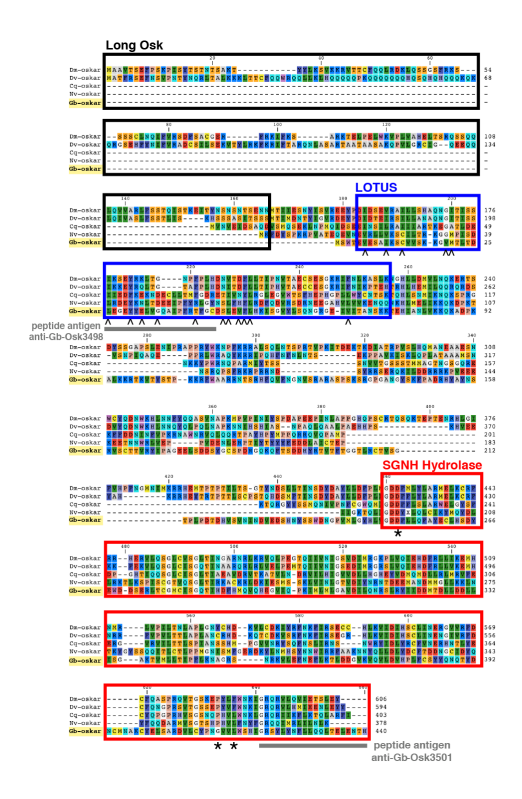
## of Germ Plasm in Insects

Ben Ewen-Campen, John R. Srouji, Evelyn E. Schwager, and Cassandra G. Extavour

### **Supplemental Inventory**

- Supplemental Data
  - o Figures S1-S4
  - o Tables S1 and S2
- Supplemental Experimental Procedures
- Supplemental References

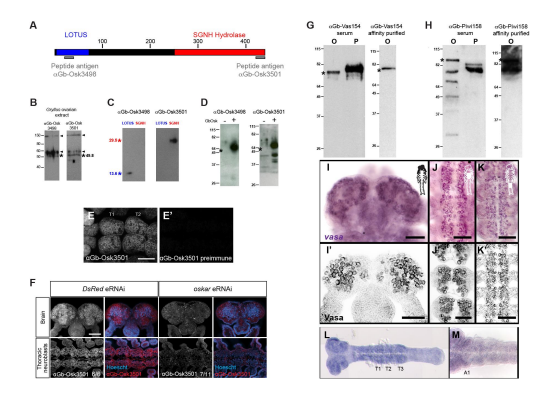
Ewen-Campen et al. Supplemental Information Page 1 of 18



Ewen-Campen et al. Supplemental Information Page 2 of 18

Figure S1, related to Figure 1. ClustalW alignment of Gb-Oskar with known oskar orthologs. Gb-Oskar contains the LOTUS (aka OST-HTH) domain (blue box) and the SGNH hydrolase domain (red box) that are characteristic of Oskar orthologs. The Long Osk domain (black box) is only present in Oskar orthologs from Drosophilid species. Residues are colored using the RasMol color scheme to indicate physicochemical properties. Structurally conserved residues within the LOTUS domain are indicated with carets [1, 2]. Indicated with astrices are the positions of the serine, aspartate, and histidine residues (notably absent in Oskar) that constitute the catalytic triad in functional hydrolases [1]. The two peptide antigens used to generate Gb-Oskar antibodies are indicated in grey. Species (accession IDs): Dm = Drosophila melanogaster (AAF54306.1), Dv = Drosophila virilis (AAA28426.1), Cq = Culex quinquefasciatus (ACB20969.1); Nv = Nasonia vitripennis (ADK94458.1), Gb = Gryllus bimaculatus (JQ434102).

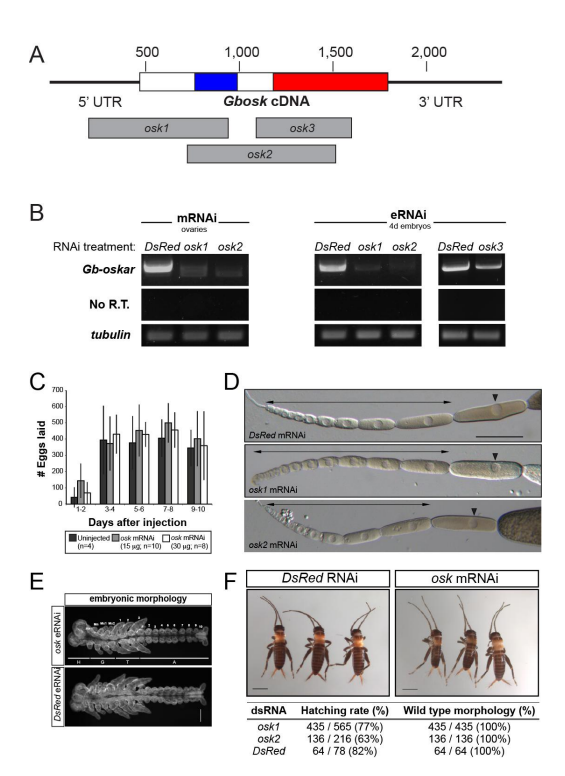
Ewen-Campen et al. Supplemental Information Page 3 of 18



Ewen-Campen et al. Supplemental Information Page 4 of 18

Figure S2, related to Figures 2, 3. Validation of rabbit  $\alpha$ Gb-Oskar,  $\alpha$ Gb-Vasa and  $\alpha$ Gb-Piwi antibodies. (A) Schematic of Gb-Oskar, showing the location of the LOTUS (blue) and SGNH hydrolase (red) domains, and the N- and C-terminal peptides used to generate αGb-Osk3498 and αGb-Osk3501 respectively (grey). (B)Western blot of Gryllus ovarian extract probed with \alpha Gb-Oskar antibodies. Bands are detected at the predicted weight of Gb-Oskar (black asterisk: 49.8 kDa). (C) Western blot of purified recombinant LOTUS and SGNH domains from Gb-Oskar probed with αGb-Oskar antibodies. αGb-Osk3498 specifically recognized the LOTUS domain (blue askterisk indicates expected size of 13.6 kDa) but not the SGNH hydrolase domain (red asterisk indicates expected size of 29.6 kDa). Conversely, \alpha Gb-Osk3501 specifically recognized the SGNH hydrolase domain but not the LOTUS domain. (D) Western blot of whole cell lysate of E. coli expressing full length Gb-Oskar. The predominant band is slightly larger than expected (black asterisk 49.8 Kda), as in (A). (-) uninduced E. coli, (+) induced E. coli.(E-E") The expression of Gb-Oskar in neuroblasts (here shown in thoracic segments T1 and T2) is only seen using the final bleed serum (E), and is not present using the preimmune serum (E'). (F) Gb-Oskar immunostaining in brain (top row) and thoracic neuroblasts (bottom row) is greatly reduced following Gb-oskar eRNAi, validating the specificity of the antiserum and the efficacy of the RNAi knockdown. Sample sizes are given at the bottom of each panel. Anterior is to the left in (E-E") and in (F) bottom row; anterior is up in (F) top row. Scale bars = 100 µM. (G) Western blots of unpurified αGbVas154 serum on Gryllus ovarian extracts (O) and purified protein (P). Serum used at 1:5,000 recognizes a band of the expected size (asterisk: 75kDa) and a smaller secondary band from ovarian extracts. Following affinity purification against recombinant Gb-Vasa protein used at 1:500, the antibody recognizes a single band of the expected size from ovarian extracts (O). (H) Western blots of unpurified αGbPiwi158 serum on Gryllus ovarian extracts (O) and purified protein (P). Serum used at 1:2,000 recognizes a band of the expected size (asterisk: 92kDa) and several smaller secondary bands from ovarian extracts. Following affinity purification agasint Protein-A, serum used at 1:500 shows reocgnition of most smaller secondary bands is abolished and the antibody strongly recognizes a band of the expected size as well as a slightly smaller secondary band from ovarian extracts (O). In situ hybridization for Gb-vasa (I-K) and antibody staining with  $\alpha$ GbVas154 (I'-K') shows expression of Gb-vasa transcript and protein in neuroblasts of the embryonic brain (I, I'), thorax (J, J') and abdomen (K, K') in Stage 9 embryos. Gb-oskar expression increases in the ectodermal neuroblast precursor territory just prior to neuroblast formation at Stage 5 (L) and in newly forming neuroblasts at Stage 6 (M). Scale bar =  $100 \mu M$ .

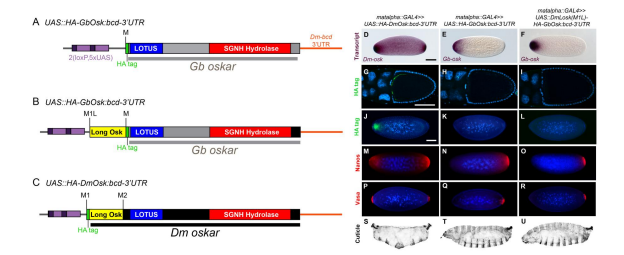
Ewen-Campen et al. Supplemental Information Page 5 of 18



Ewen-Campen et al. Supplemental Information Page 6 of 18

Figure S3, related to Figures 2, 3. Validation and phenotypes of Gb-oskar RNAi. (A) Schematic of the three dsRNA fragments used for Gb-oskar RNAi experiments. Blue and red regions indicate locations of the LOTUS and SGNH hydrolase domains respectively. Grey boxes indicate fragments used for RNAi treatments. (B) Semi-quantitative PCR analysis of Gb-oskar levels in negative control (DsRed), osk1 and osk2 RNAi treatments. Maternal RNAi (mRNAi) greatly reduces Gb-oskar transcript levels in ovaries of injected females even 10 days after injection (left panel). Near-complete reduction of Gb-oskar transcript in embryos via embryonic RNAi (eRNAi) with fragments osk1 and osk2, and a lesser reduction with fragment osk3, persists through until at least 4 days after egg laying (right panel). No R.T. = No reverse transcriptase control to ensure the absense of genomic DNA. β-tubulin serves as a loading and RNAi specificity control. (C) Average daily egg-laying rate is not significantly different between Gboskar mRNAi-injected females and controls, at two different concentrations of Gb-oskar dsRNA. Error bars indicate 95% confidence interval. osk1 and osk2 results are pooled for each concentration, as no significant differences were seen between these treatments. (D) Gb-oskar mRNAi does not disrupt oogenesis or ovarian morphology. Double-headed arrows indicate that oocytes at stages of oogenesis are present in RNAi treated ovaries as well as controls. Arrowhead indicates the posterior localization of the oocyte nucleus, which indicates normal oocyte patterning [3]. (E) Nuclear stain of Gb-oskar eRNAi embryos reveals wild type anteriorposterior patterning compared to controls. (F) Gb-oskar mRNAi hatchlings (right panel) do not display obvious morphological defects compared to controls (left panel). Hatching rate is not significantly different between Gb-oskar mRNAi embryos (right panel) and controls (left panel). Scale bars =  $100 \mu M$  in D-E, 1 mm in F.

Ewen-Campen et al. Supplemental Information Page 7 of 18



Ewen-Campen et al. Supplemental Information Page 8 of 18

Figure S4, related to Figure 4. Schematic of constructs used for transgenic *D. melanogaster*, which show that *Gb-oskar* and *DmLosk(M1L)*: *GbOskar* transgenes are transcribed and localized but not translated in *Drosophila* ovaries or embryos. Full length cDNAs of *Gb-oskar* (A), *Gb-oskar* downstream of *Dm-Losk*(M1L) (B) or full length *Dm-osk* (C) were fused to an N-terminal HA tag fused to the *Dm-bicoid* 3'UTR sequence. Constructs were cloned into the pValium22 vector for site-directed insertion via the ΦC31 targeted transgenesis system . HA-tagged *Dm-osk* (D, G, J, M, P, S), *Gb-oskar* (E, H, K, N, Q, T), or DmLosk(M1L)-*Gb-osk* (F, I, L, O, R, U) fused to the *bicoid* 3'UTR were expressed in transgenic *D. melanogaster* with a maternal GAL4 driver (matalpha). In situ hybridization for *Dm-osk* (D) or *Gb-oskar* (E, F) shows that all transcripts are localized to the anterior of early embryos. Dm-Oskar but not Gb-Oskar protein is translated in oocytes (G, H, I) and early embryos (J, K, L). Ectopic Dm-Osk causes anterior localization of Nanos (M) and ectopic Vasa-positive pole cells (P) and a bicaudal phenotype revealed in larval cuticle preparations (S). Absence of detectable ectopic Gb-Oskar (H, I, K, L) is correlated with absence of ectopic Nanos (N, O), Vasa or pole cells (Q, R) and larval morphology is wild type (T, U). In all panels anterior is to the left, scale bar = 100 μm.

Ewen-Campen et al. Supplemental Information Page 9 of 18

 $\begin{tabular}{ll} Table S1, related to Figure 2. Anterior-posterior axis phenotypes in $\it Gb-oskar$ RNAi embryos. \end{tabular}$ 

RNAi method	dsRNA injected	# embryos injected <sup>1</sup>	# developed (%)	# surviving embryos with morphological abnormalities (%)
Maternal (mRNAi)	osk1	123	75 (61.0%)	0
	osk2	52	23 (44.2%)	0
	caudal	589	486 (82.5%)	321 (66.0%)
	DsRed	135	81 (60%)	0
				·
Embryonic (eRNAi)	osk1	125	101 (80.8%)	0
	osk2	67	41 (61.2%)	0
	osk3	150	53 (35.3%)	0
	caudal	168	131 (78.0%)	110 (84.0%)
	DsRed	74	65 (87.8%)	0
	DsRed <sup>2</sup>	85	65 (76.5%)	0
	uninjected	90	64 (71.1%)	0

<sup>1.</sup> For maternal RNAi, this number refers to number of eggs dissected.

Ewen-Campen et al. Supplemental Information Page 10 of 18

Table S2, related to Figure 4. Effect of the *Gb-oskar* transgenes in *D. melanogaster* using the bicoid 3' UTR.

Maternal genotype	bicaudal phenotype / total (%)	anterior Vasa-positive cells / total (%)
matalpha >> HA-Dm-osk:bcd3'UTR	63 / 76 (82.9%)	42 / 60 (70%)
matalpha >> HA-Gb-osk:bcd3'UTR	0 / 73 (0%)	0 / 54 (0%)
matalpha >> DmLosk(M1L):HA-Gb- osk:bcd3'UTR	0 / 91 (0%)	0 / 61 (0%)
matalpha >> DmOsk(M1L):bcd3'UTR	20 / 26 (76.9%)	14 / 28 (50%) <sup>1</sup>

<sup>1.</sup> Of the 14 embryos with ectopic germ cells, six (42.8%) did not form at the anterior pole, but instead formed slightly more posteriorly.

Ewen-Campen et al. Supplemental Information Page 11 of 18

## **Supplemental Experimental Procedures**

## Gb-Oskar Identification and Phylogenetic Analysis

Gb-oskar was recovered from a de novo transcriptome generated using 454 Titanium sequencing. A 1,323 bp predicted full length Gb-oskar sequence was assembled from 695 raw reads (from a total of 4.2 million reads in the transcriptome), including additional 5' and 3' UTR sequence (462 bp and 607 bp, respectively). Our transcriptome assembly also contained a predicted alternate isoform of Gb-oskar that includes an additional 162 bp of sequence between the LOTUS and SGNH hydrolase domains. However, RT-PCR failed to amplify this larger product, and the additional sequence was not conserved in any other oskar orthologue. We therefore focused on the smaller of these two predicted isoforms, which we have confirmed is transcribed during Gryllus development by RT-PCR (Fig. 2J). Portions of Gryllus piwi and tdrd7 orthologues were also found in our Gryllus transcriptome, and Gryllus vasa and even-skipped were cloned based on published sequence (AB378065, AB120736). All new sequence data have been submitted to NCBI (GenBank Accessions JQ434102-4).

In the course of performing reciprocal best BLAST hit analysis, we found that *Nasonia vitripennis oskar* (Nv-Osk, ADK94458.1), unlike *oskar* orthologues from *D. melanogaster* (AAF54306.1), *Culex quinquefasciatus* (ACB20969.1), *Aedes aegypti* (ABC41128.1), *Anopheles gambiae* (ABC54566.1), *and Messor pergandei* (ADM07366.1), retrieved a best hit against the *Gryllus* transcriptome sequence for *Gb-tdrd7* (e-value = 5e-11). However, this BLAST hit only extended over the LOTUS domain (amino acids 14-120), which is conserved in both *oskar* and *tdrd7* genes, while the second best hit, *Gb-oskar* (e-value = 2e-10), extended over both the LOTUS and SGNH hydrolase domain (amino acids 19-106 and 183-375 of the query). Further, *Gb-oskar* retrieves Nv-Osk as its top hit against the *N. vitripennis* proteome. We therefore concluded that across the length of the gene *Nv-oskar* is more similar to *Gb-oskar* than to *Gb-tdrd7*, a result that agrees with all other known *oskar* orthologues as well as our phylogenetic analysis (Fig. 1B).

Amino acid identity conservation (i.e. percentage of sequences containing an identical amino acid at a given alignment position) and physicochemical conservation were calculated using JalView [25].

To estimate a gene tree of *oskar* and *tdrd7* genes, *Gb-oskar* and *tdrd7* were aligned to their publically available orthologues using MUSCLE [26] (Fig. S1). Although *oskar* orthologues have been bioinformatically predicted in several recently sequenced Hymenopterans (retrievable at <a href="http://hymenopteragenome.org/ant\_genomes/?q=blast">http://hymenopteragenome.org/ant\_genomes/?q=blast</a>), we only included those from *N. vitripennis* [NP\_001234884.1] and *M. pergandei* [ADM07366.1] in our phylogenetic analysis, as these have been experimentally validated as true *oskar* orthologues [3]. Regions of uncertain alignment were removed using GBlocks [27] with the least stringent settings, which produced an alignment containing 294 amino acids. A gene tree for *oskar* and *tdrd7* was estimated under both Bayesian and maximum likelihood criteria. The "mixed" model of amino acid evolution was used in MrBayes v3.1.2 [28] to choose the optimal model for amino acid evolution for both this and the maximum likelihood tree. MrBayes selected the WAG model [29] with a probability of 1.0. Two runs of four independent MCMC chains each were executed for 2 million generations, sampling trees every 1,000 generations and the first 250,000 generations discarded as burn-in. The average standard deviation of split frequencies between the two runs fell below 0.01 after 154,000 generations, indicating that the two chains had converged. Maximum likelihood analysis

Ewen-Campen et al. Supplemental Information Page 12 of 18

was estimated using RAxML-MPI v7.2.8. 2000 runs from independent starting trees were executed under the WAG model of protein evolution with gamma distribution of rate heterogeneity to simultaneously estimate the best scoring tree and perform rapid bootstrap analysis. All tree estimations were conducted on the Odyssey Cluster (Harvard University), supported by the FAS Sciences Division Research Computing group.

## **Gb-Oskar Antibody Generation**

Two rabbit polyclonal antibodies were raised against an N-terminal peptide DLEGEYYELVGQAIPFRTFGC ( $\alpha$ Gb-Osk3498) and a C-terminal peptide GRSYLYNFLLQQLTELENTH ( $\alpha$ Gb-Osk3501) from Gb-Oskar (Figs. S1, S2A) (Open Biosystems, Inc., Alabama). Rabbits were boosted on days 14, 28, 42 and 82 after initial injection, and the final bleed was performed on day 138. Specificity was tested using Western blot analysis and immunostaining (Fig. S2B-F).

To prepare *Gryllus* ovaries for Western blotting,  $\sim 30\text{-}50$  ovarioles were homogenized in 300  $\mu L$  of 5X SDS loading buffer, boiled for 5 minutes at 95-100°C, and stored at -20°C until use. 1:10 and 1:100 dilutions of this homogenate, run in adjacent wells, were used in Western blots. For purified Gb-Oskar domains, 10  $\mu g$  of purified protein were run. For whole-cell lysates containing full-length bacterially expressed Gb-Oskar, cells were grown (either with or without induction) overnight at 20°C, and OD600 readings were used to calculate equal volumes of induced versus uninduced cells.

For Western blotting, samples were separated on a 12% acrylamide SDS-PAGE gel in running buffer (25 mM Tris base, 192 mM glycine, 0.1% w/v SDS) at ~180V for approximately one hour and then transferred to a nitrocellulose membrane (0.2 μm pore size, BIO-RAD cat. # 162-0112) at ~300V for 90-120 minutes at 4° C in transfer buffer (25 mM Tris base, 192 mM glycine, 0.1% w/v SDS, 20% v/v methanol). Ponceau S staining was used to verify that equal volumes were loaded in each lane and had transferred properly. Blots were blocked in 5% BSA + 5% milk powder for at least 30 minutes, and then incubated overnight with primary antibodies (1:500 dilution). Blots were washed at least 5 x 5 minutes in TBST (20 mM Tris pH 7.5, 500 mM NaCl, 0.05% (v/v) Tween-20) and then incubated with donkey anti-rabbit HRP-coupled secondary antibody (Jackson Laboratories, 1:5,000 dilution) in block for two hours. After 5 x 5 minute washes in TBST, the secondary antibody was detected with SuperSignal West Pico Chemiluminescent substrate (ThermoScientific) following manufacturer's protocols. Signal was visualized using Kodak BioMax film, and exposures were tested at 10 seconds, 1 minutes, 5 minutes, and 10 minutes to optimize exposure.

Both antibodies recognized proteins of the predicted size (asterisk in Fig. S2B) from *Gryllus* ovary extracts, as well as additional species of larger molecular weights (arrowheads in Fig. S2B), suggesting possible post-translational modification as has been reported for *Drosophila* Oskar [30]. Western blot analysis showed that as predicted, the  $\alpha$ Gb-Osk3501 serum recognized the C-terminal SGNH hydrolase domain and not the N-terminal LOTUS domain; conversely, the  $\alpha$ Gb-Osk3498 serum recognized the LOTUS domain and not the SGNH hydrolase domain (Fig. S2C). Both antibodies also recognized bacterially expressed full length Gb-Oskar at a slightly higher MW than predicted (Fig. S2D). The staining pattern seen in whole mount immunostained *Gryllus* embryos was not detected in pre-immune serum (Fig. S2E', E''), and was abolished by *oskar* RNAi (Fig. S2F).

Ewen-Campen et al. Supplemental Information Page 13 of 18

## Gb-Vasa and Gb-Piwi Antibody Generation

Rabbit polyclonal antibodies were raised against recombinant Gryllus Vasa and Piwi protein fragments (McGill Biology CIAN facility, Canada). Full-length Gryllus vasa (amino acids 2-649) and a fragment of Gryllus piwi (774 C-terminal amino acids ending at the stop codon) were each cloned into the pET151/dTOPO expression vector (Invitrogen Cat No. K151-01), thus introducing an N-terminal 6X-polyhistidine tag, Protein expression was induced in E. coli BL21 (DE3) by addition of IPTG to a final concentration of 0.25 mM and incubated at 30°C for 4 hrs. The overexpressed proteins were highly insoluble, so the cell pellets were disrupted by sonication in non-denaturing buffer, and soluble fraction were separated by centrifugation and discarded. The pellet was dissolved in 8 M urea (crude prep). Crude preps were further purified by SDS gel-purification. Protein bands were excised from the gel after reverse staining with zinc chloride and the proteins were collected by electroelution. Acetone precipitation was performed and the precipitated protein was washed and redissolved in 8M urea. Protein concentration was adjusted to 1-2 mg/ml for injection. Rabbits were injected using a standard 80-day protocol with three boosts after the initial injection; final bleed was performed after 87 days. The serum was processed by addition of NaN3 to 0.02% (w/v). Both antibodies were then affinity purified: αGb-Vasa was purified against bacterially expressed Gb-Vasa protein, and αGb-Piwi was purified with Protein A (Primm Biotech, Cambridge MA).

The CIAN facility determined specificity prior to purification using Western blot analysis (A-B, left blots), and we repeated this analysis following affinity purification using the above protocol (Fig. S2G-H, right blots).

## In situ Hybridization and Antibody Staining

In situ hybridizations and antibody stainings were performed as described in [4]. Four different DIG-labeled fragments of *Gb-oskar* were used as in situ probes (ranging from 742 to 2,103 bp in size), all of which gave consistent staining results. *Gb-piwi* and *Gb-vasa* in situ probes were 781 bp and 1,953 bp, respectively. Probes were used at between 0.5-2  $\mu$ g/ $\mu$ l during hybridization. For immunostainings, all species-specific primary antibodies used were used at a final dilution of 1:300.  $\alpha$ Gb-Oskar was preabsorbed against Oregon R *Drosophila* mixed stage embryos for 90 minutes at room temperature prior to staining to reduce background. Secondary antibodies used were goat anti-rabbit or goat anti-mouse coupled to Alexa 555, Alexa 488 or Alexa 647 (Invitrogen) at 1:1,000. Nuclei were counterstained with Hoechst 33342 at 1:5000 of a 10mg/ml stock solution. Cy3 conjugated anti-HRP (gift of Sam Kunes, Harvard University, MA, USA) was used at 1:50.

## Injections for RNA Interference (RNAi)

Maternal and embryonic injections of dsRNA were carried out as described in [4]. To synchronize adult female age for maternal injections, final nymphal stage females were isolated and monitored daily for final molt. All females were injected with 15  $\mu$ g of dsRNA on the third day following the final molt. Because we observed that egg laying could be reduced by the continued presence of two males, each female was housed separately with two males only for days one, two and five of a 10 day experiment.

Three different fragments of *Gb-oskar* were used as template for dsRNA synthesis (Fig. S3A): (1) *osk1*: a 742bp fragment starting 262 bp upstream of the first methionine and ending at position 480; (2) *osk2*: a 789 bp fragment from position 255-1044, and (3) *osk3*: a 503 bp fragment from position 638-1141. Results from these three fragments were consistent for all

Ewen-Campen et al. Supplemental Information Page 14 of 18

phenotypic analyses. DsRNA was adjusted to a final concentration of 3  $\mu$ g/ $\mu$ l (6.1 and 5.8  $\mu$ M respectively) for fragments osk1 and osk2, and 5  $\mu$ g/ $\mu$ l for fragment osk3 (15.1  $\mu$ M). The DsRed negative control and caudal positive control dsRNAs have been described in [4] and were both used at the same final concentration as the corresponding oskar fragments used in that experiment (either 3 or 5  $\mu$ g/ $\mu$ l).

### Validating RNAi Knockdown

RNAi knockdown efficiency was estimated using both semi-quantitative RT-PCR and whole-mount antibody staining. For semi-quantitative RT-PCR (Fig. S3B), ovaries from maternal RNAi-injected females were dissected 10 days after maternal dsRNA injection, and stage 7-10 embryos laid by maternal RNAi-injected females, or resulting from zygotic RNAi injections, were dissected from 4-5 days after egg laying (AEL). All tissues were collected in Trizol (Invitrogen), and total RNA was isolated following manufacturer's protocols, followed by a 30 minute DNase treatment (Ambion) at 37°C to remove genomic DNA. Equal quantities of total RNA were used as template for first strand cDNA synthesis using SuperScript III (Invitrogen). *Gb-oskar* levels were estimated following 35 PCR cycles (98°C for 3 minutes, 35 cycles of 98°C for 30 seconds, 55°C for 30 seconds, 72° C for 2 minutes 15 seconds, followed by a 10 minute final extension at 72°C) and electrophoresis on a 1.0% agarose gel, compared to *Gb-beta-tubulin* to ensure equivalent amounts of template. Negative controls without reverse transcriptase were run in parallel and revealed no genomic DNA contamination. The *Gb-oskar* primers used (F: 5' TGGTAGTTCGAAGGGAACTTG-3'; R: 5'-CATCTTCCATTTGCCACAGA-3') amplify a band of 2,149 bp. The *Gb-beta-tubulin* primers

CATCTTCCATTTGCCACAGA-3') amplify a band of 2,149 bp. The *Gb-beta-tubulin* primers used (F: 5'-TCAGACACCGTCGTTGAACC-3'; R: 5'-GATGGTTCAGGTCGCCGTAG-3') amplify a band of 157 bp.

Knockdown appeared to be less efficient using fragment *osk3* (Fig. S3B), but this fragment still produced CNS phenotypes consistent with those produced using fragment *osk1*.

RNAi efficiency was independently assayed by staining osk eRNAi and DsRed eRNAi embryos with the  $\alpha$ Gb-Oskar3501 antibody and imaging the embryos under identical conditions (Fig. S2F).

### Scoring RNAi Phenotypes

To quantify egg laying, each injected female was individually housed with a dish of moist sand. Because we observed that Gryllus females typically only oviposit substantial numbers of eggs on alternate days, egg dishes were collected every 48 hours. Eggs were separated from sand using a 500  $\mu$ m mesh under running tap water, and spread out in a monolayer in a 15cm petri dish. An image of the eggs was captured on a flatbed scanner and eggs were manually tallied using the ImageJ Cell Counter plug-in (Fig. S3C).

Ovarian morphology was assayed on ovaries of injected females 10 days after dsRNA injection. Wild type A-P patterning was assessed by the asymmetric localization of the oocyte nucleus in mid-stage oocytes [3]. We also tested for oogenesis phenotypes by looking for the presence of oocytes at all stages of development (Fig. S3D).

Embryonic morphology was visualized at stages 7-10 using both Nomarski optics and fluorescent nuclear staining (Fig. S3E). The presence of embryonic germ cells was assayed using in situ hybridization against *piwi* (not shown), as well as antibody staining for both Gb-Vasa and Gb-Piwi (Fig. 2G-H). The presence of functional ovarioles was also assayed in *osk* eRNAi embryos that were grown to adulthood.

Ewen-Campen et al. Supplemental Information Page 15 of 18

Axonal scaffolds were visualized with anti-HRP immunostaining followed by confocal microscopy of the entire embryonic nervous system. Before scoring axonal defects, stagematching of embryos was performed by comparison of morphogenetic progress in the thoracic appendages, antennae, abdominal segmentation and terminal cerci, rather than by developmental progression of the nervous system. Only embryos at stages 8.5-10 were used for this analysis, since formation of the global CNS axonal scaffold is complete by this time. Axonal patterning of Gb-oskar eRNAi embryos, DsRed eRNAi embryos, and uninjected embryos at comparable developmental stages were scored blind (embryos were grouped by matched developmental stages, but whether they were wild type, DsRed eRNAi or Gb-oskar eRNAi embryos was hidden) for consistent and symmetrical thickness and continuity of longitudinal connectives, timing/morphology of anterior commissure formation relative to central neuron mitoses, and timing/morphology of posterior commissure formation relative to anterior commissure formation. Two independent blind scoring rounds were performed and the results were consistent. After scoring was completed, the nature of the embryonic treatment was revealed, and data from embryos of the same treatment (uninjected, DsRed eRNAi or Gb-oskar eRNAi) were grouped for statistical analysis (Chi-squared test).

To test for defects in aCC/pCC neurons, we performed in situ hybridization on stage 9 embryos using a 528 bp probe against *Gryllus even-skipped*. Samples were scored blind for the absence of any of the four aCC/pCC neurons in each segment. Two independent knockdowns using two dsRNA fragments (fragments *osk1* and *osk3* described above) gave comparable results (5/11 and 5/18 embryos contained defective aCC/pCC neurons in the two experiments respectively). To reliably distinguish between aCC/pCC neurons and the nearby U/CQ neurons, we used several approaches. First, we noted that the U/CQ neurons are clearly distinguishable from aCC/pCC by their position in the dorsal-ventral axis and their relationship to the axonal scaffold. Specifically, aCC/pCC are the dorsal-most *eve-*positive cells while U/CQ are more ventral [see e.g. 3; their Figure 4a-h]. Second, aCC/pCC are reliably located in the corner where the longitudinal connectives meet the posterior commissure [see e.g. 4; their Figure 2d], and in the same focal plane. Third, in our analysis we examined optical sections through the entire dorso-ventral axis of every affected segment to ensure that apparently missing aCC/pCC cells were not visible in any other focal plane, and that U/CQ could be identified in more ventral focal planes.

## Transgenic Constructs

To target HA-tagged Dm-Oskar or Gb-Oskar to the *D. melanogaster* oocyte anterior, we cloned Gb-Oskar, Dm-Losk(M1L):Gb-Oskar, or Dm-Osk-bcd3'UTR [5] into pValium22 [6], which contains attB sites for PhiC31-based targeted integration and is optimized for expression in the female germ line. We used a previously described bcd3'UTR that has a putative NRE replaced by a 10bp linker to prevent potential transcript clearing by *nanos* [5, 7]. Cloning was performed using the circular polymerase extension cloning method [8] with primers containing an N-terminal HA-tag (pValium22 was amplified with primers F: 5'-AGAACTAGAGCCGCG-3', R: 5'GCCCGAGCTTAAGACT-3'; Dm-Osk-bcd3'UTR was amplified with primers F: 5'-GCCAGTCTTAAGCTCGGGCATG

TACCCATACGATGTTCCGGATTACGCTGCCGCAGTCACAAGT-3' and R: 5'-CAATTCCGCGGCTCTAGTTCTGGATCCACCCGAGTA-3'; Gb-Oskar was amplified with primers F: 5'-GCCAGTCTTAAGCTCGGGCATG

TACCCATACGATGTTCCGGATTACGCTAGTTGGACTGAGGTT-3' and R: 5'-

Ewen-Campen et al. Supplemental Information Page 16 of 18

CCTCTCATCCAGGCTCGAGCGCCGGCGTCAATGTGTGTTTTC-3'; to amplify the bcd3'UTR for fusion with Gb-Oskar, primers F: 5'-CTCGAGCCTGGATGA-3' and R: 5'-CAATTCCGCGGCTCTAGTTCTGGATCCACCCGAGTA-3' were used). The transgenes for GbOsk:bcd3'UTR and DmOsk:bcd3'UTR were inserted into the attp40 site on 2L, and the DmLosk(M1L):GbOsk:bcd3'UTR transgene was inserted into the attp2 site on 3L [9] both of which show high levels of specific expression. Injection to create transgenic flies was carried out by Genetic Services, Inc. (Cambridge, MA). Transgenic progeny were identified and maintained using standard genetic crosses.

## Drosophila Stocks

The maternal driver line  $w^*$ ;  $P\{w^{+mC}=matalpha4-GAL-VP16\}V2H$  was obtained from the Bloomington *Drosophila* Stock Center (stock number 7062). y v; Sco/CyO and y Sc v; Dr e/TM3,Sb were obtained from the Perrimon lab (Harvard University).

### Imaging and Image Analysis

Images were captured with AxioVision v.4.8 driving a Zeiss Stereo Lumar equipped with an AxioCam MRc camera, or a Zeiss Axio Imager equipped with an AxioCam MRm camera, using epifluorescence either with or without an Apotome. Confocal microscopy was performed with a Zeiss LSM 710 confocal, using comparable gain, offset, and averaging parameters for all samples. Image analyses were performed with AxioVision v.4.8, Zen 2009 (Zeiss), and figures were assembled in Photoshop CS4 or Illustrator CS4 (Adobe). For the confocal images shown in Fig. 2 (G-N), a maximum-intensity projection of the antibody staining was superimposed over a single z-plane of the nuclear counterstain for visual clarity.

Author Contributions: BE-C and CGE designed research, analysed data and wrote the paper; experiments were carried out by JRS (recombinant Gb-Oskar purification), EES (cloning *Gb-vasa* and *Gb-piwi*), CGE (axonal scaffold phenotypic analysis) and BE-C (all other experiments); CGE obtained funding for the research.

Ewen-Campen et al. Supplemental Information Page 17 of 18

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Ewen-Campen et al. Supplemental Information Page 18 of 18

## CHAPTER 3

## **Article**

# Germ Cell Specification Requires Zygotic Mechanisms Rather Than Germ Plasm in a Basally Branching Insect

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### Summary

Background: Primordial germ cell (PGC) specification is a universal process across animals, but the molecular mechanisms specifying PGCs are remarkably diverse. In *Drosophila*, PGCs are specified by maternally provided, asymmetrically localized cytoplasmic factors (germ plasm). In contrast, historical literature on most other arthropods reports that PGCs arise from mesoderm during midembryogenesis, suggesting that an arthropod last common ancestor may have specified PGCs via zygotic mechanisms. However, there has been no direct experimental evidence to date for germ plasm-independent arthropod PGC specification.

Results: Here we show that in a basally branching insect, the cricket *Gryllus bimaculatus*, conserved germ plasm molecules are ubiquitously, rather than asymmetrically, localized during oogenesis and early embryogenesis. Molecular and cytological analyses suggest that *Gryllus* PGCs arise from abdominal mesoderm during segmentation, and *twist* RNAi embryos that lack mesoderm fail to form PGCs. Using RNA interference we show that *vasa* and *piwi* are not required maternally or zygotically for PGC formation but rather are required for primary spermatogonial divisions in adult males.

Conclusions: These observations suggest that *Gryllus* lacks a maternally inherited germ plasm, in contrast with many holometabolous insects, including *Drosophila*. The mesodermal origin of *Gryllus* PGCs and absence of instructive roles for vasa and piwi in PGC formation are reminiscent of mouse PGC specification and suggest that zygotic cell signaling may direct PGC specification in *Gryllus* and other Hemimetabola.

### Introduction

Of the many specialized cell types that comprise an animal's body, only one is capable of contributing genetic information to the next generation: the germ cells. The restriction of reproductive potential to a small subset of cells is a universal process across sexually reproducing animals and represents a profound evolutionary novelty likely required for the evolution of multicellularity [1]. The molecular mechanisms that specify these cells, however, are remarkably diverse between taxa [2–5] and only well understood in a handful of model organisms.

Primordial germ cell (PGC) specification mechanisms have been categorized into two modes: cytoplasmic inheritance and zygotic induction [3, 4, 6]. Cytoplasmic inheritance (e.g.,

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in *Drosophila melanogaster*) involves the localization of maternal mRNAs and proteins (germ plasm) to a subcellular region of the oocyte. Germ plasm is necessary and sufficient to induce PGC fate. In zygotic induction (e.g., in *Mus musculus*), by contrast, there is no germ plasm, and PGCs instead form in response to inductive signals from neighboring somatic cells [7].

Within insects, cytoplasmic inheritance appears to be a derived character confined primarily to the holometabolous insects [8] (Figure 1A; see also Table S1 available online), where germ plasm has been demonstrated experimentally in many species (Table S1). Histological studies of insects branching basally to Holometabola (the Hemimetabola), in contrast, have reported the absence of both germ plasm and pole cells in nearly all of these taxa [3, 6] (Figure 1A; Table S1). Studies of molecular markers for PGCs in hemimetabolous insects have been limited to the highly atypical parthenogenetic embryos of the pea aphid *Acyrthosiphon pisum*, a milkweed bug, and several orthopteran species (Table S1), yet there is no conserved pattern of PGC origin across these taxa.

In this study, we use multiple conserved molecular markers and RNAi to characterize PGC formation in the cricket *Gryllus bimaculatus* (Orthoptera), a hemimetabolous model species for studying the development of basally branching insects [9]. We provide several lines of evidence that *Gryllus* PGCs form from the abdominal mesoderm via inductive signaling and discuss the implications of these results for the evolution of germ plasm and the possibility of an ancient relationship between bilaterian PGCs and mesoderm.

### Results

## Gryllus Germ Cells Express a Suite of Conserved Genes

Within the Orthoptera, neither germ plasm nor pole cells have been reported (Figure 1A; Table S1). Histological examinations of orthopteran embryos conducted by William Wheeler over a century ago [10] suggested that PGCs arise from or among abdominal mesoderm cells during abdominal segmentation (Figure 1C), consistent with reports of germline origin both in other Hemimetabola and in most arthropods [3, 6]. However, conserved molecular markers can reveal a cryptic germ plasm that eludes histological examinations [11-13]. We therefore examined the expression of several conserved molecular PGC markers (vasa [14], piwi [14], tudor, boule, and germ cell-less) and three additional PIWI family genes (Figures S1A and S1B) in Gryllus ovaries and embryos. Because some germ plasm components localize as proteins rather than transcripts (see for example [15, 16, 17]), we also examined the expression of Vasa and Piwi proteins [14].

In fully segmented (stage 9) *Gryllus* embryos (Figures 1B and 1C), we identified cells matching Wheeler's description [10] that express both mRNA and protein of *piwi* and *vasa*, as well as *bol* and *gcl* transcripts (Figures 1D, 1E, and S1C). These cell clusters were found in abdominal segments A2–A3 in all embryos, and in A4–A5 in 45% of embryos (Figures 1D and 1E, arrowheads). Clusters were located on the dorsal medial face of mesodermal structures termed "coelomic pouches," which are present in every gnathal, thoracic, and abdominal

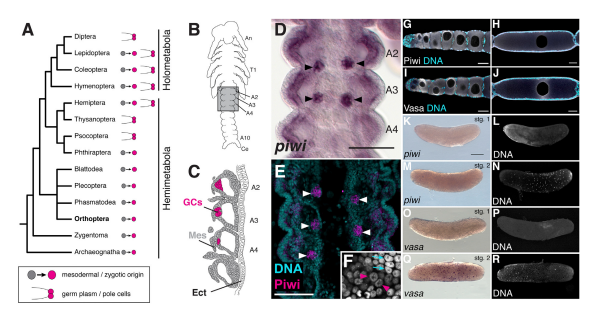


Figure 1. Molecular Markers Suggest Absence of Germ Plasm in Gryllus

- (A) Phylogenetic distribution of reported PGC specification mechanisms across insects (see Table S1).
- (B) Schematic of a stage 9 Gryllus embryo, highlighting the region enlarged in (D)-(E) (gray box). A2-A4, abdominal segments 2-4.
- (C) Tracing of Wheeler's description of orthopteran PGCs at the earliest time point they could be identified [10]. GCs, germ cells (magenta); Mes, mesoderm (gray); Ect, ectoderm (white).
- (D and E) Gryllus PGCs (arrowheads) express piwi transcripts (D) and protein (E).
- (F) PGCs (arrowheads) display nuclear morphology distinct from somatic cells (arrows).
- (G-J) Piwi (G and H) and Vasa (I and J) proteins do not localize asymmetrically in the ooplasm.
- (K and M) piwi transcripts are undetectable during stages 1-2.
- (O and Q) vasa transcripts are undetectable at stage 1 (O) and associated with all energid nuclei at stage 2 (Q).
- (L, N, P, and R) Corresponding nuclear stains.

Scale bar represents 100 μm in (D) and (E); 50 μm in (G)–(J); 200 μm in (K)–(R). Anterior is up in (B)–(F), left in (G)–(R). See also Figures S1, S2, and Table S1.

segment. These cells possessed universal PGC characteristics [3] of large nuclei with diffuse chromatin and a single large nucleolus (Figure 1F). Based on these gene expression, nuclear morphology, and embryonic location data, we conclude that these cells are *Gryllus* PGCs. We also examined the expression of four additional putative candidate PGC marker genes (*tudor*, *piwi-2*, *AGO3-A*, and *AGO3-B*) but found that they were not specific PGC markers in *Gryllus* embryos (Figures S1B and S1F).

# **Gryllus Germline Markers Do Not Localize within Oocytes or Reveal PGCs in Early Embryos**

We next examined the expression of *Gryllus* PGC markers during earlier stages of embryogenesis and oogenesis to test whether they revealed the presence of germ plasm in oocytes or PGCs in early embryos. All genes tested were consistently ubiquitous throughout oogenesis and never localized asymmetrically within the ooplasm (Figures 1G–1J, S1D, and S1F), although Vasa and Piwi proteins were enriched around the oocyte nucleus (Figures 1G–1J). In blastoderm-stage embryos (stages 1–3) and early germband-stage embryos (stage 4), *piwi* (Figures 1K–1N, S1E, and S2G–S2P'), *vasa* (Figures 10–1R, S1E, and S2Q–S2Z'), *bol*, and *gcl* (Figures S2A–S2F) were expressed ubiquitously at low levels and showed no asymmetric localization within the embryo. These results are in stark contrast to the posterior accumulation of PGC determinants in *Drosophila* oocytes and early embryos [16, 18–20] and

suggest an absence of germ-plasm-driven PGC specification in *Gryllus*.

## Gryllus PGCs Arise De Novo during Midembryogenesis

To determine the embryonic origin of Gryllus PGCs, we examined the expression of piwi and vasa transcripts and proteins throughout abdominal elongation and segmentation. During early germband stages (stage 4), we detected low-level ubiquitous expression of both genes in all ectodermal and mesodermal cells (Figures 2A-2B' and S3A-S3B'). It was not until thoracic limb bud enlargement began (stage 5) that piwi transcripts were detected at higher levels in two subsets of cells in abdominal segments A2-A4 among the lateral abdominal mesoderm (Figures 2C and 2C'). As appendage elongation began (stage 6), piwi-positive cells split into distinct groups along the anterior-posterior axis (Figures 2E and E'), and Piwi protein levels rose in these cells (Figures 2E" and 2E""). During morphological segmentation of the abdomen (stages 7-9) these cell groups coalesced into four to six distinct clusters adjacent and dorsal to the coelomic pouches in segments A2-A4 and continued to express high levels of piwi transcripts and protein (Figures 2F-2H" and S3D-S3F"). vasa transcript and protein expression was similar to that of piwi, but vasa became enriched in PGCs slightly later than piwi and showed higher expression levels in the soma (Figure S3).

Interestingly, hallmarks of active transcription were observed in PGCs throughout all stages examined (Figures

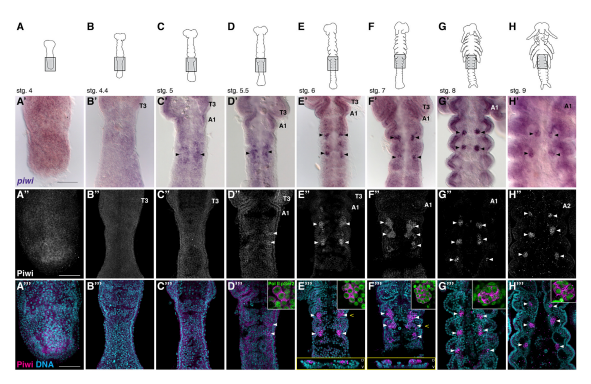


Figure 2. Gryllus PGCs Arise during Early Segmentation Stages

Drawings of *Gryllus* embryogenesis (A–H), highlighting the region depicted below (gray boxes). *piwi* transcripts are ubiquitous in stages 4 and 4.4 embryos (A' and B'), but beginning at stage 5 (C'), two bilateral groups of *piwi*-positive cells arise on the dorsal surface of the embryo, then resolve into clusters during later stages in abdominal segments A2–A4 (D'–H'). Piwi protein shows a similar expression pattern to that of *piwi* transcript but is enriched slightly later in development (A"–H"). Piwi (magenta) overlaid on nuclear stain (cyan) reveals that PGCs arise prior to coelomic pouch formation and ultimately reside medial to these mesodermal structures (A""–H"). Yellow-framed insets in (E"') and (F"') show orthogonal projections at the position of the caret, illustrating the dorsal location of PGCs. Insets in (D")–(H"') show expression of RNA polymerase II pSer2 (green) in PGCs. T3, thoracic segment 3; A1 and A2, abdominal segments 1 and 2.

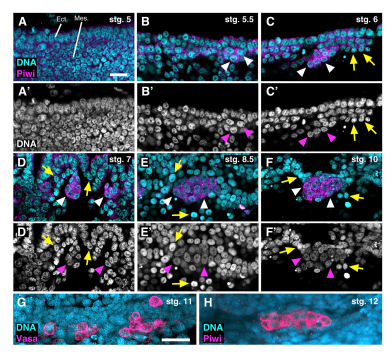
Scale bar represents 100  $\mu\text{m}.$  Anterior is up in all panels. See also Figures S2 and S3.

2D"'-2H"', insets). This is consistent with *Gryllus* PGC formation via active transcriptional response to inductive signaling between cells rather than PGCs being a transcriptionally quiescent subpopulation of early-segregated cells as seen in *Drosophila* and other species with germ plasm [21].

Consistent with a conversion of presumptive mesoderm cells to PGCs beginning at stage 5, the nuclear morphology of mesodermal cells correlated with the relative levels of Piwi expression throughout development. At stage 4, all mesoderm cells had uniform Piwi expression and nuclear morphology, relatively compact chromatin, and multiple nucleoli (Figures 3A and 3A'). As Piwi expression increased in presumptive PGCs, their nuclei became larger with increasingly diffuse chromatin, whereas nuclei of neighboring Piwipoor cells decreased in size, and their chromatin became compact as they progressed through mesoderm differentiation (Figures 3B-3D'). By stages 8-9, PGCs were clearly distinguished by their high nuclear-cytoplasmic ratio, diffuse chromatin, and single large nucleolus (Figures 3E-3F'), criteria used to identify PGCs in historical studies of Orthoptera and other animals [3]. Following stage 10, PGC clusters merged via short-range cell migration (Figure 3G) and coalesced into two bilateral gonad primordia (Figure 3H) located in segments A3-A4. Thus, Gryllus PGCs do not undergo long-range migration, as they do in many other species including *Drosophila* [22], but rather arise near the location of the embryonic gonad.

# Knockdown of *Gryllus piwi* or *vasa* Does Not Disrupt PGC Formation or Maintenance

We knocked down vasa and piwi function using both maternal and zygotic RNAi (mRNAi and eRNAi, respectively) and confirmed knockdown using qPCR and immunostaining (Figures 4A, 4B, 4E, and 4H). In contrast to Drosophila, in which vasa and piwi are required maternally for embryonic PGC formation [15, 18], mRNAi against vasa and piwi did not disrupt PGC formation in Gryllus embryos (Figures 4C-4H), and there was no significant difference in the number of PGCs in either vasa or piwi mRNAi or eRNAi embryos relative to controls (Figures 4I and 4J). Furthermore, female embryos laid by mothers injected with vasa or piwi doublestranded (ds)RNA ultimately grew into fertile adults with fully functioning ovaries (Figure 4K-4M). In contrast to the Drosophila requirement for vasa and piwi in oogenesis and axial patterning [15, 23], Gryllus females injected with vasa or piwi dsRNA displayed no defects in egg laying, oogenesis, or axial patterning (Figures S4A-S4C). Moreover, double knockdown of vasa + piwi maternally or zygotically did not



disrupt PGC formation or axial patterning (Figures S4C–S4E), indicating that these genes do not act redundantly to direct PGC specification.

### vasa and piwi Play Roles in Gryllus Spermatogenesis

In mice, which lack germ plasm and specify PGCs from presumptive mesoderm via signaling, vasa and piwi are not required for PGC specification but do mark established PGCs of both sexes and play roles in adult spermatogenesis [24, 25]. We tested whether these genes were required for adult spermatogenesis in Gryllus by injecting adult males with dsRNA for vasa or piwi to achieve paternal RNAi (pRNAi). Gryllus testes comprise 200-300 testioles (sperm tubules) [26], within which spermatogenesis proceeds from anterior to posterior (Figures 5A and S5A-S5G). The anterior region of each testiole expresses Vasa and Piwi proteins (Figures S5T and S5U) and contains primary and secondary spermatogonia (Figure 5A). Knockdown of vasa or piwi via pRNAi severely reduced spermatogonial region length (Figure 5H). In both vasa and piwi pRNAi testes, meiotic spermatocytes were found in the anterior region of testioles, in some cases almost abutting primary spermatogonia (Figures 5C, 5D, 5F, and 5G, yellow arrowheads), and secondary spermatonial cysts were reduced (Figures 5C, 5D, and 5G, red arrows) or absent (Figure 5F), suggesting that the mitotic divisions of primary spermatogonia were affected. The misregulation of primary spermatocyte divisions was not due to absence of the germline stem cell niche (apical cell), which was present in piwi and vasa pRNAi testes (Figures 5E-5G, asterisks, and S5B, S5H, and S5N). Postspermatogonial stages of spermatogenesis appeared unaffected (Figures S5I-S5M and S5O-S5S). These data indicate that, as in mice and other animals (see Discussion), piwi and vasa play a role in Gryllus gametogenesis in adult males.

Figure 3. Piwi Expression Correlates with Acquisition of PGC Nuclear Morphology in Abdominal Mesoderm Cells

(A–F') Between stages 5 and 10, Piwi expression increases in PGCs (A–F, arrowheads), and nuclear morphology of Piwi-enriched cells changes accordingly (A′–F', arrowheads). Late stage 5 PGCs have chromatin compaction and multiple nucleoli similar to neighboring mesodermal cells (B and B'). As Piwi enrichment in PGCs increases (arrowheads), their chromatin becomes more diffuse and nuclear size increases (yellow arrows) (C and C'). In subsequent stages chromatin morphology differences become more pronounced (D–F').

(G and H) At stage 11 (G), PGCs commence shortrange migration along the anterior-posterior axis toward the intersegmental region of A3–A4 to form a single gonad primordium on each side of the embryo by stage 12 (H).

Scale bars represent 50  $\mu m$  in (A) (applies to A'-F') and (G) (applies to H). Anterior is to the left.

## Mesoderm Is Required for *Gryllus* PGCs

Our observations thus far suggested that PGCs arise from among mesodermal cells during abdominal segmentation. To test this hypothesis, we took advantage of the conserved role of the

twist gene in mesoderm development [27] to ask whether PGCs could form if mesoderm development was compromised. Gryllus twist is expressed in the abdominal mesoderm beginning during axial elongation, including in cells of the region where PGCs arise (Figures S6A-S6D2'). In Drosophila, twist mutants display gastrulation defects [28], yet PGCs form normally because PGC specification occurs via germ plasm well before gastrulation (Figures 6A and 6E). In Gryllus, twist eRNAi similarly causes disorganization or loss of major mesodermal structures within all body segments (Figures 6F and 6G, compare to 6B and 6C). In contrast to Drosophila, however, 49% of Gryllus twist eRNAi embryos lack PGCs, compared to 0% of controls (p < 0.01, Figures 6D and 6G'), and those twist eRNAi embryos that specify PGCs have fewer than controls (p = 0.05, Figure 6H). These results are consistent with the hypothesis that PGCs form from a subset of abdominal mesoderm. Alternatively, PGCs may be formed normally at stage 5 but fail to be maintained due to absent or compromised mesodermal surroundings.

### Discussion

We have shown that neither vasa nor piwi are required maternally or zygotically for the formation of functional PGCs (Figures 4 and S4) but instead play a role in spermatogonial divisions in adult males. Our results differ from those of analogous experiments in D. melanogaster [15, 18], indicating that the functions of these genes have diverged between Gryllus and Drosophila. Although these genes are not required for Gryllus PGC formation, we propose that, together with gcl and boule expression (Figure S2) and the transition from mesodermal to PGC-like morphology in situ (Figure 3), vasa and piwi are nevertheless informative Gryllus PGC markers, despite their pleiotropic roles in other developmental

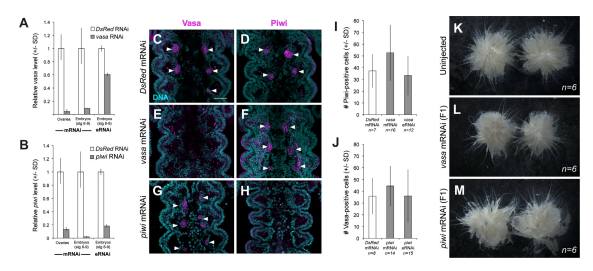


Figure 4. vasa and piwi Are Not Required for Gryllus PGC Specification

(A and B) qPCR validation of vasa and piwi knockdown following mRNAi and eRNAi.

(C–H) Piwi-positive PGCs (arrowheads) form in vasa RNAi embryos, and Vasa-positive PGCs form in piwi RNAi embryos. Consistent with qPCR results, vasa mRNAi (E; 100%, n = 9) and piwi mRNAi (H; 60%, n = 10) abolished respective protein expression. eRNAi produced similar results (not shown). (I and J) PGC quantification confirms that PGC formation is not reduced (Student's t test: vasa mRNAi p = 0.07; vasa eRNAi p = 0.57; piwi mRNAi p = 0.24; piwi eRNAi p = 0.77).

(K–M) Ovaries from adult offspring of vasa and piwi pRNAi mothers (L–M) are indistinguishable from uninjected controls (K). Scale bar represents 50 μm in (C)–(H).

See also Figure S4.

processes. We cannot eliminate the possibility that untested marker genes might show an earlier PGC specification event than the one we identify in stage 5 (Figure 2C). However, given the conserved coexpression of the tested genes in PGCs of

multiple metazoans, we believe it unlikely that all four would be absent from *Gryllus* PGCs at the time of their specification.

Evidence from multiple systems suggests that functional divergence of vasa and piwi is widespread. In

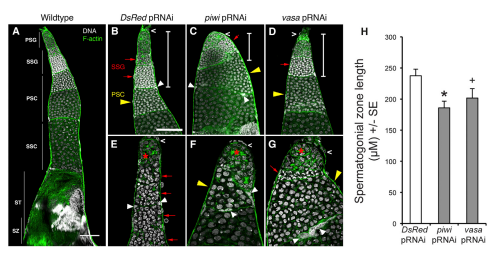


Figure 5. piwi and vasa pRNAi Causes Defects in Spermatogonial Proliferation

(A) Wild-type  ${\it Gryllus}$  testiole showing the stages of spermatogenesis.

(B–G) White bars in (B)–(D) indicate the spermatogonial zone containing secondary spermatogonia (SSG, red arrows). The zone of primary spermatocytes (PSC, yellow arrowheads) nearly abuts the primary spermatogonial zone in piwi (C and F) and vasa (D and G) pRNAi testes because of the shortened SSG zone but is absent from the anterior region of control testioles that have extensive SSG populations (B and E). Higher magnification (E–G) is shown of anterior testiole regions in control (E), piwi RNAi (F), and vasa RNAi (G) testes.

(H) vasa or piwi paternal RNAi results in a shortened spermatogonial zone compared to controls (Student's t test:  $^+p < 0.01$ ,  $^+p = 0.06$ ). Scale bar represents 100  $^{\mu}m$  in (A) and 50  $^{\mu}m$  in (B) (applies also to C–G). Anterior is up in (A)–(G). See also Figure S5.

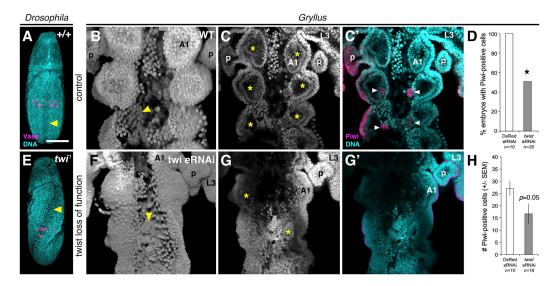


Figure 6. twist eRNAi Disrupts Gryllus PGC Formation

Drosophila twist<sup>1</sup> embryos display gastrulation defects (A and E; arrowhead indicates ventral furrow) but specify PGCs (magenta) properly. In comparison to wild-type (B-C'), Gryllus twist eRNAi embryos (F-G') have disorganized mesoderm (F, arrowhead), show absent or rudimentary coelomic pouches (G, asterisks), and fail to specify PGCs (G') significantly more often than controls (D; Fisher's exact test, \*p < 0.01). When they do specify PGCs, Gryllus twist eRNAi embryos have fewer PGCs than controls (H; Student's t test, p = 0.05).

Scale bar represents 50 µm and applies to all panels. L3, third thoracic leg; A1, first abdominal segment; p, pleuropodia. Anterior is up. See also Figure S6.

D. melanogaster, where both genes were first discovered, mutations in vasa or either of the two piwi orthologs (piwi and aubergine) cause defects in germ plasm formation, oogenesis, PGC specification, and posterior patterning [15, 18, 19, 29]. Similarly, vasa and piwi orthologs are required for PGC specification, development, and oogenesis in C. elegans, D. rerio [see 2], and medaka [30]. In mice, however, vasa is expressed in embryonic PGCs of both sexes, but vasa<sup>-/-</sup> homozygotes display no discernable defects in PGC specification or oogenesis and instead show a male-specific defect in spermatogenesis [25]. Similarly, knockout mice for any of the three PIWI family homologs display spermatogenic defects only, with no defects in females [24, 31, 32]. Our data therefore suggest that the roles of Gryllus vasa and piwi are similar to those of their mouse homologs. Functional genetic and gene expression data from insects (Table S1) suggest that, in this clade, an instructive role for these genes in PGC formation may be restricted to the Holometabola, perhaps concomitant with the co-option of oskar to the top of the PGC specification pathway [14]. Consistent with this hypothesis, vasa is dispensable for PGC formation in another hemimetabolous insect, the milkweed bug, Oncopeltus fasciatus [33].

Our data indicate that a zygotic mode of PGC specification is likely present in *Gryllus*, whereby PGCs appear to arise from presumptive mesoderm. Because *twist* is expressed broadly in mesodermal cells (Figure S6), our *twist* RNAi results could indicate either that mesoderm gives rise to PGCs directly or that mesoderm is required to maintain PGCs (we note that hese interpretations are not mutually exclusive). However, our morphological (Figure 3) and gene expression (Figures 2, S3, and S6) analyses strongly suggest that cells convert from mesodermal to PGC fate in situ. Alternatively, an undifferentiated population of PGC precursors could exist that do not express any of the tested PGC marker genes but are induced

to adopt PGC fate by adjacent mesodermal cells. If this is the case, however, we note that such pluripotent precursors cannot require maternal provision of *vasa* or *piwi* and would most likely be specified by zygotic mechanisms.

Several lines of evidence suggest that a cell lineage relationship between mesoderm and the germline may be a cell type association predating the emergence of Bilateria. Bilaterian germ cells are strikingly similar in gene expression and cytological characteristics to endomesodermally derived stem cells in bilaterian outgroups. Whereas nonbilaterians do not have a dedicated germline per se, their pluripotent stem cell populations serve the function of the germline (reviewed in [34]), and cnidarian pluripotent stem cells are derived from endomesoderm during embryogenesis [35-39]. Within bilaterians, gametogenic cells are consistently described as arising from gonadal epithelia of mesodermal origin in most arthropods and many marine invertebrates (reviewed in [3]). In many spiralians, cytological, cell lineage, and molecular data indicate that PGCs originate from a multipotent mesodermal precursor or precursors (see also [3, 40-47]). Recent studies suggest that mouse PGCs may default to a mesodermal specification program if germline induction signals are absent [48, 49]. The work presented here illuminates broad similarities between PGC specification and vasa function in Gryllus and in the mouse. Future work will be required to explore this apparent similarity in greater depth and to determine the extent of conservation in the developmental and molecular processes involved in specifying the germline across Bilateria.

### **Experimental Procedures**

Gryllus husbandry, gene expression analysis, mRNAi, eRNAi, and phenotypic analysis were carried out as previously described [50]. For pRNAi, 5 µl of 3 µg/ml dsRNA was injected into the coelomic cavity of adult males

1–3 days after the final molt to sexual maturity, and testes of injected males were dissected for analysis 7 days after injection (details in Supplemental Experimental Procedures).

#### Accession Numbers

Sequences have been deposited in GenBank (accession numbers KC242803-KC242808).

### **Supplemental Information**

Supplemental Information includes six figures, Supplemental Experimental Procedures, and one table and and can be found with this article online at http://dx.doi.org/10.1016/j.cub.2013.03.063.

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Current Biology, Volume 23 Supplemental Information

Germ Cell Specification Requires Zygotic Mechanisms Rather Than Germ Plasm in a Basally Branching Insect

Ben Ewen-Campen, Seth Donoughe, Donald Nat Clarke, and Cassandra G. Extavour

## **Author Contributions**

B.E.-C. and C.G.E. designed research, analyzed data, and wrote the paper; experiments were carried out by S.D. (*twist* eRNAi experiment), D.N.C. (pRNAi egg-laying and embryonic survival scoring, PGC migration analysis), C.G.E. (nuclear morphology analysis, *twist* in situ hybridization analysis, spermatogenesis analysis of pRNAi experiments), and B.E.-C. (all other experiments); C.G.E. obtained funding for the research.

## Supplemental Inventory

- Supplemental Figures S1-S6
- Table S1, Related to Figure 1. Data on Insect PGC Origin during Embryogenesis
  Please see accompanying Excel file. Only studies directly addressing the mechanism
  and/or description of the first embryonic appearance of PGCs are referenced. Numbered
  references in the Excel file are listed at the end of this Supplemental Information
  document.
- Supplemental Experimental Procedures
- Supplemental References

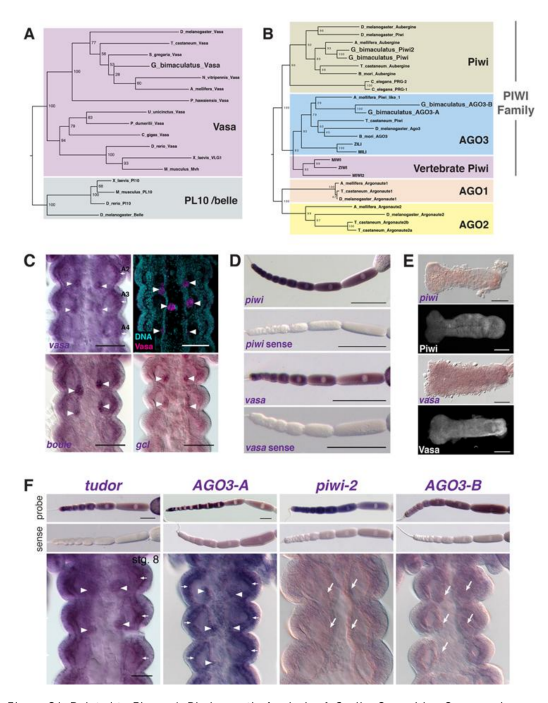


Figure S1, Related to Figure 1. Phylogenetic Analysis of *Gryllus* Germ Line Genes and Expression Patterns of Additional Germ Line Marker Candidates

- (A and B) Maximum-likelihood phylogenetic reconstruction of Vasa and Piwi amino acid sequences. As previously reported [1], *Gryllus* Vasa falls clearly within other insect *Vasa* genes, not the PL10/Belle class of RNA helicases (A). *Gryllus* possesses two *piwi*-like genes and two AGO3-like genes, both of which represent species-specific duplications (B). As only the first identified *piwi*-like gene [2] was enriched in *Gryllus* PGCs, we focus the present analyses on this orthologue, which we refer to here simply as *piwi* as it is clearly orthologous to other animal *piwi* genes. Note that *aubergine* is a *Drosophila*-specific duplication of *piwi*.
- (C) *Gryllus* PGCs (arrowheads) express high levels of Vasa protein and transcripts of *Vasa*, *boule* and *germ cell less*. All genes are also expressed at lower levels throughout the somatic tissues of the embryo.
- (D) *piwi* and *vasa* transcripts are expressed ubiquitously during all stages of oogenesis and do not localise to the posterior ooplasm.
- (E) *piwi* and *vasa* transcripts and protein products are expressed ubiquitously in stage 4 embryos. The apparent increased expression levels at the germ band posterior are an artifact of tissue thickness. (F) Top row: *tudor*, *AG03-A*, *piwi-2*, and *AG03-B* transcripts are not localized asymmetrically in
- oocytes. Bottom row: these genes do not specifically label *Gryllus* PGCs. In the PGC-containing region (Figure 1B-E) at stage 9, *tudor* and *AGO3-A* are detectable in PGCs (arrowheads), but are also expressed throughout the somatic tissues of the embryo (arrows). *piwi-2* and *AGO3-B* are not detected above background levels in stage 8-9 embryos. Arrows mark PGCs recognisable based on morphology and anatomical position independent of gene expression. Anterior is to the left in D and top two rows row of E, and up in C and bottom row of E. Scale bars = 100  $\mu$ M in C, 500  $\mu$ M in D; 200  $\mu$ M in E and top two rows of F; 50  $\mu$ M bottom row of F.

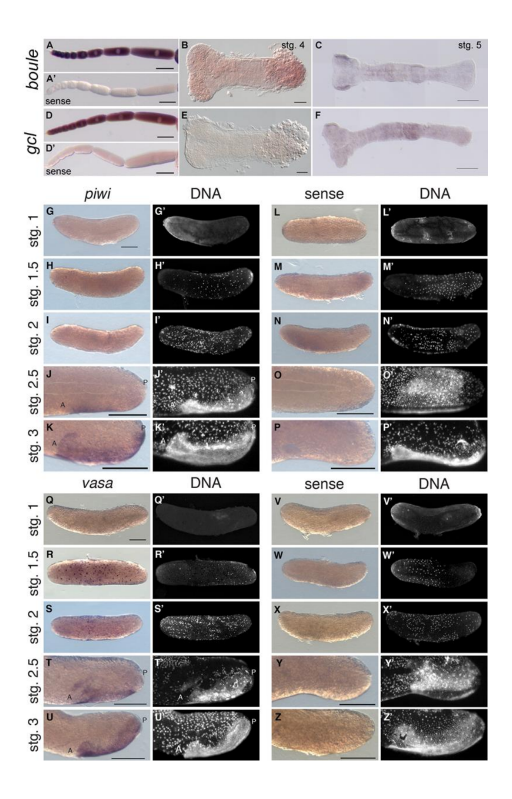


Figure S2, Related to Figures 1 and 2. Additional Gene Expression Data in Support of the Absence of Germ Plasm in *Gryllus* 

(A–F) Expression of *boule* and *gcl*, which mark PGCs in stage 9 embryos (Figure 1) during oogenesis and early embryogenesis. Neither gene is asymmetrically localized in oocytes (A and D; A' and D' show sense controls). Both genes are expressed ubiquitously during stages 4 (B and E) and 5 (C and F), and do not reveal any segregated PGCs during these stages. (G–Z) *piwi* (G–K) and *vasa* (Q–U) expression during blastoderm stages. (L-P and V-Z) Corresponding sense controls. (G'–Z') Nuclear stains of adjacent panels. *piwi* transcripts are undetectable during blastoderm stages (G–I), and are found ubiquitously at low levels as the germ band condenses (J and K). *vasa* transcripts are undetectable in just-laid eggs (Q), and energids are associated with all nuclei along the A–P axis as they populate the blastoderm surface (R). During subsequent blastoderm divisions (S), *vasa* expression is not localized to any specific subset of nuclei. As the germ band condenses at the posterior of the egg (T, U), *vasa* expression is detected at similar, low levels throughout the germ band but not enriched at the posterior or in any other specific region. A = germ band anterior, P = germ band posterior. Scale bars = 200  $\mu$ M in (A)–(A') and (D)–(D'); 100  $\mu$ M in (B) (applies also to E); 200  $\mu$ M in (C) (applies also to F), (G), and (Q) (applies also to H–

I and R-S, respectively), and (J)-(K), (O)-(P), (T)-(U), and (Y)-(Z). Anterior is to the left in all

panels.

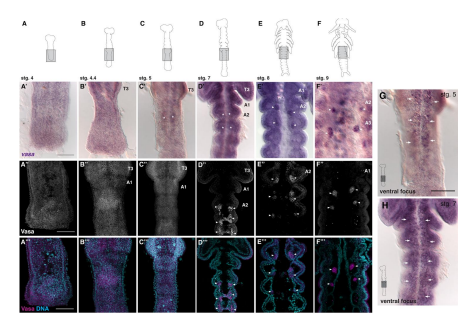


Figure S3, Related to Figure 2. Expression of *vasa* Transcript and Protein throughout Abdominal Segmentation (A–F) Schematic drawings of *Gryllus* mid-staged embryos of the stages shown here; boxed grey areas indicate regions shown in panels below. *Vasa* transcripts are expressed ubiquitously during stages 4–4.4 (A and B), and do not reveal the presence of PGCs at this stage, consistent with *piwi* expression. (C and D) During stages 5 and 7 *vasa* transcripts do not reveal the presence of PGCs. Astrices in (C' and D') denote out-of-focus staining in the ventrally located nervous system, which is shown in focus in (G–H). *vasa* transcripts are detected in PGCs during stage 8 and 9 (E and F). Vasa protein is ubiquitously expressed during stages 4-5 (A'–C'). In stage 7 embryos (D' and D''), Vasa protein is strongly enriched in PGCs, and this expression continues in stage 8 and 9 (E'–F' and E"–F"). Arrowheads indicate PGC clusters. T3 = thoracic segment 3; A1, A2 = abdominal segments 1 and 2. Scale bar = 100 μM. Anterior is up in all panels.

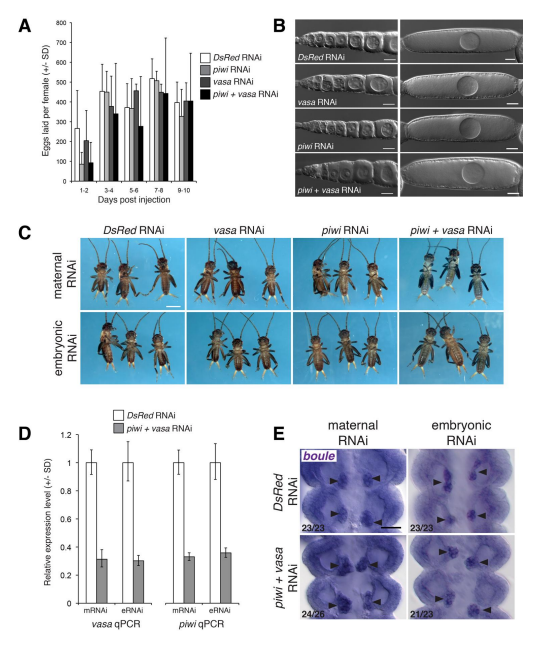


Figure S4, Related to Figure 4. Phenotypic Analysis of *vasa* and *piwi* mRNAi and *vasa* + *piwi* Double eRNAi and mRNAi, in Ovaries and Embryos
(A) Females injected with dsRNA against *vasa* or *piwi* lay numbers of eggs that do not differ

significantly from controls (student's t-test, p>0.05 in every pairwise comparison of vasa or piwi RNAi with DsRed RNAi on the indicated days post-injection).

## (B) Ovaries dissected from *vasa* or *piwi* mRNAi females 10 days after injection are morphologically

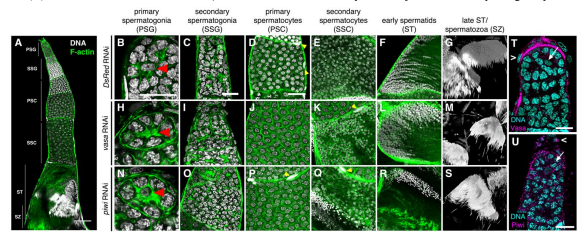


Figure S5, Related to Figure 5. *vasa* or *piwi* pRNAi Does Not Disrupt Postspermatogonial Stages of Spermatogenesis in *Gry/lus* Wild type *Gry/lus* testiole showing the stages of spermatogenesis (A). Primary spermatogonia (PSC) undergo self-renewing divisions, which are thought to occur under the influence of a single apical cell (red arrowheads in B, H, N) that provides a "stem cell niche" analogous to the hub of *Drosophila* testes [3, 4]. Following seven mitotic divisions by cysts of secondary spermatogonia (SSG) enclosed by somatic cell sheaths (yellow arrowheads in D, K, P–Q), the resulting 128 primary spermatocytes (PSC) undergo meiosis (secondary spermatocytes: SSC) to produce 512 clonally related spermatids (ST), which undergo synchronous spermeiogenesis to produce bundles of mature spermatozoa (SZ) [5]. Although *vasa* and *piwi* pRNAi testes display a reduction in the number of secondary spermatogenesis (D–G, I–M, O–S), which are surrounded by somatic sheath cells (arrowheads), and cysts proceeding normally through all stages of spermatogenesis (D–G, I–M, O–S), which are surrounded by somatic sheath cells (arrowheads) as in controls. Vasa (T) and Piwi (U) proteins are expressed in the anterior region of testioles, which contains primary spermatogonia (arrows) and somatic sheath cells (arrowheads), and is enclosed by a cellular peritoneal sheath (carets). Scale bars = 100 μm in A; 50 μm in (D) (applies also to E–G, J–M, P–S); 25 μm in (B) (applies also to I and O); 20 μm in (T)–(U).

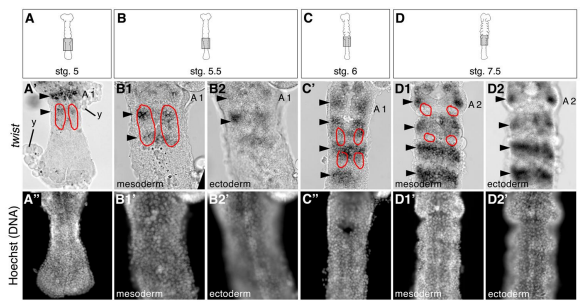


Figure S6, Related to Figure 6. *twist* Expression in the Abdominal Mesoderm where *Gryllus* Germ Cells Arise (A–D) Schematic drawings of progressive stages of *Gryllus* embryogenesis; grey box indicates region shown in panels below. *twist* transcripts accumulate in an anterior to posterior progression in abdominal segments, indicated by black arrowheads in (A'), (B1)–(B2), (C'), and (D1)–(D2). Red outlines in (A'), (B1), (C'), and (D1) indicate the regions that become enriched for *piwi* expression at these stages, suggesting that these are the sites of PGC origin and showing that these regions express *twist* at the proposed onset of PGC specification (stage 5). Bottom row shows nuclear staining of corresponding bright field images in the row above. Anteriormost abdominal segment is labeled in each panel. *y* = yolk. *twist* expression is confined to the mesoderm and absent from the ectoderm, as shown in micrographs of mesodermal focal planes in (B1 and D1), and ectodermal focal planes in (B2 and D2).

## Supplemental Experimental Procedures

## Insect Cultures and Embryonic Staging

*Gryllus bimaculatus* cultures were maintained as previously described [139] and embryos were staged according to [140]. *Drosophila Oregon R* and *twist*<sup>1</sup> stocks were obtained from the Bloomington Drosophila Stock Center (#5, #2381).

## Cloning and Phylogenetic Analysis

Orthologues of *boule*, *tudor*, *germ cell-less*, an additional *piwi*-like gene, and two *AGO3*-related genes were identified in a *Gryllus* developmental transcriptome via reciprocal best BLAST hit analysis against the *Drosophila melanogaster* proteome. *Gryllus twist* was a gift of S. Roth (University of Cologne, Germany).

To resolve the orthology of the four *Gryllus* PIWI family proteins, we used maximum-likelihood based phylogenetic reconstruction as implemented by RAxML v 7.2.8 [141, 142] on the Odyssey Cluster, maintained by the FAS Sciences Division Research Computing Group (Harvard University). The alignment was produced using Muscle [143] and trimmed using Gblocks [144] under the least stringent settings. The best tree and rapid bootstrap analysis were conducted from 2000 independent runs under the WAG model of protein evolution with a gamma distribution of rate heterogeneity.

## In Situ Hybridization

DIG-labeled probes were hybridised at 68° C following standard protocols [139], with 50% polyvinyl alcohol included in the NBT/BCIP development step. Probe lengths were as follows: vasa: 1,953 bp; piwi-1: 781 bp; piwi-2: 821 bp; AGO3-A: 760 bp; AGO3-B: 832 bp; germ cellless: 1,691 bp; boule: 995 bp; tudor: 1,707 bp. Our results for vasa expression (both mRNA and protein) in *Gryllus* differ from those reported by Mito et al. [1], who failed to identify the germ cell clusters that we observed beginning at stages 6/7 (Figures 1, 4, S1, S4). This discrepancy may be due to the strong nervous system expression of *Vasa* that can obscure the relatively weaker PGC expression (their Figures 3I, 3J, 4A-D), and to our use of a species-specific Vasa antibody [2] as opposed to the cross-reactive antibody [145] used by Mito et al. Mito et al. also reported detection of transient *vasa* mRNA staining at the posterior of stage 4 embryos (their Figure 3C-D), and interpreted it as consistent with Heymons' 1895 claim that germ cell precursors arose among the posterior germ band mesoderm shortly after gastrulation [125]. However, we found that this apparently stronger expression is due to the thickness of the posterior germ band tissue at these stages. In three *in situ* hybridization replicates and  $\ge 30$  early stage embryos, Vasa did not show consistent enrichment in any specific embryonic region before stage 5. We therefore conclude that germ cells are not specified until this stage, in agreement with the majority of previous authors on orthopteran germ cell origin [131-133].

## **Immunohistochemistry**

Primary antibodies used were rabbit anti-Gb-Vasa and anti-Gb-Piwi [2] at 1:300, mouse anti-RNA polymerase II pSer 6 Mab H5 (Covance MMS-129R) 1:100, FITC-conjugated anti-alpha Tubulin (Sigma F2168) 1:100 and rabbit anti-Drosophila Vasa 1:500 (gift of P. Lasko) following standard procedures. Goat anti-rabbit secondary antibodies conjugated to Alexa 488, Alexa 555 or Alexa 568 (Invitrogen) were used at 1:500 or 1:1000. Counterstains were Hoechst 33342 (Sigma B2261) 0.1 to 0.05 mg/ml and FITC-conjugated phalloidin (Sigma P5282) 1 U/ml.

### RNA Interference

dsRNA injection into adult females (maternal RNAi = mRNAi) and newly laid embryos (embryonic RNAi = eRNAi) was conducted as previously described [2]. dsRNA fragments for *vasa* and *piwi* were 541 bp and 646 bp, respectively. For eRNAi double knockdown experiments, equal volumes of *vasa* and *piwi* dsRNA were mixed prior to injection. For mRNAi double knockdown experiments, twice the volume of dsRNA as that used for single RNAi experiments (15  $\mu$ g each of *vasa* and *piwi* dsRNA, or 30  $\mu$ g of the DsRed control dsRNA) was injected into adult females. dsRNA was used at a concentration of 3  $\mu$ M (mRNAi) and 5  $\mu$ M (eRNAi and pRNAi).

## qPCR Analysis of Knockdown

qPCR was used to verify RNAi efficacy as follows: total RNA was extracted from RNAi-treated ovaries or stage 8-9 (day 4) embryos using TRIzol (Invitrogen) and including a 30-minute DNase digestion at 37° C to remove genomic contamination. Equal volumes of RNA were used as template for first strand cDNA synthesis using SuperScript III (Invitrogen) including a no reverse transcriptase control. cDNA was diluted 1:5 prior to qPCR. qPCR was conducted using PerfeCta SYBR Green SuperMix (Low ROX, Quanta Biosciences) in a Stratagene MxP3005 machine. Primers amplifying single amplicons of *piwi* (129 bp; F:

TTCGGCCAACTACTTCAAGC; R: AGAGTTTCCCGATGAACACG), *vasa* (150 bp; F: GAACATTGTGAGCCTCATGC; R: TTGCTGAGCCTGGTGGTAT) and beta-tubulin (166 bp; F: TGGACTCCGGTCAGGC; R: TCGCAGCTCTCGGCCTCCTT) were used. Each reaction was conducted in triplicate, and fluorescence measurements were normalised and background-subtracted using the ROX dye present in the PCR reactions.

 $C_t$  values were used to calculate fold change compared to *DsRed*-injected controls using the  $2^{-\Delta\Delta Ct}$  method [146]. Triplicate  $C_t$  values were averaged and the standard deviation was propagated using standard methods.

## Imaging and Image Analysis

Micrographs were captured with AxioVision v.4.8 driving a Zeiss Stereo Lumar equipped with an AxioCam MRc camera, Zen Blue 2011 driving a Zeiss Stereo Zoom equipped with an AxioCam HRc camera, a Zeiss Axio Imager equipped with an AxioCam MRm camera using epifluorescence either with or without an Apotome, or an Olympus IX71 equipped with a Hamamatsu C10600-108 camera. Confocal microscopy was performed with a Zeiss LSM 710 or 780 confocal, using comparable gain, offset, and averaging parameters for all samples. Image analyses were performed with AxioVision v.4.8, Zen 2009 or Zen 2011 (Zeiss), and figures were assembled in Photoshop CS4, InDesign CS4, or Illustrator CS4 (Adobe). For confocal images shown in Figures 1E; 2A"-H"; 4C-H; 6C' and G'; S3A"'-F"', a maximum-intensity projection of multiple optical sections of the antibody staining was superimposed over a single optical section of the nuclear counterstain for visual clarity. All other confocal micrographs are maximum intensity projections (Figures 2A"-H'; 3G and H; S3A"-F"), three-dimensional projections (Figure 6B, F) or single optical sections (all other confocal micrographs).

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# **Supplementary Table 1**

Accompany familiary   State	Insect Order	Species[1]		Presumptive PGC Origin	Mode of PGC Specification[2]	PGC Identification Criteria[3]	Functional or Experimental Evidence[4]	Referenc
Annihological forcidada			n.d.					
March Stager	Diptera							
Angeles personal Basis from Pine office GP UM IN IN 19. Angeles personalered Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijania Statistics Pine office GP UM IN IN 19. California optimizacijani Statistics Pine office GP UM IN IN IN 19. California opt		Acricotopus lucidus	Blastoderm	Pole cell[7]	GP	LM, LSM	N	[6]
Augustes maceleonics   Basicionis   Pies de Che   Circ   N. N.   1.		Aedes aegypti	Blastoderm	Pole cells	GP	LM, MM	N	[7, 8]
Calphone opmonephole   Calphone		Anopheles gambiae	Blastoderm	Pole cells	GP	MM	N	[8]
Canthors apprincepable  Control congress of the State of the Pice or the Cipe		Anopheles maculipennis	Blastoderm	Pole cells	GP	LM	N	[7]
Cultific appliant		Calliphora crythrocephala	Blastoderm	Pole cells	GP	LM	N	[9]
Policy Composed as   Bastedom   Policy of the 100		Calliphora erythrocephala	Blastoderm	Pole cells	GP	LM	N	[10]
Compress provided   Particular   Particula		Ceratitis capitata	Blastoderm	Pole cells	GP	LM, TEM, LSM	N	[11]
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Companies accurated   Balactoom   Pine oath   OP		Clogmia albipunctata	nd[8]	nd	nd	LM	N	[14]
Clark program		Compisura concinnata		Pole cells	GP	LM	N	[15]
Clear gromperfections		Culex fatigans	Blastoderm	Pole cells	GP	LM	N	
Care care-participation			Blastoderm	Pole cells	GP	LM	Υ	[7, 17]
Decay print   Baskoom							N	
Description prolonger   Bistochem								
Memorance programs		-						
Luciole agrams								
Licale serioraza Mesperagno rovia Mesper								
Master mereleane   Balestoderm   Pule cell   CP								
Master ammiricans								
Master mersebase   Bistrictoms								[31]
Masse domested   Mass								[32]
Marcar combon   Bilandorm   Pries cells   GP								[33, 34
Mycophia gapper  Bistocom					GP			[13, 35
Processing servicials   Blastodorm   Pule calls   CP		Musca vomitora	Blastoderm	Pole cells	GP	LM	N	[36, 37
Phonesina sericatal   Bisatoderim   Pide cells   GP		Mycophila speyeri	Blastoderm	Pole cells	GP	LM	N	[38]
Price cate   Pri			Blastoderm	Pole cells	GP	LM	N	[39]
Prince   P		Phormia regina	Blastoderm	Pole cells	GP	LM	N	[40]
Science coproposable   Blastodorm   Pote cells   GP   LM   N   1.65		Rhynchosciara americana	Blastoderm			MM		[41]
Simple proper   Biastodorm   Pule cells   GP   LM   N   145			Blastoderm		GP	LM	N	
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Ephestis kuterhiella         Germ band         PGB primary ectoderm         MZ         LM, MM         N         [59, 169, 169, 169, 169, 169, 169, 169, 16		Endromis versicolora	Germ band	PGB primary ectoderm	MZ	LM	N	[58]
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Euvanessa antiopa Blastoderm Ventral cellular blastoderm MZ LM N [5] Hellothia zea Gern band Gern band dem band midline MZ LM N [6] Luehdorila japonica Gern udiment Ventral gern udiment MZ LM N [5] Neomicropteryx ripponensis Gern band CP Mesoderm MZ LM N [5] Peringhora gosspiella Blastoderm Ventral gern udiment MZ LM N [5] Peringhora gosspiella Blastoderm Pole cells GP LM, TEM N [5] Priest rapse Gern band Mesoderm MZ LM N [6] Solenobla friquetrella Gern band Mesoderm MZ LM N [6] Solenobla friquetrella Gern band Mesoderm MZ LM N [6] Solenobla friquetrella Gern band PGB primary ectoderm MZ LM N [6] Callandra oryzae Blastoderm Pole cells GP LM N [6] Callandra oryzae Blastoderm Pole cells GP LM N [6] Callandra oryzae Blastoderm Pole cells GP LM N [7] Callandra oryzae Blastoderm Pole cells GP LM N [7] Callandra oryzae Blastoderm Pole cells GP LM N [7] Callandra pranaira Blastoderm Pole cells GP LM N [7] Callandra oryzae Blastoderm Pole cells GP LM N [7] Callandra oryzae Blastoderm Pole cells GP LM N [7] Callandra oryzae Blastoderm Pole cells GP LM N [7] Callandra callosa Blastoderm Pole cells GP LM N [7] Callandra callosa Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis Blastoderm Pole cells GP LM N [7] Corynodes pusis GP LM N [		Epiphyas postvittana		Germ band midline	MZ			
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Luehdorfia japonica         Germ nudiment         Ventral germ rudiment         MZ         LM         N         155           Neomicrophery napponensis         Germ band         CP Mesonderm         MZ         LM         N         164           Parassius glaciales         Germ band         Ventral germ undiment         MZ         LM         N         155           Pectinophora gossypiella         Blastoderm         Pole cells         GP         LM, TEM         N         166           Specification in graph and problem of the pr								
Neomicropteryx nipponensis Germ band CP Mesoderm MZ LM N [56]  Parnassius glacialis Germ udiment Ventral germ udiment MZ LM N [55]  Pectrophora gossypiella Blastoderm Pole cells GP LM, TEM N [66]  Pieris rapae Germ band Mesoderm MZ LM N N [66]  Solenolis triquetrella Germ band PGB primary ectoderm MZ LM N N [66]  Calendra granaria Blastoderm Pole cells GP LM N N [68]  Calendra orgaze Blastoderm Pole cells GP LM N N [68]  Calendra callosa Blastoderm Pole cells GP LM N N [70]  Calligrapha multipunctata Blastoderm Pole cells GP LM N N [70]  Calligrapha multipunctata Blastoderm Pole cells GP LM N N [70]  Corynodes pusis Blastoderm Pole cells GP LM N N [70]  Corynodes pusis Blastoderm Pole cells GP LM N N [70]  Pophybrar deceminenta Germ band PCB MZ LM N N [71]  Hydrophilus piceus Germ band PCB MZ LM N N [72]  Expinotarsa deceminenta Blastoderm Pole cells GP LM N N [73]  Rhapophthalmus ohbai Germ Sand POB MZ LM N N [73]  Rhapophthalmus obbai Germ Band PCB MZ LM N N [73]  Rhapophthalmus obbai Germ Band PCB MZ LM N N [73]  Rhapophthalmus obbai Germ Band PCB MZ LM, SEM N [73]  Rhapophthalmus obbai Germ Band PCB MZ LM, SEM N [73]  Rhapophthalmus obbai Germ Band PCB MZ LM, SEM N [74]  Rhapophthalmus obbai Blastoderm Pole cells GP LM N N [73]  Rhapophthalmus obbai Blastoderm Pole cells GP LM N N [74]  Rhapophthalmus obbai Blastoderm POle cells GP LM N N [74]  Rhapophthalmus obbai Blastoderm Pole cells GP LM N N [74]  Rhapophthalmus obbai Blastoderm Pole cells GP LM N N [78]  Rhapophthalmus obbai Blastoderm Pole cells GP LM N N [78]  Rhapophthalmus obbai Blastoderm Pole cells GP LM N N [78]  Rhapophthalmus obbai Blastoderm Pole cells GP LM N N [78]  Rhapophthalmus obbai Germ Band Mesoderm MZ EEM, LM, SEM, MM N [78]  Rhapophthalmus obbai Germ Band Mesoderm GP LM N N [78]  Rhaporterous pulgiandis Blastoderm Pole cells GP LM N N [78]  Rhapophthalmus obbai Germ Band Mesoderm MZ EEM, LM, SEM, MM N [78]  Rhapophthalmus obbai Germ Band Mesoderm MZ EEM, LM, SEM, MM N [78]  Rhapophthalmus obbai Germ Band Mesoderm MZ EEM, LM, SEM, M								
Parnassius glacialis Gern rudiment Ventral gern rudiment MZ LM N 155 Pectinophora gossypella Blastoderm Pole cells GP LM, TEM N 165 Pieris rapea Gern band Mesoderm MZ LM N N 165 Solenoba triquethelia Gern band PGB primary ectoderm MZ LM N N 165 Calendra granaria Blastoderm Pole cells GP LM N N 166 Calendra oriyzae Blastoderm Pole cells GP LM N 166 Calendra callosa Blastoderm Pole cells GP LM N 166 Calendra callosa Blastoderm Pole cells GP LM N 166 Calendra callosa Blastoderm Pole cells GP LM N 172 Corynodes pusis Blastoderm Pole cells GP LM N 172 Corynodes pusis Blastoderm Pole cells GP LM N 172 Doryphora decemineata Gern band PGB MZ LM N 173 Hydrophilus picues Gern band PGB MZ LM N 173 Captionara decemineata Gern band PGB MZ LM N 174 Captionara decemineata Gern band PGB MZ LM N 174 Captionara decemineata Gern band PGB MZ LM N 174 Captionara decemineata Blastoderm Pole cells GP LM N 174 Captionara decemineata Gern band PGB MZ LM N 174 Captionara decemineata Blastoderm Pole cells GP LM N 174 Captionara decemineata Blastoderm Pole cells GP LM N 175 Captionara decemineata Blastoderm Pole cells GP LM N 175 Captionara decemineata Blastoderm Pole cells GP LM N 175 Captionara decemineata Blastoderm Pole cells GP LM N 175 Captionara decemineata Blastoderm Pole cells GP LM N 175 Captionara decemineata Blastoderm Pole cells GP LM N 175 Captionara decemineata Blastoderm Pole cells GP LM N 175 Captionara decemineata Blastoderm Pole cells GP LM N 175 Captionara decemineata Blastoderm Pole cells GP LM N 183. Captionara decemineata Blastoderm Pole cells GP LM N 183. Captionara decemineata Blastoderm Pole cells GP LM N 183. Captionara decemineata GP LM N 183. Captionara dec								
Pectinophora gassypiella   Biastoderm   Pole cells   GP								
Perin rapae   Germ band   Mesoderm   MZ								[55]
Solenobla triquetrella         Germ band         PGB primary ectoderm         MZ         LM         N         167           coptera           Callandra granaria         Blastoderm         Pole cells         GP         LM         N         168           Caliandra oryzae         Blastoderm         Pole cells         GP         LM         N         176           Calligrapha multipunctata         Blastoderm         Pole cells         GP         LM         N         171           Corynodes pusis         Blastoderm         Pole cells         GP         LM         N         172           Doryphora decemtineata         Germ band         PGB         MZ         LM         N         173           Hydrophilus picuss         Germ band         CP mesoderm         MZ         LM         N         173           Hydrophilus picuss         Germ disc         GP         LM         N         173           Rhapophthalmus chbai         Germ disc         GP         LM         N         173           Rhapophthalmus chbai         Germ disc         GB         MZ         LM, SEM         N         175           Tanchoiro molitor         Germ band         PGB mesoderm         MZ						,		[65]
Calandra granaria   Blastoderm								[66]
Calandra granaria         Blastoderm         Pole cells         GP         LM         N         [68]           Calendra orgaze         Blastoderm         Pole cells         GP         LM         N         [66]           Calendra calicas         Blastoderm         Pole cells         GP         LM         N         [77]           Caligrapha multipunctata         Blastoderm         Pole cells         GP         LM         N         [72]           Corynodes pusis         Blastoderm         Posterior         GP         LM         N         [72]           Doryphra decemilneata         Germ band         PGB         MZ         LM         N         [73]           Hydrophilus piceus         Germ band         CP mesoderm         MZ         LM         N         [73]           Rhagophthelmus obbai         Germ disc         Posterior germ         MZ         LM         N         [75]           Tenebrio molitor         Germ band         PCB mesoderm         MZ         LM         N         [75]           Tribolum castaneum         Germ band         PCB mesoderm         MZ         TEM, LM, SEM, MM         N         [78]           Aparteles glomeratus         Blastoderm         Pole cells <td< td=""><td></td><td>Solenobia triquetrella</td><td>Germ band</td><td>PGB primary ectoderm</td><td>MZ</td><td>LM</td><td>N</td><td>[67]</td></td<>		Solenobia triquetrella	Germ band	PGB primary ectoderm	MZ	LM	N	[67]
Calendra oryzae         Blastoderm         Pole cells         GP         LM         N         150           Calendra calicaa         Blastoderm         Pole cells         GP         LM         N         177           Calligraphe multipunctata         Blastoderm         Pole cells         GP         LM         N         171           Corynodes pusis         Blastoderm         Postarior         GP         LM         N         172           Doryphora decemlineata         Germ band         PGB         MZ         LM         N         173           Hydrophilus picuss         Germ band         CP mesoderm         MZ         LM         N         173           Leptinotars a decemlineata         Blastoderm         Pole cells         GP         LM         N         173           Rhappohthalmus orbai         Germ disc         disc         MZ         LM, SEM         N         175           Tanchoiro molitor         Germ band         PGB mesoderm         MZ         LM         N         176.5           Tibolium castaneum         Germ band         PGB mesoderm         MZ         LM         N         178.8           Xylebous terrugineus         Blastoderm         Pole cells         GP	eoptera							
Calendria callosa         Blastoderm         Pote cells         GP         LM         N         177           Caligrapha multipunctata         Blastoderm         Pote cells         GP         LM         N         177           Corynodes pusis         Blastoderm         Potestrior blastoderm         GP         LM         N         172           Doryphora decemineata         Gern band         PGB         MZ         LM         N         173           Hydrophilus pieus         Germ band         CP nesoderm         MZ         LM         N         174           Leptinatarsa decemilineata         Blastoderm         Pole cells         GP         LM         N         173           Rhappophthalmus ohbai         Germ disc         Posterior germ         MZ         LM, SEM         N         175           Tenebrio molitor         Germ band         PGB mescoderm         MZ         LM, SEM, MM         N         178.           Tibolium castaneum         Germ band         PGB mescoderm         MZ         LM, SEM, MM         N         178.           Ayiebous terrugineus         Blastoderm         Pole cells         GP         LM         N         182.           Nemoriteter         Apia mellifera         G		Calandra granaria	Blastoderm	Pole cells	GP	LM	N	[68]
Calendra callosa         Blastoderm         Pole cells         GP         LM         N         T/C           Calligrapha multipunctata         Blastoderm         Pole cells         GP         LM         N         1/7           Corynodes pusis         Blastoderm         Posterior blastoderm         GP         LM         N         1/2           Doryphora decemilneata         Germ band         PGB         MZ         LM         N         1/3           Hydrophilus pieus         Germ band         CP mesoderm         MZ         LM         N         1/7           Rhappophthalminus ohbai         Germ dac         Posterior germ         MZ         LM         N         1/7           Tenebrio molitor         Germ band         PGB mesoderm         MZ         LM         N         1/7           Tenebrio molitor         Germ band         PGB mesoderm         MZ         LM         N         1/7           Tenebrio molitor         Germ band         PGB mesoderm         MZ         LM         N         1/7           Tenebrio molitor         Germ band         PGB         MZ         TEM, LM, SEM, MM         N         1/7           Tenebrio molitor         Germ band         PGB         MZ		Calandra oryzae	Blastoderm	Pole cells	GP	LM	N	[69]
Caligraphe multipunctata         Blastoderm         Pole cells         GP         LM         N         171           Corynodes pusis         Blastoderm         Posterior person         GP         LM         N         172           Doryphror decemilneata         Germ band         PGB         MZ         LM         N         173           Hydrophilus piceus         Germ band         CP mesoderm         MZ         LM         N         174           Leptinotarsa decemilneata         Blastoderm         Pole cells         GP         LM         N         173           Rhappothriamus orbai         Germ disc         Posterior germ         MZ         LM, SEM         N         173           Tenebrio molitor         Germ band         PCB mesoderm         MZ         LM, SEM         N         175           Tribolium castaneum         Germ band         PCB mesoderm         MZ         TEM, LM, SEM, MM         N         178.8           Xyleborus terrugineus         Blastoderm         Pole cells         GP         LM         N         183.2           Aparteles glomeratus         Blastoderm         Pole cells         GP         LM         N         183.4           Aparteles glomeratus         Blastoderm		Calendra callosa	Blastoderm	Pole cells	GP	LM	N	[70]
Corynodes pusis   Blastoderm   Posterior   GP		Calligrapha multipunctata	Blastoderm	Pole cells	GP	LM	N	[71]
Darphora deceminesta   Germ band   POB   MZ   LM   N   173				Posterior			N	
Hydrophilus piceus   Germ band   CP mesoderm   MZ								
Leptinotara decemilineata   Blastoderm   Pole cells   GP   LM   N   173								[73]
Rhagophthalmus chbai   Germ disc   Postenico germ   MZ								[74]
Trebopt motifor   Germ band   PGB mesoderm   MZ   LM   N   [76.]		Leptinotarsa decemlineata	Blastoderm		GP	LM	N	[71]
Tenebrio molifor   Germ band   PGB mesoderm   MZ   LM   N   [76.1]		Rhagophthalmus ohbai	Germ disc	Posterior germ disc	MZ	LM, SEM	N	[75]
Tribolium castaneum Germ band PGB MZ TEM.LIM.SEM,MIM N [78-6 Xyleborus ferragineus Blastoderm Pole cells GP LIM N [83.1 K]  Aparteles glomeratus Blastoderm Pole cells GP LIM N [83.1 K]  Apis mellifera Germ band Mesoderm MZ SEM,MIM N [78-85 Copidosoma sp. 2 2" desvage B4 at four cell stage[9] GP LIM N [91-1 K]  Haborbacon juginals Blastoderm Pole cells GP LIM N [91-1 K]  Momoniella (Nasonia) vitripennis Blastoderm Pole cells GP LIM N [100-1 K]  Pimpla turrionellae Blastoderm Pole cells GP LIM N [100-1 K]  Pimpla turrionellae Germ Pole cells GP LIM N [100-1 K]  Pitronicea ribesii Germ band PGB MZ LIM N [100-1 K]  Tiribogaramae evanessons Blastoderm Pole cells GP LIM N [100-1 K]  Tiribogaramae evanessons Blastoderm Pole cells GP LIM N [100-1 K]  Tiritneptis dipronis Blastoderm Pole cells GP LIM N [100-1 K]		Tenebrio molitor	Germ band		MZ	LM	N	[76, 77
Xyleborus terrugineus Blastoderm Pole cells GP LM N 182  Apanteles glomeratus Blastoderm Pole cells GP LM N 183,4  Apanteles glomeratus Blastoderm Pole cells GP LM N 183,4  Apis mellifera Germ band Mesoderm MZ SEM,MM N [78,88  Copidosoma sp. 2" cleavage B4 at four cell stage[9] GP LM M Y [91-6]  Habobracon juglands Blastoderm Pole cells GP LM N 193  Mormoniella (Nasonia) vitripennis Blastoderm Pole cells GP LM N 193  Pirmpla turionellee Blastoderm Pole cells GP LM N 103,3  Pirmoniea ribessi Germ band PGB MZ LM N 100,3  Tirkogarmane evanescens Blastoderm Pole cells GP LM N 100,1  Tirkogarmane evanescens Blastoderm Pole cells GP LM N 100,1  Tirkogarmane evanescens Blastoderm Pole cells GP LM N 100,1								
Aparteles glomeratus         Blastoderm         Pole cells         GP         LM         N         [83, 6]           Apis mellifera         Germ band         Mesoderm         MZ         SEM, MM         N         [78, 88]           Copidosoma sp.         2°d cleavage         B4 at four cell stage[9]         GP         LM         M         Y         [91-6]           Habobraccon juglandis         Blastoderm         Pole cells         GP         LM         N         [90]           Mormoniella (Nasonia) vitripennis         Blastoderm         Pole cells         GP         LM         N         [100-7]           Pilmpla turionellae         Blastoderm         Pole cells         GP         LM         N         [100-7]           Trichogramma evanescens         Blastoderm         Pole cells         GP         LM         N         [106, 106]           Tritheptis dipronis         Blastoderm         Pole cells         GP         LM         N         [106, 106]	enontera	,		Fore cells	5.			[04]
Apis mellifera         Germ band         Mesoderm         MZ         SEM, MM         N         [78,88           Copidosoma ga.         2° delavage         B4 at four cell stage[9]         GP         LM, MM         Y         [91-18]           Habobracon juglandis         Blastoderm         Pole cells         GP         LM         N         [99]           Mormoniella (Nasonielle         Blastoderm         Pole cells         GP         LM         N         [100-19]           Pimpla turionelle         Blastoderm         Pole cells         GP         LM         N         [103,19]           Pricopida ribosii         Germ band         PGB         MZ         LM         N         [100,19]           Trichogramma evanescens         Blastoderm         Pole cells         GP         LM         N         [106,10]           Tritheptis dipronis         Blastoderm         Pole cells         GP         LM         N         [106,10]	-choptera	Apartolas alamentus	Plantadorm	Dala sella	GD.	114	<b>b</b> 1	102 0
Copidosoma sp.         2 <sup>rd</sup> cleavage         B4 at four cell stage[9]         GP         LM, MM         Y         [91:6]           Habobriscon juglandis         Blastoderm         Pole cells         GP         LM         N         199           Mormoniale (Nasonia) vitripennis         Blastoderm         Pole cells         GP         LM         N         100-1           Pimpla turionellae         Blastoderm         Pole cells         GP         LM         N         100-1           Peteronidea ribesii         Gem band         Pole cells         GP         LM         N         110-1           Trichogramme evanessons         Blastoderm         Pole cells         GP         LM         N         1106-1           Trichogramme evanessons         Blastoderm         Pole cells         GP         LM         N         1106-1								
Habobracon juglandis         Blastoderm         Pole cells         GP         LM         N         199           Momnoniella (Nasonia) virippennis         Blastoderm         Pole cells         GP         LM         N         100-2           Pimpla turionellae         Blastoderm         Pole cells         GP         LM         **         100-3           Pteronidea ribesi         Germ band         PGB         MZ         LM         N         100           Trichogramma evanescens         Blastoderm         Pole cells         GP         LM         N         106           Tritneptis dipronis         Blastoderm         Pole cells         GP         LM         N         100								
Mormoniella (Nasonia) vitripennis         Blastoderm         Pole cells         GP         LM         N         [100-2]           Pirmpia turionellare         Blastoderm         Pole cells         GP         LM         **         [103, 3]           Pteronidea ribeasi         Germ band         PGB         MZ         LM         N         [100]           Trichogramma evanescens         Blastoderm         Pole cells         GP         LM         N         [106, 3]           Tritheptis dipronis         Blastoderm         Pole cells         GP         LM         N         [100]								[91-9
Pimpla turioneliae   Blastoderm   Pote cells   GP   LM   *+*   [103.								[99]
Pteronidea ribesi         Germ band         PGB         MZ         LM         N         [10]           Trichogramma evanescens         Blastoderm         Pole cells         GP         LM         N         [106, 1]           Tritneptis dipronis         Blastoderm         Pole cells         GP         LM         N         [10]		Mormoniella (Nasonia) vitripennis	Blastoderm	Pole cells	GP	LM		[100-1
Trichogramme evanescens Blastoderm Pole cells GP LM N [106, 7 inteptis dipronis Blastoderm Pole cells GP LM N [10]		Pimpla turionellae	Blastoderm	Pole cells	GP	LM	*+*	[103, 10
Trichogramme evanescens Blastoderm Pole cells GP LM N [106, 1 Tritneptis dipronis Blastoderm Pole cells GP LM N [10]		Pteronidea ribesii	Germ band	PGB	MZ	LM	N	[105
Tritneptis dipronis Blastoderm Pole cells GP LM N [10]		Trichogramma evanescens	Blastoderm	Pole cells	GP	LM	N	[106, 10
		Tritneptis dipronis	Blastoderm	Pole cells	GP	LM	N	[108
	iptera							
Acyrthosiphon pisum Blastoderm Pole cells GP LM, MM N [109-1								[109-1

### **Supplementary Table 1 (Continued)**

	Aphis pelargonii	Blastoderm	PGB	MZ	LM	N	[114]
	Aphis plantoides	Blastoderm	PGB	MZ	LM	N	[115]
	Aphis rosea	Blastoderm	PGB	MZ	LM	N	[33]
	Oncopeltus fasciatus	Blastoderm	PGB	MZ	LM, TEM, MM	Y	[116-118]
	Pyrrhocoris apterus	Blastoderm	PGB	MZ	LM	N	[119]
	Rhodnius prolixus	Blastoderm	PGB	GP	LM	N	[120, 121]
Thysanoptera							
	Haplothrips verbasci	Blastoderm	Pole cells	GP	LM	N	[122]
socoptera							
	Liposcelis divergens	Blastoderm	PV blastoderm	MZ	LM	N	[123]
Phthiraptera							
		Blastoderm	Pole cells	GP	LM	N	[cited in 122,
Blattodea							124]
	Periplaneta orientalis	Germ hand	Mesoderm	M7	LM	N	[125]
	Phyllodromia (Blatta) germanica	Germ band	Mesoderm	MZ	LM	N	[125]
Plecoptera	r nyilodromia (biatta) germanica	Geriii baliu	Wesouchin	IVIZ	LM	14	[123]
recoptera	Pteronarcys proteus	Germ hand	Mesoderm	M7	LM	N	[126]
Phasmatodea	r teronarcys proteus	Geriii baliu	Wesouth	IVIZ	LM	14	[126]
riiasiliatouea	Carausius morosus	Germ band	Mesoderm	MZ	LM. TEM	Y	[127-130]
Orthoptera	Carausius morosus	Geriii banu	Wesouth	IVIZ	LW, TLW		[127-130]
Orthoptera	Gryllus bimaculatus	Germ hand	Mesoderm	M7	LM. MM	Y	This study
	Gryllus campestris	Germ band	PGB	MZ	LM.	N N	[125]
	Gryllus domesticus	Germ band	PGB	MZ	LM	N	[125]
		Germ band	CP Mesoderm	MZ	LM	N	[131]
	Locusta migratoria	Germ band	Lateral abdominal ectoderm	MZ	LM	N	[131]
	Melanoplus differentialis  Xiphidium enisterum	Germ band Germ band	Mesoderm	MZ	LM	N	[133]
Zygentoma	Alphidiam enisteram	Geriii banu	Mesodellii	IVIZ	LM	N N	[133]
Lygentoma			0044				[40.4]
	Ctenolepisma lineata	Germ band	CP Mesoderm	MZ	LM	N	[134]
	Lepisma saccharina	Germ band	PGB CP Mesoderm	MZ	LM TEM	N	[135]
	Thermobia domestica	Germ band	CP Mesoderm	MZ	LM, TEM	N	[134, 136]
Archaeognatha							
	Pedetontus unimaculatus	Blastoderm	PGB	MZ	LM	N	[137]
	Petrobius brevistylis	Germ band	CP Mesoderm	MZ	TEM	N	[138]

 $\begin{tabular}{l} [1] Species name is shown as reported in the primary data reference, listed alphabetically within an insect order. \end{tabular}$ 

(1) Species name is shown as regorded in the primary data reference, listed alphabetically within an insect order.

[2] M2 = mesodermal/papotic origin, GP = perm plasm/poole cells

[3] M3 = species in the processor is biotological analysis, of either whole mounts or sections, TTM = transmission electron microscopy, STM = scanning electron microscopy. [M = enzymatic markers, LM = molecular markers, LB = cell ineage studies.

[4] X = yes, X = no, included state derived from functional genetic evidence and/or physical perturbation/abbitton.

[3] Since direct comparison of the duration of states of development in different species is not appropriate due to differences in culture conditions, we describe here developmental stages rather than absolute time.

[3] Find cells | O = permission of the duration of states of development most processor in the literature.

[3] Ind. no date, for Comma abbunctats pole cells have been specifically noted as absent, but no hypotheses on PGC origin have been reposed in the literature.

[3] Conformed information in the interaction processor in the transport of processor in the processor in the processor in the literature.

[9] Copidosoma floridanum is unusual among insects in that it displays holoblastic cleavage rather than syncytial cleavage in early development. B4 is the name given to the small blastomere formed at second cleavage, which inherits the cosome (germ plasm) and is the primordial germ cell.

# Evidence against a germ plasm in the milkweed bug Oncopeltus fasciatus, a hemimetabolous insect

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#### Summary

Primordial germ cell (PGC) formation in holometabolous insects like Drosophila melanogaster relies on maternally synthesised germ cell determinants that are asymmetrically localised to the oocyte posterior cortex. Embryonic nuclei that inherit this "germ plasm" acquire PGC fate. In contrast, historical studies of basally branching insects (Hemimetabola) suggest that a maternal requirement for germ line genes in PGC specification may be a derived character confined principally to Holometabola. However, there have been remarkably few investigations of germ line gene expression and function in hemimetabolous insects. Here we characterise PGC formation in the milkweed bug Oncopeltus fasciatus, a member of the sister group to Holometabola, thus providing an important evolutionary comparison to members of this clade. We examine the transcript distribution of orthologues of 19 Drosophila germ cell and/or germ plasm marker genes, and show that none of them localise asymmetrically within Oncopeltus oocytes or

early embryos. Using multiple molecular and cytological criteria, we provide evidence that PGCs form after cellularisation at the site of gastrulation. Functional studies of vasa and tudor reveal that these genes are not required for germ cell formation, but that vasa is required in adult males for spermatogenesis. Taken together, our results provide evidence that Oncopeltus germ cells may form in the absence of germ plasm, consistent with the hypothesis that germ plasm is a derived strategy of germ cell specification in insects

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Key words: vasa, tudor, boule, Germ line, RNA interference, Spermatogenesis

#### Introduction

In sexually reproducing animals, only germ cells contribute genetic information to future generations. The germ line/soma separation is a cell fate decision shared across Metazoa (Buss, 1987). Despite the fundamental commonality of germ cell function in animals, the molecular mechanisms underlying germ cell specification are remarkably diverse across different taxa (Extavour and Akam, 2003; Extavour, 2007; Ewen-Campen et al., 2010; Juliano et al., 2010).

Primordial germ cells (PGCs) can be specified via different developmental mechanisms; here we call these "cytoplasmic inheritance" and "zygotic induction." (We and others have previously referred to these mechanisms as "preformation" and "epigenesis" respectively (Nieuwkoop and Sutasurya, 1981; Extavour and Akam, 2003; Extavour, 2007). However, these terms can hold different meanings in other contexts of the history and philosophy of biology (e.g. Callebaut, 2008). We therefore avoid them here in favour of more mechanistically descriptive terms.) Cytoplasmic inheritance is characterised by the asymmetric formation of a specialised cytoplasmic region within the oocyte or early embryo, termed "germ plasm." Germ plasm contains maternally provided mRNAs and proteins that are individually necessary and collectively sufficient for PGC formation. Cells that inherit germ plasm during embryogenesis acquire germ line fate. The best understood

example of cytoplasmic inheritance occurs in *Drosophila melanogaster*, where germ plasm is maternally synthesised, localised to the posterior of the oocyte during oogenesis, and subsequently incorporated into PGCs (pole cells) during cellularisation. Removing pole cells after their formation, or compromising the molecular components of germ plasm, leads to loss of PGCs and sterility in adulthood (reviewed by Mahowald, 2001). In contrast, zygotic induction of PGCs takes place later in development and requires signalling from neighbouring somatic cells to induce germ line fate. This mode of PGC development is exemplified by *Mus musculus*, wherein PGCs develop from a subset of presumptive mesodermal cells after the segregation of embryonic and extraembryonic tissues in response to local signalling (reviewed by Magnúsdóttir et al., 2012).

Across Insecta, germ plasm has been almost exclusively reported in taxa nested within Holometabola ("higher" insects, which undergo complete metamorphosis) including *D. melanogaster* (reviewed by Kumé and Dan, 1968; Anderson, 1973; Nieuwkoop and Sutasurya, 1981), and in only three species belonging to the sister assemblage to the Holometabola (see below). Thus, although the vast majority of our knowledge of insect germ cell development comes from studies of germ plasm in *D. melanogaster*, this mode of germ cell specification is likely a derived feature of Holometabolous insects and their close sister taxa.

Our present knowledge of PGC specification in basally branching insects (Hemimetabola) is based almost entirely on classical histological studies of insect development conducted over the past 150 years. Nearly all of these report that PGCs arise late in embryogenesis, raising the possibility that they may be specified through inductive mechanisms (Wheeler, 1893; Heymons, 1895; Hegner, 1914; Nelsen, 1934; Roonwal, 1937). Experimental approaches to discovering germ plasm in Hemimetabola are limited, but a study involving destruction of the germ rudiment via irradiation in the cricket Gryllus domesticus (Schwalm, 1965) showed that no specific region of early embryos in this species contains a germ line determinant. Functional tests of genes that may specify germ cells in Hemimetabola have been performed in only one insect, the cricket Gryllus bimaculatus. In this cricket, the conserved germ line markers vasa and piwi are dispensable maternally and zygotically for PGC formation (Ewen-Campen et al., 2013). Most evidence available for the Hemimetabola therefore suggests the absence of germ plasm and the operation of zygotic PGC specification mechanisms.

Exceptions have been reported, however, in some members of the Paraneoptera, an assemblage of insect orders (including Hemiptera [true bugs], Psocoptera [book lice], and Thysanoptera [thrips]) that collectively form the sister group to Holometabola (Yeates et al., 2012). Cytological studies of three paraneopteran species, a book louse (Psocoptera (Goss, 1952)), a thrip (Thysanoptera (Heming, 1979)) and an aphid (Hemiptera (Chang et al., 2009)) suggested the presence of germ plasm in oocytes or early embryos, as did expression studies of vasa, piwi and nanos expression during asexual development of the pea aphid Acyrthosiphon pisum (Chang et al., 2006; Chang et al., 2007; Chang et al., 2009; Lu et al., 2011). However, A. pisum embryogenesis is highly modified relative to that of other hemimetabolous insects and even relative to other members of the same order (Miura et al., 2003). Studies of embryogenesis in most other hemipterans describe absence of germ plasm and PGC origin after cellularisation from the blastopore region at gastrulation stages (Metschnikoff, 1866; Witlaczil, 1884; Will, 1888; Seidel, 1924; Mellanby, 1935; Butt, 1949; Sander, 1956; Kelly and Huebner, 1989; Heming and Huebner, 1994). We therefore wished to examine the expression and function of germ line genes in a hemipteran displaying embryological characteristics more representative of the order.

Here we characterise germ cell formation and migration in the milkweed bug *Oncopeltus fasciatus* (Hemiptera). We examine the expression of 19 molecular markers including *vasa*, *nanos*, and *piwi*, and test the germ cell function of three of these using RNA interference. We show that in striking contrast to *Drosophila*, transcripts of none of these genes localise asymmetrically within *Oncopeltus* oocytes or early embryos. We identify PGCs using multiple criteria, and show that neither *vasa* nor *tudor* are required for PGC specification or oogenesis in this species, but that *vasa* is required for spermatogenesis in adult males. These data show that the PGC specification role of *vasa* has diverged between *Oncopeltus* and the Holometabola, and suggest that *Oncopeltus* PGCs may form in the absence of maternally supplied germ plasm.

#### Results

Putative germ cells are first detectable in the late blastoderm stage

In contrast to *D. melanogaster*, classical studies of *Oncopeltus fasciatus* embryogenesis have not revealed a germ plasm in

oocytes or early embryos, and instead first identify cells with cytological characteristics of PGCs at the posterior of the embryo at the end of the cellular blastoderm stage (Butt, 1949). We used semithin plastic sectioning and fluorescence microscopy to confirm these observations, and traced the development of these putative PGCs throughout gastrulation and germ band elongation (supplementary material Fig. S1). Our observations of putative PGC formation in Oncopeltus were consistent with historical studies (Butt, 1949), showing that these cells first arise at the blastoderm posterior immediately prior to gastrulation (supplementary material Fig. S1). Unlike pole cells in D. melanogaster, presumptive PGCs in Oncopeltus arise on the basal side of the blastoderm surface, adjacent to the yolk (supplementary material Fig. S1G-H). In order to obtain further evidence that these cells were PGCs and test for the presence of a maternally supplied germ plasm, we examined the expression of conserved germ line markers.

#### Cloning Oncopeltus germ line markers

We cloned fragments of *vasa*, *nanos*, and *piwi* (Ewen-Campen et al., 2010) and confirmed that each was the best reciprocal BLAST hit to its respective orthologue in *D. melanogaster. vasa* was cloned using degenerate primers (supplementary material Table S1). *nanos* and a single *piwi* gene were recovered from the *Oncopeltus* transcriptome, in addition to a single AGO-3 orthologue (an additional PIWI family protein belonging to a separate sub-family; not shown). We believe it is unlikely that *Oncopeltus* possesses additional orthologues of these genes because (1) the *Oncopeltus* ovarian and embryonic transcriptome, which is nearly saturated for gene discovery and has an average coverage of 23× (Ewen-Campen et al., 2011), contained only one orthologue of each gene; and (2) degenerate PCR for *vasa* using primers flanking the conserved DEAD box helicase domain (Rocak and Linder, 2004) recovered only a single *vasa* orthologue.

Phylogenetic reconstruction confirmed that *Oncopeltus vasa* is nested within other insect *vasa* genes (supplementary material Fig. S2A), and that *Oncopeltus piwi* belongs to the PIWI sub-family containing the *Drosophila* genes *piwi* and *aubergine* (which are *Drosophila*-specific duplications) (supplementary material Fig. S2B). The portion of animal Nanos proteins with conservation sufficient for confident alignment (48 amino acids) is too short to yield significant phylogenetic signal (supplementary material Fig. S2C, note low support values), but *Oncopeltus* Nanos does contain the diagnostic 2×(CCHC) zinc finger domain found in all Nanos orthologs (supplementary material Fig. S2D).

Our analysis of the Oncopeltus nanos sequence produced an unexpected result: we found that a stop codon is present 771 bp upstream of the first CCHC zinc finger domain, although no methionine is found anywhere in this region. This is unlikely to be a sequencing error, as it was identified with high coverage (22 reads/bp at this position) in the transcriptome (Ewen-Campen et al., 2011) and confirmed using Sanger sequencing of independent clones generated from a different cDNA pool than that used to generate the transcriptome. Furthermore, repeated attempts at 5' RACE using a third independent DNA pool failed to amplify a start codon. Several lines of evidence confirm that this Oncopeltus nanos sequence represents a highly expressed mRNA and is therefore unlikely to be a pseudogene: it was recovered from a transcriptome made solely from poly(A)-RNA, and is detected via both RT-PCR (Ewen-Campen et al., 2011) and in situ hybridisation (see below). We hypothesise that a large, unspliced intron downstream of the start codon may have been

present in our mRNA preparations. Alternatively, given that the length of the predicted translated region upstream of the first CCHC zinc finger domain (266 amino acids) is within the range of known arthropod Nanos orthologues (95 to 332 amino acids) (Wang and Lehmann, 1991; Curtis et al., 1995; Calvo et al., 2005; Lynch and Desplan, 2010), it may be that the *Oncopeltus* Nanos N terminus has a non-methionine start codon. Although rare, eukaryotic non-AUG translation initiation can occur in nuclear-encoded genes, including developmentally relevant genes (Hellen and Sarnow, 2001), and can be recognized by insect ribosomes (Sasaki and Nakashima, 2000; Jan et al., 2001). In the absence of a complete genome sequence we cannot distinguish between these hypotheses. Despite this uncertainty, we report *nanos* transcript expression here for the sake of completeness.

## vasa, nanos, and piwi transcripts do not localise asymmetrically in ovaries

The distinct cytoplasm inherited by early-specified PGCs in multiple organisms, including *D. melanogaster*, *Caenorhabditis elegans*, *Xenopus laevis* and *Danio rerio*, contains transcripts of the highly conserved *piwi*, *vasa* and *nanos* gene families. Of these, only *nanos* mRNA is asymmetrically localised to *D. melanogaster* germ plasm, while *piwi* and *vasa* transcripts are ubiquitous throughout the fly oocyte and embryo. However, in several other organisms *vasa* orthologue transcripts are asymmetrically localised germ plasm components (reviewed by Ewen-Campen et al., 2010).

To test whether any of these transcripts were asymmetrically localised to putative germ plasm in Oncopeltus oocytes, we conducted in situ hybridisation on adult ovaries. The structure of Oncopeltus ovaries is typical of Hemiptera and several other insect orders but differs remarkably from that of Drosophila (Fig. 1A) (Büning, 1994). Rather than each oocyte developing together with its own complement of 15 nurse cells as in Drosophila, all oocytes in Oncopeltus ovarioles share a common pool of syncytial nurse cells located at the anterior of each ovariole in a region termed the "tropharium" (Fig. 1A1). The nurse cell syncytium connects to all oocytes via elongated, microtubule-rich tubes called "nutritive tubes" (Hyams and Stebbings, 1979; Harrison et al., 1991) through which maternal factors, including mRNA, proteins and mitochondria, are transported to developing oocytes (Fig. 1A2,A3) (Stebbings et al., 1985; Stebbings and Hunt, 1987; Anastas et al., 1991; Hurst et al., 1999; Stephen et al., 1999).

vasa, nanos, and piwi were expressed at high levels in Oncopeltus nurse cells and oocytes of all stages, but at no stage of oogenesis did any of these three transcripts localise asymmetrically within oocytes (Fig. 1B–D). Expression was detected in nurse cells, resting oocytes, nutritive tubes, and developing oocytes, suggesting that these transcripts are synthesised in the nurse cells and subsequently transported to oocytes via nutritive tubes (Fig. 1B–D). nanos and piwi were expressed throughout the tropharium (Fig. 1C,D), in contrast to vasa, whose expression was primarily in nurse cells of the posterior tropharium, resting and developing oocytes (Fig. 1B). In late stage oocytes, expression remained ubiquitous (not shown), similar to the expression in just-laid eggs (see below).

# In situ screen of conserved *Drosophila* germ plasm markers fails to reveal a germ plasm in *Oncopeltus*

The expression of piwi, vasa and nanos suggests that a maternally localised germ plasm containing transcripts of these genes is not

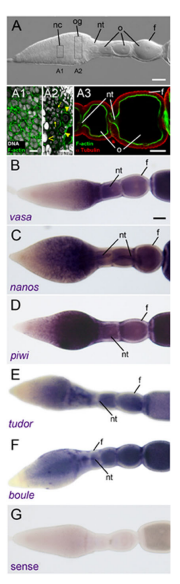


Fig. 1. Germ cell markers do not localise asymmetrically during oogenesis.

(A) Overview of a single Oncopeltus ovariole. nc: nurse cells; o: oocytes; nt: nutritive tubes; f: follicle cells. Boxed regions are enlarged in (A1–A3). (A1) Nurse cell syncytium containing polyploid nurse cell nuclei (white) connected by cytoplasmic bridges (green). (A2) Posterior tropharium containing oogonia (arrows) and resting oocytes (arrowheads). Caret indicates polyploid nurse cells in the anterior of this region. (A3) Nutritive tubes (nt) are actin-rich at the end that enters the anterior of each oocyte. Transcripts of vasa (B), nanos (C), pivi (D), tudor (E) and boule (F) are detected in nurse cells, nutritive tubes, and uniformly in oocytes. (G) A representative sense control (for vasa) is shown; sense controls for other genes were similar. Scale bars: 100 µm in A,A3,B (applies to C–G): 25 µm in A1.A2. Anterior to the left in all panels.

present in *Oncopeltus* oocytes. However, a functional germ plasm that contains gene products other than those encoded by these three genes could be present in oocytes or early embryos. To

explore this possibility, we examined the expression of 14 additional genes whose transcripts are enriched in the germ plasm and germ cells of *Drosophila* (supplementary material Table S2) (Tomancak et al., 2002; Lécuyer et al., 2007; Tomancak et al., 2007) that were also recovered from the *Oncopeltus* ovarian and embryonic transcriptome (Ewen-Campen et al., 2011) based on best reciprocal BLAST hit analysis with the *Drosophila* proteome (Zeng and Extavour, 2012). Although several of these genes do not have documented mutant phenotypes for germ cell formation in *Drosophila* (supplementary material Table S2), all are expressed at high levels in germ plasm and/or pole cells and are therefore molecular markers for germ plasm in *Drosophila*. We reasoned that if *Oncopeltus* possessed germ plasm it would likely be revealed by at least one of these genes.

In addition, we examined the expression of *boule* and *tudor*, which have widely conserved functions in germ cells across Metazoa (Eberhart et al., 1996; Ewen-Campen et al., 2010; Shah et al., 2010). *tudor* is one of 23 Tudor domain-containing proteins

in *Drosophila* (Ying and Chen, 2012), but there is no evidence that loss of function of other Tudor domain-containing genes have grandchildless phenotypes in *Drosophila* (Handler et al., 2011; Pek et al., 2012). We therefore focus only on the expression and function of the orthologue of *Drosophila tudor* (CG9450). We examined *boule* and *tudor* transcript expression throughout oogenesis and embryogenesis through mid-germ band stages.

None of these 16 transcripts localised asymmetrically in ovaries (Fig. 1E,F; supplementary material Fig. S3). Instead, like vasa, piwi and nanos (Fig. 1B–D), all of these genes were expressed ubiquitously throughout oogenesis. Half of the genes examined (sra, CycB, Bsg25D, Uev1A, CG16817, Unr, mael and tud) were expressed, like vasa (Fig. 1B), in nurse cells adjacent to resting oocytes, as well as in the resting and early oocytes themselves (Fig. 2E; supplementary material Fig. S3B–H). Five genes (Gap1, elF5, bel, orb and bol) were, like piwi and nanos (Fig. 1C,D), strongly expressed in all nurse cells of the

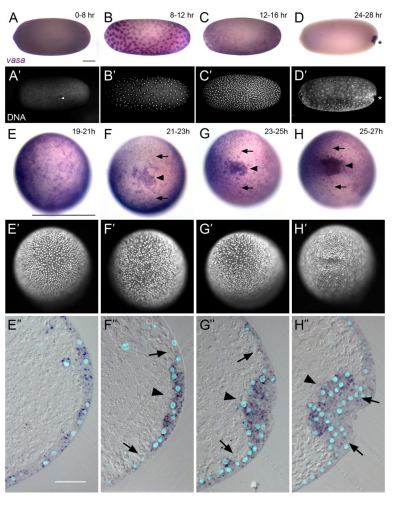


Fig. 2. vasa transcript expression first labels PGCs at late blastoderm stages. (A-D) vasa transcript expression. (A'-D') Corresponding images of nuclear stains. (A,A') Immediately following fertilisation vasa is detected ubiquitously. Arrowhead: polar body.  $(B,\!B^\prime)$  In early cleavage stages vasa transcripts are associated with all energid nuclei. (C,C') During early syncytial blastoderm stages, vasa expression remains ubiquitous. (D,D') At cellular blastoderm stages (24-28 h AEL), vasa marks putative PGCs at the posterior pit (asterisk). (E-H) End-on perspective of the posterior of Oncopeltus embryos showing vasa expression during PGC formation. (E'-H') Corresponding images of nuclear stains. (E"-H") Medial sections of vasa-(purple) and nuclear- (cyan) stained embryos at corresponding time points. (E-E") In late syncytial blastoderm stages, vasa is expressed ubiquitously. (F-F") In early cellular blastoderm embryos, *vasa* expression increases in some posterior cells (arrowheads in F,F") while levels in the remainder of the blastoderm decrease (arrows in F,F"). (G-G") At posterior germ band invagination vasa-positive cells (arrowheads) are the first cells to enter the yolk; vasa transcripts continue to be cleared from somatic tissue (arrows). (H-H") As invagination proceeds vasa expression is largely restricted to PGCs (arrowhead) and nearly cleared from the soma (arrows). Scale bars: 100 µm in A (applies to B-D,A'-D'); 500 μm in E (applies also to F-H'); 50 μm in E" (applies also to F"-H"). Anterior is to the left in A-D' and E"-H".

tropharium (Fig. 2F; supplementary material Fig. S3I–L). Two genes (*cta* and *Tao*) were expressed in resting and early oocytes but barely at all in the tropharium (supplementary material Fig. S3M,N), suggesting that these genes may be transcribed by resting oocyte nuclei rather than by nurse cells. Finally, *aret* (aka *bruno*), which is a translational regulator of Oskar in *Drosophila* (Kim-Ha et al., 1995; Webster et al., 1997), was expressed in nurse cells of the posterior tropharium and in early stages of oogenesis but excluded from resting oocytes (supplementary material Fig. S3O), suggesting that it is transcribed by oocyte nuclei after the onset of oogenesis. In summary, although transcripts of most of these genes are likely to be supplied maternally to oocytes, they are not asymmetrically localised within oocytes of any stage.

# vasa, boule and tudor transcripts mark PGCs throughout embryogenesis but are not asymmetrically localised in early embryos

Although none of the genes examined showed asymmetric localisation during oogenesis or early embryogenesis, at late blastoderm stages many of the genes appeared enriched at the posterior pit, where PGCs had previously been identified based on cytological criteria (supplementary material Fig. S1) (Butt, 1949). However, because at this stage of development gastrulation begins at the posterior, this region of the blastoderm is multilayered. Upon close examination, we found that the apparent transcript enrichment was an artifact of tissue thickness for all genes except *vasa*, *tudor* and *boule*, whose transcripts appeared truly enriched in putative PGCs at late blastoderm/early gastrulation stages (Fig. 2D–H", Fig. 4N,S).

Strikingly, we found that vasa, tudor and boule marked PGCs from the time of their formation at cellular blastoderm stages, but

that none of these genes' transcripts were asymmetrically localised prior to PGC formation. Immediately after egg laying, vasa transcripts were not localised asymmetrically but rather were ubiquitously distributed throughout the embryo (Fig. 2A). As energid nuclei reached the embryonic surface (Fig. 2B), cytoplasmic islands enriched with these transcripts were distributed evenly across the embryonic surface, remaining there as these energids divided to form the uniform blastoderm (Fig. 2C). Prior to posterior pit formation, vasa expression became restricted to putative PGCs at the embryonic posterior (Fig. 2D).

To visualise *vasa* expression in the developing PGCs in greater detail, we collected staged embryos in two-hour intervals over the period during which PGCs arise (19 to 27 hours after egg laying (AEL)), performed in situ hybridisation for vasa (Fig. 2E-H), and sectioned the embryos in plastic resin (Fig. 2E"-H"). During this eight-hour period, the blastoderm nuclei undergo two concurrent, dynamic processes: continuing cell divisions increase the nuclear density throughout the blastoderm, and the blastoderm nuclei move towards the posterior pole and ultimately into the yolk (Butt, 1949; Liu and Kaufman, 2004) (Fig. 2E'-H'). From 19-21 hours AEL, the ubiquitous vasa expression seen in early embryos remained unchanged (Fig. 2E-E"). However, from 21-23 hours AEL vasa expression became enriched in a subset of cells at the blastoderm posterior (Fig. 2F-F"). From 23-25 hours AEL, vasa-positive cells increased in density at the blastoderm posterior and began to move into the yolk (Fig. 2G-G"). This movement appeared passive, due to the formation of the posterior pit by invagination of the germ rudiment. However, in the absence of time-lapse data we cannot rule out the possibility of active PGC movement out of the blastoderm epithelium and

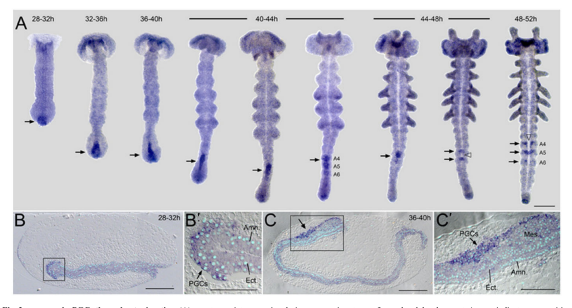


Fig. 3. vasa marks PGCs throughout migration. (A) vasa transcript expression during progressive stages of germ band development. Arrows indicate vasa-positive PGCs. (B) Medial section of an embryo at 28–32 h AEL, showing vasa in situ hybridisation (purple) and nuclear stain (cyan). Boxed region enlarged in (B') shows PGCs in contact with ectoderm (Ect.) and the amnion (Amn.). (C) Medial section of an embryo at 36–40 h AEL, when PGCs (arrow) iritiate migration along the mesoderm (Mes.). Boxed region enlarged in (C'). Scale bars: 200 μm in A-C; 100 μm in B',C'. Anterior is up in A, left in B-C'.

towards the yolk. From 25–27 hours AEL, as the germ rudiment began its invagination into the yolk, *vasa*-positive cells formed a distinct mesenchymal clump within the yolk at the posterior of the embryo (Fig. 2H–H"). During this and all following stages, in addition to the marked enrichment in PGCs, *vasa* transcripts were additionally observed ubiquitously at low levels throughout somatic tissue (Figs 2, 3).

Throughout all subsequent stages of germ band elongation and patterning, vasa continued to mark PGCs (Fig. 3). During early stages of germ band elongation prior to limb bud formation (~28-32 hours AEL) vasa-positive PGCs remained at the embryonic posterior on the dorsal surface of the newly forming mesoderm (Fig. 3A,B,B'). The PGC cluster then became pearshaped from 32-42 hours AEL, as the anteriormost PGCs began to move towards the anterior of the embryo (Fig. 3A,C,C'). As the head lobes enlarged (36-40 h AEL), PGCs began to migrate anteriorly on the dorsal surface of the embryo and continued their migration during limb bud stages (40-44 h AEL) (Fig. 3A). During appendage elongation stages (44-48 h AEL) PGCs split into distinct clusters spanning the midline in abdominal segments A4-A6, one cluster per segment. As appendage segmentation became morphologically distinct (48-52 h AEL), the segmental clusters split along the ventral midline into bilateral clusters in

tudor and boule were also expressed in PGCs at all stages in a pattern indistinguishable from that of vasa (Fig. 4), providing further evidence that the vasa-positive cells are Oncopeltus PGCs. None of the other genes we examined (supplementary material Fig. S3), including nanos and piwi (Fig. 4), were enriched in PGCs at any stage of embryogenesis.

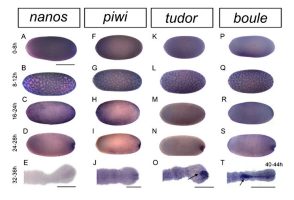


Fig. 4. Oncopeltus PGCs express tudor and boule, but not nanos or piwi. In early embryos, expression of all four genes remains ubiquitous during energid proliferation (A,F,K,P) and blastoderm formation (B,C,G,H,L,M,Q,R). During posterior pit formation nanos is expressed throughout the length of the embryo (D), whereas piwi expression is reduced in the presumptive extraembryonic serosal tissue in the anterior of the embryo (I). Apparent posterior staining in (D) and (I) is the result of tissue thickness in that location, and is not specific to PGCs. tudor (N) and boule (S) transcripts become restricted to presumptive PGCs at the time of their specification. In germ band stage embryos, while tudor (O) and boule (T) mark presumptive PGC clusters, nanos is not detected (E), while piwi expression is ubiquitous (J). Scale bars: 500 µm in A (applies also to B–D,F–I,K–N,P–S); 100 µm in E,J; 200 µm in O,T. Anterior to the left.

Neither vasa nor tudor are required for PGC formation

Our gene expression analysis demonstrates that *vasa*, *tudor*, and *boule* are specifically expressed in PGCs beginning at the putative time of their specification at the embryonic posterior just prior to gastrulation. To determine whether these genes were required for PGC formation or development in *Oncopeltus*, we performed maternal RNAi (mRNAi) for each gene. We confirmed that mRNAi effectively reduced zygotic transcript levels in our experiments using RT-PCR (Fig. 5E). PGC presence or absence was determined with in situ hybridisation against PGC markers at ~40–54 hours AEL, when germ cells are visible on the dorsal mesoderm.

RNAi knockdown of vasa or tudor did not disrupt embryonic patterning or germ band development (supplementary material Table S3), despite the widespread expression of these genes at early blastoderm stages (Figs 2, 4), and their persistent low levels of expression in somatic cells even after PGC formation (Figs 3) 4). Strikingly, germ cells were clearly present in both vasa (93.8%, n=16) and tudor (100%, n=20) knockdowns, suggesting that neither of these genes is required for PGC specification (Fig. 5A-C'). It is formally possible that residual vasa or tudor transcripts that may have escaped destruction by mRNAi could be sufficient to play an instructive role in PGC formation. However, we note that transcript levels of both genes in the progeny of injected mothers were barely detectable in the case of vasa, and undetectable in the case of tudor, when assessed with RT-PCR even as late as 4 days AEL (Fig. 5E). Moreover, even hypomorphic alleles of tudor (Schüpbach and Wieschaus, 1986) and vasa (Lasko and Ashburner, 1990; Schüpbach and Wieschaus, 1991; Liang et al., 1994) lead to loss of PGCs in Drosophila. We therefore hypothesise that in Oncopeltus, vasa and tudor are required neither maternally nor zygotically for germ cell specification, although they are expressed in the cells specified as PGCs.

To address the possibility of redundancy between these two genes, we performed double knockdown of *vasa* and *tudor*, which reduced transcripts of both genes to undetectable levels (Fig. 5E). Eggs laid by *vasa* + *tudor* double RNAi females had an increased rate of embryonic lethality relative to controls (supplementary material Table S3; 47.4%, *n*=19 vs 10.3%, *n*=39), which may mean that these genes work together to play roles in somatic development. However, embryos that escaped this lethality still had PGCs (100%, *n*=10) (Fig. 5D,D').

None of the knockdowns caused any qualitative or quantitative change in egg laying by injected females compared to controls, and ovaries of injected females showed neither morphological abnormalities nor signs of disrupted oogenesis (not shown). This indicates that, in contrast to *Drosophila* (Schüpbach and Wieschaus, 1991; Styhler et al., 1998; Tomancak et al., 1998; Johnstone and Lasko, 2004), *vasa* is not required individually or redundantly with *tudor* for *Oncopeltus* oogenesis or egg laying.

# boule is necessary for *Oncopeltus* oogenesis and embryonic survival

boule mRNAi caused a complete cessation of egg laying by injected females after four to five clutches (one clutch is laid every one to two days). In contrast, vasa, tudor and control mRNAi females continued to lay up to 12 clutches. Ovaries of boule dsRNA-injected females possessed only a few oocytes at early stages of oogenesis, and few or no mature oocytes (not shown), indicating a requirement for boule in the progression of

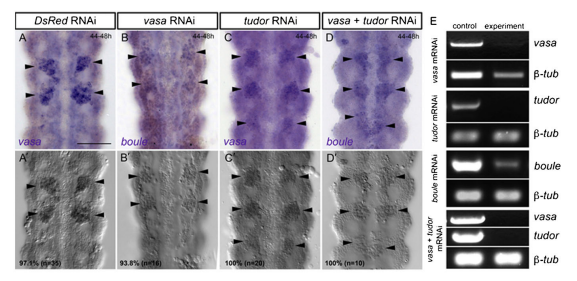


Fig. 5. vasa and tudor are not required for PGC specification in Oncopeltus. (A–D) Bright field images of in situ hybridisations for PGC markers in different RNAi conditions; numbers indicate sample sizes and % of embryos with PGCs. Arrowheads indicate PGC clusters in abdominal segments A4–A6. (A′–D′) DIC images of the same embryos shown in (A–D) showing distinct PGC cluster morphology. (A,A′) In control embryos vasar-positive PGCs are visible on the dorsal surface of abdominal segments 4–5. PGCs are present in vasa RNAi (B,B′), tudor RNAi (C,C′), and double vasa + tudor RNAi (D,D′) embryos. (E) RT-PCR validation of RNAi knockdown. Controls are animals injected with DsRed dsRNA. Expression of β-tubulin was analysed to confirm cDNA integrity and allow comparison of amounts of template per lane. Scale bar: 100 μm. Anterior is up in A–D.

oogenesis. Eggs laid by boule RNAi females displayed nearly complete embryonic lethality (81.8%, n=22) in all but the first clutch laid. (The first clutch of Oncopeltus eggs laid following mRNAi typically displays no abnormalities, as these eggs have developed their chorion by the time of injection and are therefore impervious to dsRNA (Liu and Kaufman, 2004).) This was a striking increase in embryonic lethality compared to DsRed controls (26.8%, n=190), vasa knockdowns (23.2%, n=198) and vudor knockdowns (5.6%, n=54). The oogenesis requirement for vudor boule and resulting embryonic lethality thus prevented us from determining whether vudor boule is required for germ cell specification in vudor vudor

#### vasa is required for Oncopeltus spermatogenesis

Given that in contrast to *Drosophila*, vasa is not required for germ cell specification or oogenesis in *Oncopeltus*, we wished to test for other possible functions of this gene. In mice, despite its expression in the embryonic PGCs of both sexes once they reach the genital ridge (Fujiwara et al., 1994; Diez-Roux et al., 2011). vasa is required not for PGC specification, but rather for gametogenesis in males (Tanaka et al., 2000). Similarly, we recently showed that vasa plays a role in spermatogenesis in the cricket *G. bimaculatus* (Ewen-Campen et al., 2013). We therefore asked whether vasa also functions during spermatogenesis in *Oncopeltus*.

The testes of *Oncopeltus* show an organisation typical of insect testes (Dumser, 1980), with stages of spermatogenesis located in an anterior–posterior progression (supplementary material Fig. S4). Unlike *Drosophila*, which has a single sperm tubule (testiole) per testis (Hardy et al., 1979), each *Oncopeltus* testis

comprises seven testioles (Bonhag and Wick, 1953). In situ hybridisation for vasa showed that it is strongly expressed in secondary spermatogonia of each testiole, and at lower levels in early primary spermatocytes and post-spermatocyte stages, but not in primary spermatogonia or somatic cells (Fig. 6A). Adult males injected with dsRNA against vasa displayed multiple abnormalities in spermatogenesis. Testioles of vasa RNAi males lacked clearly defined cysts and contained large numbers of small, dense nuclei in the anterior region (Fig. 6H,I,I'), which in controls contained only spermatocytes with large, pale nuclei (Fig. 6C,D,D'; supplementary material Fig. S4E). The primary spermatogonial region of vasa RNAi testioles contained cysts of irregular size (Fig. 6I, arrowheads) with poorly defined cytoplasmic bridges (Fig. 6I', arrows). In the spermatocyte region vasa RNAi testioles contained large, poorly defined clusters of several hundred cells (Fig. 6J, arrowheads) at varying stages of spermatogenesis (Fig. 6J'). The nuclear morphology of cells in these cysts corresponded to spermatocyte (Fig. 6E; supplementary material Fig. S4E) or early spermatid (Fig. 6F) stages, as well as shell stage-like nuclei (Fig. 6K,K') typical of the mid-stage spermatids of controls (Fig. 6F'). Cysts of wild type shell stage spermatids are no longer syncytial as the actinrich cytoplasmic bridges disappear during spermatocyte stages (supplementary material Fig. S4G). In contrast, the anterior shell stage-like nuclei in vasa RNAi testioles remained connected by cytoplasmic bridges (Fig. 6K', red arrows), consistent with precocious spermatid differentiation. Moreover, although they displayed clear shell stage nuclear morphology (Fig. 6K,K', red arrowheads), they were larger than wild type shell stage nuclei (Fig. 6F', red arrowheads), suggesting that they had begun spermatid differentiation as syncytial diploid cells without first

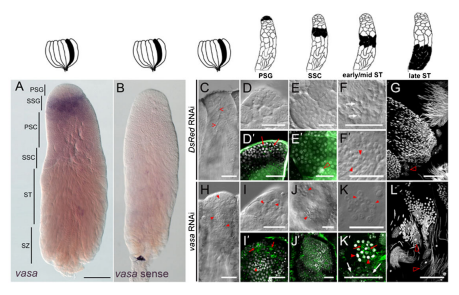


Fig. 6. vasa is expressed in adult testes and required for spermatogenesis in Oncopellus. Schematics indicate the region of the testis (A–C) or testiole (D–L) shown in each column. (A) vasa in situ of an adult vid-type testiole showing expression in the secondary spermatogonia. PSC: primary spermatogonia; SSC: secondary spermatogenia; PSC: primary spermatocytes; SSC: secondary spermatogonia psermiogenesis; SZ: spermatozoa. (B) vasa sense control probe. DIC optics (C–F,F') and F-actin (green) and nuclear staining (white) (D',E') of control testioles reveals distinct, synchronized spermatogenic cysts separated by clear cyst boundaries (carets) (C), small cysts of spermatogonia (PSG) at the apex (D,D'), larger cysts of secondary spermatocytes (SSC) posterior to the apex (E,E'; arrowhead in (E') indicates somatic sheath cells associated with cysts of germ cells), early spermatids with round prominent nuclei (early ST) (F) and mid-stage spermatids with smaller, compact round nuclei (mid ST) (F'). (G) Late spermatid cysts in controls are synchronized in spermiogenesis; hollow arrowheads indicate somatic sheath cells. vasa RNAi testioles contain large masses of cells with heterogeneous nuclear morphologies (H; arrowheads). (I) PSG cysts are abnormal in shape and size, contain nuclei of multiple sizes (arrowheads), and (I') have filamentous actin masses interspersed between nuclei (arrows) arther than clearly defined cytoplasmic bridges (compare with D', arrows). (J,J') Abnormal cysts contain clusters of small dense nuclei (arrowheads). (K,K'). Aberrant cysts retain cytoplasmic bridges at spermatid stages (red arrows), and contain nuclei with morphologies corresponding to different spermatogenic stages, including both early (white arrows; compare with F) and mid ST (red arrowheads; compare with F') stages. (L) vasa RNAi late spermatid cysts are asynchronous, comprising multiple late spermatid and spermatozoon differentiation stages within a single cyst; cysts remain associated with sheath cells (hollow arrowheads). Sc

proceeding through meiosis as in wild type. Finally, the posteriormost region of vasa RNAi testioles contained irregular groups of cells at mixed stages of late spermatid and spermatozoon differentiation (Fig. 6L), rather than the perfectly synchronised cysts of late spermiogenic stages seen in controls (Fig. 6G). These defects were observed in testes examined 28–29 days following injection of adult males, but are not artefacts of age, as testes of 10 week old wild type adult males showed normal progression through all stages of spermatogenesis (supplementary material Fig. S4B).

Taken together, these data suggest that *vasa* is required for the maintenance of synchrony within cysts at multiple stages of spermatogenesis. In addition, *vasa* may be required for secondary spermatogonia to enter correct meiotic progression as spermatocytes, in the absence of which germ cells are nonetheless able to continue with subsequent stages of spermatogenesis.

#### Discussion

#### Oncopeltus germ cell formation

In several cases, analyses of molecular markers such as *vasa* mRNA have revealed the presence of a cryptic germ plasm that had eluded prior histological studies (Yoon et al., 1997; Tsunekawa et al., 2000; Wu et al., 2011). In *Oncopeltus*, we

have shown that none of the transcripts of an extensive suite of conserved germ cell markers localise asymmetrically within oocytes or in early embryos (Figs 1, 2, 4; supplementary material Fig. S3), including transcripts of genes that localise to and are required for the function of germ plasm in Drosophila. Gene products of at least one of these conserved germ line markers have been found in the germ plasm of every species where a germ plasm is known to exist (Ewen-Campen et al., 2010), although we note the important caveat that in Drosophila, several of these genes (vasa, piwi, and tudor) are localized as proteins rather than mRNAs. Thus, the lack of localisation of transcripts of any of these 19 genes during oogenesis or early embryogenesis suggests that Oncopeltus lacks germ plasm. Instead, our data support the hypothesis that Oncopeltus germ cells form in the absence of germ plasm, and are not present prior to the onset of posterior invagination at the end of the cellular blastoderm stage. We cannot, however, rule out the possibility that untested molecular markers, including protein products of the genes examined here, could be asymmetrically localised to a putative germ plasm in Oncopeltus.

While we provide multiple markers of PGCs, further experiments could be useful to confirm the identity of these cells. However, demonstration that these cells are functional PGCs via ablation experiments is complicated by the fact

that they arise at the inner face of the blastoderm at the gastrulation center, so that their physical disruption would likely also compromise mesoderm formation and subsequent embryogenesis. Moreover, we note that while pole cell removal experiments in Drosophila result in sterility, pole cell removal in another insect with germ plasm, the wasp Pimpla turionellae, yields fertile adults despite the fact that these pole cells are bona fide PGGs in wild type embryos (Bronskill, 1959; Achtelig and Krause, 1971; Fleischmann, 1975). Further, we are currently unable to genetically ablate these cells and determine their effect on fertility, as our vasa, tudor and vasa + tudor RNAi double RNAi experiments do not disrupt their formation (Fig. 5). Lineage tracing techniques that would permit tracking of the putative PGCs over the six-week period between PGC formation and sexual maturity are not currently available for Oncopeltus. These caveats notwithstanding, the molecular and morphological evidence that the cells we identify in this report are bona fide Oncopeltus PGCs is comparable to that available for PGC identification in most studied animal species: (1) three conserved germ line genes, vasa, tudor, and boule, are specific germ cell markers in Oncopeltus (Figs 1-4); (2) transcripts of these genes first become enriched in germ cells specifically at the time that these cells were previously reported to arise based on morphological and cytological criteria (Figs 2-4) (Butt, 1949); and (3) cells with these molecular markers undergo migration and primordial gonad occupation (supplementary material Fig. S1; Figs 3, 4) consistent with the well-documented behavior of PGCs in many other hemipterans (Seidel, 1924; Mellanby, 1935; Butt, 1949; Sander, 1956; Kelly and Huebner, 1989; Heming and Huebner, 1994).

Although the posterior location of germ cells at the time of their specification is superficially similar to that of pole cells in Drosophila and other Diptera, PGC specification and development in Oncopeltus differs in several important ways. First, while Drosophila pole cells form on the exterior of the posterior syncytial blastoderm before somatic cellularisation, Oncopeltus germ cells appear on the yolk side of the cellular blastoderm. Second, while Drosophila pole cells are the first cells in the embryo to cellularise (Huettner, 1923), Oncopeltus germ cells arise after blastoderm cellularisation is complete (Butt, 1949). Third, because Oncopeltus is an intermediate-germ insect, only the gnathal and thoracic segments have been specified at the time that germ cells arise (Liu and Kaufman, 2004), whereas in the long-germ insect Drosophila, pole cells form posterior to the abdominal embryonic segments. Lastly, Oncopeltus germ cells form on the dorsal surface of the embryo, and remain on the yolkfacing surface of the mesoderm during their migration to the gonad primordium in anterior abdominal segments (Fig. 3). As a result, they do not undergo a transepithelial migration through the hindgut epithelium as in Drosophila (reviewed by Richardson and Lehmann, 2010).

#### The function of "germ line genes" in Oncopeltus

Our functional analysis led to the surprising discovery that neither vasa nor tudor play instructive roles in germ cell specification in Oncopeltus. Both of these genes are required for germ cell specification in Drosophila (Boswell and Mahowald, 1985; Schüpbach and Wieschaus, 1986) and other species (Sunanaga et al., 2007; Spike et al., 2008). However, vasa has widely divergent roles across Metazoa (reviewed by Yajima and Wessel, 2011), and in many cases is dispensable for PGC

specification (Tanaka et al., 2000; Braat et al., 2001; Li et al., 2009; Özhan-Kizil et al., 2009). In several organisms it plays a role in adult gametogenesis (Tanaka et al., 2000; Ohashi et al., 2007; Salinas et al., 2007; Fabioux et al., 2009; Salinas et al., 2012; Ewen-Campen et al., 2013).

Intriguingly, we find that similar to the mouse and the cricket, vasa is required for spermatogenesis in adult Oncopeltus (Fig. 6), but not for oogenesis. This sex-specific function may relate to a putative role in stem cell function. As in other hemimetabolous insects (Büning, 1994), in Oncopeltus germ line stem cells are likely active in the apex of the testes (Schmidt and Dorn, 2004) but are not thought to be present in adult ovaries. One caveat to this hypothesis is that vasa transcript was not detected by in situ hybridisation in the primary spermatogonia (Fig. 6A), although it may be present at very low levels in those stem cells. Alternatively, given its strong expression in secondary spermatocytes and the defects in cyst integrity and synchrony caused by vasa RNAi (Fig. 6), Oncopeltus vasa may play a malespecific role in the onset or synchrony of meiosis. Consistent with a conserved role for *vasa* in bilaterian meiosis, male germ cells in vasa knockout mice arrest just prior to meiosis onset (Tanaka et al., 2000), and in human stem cell-derived germ cells vasa overexpression enhances meiotic progression (Medrano et al., 2012). Oncopeltus vasa RNAi leads to premature spermatid differentiation by some diploid secondary spermatocytes within a cyst, resulting in cyst asynchrony. In Drosophila, mutations are known that disrupt meiosis but do not prevent sperm formation (Davis, 1971), consistent with the hypothesis that spermiogenesis can be decoupled from meiotic status.

#### The evolutionary origins of germ plasm in insects

Together with recent molecular and classical histological data on germ cell specification in other insects, our results are consistent with the hypothesis that germ plasm is a derived mode of germ cell specification that arose in the ancestor to holometabolous insects (Fig. 7) (Lynch et al., 2011; Ewen-Campen et al., 2012). The only other functional genetic analysis of germ line specification in a hemimetabolous insect to date (Ewen-Campen et al., 2013) has also provided evidence that maternally supplied posterior germ plasm is absent, and that *vasa* is dispensable maternally and zygotically for germ cell formation. Our data thus support the notion that germ plasm-driven germ cell specification mechanisms operative in *Drosophila melanogaster* and *Nasonia vitripennis* are derived relative to the Hemimetabola (Fig. 7).

The ubiquitous distribution of germ cell markers in early *Oncopeltus* embryos and their subsequent enrichment in presumptive germ cells at the blastoderm posterior is reminiscent of *vasa* expression in the beetle *Tribolium* (Fig. 7) (Schröder, 2006; C. von Levetzow, Konservierte und divergente Aspekte der *twist-*, *snail-* und *concertina-*Funktion im Käfer *Tribolium* castaneum, PhD thesis, Universität zu Köln, 2008). Further taxonomic sampling, and functional studies in *Tribolium*, will be needed to determine whether the PGC specification mechanisms in these two species may be the result of common ancestry (Fig. 7).

A large number of transcripts that localise to germ plasm in *Drosophila* are expressed ubiquitously in *Oncopeltus* oocytes and early embryos. This suggests that the evolution of germ plasm in Holometabolous insects involved a large-scale change in the localisation of many transcripts in the oocyte. We propose that this likely resulted from a change in the localisation of an upstream component capable of recruiting many downstream

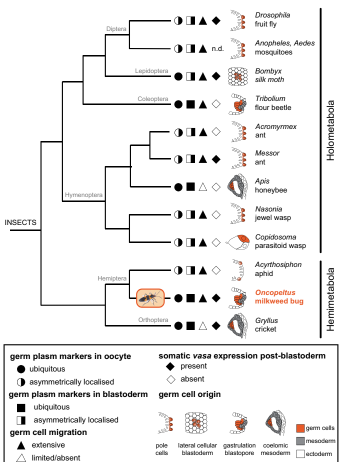


Fig. 7. Phylogenetic distribution of germ cell specification mechanisms and migration patterns across insects. Species shown are those for which data on the expression and/or function of molecular markers for germ cells during oogenesis and embryogenesis are available. Molecular data suggest absence of germ plasm in oocytes (circles) and early embryos (squares) of some Holometabola (Tribolium, Apis) and Hemimetabola (Oncopeltus and Gryllus), and somatic expression of vasa at post-blastoderm stages of development (diamonds) is not uncommon. In most species, PGCs undergo extensive migration from the site of specification to the gonad primordia (triangles). Data from this study (also Nakao, 1999; Mahowald, 2001; Donnell et al., 2004; Zhurov et al., 2004; Chang et al., 2006; Dearden, 2006; Juhn and James, 2006; Nakao et al., 2006; Schröder, 2006; Chang et al., 2007; Juhn et al., 2008; Zhao et al., 2008; Chang et al., 2009; Khila and Abouheif, 2010; Lynch et al., 2011; Ewen-Campen et al., 2013; C. von Levetzow, Konservierte und divergente Aspekte der twist-, snail- und concertina-Funktion im Käfer Tribolium castaneum, PhD thesis, Universität zu Köln, 2008). Phylogenetic relationships from Yeates et al. (Yeates et al.,

transcripts, rather than via the sequential evolution of distinct localisation mechanisms for individual transcripts. Studies on the genetic mechanism of evolutionary redeployments of multiple downstream genes have largely focused on transcription factors, as individual transcription factors are capable of regulating large numbers of target genes (Hoekstra and Coyne, 2007; Moczek, 2008; Stern and Orgogozo, 2008; Craig, 2009). Interestingly, in the case of germ plasm, transcription factors are unlikely to have been key players in the mechanisms of evolutionary change for a number of reasons. First, regulation of germ line determinants is largely post-transcriptional (Arkov and Ramos, 2010; Richter and Lasko, 2011; Sengupta and Boag, 2012; Nousch and Eckmann, 2013). Second, germ plasm transcript function relies on their subcellular localisation (often mediated via signals in their 3'UTRs) rather than their presence or absence (Rangan et al., 2009). Finally, unlike key transcription factors identified as largely sufficient to induce specific somatic cell fates (e.g. Akam, 1998; Kozmik, 2005; Baena-Lopez and García-Bellido, 2006), there is no single conserved gene that is sufficient to confer germ cell fate across metazoans. The evolution of germ plasm may therefore serve as an example of how a novelty (asymmetrically

localized germ plasm in the oocyte) arose via changes in RNA localisation rather than transcriptional regulation.

#### **Materials and Methods**

#### Animal culture

Oncopeltus fasciatus were cultured at 28°C as previously described (Ewen-Campen et al., 2011). Timing of embryonic events reported here may differ from that reported in other studies using lower rearing temperatures (e.g. Liu and Kaufman, 2004).

#### Cloning and phylogenetic analysis

Total RNA was extracted from mixed-stage embryos and ovaries using TRIzol (Invitrogen) and used for first strand cDNA synthesis with qScript cDNA SuperMix (Quanta BioSciences). An Oncopeltus vasa fragment was cloned using degenerate primers (supplementary material Table S1). nanos and piwi fragments were obtained from the Oncopeltus transcriptome (Ewen-Campen et al., 2011). Fragments were extended using RACE PCR (SMART RACE cDNA kit, Clontech), and used as templates for DIG-labeled in situ probes and dsRNA fragments following sequence verification (supplementary material Table S1). Genes for the in situ hybridisation screen (supplementary material Tables S1, S2) were obtained from the Oncopeltus transcriptome (Ewen-Campen et al., 2011; Zeng and Extavour, 2012) and amplified using primers containing linker sequence (5'-CCCGGGGC-3') enabling direct addition of a T7 site to the 3' end in a subsequent PCR reaction. Extended sequences are available from ASGARD (http://asgard.rc.fas.harvard.edu) (Zeng and Extavour, 2012). All coding sequences

reported in this study have been submitted to GenBank [accession numbers KC261571-KC261587] except for orb and UevIA, for which we obtained only 3' UTR sequence

Maximum-likelihood phylogenetic analysis was performed for vasa, piwi, and nanos as previously described (Ewen-Campen et al., 2012).

#### Tissue fixation and gene expression analysis

Embryos were fixed and stained as previously described (Liu and Kaufman, 2004; Erezyilmaz et al., 2009; Kainz et al., 2011). Adult gonads were dissected in 1× PBS and fixed in 4% formaldehyde in 1× PBS for at least one hour. Antibodies used were mouse anti-alpha tubulin DM1A (Sigma) 1:50 and goat anti-mouse Alexa Fluor 568 (Invitrogen) 1:500–1:1000, and counterstains were FITC-phalloidin (Invitrogen) 0.5–1  $\mu l$  and Hoechst 33342 (Sigma) 0.1–0.5  $\mu g/ml$ .

#### Plastic sectioning

In situ hybridisation and/or Sytox Green (Invitrogen) staining were performed prior to embedding embryos in Durcupan ACM Fluka (Sigma), mixed at a ratio of 32:27:1:0.6 = components A:B:C:D. Embryos were dehydrated through 10-minute washes in each of 50%, 70%, 90%, 2× 100% ethanol and 100% acetone, transferred to a 1:1 mixture of acetone: catalysed Durcupan, and left uncovered in a fume hood overnight. Embryos were individually transferred to fresh Durcupan in silicone molds (Electron Microscope Sciences NO. 70903) and oriented following a 30-minute initial hardening at 65°C. Resin blocks were baked for 24 hours at 65℃.

Block fronts were trimmed with a razor blade and sectioned at 5-6  $\mu m$  on a Leica RM2255 microtome with a high-profile knife holder using High-Profile disposable "diamond-edge" steel knives (C.L. Sturkey NO. D554D50). Sections were collected on water droplets on charged slides, dehydrated on a heat block, and mounted in Permount (Fisher Scientific).

#### Parental RNAi

dsRNA for all genes (supplementary material Table S1) was prepared as previously described (Kainz et al., 2011) and resuspended in injection buffer (5 mM KCl, 0.1 mM NaH<sub>2</sub>PO<sub>4</sub>) to a concentration of 2 μg/uL. Male and female adults were injected three days after final molt with 5  $\mu$ L of 2  $\mu$ g/uL dsRNA using a Hamilton syringe and size 26 needles. Testes were collected from injected males 27-29 days after injection.

#### Reverse-transcription PCR

Half of each clutch laid by injected females was fixed for in situ hybridisation, and the other half was homogenised in TRIzol (Invitrogen) and stored at -80 °C before isolation of total RNA. RNA was isolated separately from late blastoderm (24-29 hours AEL), early germ band (24-48 hours AEL) and late germ band (72-96 hours AEL) embryos laid by injected mothers. Genomic DNA was treated with Turbo DNase (Ambion) at 37°C for 30 minutes, followed by DNase heat-inactivation and phenol/chloroform extraction. cDNA was synthesised from 120 ng of each RNA sample using Superscript III Supermix (Invitrogen), PCR was performed using Advantage 2 DNA Polymerase from 1 µL of cDNA template and primers indicated in supplementary material Table S1 at 60°C annealing temperature with 35 PCR cycles. RT-PCR results for samples of all three embryonic ages tested yielded indistinguishable results, indicating that maternal RNAi was effective at reducing zygotic transcripts in embryos at least up to four days AEL.

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#### **Author Contributions**

B.E.-C. and C.G.E. designed the research; B.E.-C., T.E.M.J. and C.G.E. performed experiments, collected and analysed data and wrote the manuscript; C.G.E. obtained funding for the research.

#### Competing Interests

The authors have no competing interests to declare.

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#### **Supplementary Material**

Ben Ewen-Campen et al. doi: 10.1242/bio.20134390

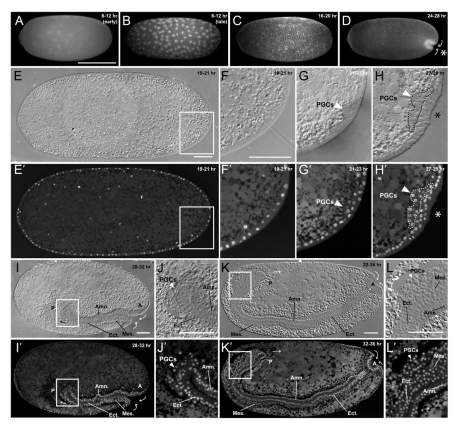


Fig. S1. Early embryogenesis of *Oncopeltus fasciatus* and morphological identification of putative PGCs. (A–C) The *Oncopeltus* syncytial blastoderm forms as a single layer of nuclei spread evenly across the surface of the yolk. (A) 8–12 hours after egg laying (h AEL) syncytially dividing nuclei are visible beneath the yolk surface. (B) Energid nuclei populate the yolk surface within 12 hours, and (C) undergo repeated mitosis and cellularisation to form a uniform cellular blastoderm by approximately 20 hours. Consistent with previous reports (Butt, 1949), we did not detect pole cell-like cells at any time during syncytial blastoderm or early cellular stages. (D) By 24–28 h AEL the posterior of the embryo has begun to invaginate into the yolk (arrows indicate direction of embryonic movements), forming the posterior pit (asterisk) where gastrulation takes place. This embryonic invagination is the beginning of the axial elongation process that will create the abdominal segments (Liu and Kaufman, 2004; reviewed by Panfilio, 2008). Immediately before posterior pit formation (~21 hours AEL), we observed putative PGCs on the inner surface of the blastoderm adjacent to the yolk (E–H'). (E) Medial section of a 19–21 h AEL embryo viewed with DIC optics and (E') stained with Sytox Green to reveal nuclei. Boxed region is enlarged in (F–H) and (F'–H'). (F,F') In 19–21 h AEL embryos, the early blastoderm is single-layered. (G,G') Between 21–23 h AEL, the embryonic posterior becomes multilayered, and the first cells visible within the yolk mass are the presumptive PGCs (arrowheads). (H,H') By 27–29 h AEL the putative PGCs (arrowhead) have fully entered the yolk. These putative PGCs are visible as a mesenchymal cluster with large round, centrally located nuclei, directly adjacent to the epithelialized somatic cells of the posterior blastoderm (asterisk), which are columnar in shape with smaller, basally located nuclei, (I,I') As the germ band elongates and its posterior end invaginates into the yolk at 28–32 h AEL (arrows ind

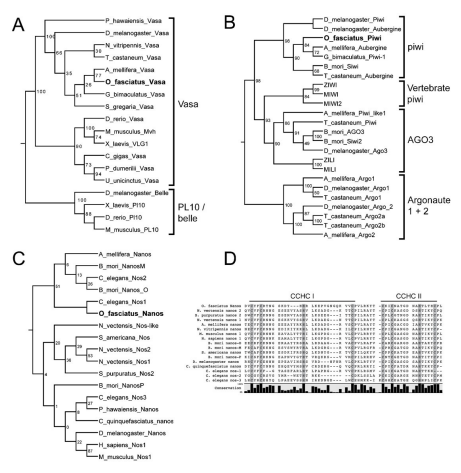


Fig. S2. Phylogenetic analysis of vasa, piwi, and nanos. Best-scoring maximum likelihood cladograms are shown with bootstrap values from 2000 replicates at nodes. (A) Oncopeltus Vasa is a member of the Vasa family of RNA helicases, not the closely related PL10/belle proteins. (B) Oncopeltus Piwi is a member of the piwi clade of PIWI proteins, closely related to Drosophila Piwi and Aubergine. (C) Phylogenetic reconstruction fails to resolve the internal relationships of nanos genes, because the conserved region of these proteins suitable for alignment (48 amino acids) is too short to provide sufficient phylogenetic signal. (D) A protein alignment of Oncopeltus Nanos protein with known orthologues demonstrates the presence of the diagnostic 2×(CCHC) zinc finger domain.

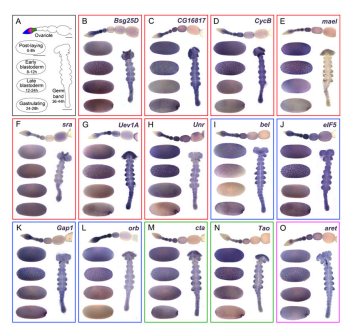


Fig. S3. Expression of additional germ plasm candidate markers in Oncopeltus ovaries and embryos. Transcripts were chosen for analysis based on their expression in the germ plasm and PGCs of Drosophila melanogaster (supplementary material Table S2). (A) Schematic figure showing the tissues depicted in subsequent panels. Embryonic ages shown in hours AEL. Coloured shading in ovariole schematic indicates spatial expression pattern of genes shown in boxes outlined in the corresponding colours. Blue = throughout entire tropharium in all nurse cells, as well as oogonia and resting oocytes; red = posterior nurse cells, oogonia and resting oocytes; green oogonia and resting oocytes but absent from or very low in only posterior nurse cells of tropharium; magenta = posterior nurse cells of tropharium but not oogonia or resting oocytes. (B-O) Expression patterns of genes studied in ovaries (top of each panel), blastoderm stages from 0-28 hours AEL (arranged vertically along the left of each panel), and in mid-germ band stages (to right of each panel), when PGCs are easily discernable in embryos stained for vasa, tudor or boule (Figs 3, 4). None of the 14 genes shown here were asymmetrically localised within oocytes, or to PGCs in later stages of development. (O) aret was strongly expressed in a population of cells located at the posterior, dorsal surface of the head at germ band stages, perhaps implicating this gene in foregut development. Scale bars: 500 µm for ovarioles and non-germ band embryos, 200 µm for germ band embryos.

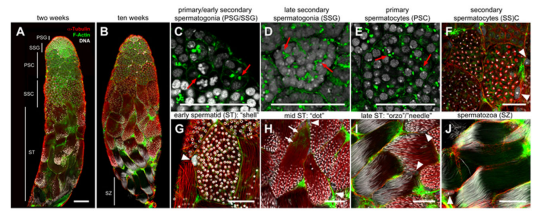


Fig. S4. Spermatogenesis in wild type *Oncopeltus* adult testioles. (A) Sperm tubule (testiole) from a two-week old male. Germ line stem cells (primary spermatogonia) and their putative niche are located at the anterior apex of each testiole. Cysts of clonally related secondary spermatogonia, spermatogytes, spermatids and spermatozoa are arranged in order posterior to the niche. All cells of a given cyst proceed synchronously through all stages of spermatogenesis, and all cysts at the same position along the anterior-posterior axis of the testiole are also roughly synchronised with each other (Economopoulos and Gordon, 1971). PSG: primary spermatogonia; PSG: secondary spermatogonia; PSG: secondary spermatogonia; PSG: primary spermatogonia; PSG: secondary spermatogonia; PSG: primary spermatogonia; PSG: p

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Table S1. Primers used for degenerate PCR, in situ hybridisation probe preparation, dsRNA preparation for RNAi, and RT-PCR. Sequence lengths are in nucleotides. F = forward; R = reverse.

				1	- 1						
Gene Name	Þ	degenerate primers	In situ probe length	Primers for men	Primers for in situ probe fragment (5' to 3')	dsRNA length	dsRNA	dsRNA primers (5' to 3')	RT-PCR ampli- con length	RT-PCI	RT-PCR primers (5' to 3')
vasa	ഥ	CCGATCGCATGC-	1403	Ħ	AAAAGGACTG-	109	Ħ	TGAGAGTA-	1392	ഥ	GGAAGAGAAA-
	2	GGTGCGGCCGA-		×	AAACCTGGA-		ĸ	TCCCGTCTGT-		×	TCCCGTCTGTT-
		TGCKRTGNACRTA			TCCCCAAATTC			TCAAGAATCC			CAAGAATCC
boule			1287	ш	ATTGAGGCAC-	716	ц	AGCCTCACCA-	614	ΙΉ	ATTGAGATGAA-
				ĸ	AGGGTGCCTA-		ĸ	AGGGTGCCTA-		×	AGTTCAGTGC-
					GGATTGGACT			GGATTGGACT			CTCAGGGAAA
tudor			1215	ш	GGTTAGCAAG-	802	ц	CCGAGAGTGC-	1876	щ	TGCTTCCAGT-
				×	CACACCTGTT-		×	AACTTGGTAC-		×	CCACCAAAAT-
				ı	GCCATAATCG			GCCTGTGGTC			CGCTTCTCATT
nanos			843	т,	GAAGGAACC- CGTAGGGAA						
				В	ATAATCCCTG-						
piwi			1023	ш	TGAAGAAGT-						
				~	GATTGAGAG-						
				:	ACGAGAAAG-						
arrest (aka bruno)			703	Ā	ATGTTCACTG-						
				×	TACAGTGCCA-						
- 11			300	Ē	TACGGTTGGA						
pelle			7.25	т,	GGGTTGAGGA- GCAAGACAAA						
				×	GCTCTGGtTT-						
beta-tubulin									199	Ħ	TGATCCCTTAC-
										~	CTCGACACA
										4	AAATCACTCT
Blastoderm- specific - gene 25D (aka Rvo 25D)			821	Ľ	AGCTGGTGGA- ACTCCAGAGA						
(1120)				×	TCAGTTCCTC-						
CG16817			717	ш	GCCATAGCTG-						
				В	GGGTCGAGGT-						
concertina			795	ш	GGGAAGTCAA-						
				м	CGITCCICAA TTGAAGAAGA-						
Cyclin B			913	Ħ	GTAAGGGAG-						
				×	TGTATATGAG- GTGTAGTGG						

Table S1. Continued.

Gene Name	degenerate primers	In situ probe length	Primers for	Primers for in situ probe fragment (5' to 3')	dsRNA length	dsRNA primers (5' to 3')	RT-PCR amplicon length	RT-PCR primers (5' to 3')
elongation initiation tion factor 5 (aka		790	ш	GGTTCTCCCA- AATCTGACGA				
(с.пэ			×	TGTCCAGGTC-				
GTPase-activating protein 1 (aka		748	Ħ	AAAGTGGCCA- TTAAGCGAGA				
(r.dan)			~	AGCCATGGTG-				
maelstrom		543	ш	GCAGTACGTT- GTGCTTTGGA				
			~	AACAGGATCG-				
oo18 RNA-binding protein (aka orb)		772	ш	TCTCGGTGAT- GGTGTTTTCA				
			~	TGCTGTGAGG-				
sarah		351	ш	TAAACCGGAA-				
			×	GATGTTGGCA-				
Tao		746	щ	TCAGGCTCTC-				
			~	AGCTCCCATG-				
UevIA		792	Г	TTTCTGCCAT-				
			~	AGCCCATTTT GCAACAAGTT-				
Upstream of N-ras		848	ц	TCCAGATGTCC AAACCCATGA-				
			×	GCCTTCACAG GCCCTCTCAA- TGAATCCAAA				

Table S2. Genes included in Oncopeltus germ plasm in situ screen.

Transcript Expression
in Drosophila Germ
Plasm and PGCs

			Pla	sm and P	GCs			
Drosophila Gene Name	Drosophila gene symbol	Drosophila CG #	Germ Plasm	Pole Cells	Stage 9 PGCs	Drosophila germ line mutant phenotype (Molecular function)	Functional conservation outside <i>Drosophila</i>	References
Boule	bol	CG4760	No	No	No	Spermatogenesis defects (RNA binding)	Germ line expression/func- tion across bilateria, often specific to spermatogenesis	[Lécuyer et al., 2007; Shah et al., 2010]
tudor	tud	CG9450	No*	No*	N.D.	Pole cell formation defects (tudor domain protein)	Germ line expression/ function across bilateria	[Golumbeski et al., 1991; Bardsley et al., 1993; Ewen-Campen et al., 2010]
orb	orb	CG10868	Yes	Yes	No	Oogenesis defects (RNA binding)	N.D.	[Lécuyer et al., 2007]
sarah	sra	CG6072	Yes	Yes	No	Oogenesis defects (Calcineurin regulation)	N.D.	[Lécuyer et al., 2007]
Cyclin B	CycB	CG3510	Yes	Yes	N.D.	Fertility defects in both sexes (Cyclin protein)	N.D.	[Lécuyer et al., 2007]
arrest (bruno)	aret	CG31762	Yes	Yes	Yes	Oogenesis defects (RNA binding)	N.D.	[Lécuyer et al., 2007]
concertina	cta	CG40010	Yes	Yes	Yes	No germ line phenotype reported (G-protein alpha subunit)	N.D.	[Lécuyer et al., 2007]
Gap1	Gap1	CG6721	Yes	Yes	Yes	No germ line phenotype reported (PH & C2-domain, Ras GTPase activation)	N.D.	[Lécuyer et al., 2007]
eIF5	eIF5	CG9177	Yes	Yes	Yes	No germ line phenotype reported (translation initiation)	N.D.	[Lécuyer et al., 2007]
Blastoderm- specific gene 25D	Bsg25D	CG14025	Yes	Yes	Yes	No germ line phenotype reported (Unknown)	N.D.	[Lécuyer et al., 2007]
Uev1A	Uev1A	CG10640	Yes	Yes	Yes	No germ line phenotype reported (ubiquitin-protein ligase)	N.D.	[Lécuyer et al., 2007]
CG16817	-	CG16817	Yes	Yes	Yes	No germ line phenotype reported (Unknown)	N.D.	[Lécuyer et al., 2007]
Tao	Tao	CG14217	Yes	Yes	Yes	No germ line phenotype reported (Protein S/T kinase)	N.D.	[Lécuyer et al., 2007]
Upstream of N-ras	Unr	CG7015	Yes	Yes	N.D.	No germ line phenotype reported (RNA and protein binding)	N.D.	[Lécuyer et al., 2007]
Belle	bel	CG9748	N.D.	Yes	No	Oogenesis and spermato- genesis defects (ATP- dependent RNA helicase)	N.D.	[Tomancak et al., 2007]
maelstrom	mael	CG11254	No	Yes	Yes	Oogenesis and spermato- genesis defects (HMG- box DNA binding )	Germ line function in mouse	[Lécuyer et al., 2007]

<sup>\*</sup>Tudor protein is localised to both pole plasm and pole cells in *Drosophila*. N.D. = no data available.

Table S3. Effects of RNAi on Oncopeltus PGC formation.

RNAi	Total # Scored	# embryos with non-specific defects* (%)	# surviving embryos with PGCs (%)
DsRed	39	4 (10.3%)	34 (97.1%)
vasa	16	0	15 (93.8%)
tudor	20	0	20 (100%)
vasa and tudor	19	9 (47.4%)	10 (100%)
boule	22	18 (81.8%)	4 (100%)

<sup>\*&</sup>quot;Non-specific defects" includes failure to develop a germ band as well as the formation of grossly defective germ bands, both of which ultimately resulted in embryonic lethality before 40–54 hours AEL, the time when we scored for PGCs. These embryos were not scored for PGC presence/absence.

#### **CHAPTER 5**

oskar functions in adult neural stem cells to influence long-term memory formation in the cricket Gryllus bimaculatus

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#### **ABSTRACT**

Although *oskar* was first described for its role in the *Drosophila* germ line, recent studies have shown that this gene has additional functions in the nervous system of larval *Drosophila* and in the embryos of a basally-branching insect, the cricket *Gryllus bimaculatus*. However, the specific molecular functions of *oskar* in the nervous system of either species remain unclear. In this study, we show that RNAi against *Gb-oskar* impairs long-term olfactory memory, but not short-term memory, in the cricket. We then show that *Gb-oskar* is expressed in the adult brain in a cluster of neuroblasts that are responsible for adult neurogenesis in the mushroom body, the anatomical substrate of olfactory memory in insects. Previous studies have shown that killing these mushroom body neuroblasts specifically impairs olfactory learning. Thus, our results are consistent with the hypothesis that *Gb-oskar* is involved in adult neuroblast function (i.e. proliferation, maintenance, or survival), and that its role in long-term olfactory memory is mediated through these cells. We discuss these results in both a functional and an evolutionary

context, and propose necessary additional experiments to directly test the role of *Gb-oskar* in adult neuroblasts.

#### INTRODUCTION

The *oskar* gene was first discovered in *Drosophila melanogaster*, where it is both necessary and sufficient to specify embryonic germ cells and recruit the posterior determinant *nanos* (Ephrussi et al., 1991; Ephrussi and Lehmann, 1992; Lehmann and Nüsslein-Volhard, 1986). Interestingly, *oskar* and several other genes originally identified as *Drosophila* germ line genes, including *nanos*, *pumilio*, *staufen*, *orb*, and the PIWI proteins *aubergine* and *AGO3*, have since been shown to have a variety of roles in the nervous system (Dubnau et al., 2003; Pai et al., 2013; Perrat et al., 2013; Wharton et al., 1998; Xu et al., 2013; Ye et al., 2004). Furthermore, a role for *oskar* in the nervous system, but not in the germ line, is conserved between *Drosophila* and a basally-branching hemimetabolous insect, the cricket *Gryllus bimaculatus*, suggesting that a neural function of *oskar* may in fact be ancestral (Ewen-Campen et al., 2012). However, the precise role(s) of *oskar* in the nervous system remain largely unknown.

The biochemical functions of some of the *Drosophila* "germ line genes" are well characterized, allowing for relatively detailed models of their function in neurons. For example, studies of Nanos (Nos) and Pumilio (Pum) during *Drosophila* embryogenesis have revealed that Pum binds to specific sequences in the 3'UTR of target mRNAs via the highly conserved PUF domain, and subsequently recruits Nos to form a translational repression complex (Murata and Wharton, 1995; Sonoda and Wharton, 1999; Zamore et al., 1997). In the *Drosophila* nervous system, Pum has been shown to function at neuromuscular junctions as a translational regulator (Mee et al., 2004; Menon et al., 2004) and in long-term memory formation (Dubnau et al., 2003), and to act together with Nanos in larval dendrite morphogenesis (Xu et al., 2013; Ye et al., 2004). A variety of relevant target neuronal mRNAs regulated by Pum have been identified at the neuromuscular junction, including the voltage-gated sodium channel *paralytic* (Mee et al., 2004) and the translation factory *eIF-4E* (Menon et al., 2004), and in the adult brain, including a membrane-associated guanylate kinase, *dlg1* (G. Chen et al., 2008). There is also evidence that

the neural roles of Nos and Pum are phylogenetically widespread: a Pum ortholog, Pumilio-2, has been shown to function mammalian neurons (Driscoll et al., 2013), and a role for Nos in the development of neurons has been functionally demonstrated in a cnidarian (Kanska and Frank, 2013).

In contrast to Nos and Pum, the biochemical function of Oskar remains largely unknown, despite decades of research (reviewed in Ewen-Campen et al., 2010). It is thus unclear how this protein may function in neurons. *oskar* is a novel, insect-specific gene, and contains two predicted protein domains, both of unknown function: a LOTUS (aka OST-HTH) domain and a SGNH hydrolase domain (Anantharaman et al., 2010; Callebaut and Mornon, 2010; Ewen-Campen et al., 2012; Lynch et al., 2011). The LOTUS/OST-HTH domain has been predicted to bind RNA (Anantharaman et al., 2010; Callebaut and Mornon, 2010), although Oskar has never been directly shown to bind RNA. The SGNH hydrolase domain belongs to a large family of lipid-processing enzymes, but, enigmatically, this domain in Oskar is predicted to be catalytically inactive (Anantharaman et al., 2010). A third domain, termed Long Osk, is found only in *Drosophilid* insects, and also has unknown biochemical function (Ewen-Campen et al., 2012; Lynch et al., 2011). The current model is that Oskar serves as a scaffolding protein, facilitating the assembly of ribonucleoprotein complexes that include a variety RNA-processing components (Jones and Macdonald, 2007; Suyama et al., 2008).

Despite the unknown molecular function of Oskar, there is evidence that this gene functions in the nervous system of *Drosophila* as well as in a basally branching hemimetabolous insect, the cricket *Gryllus bimaculatus*. In *Drosophila*, larvae that are mutant for *oskar* or express neuron-specific *oskar* RNAi display defects in *nanos* localization, ultimately leading to a defect in dendrite morphogenesis and an associated defect in motor response to mechanical stimulation (Xu et al., 2013). In addition, Dubnau *et al.* (2003) have reported a role for *oskar* in long-term memory formation based on an insertional pGal4 mutant upstream of *oskar* (the *norka* mutant), but it is unclear if *norka* is a *bona fide* allele of *oskar* (Xu et al., 2013) (See also Appendix 2 of

this dissertation). In the cricket, we have previously shown that *Gb-oskar* is expressed in embryonic neuroblasts, and that RNAi against *oskar* disrupts the development of the central nervous system (Ewen-Campen et al., 2012). Thus, although *oskar* functions in the nervous system of both species, it is unclear if it plays similar or divergent roles in each.

In the present study, we address whether *Gb-oskar* has an additional neural function in the adult brain of the cricket. We first show that RNAi against *Gb-oskar* in adult crickets impairs long-term memory formation in an olfactory associative learning assay. Next, we show that *Gb-oskar* is expressed in a well-characterized group of mushroom body neuroblasts responsible for adult neurogenesis. Given that these mushroom body neuroblasts have previously been implicated in olfactory learning in related species (Scotto-Lomassese et al., 2003), we hypothesize that *Gb-oskar* is involved in the survival and/or proliferation of neuroblasts in the adult brain. We propose future experiments to directly test whether the role of *Gb-oskar* in olfactory learning involves adult neurogenesis.

#### **METHODS**

Gryllus husbandry

For behavior experiments, *Gryllus bimaculatus* were maintained in the Mizunami laboratory at 27°C on a 12:12 light cycle, with a diet of insect food pellets, as previously described (Matsumoto and Mizunami, 2000). For gene expression analysis, qPCR, and cell proliferation experiments, crickets were maintained in the Extavour laboratory at 28°C on a 12:12 light cycle, with a diet of grain and cat food, as previously described (Kainz et al., 2011).

RNAi

Adult male crickets within one week of their final moult were injected 2 μL of 10μM dsRNA through a hole pierced in the median ocellus using a 10 μL syringe fitted with 26S gauge tip (Hamilton Inc., Nevada, USA). Behavioral tests were repeated using two non-overlapping fragments of *Gb-oskar* (742 bp and 503 bp) with a 678 bp fragment of DsRed, as a negative control (Ewen-Campen et al., 2012).

qPCR

Two days after dsRNA injection, brains were dissected in ice-cold PBS, then immediately homogenized in TRIzol (Life Technologies). Total RNA was extracted from a total of six brains per treatment, following the manufacturer's instructions, including a 30 minute DNAse treatment. 1 µg of total RNA was used as template for cDNA synthesis using SuperScript III (Life Technologies) with oligo-dT primers. cDNA was diluted 1:10 prior to qPCR, and 6 µL of template was used per 25 µL qPCR reaction. (PerfeCta SYBR Green SuperMix, Low ROX, Quanta Biosciences). qPCR reactions were conducted in triplicate, and fold change was calculated using the delta-delta Ct method (Livak and Schmittgen 2001), with standard deviation propagated following standard methods. Beta-tubulin was used as a reference gene (Kainz et al., 2011). Primers amplifying a 234 fragment of *Gb-oskar* (F: TTGTTGACCATTCCCTTCCT, R: ACTCCACAACACCACTCC) and a 166 bp fragment of Beta-tubulin (F: TGGACTCCGTCCGGTCAGGC, R: TCGCAGCTCTCGGCCTCCTT) (Kainz et al., 2011) were used.

#### Behavioral analysis

Adult male crickets at one week after final moult were used in the experiments. Three days before conditioning, individual crickets were separated into 100-mL beakers and deprived of

drinking water to enhance their motivation to search for water. Two days before a conditioning, each cricket was injected with dsRNA as described above.

Two days after dsRNA injection, each cricket was subjected to an odor preference test, in which the animal was allowed to freely visit peppermint and vanilla odors (Matsumoto and Mizunami, 2002). The time spent at each of the peppermint and vanilla source was measured cumulatively to evaluate relative odor preference (Matsumoto and Mizunami, 2002).

Crickets were subjected to 4-trial conditioning, in which an odor was paired with water reward, with inter-trial interval of 5 min (Takahashi et al., 2009). For conditioning, a small filter paper was attached to the needle of a hypodermic syringe. The syringe was filled with water reward (unconditioned stimulus: US<sup>+</sup>), and the filter paper was soaked with peppermint essence (conditioned stimulus: CS).

At one hour and one day after the end of the conditioning, each cricket was subjected to an odor preference test. Relative odor preference of each animal was measured using the preference index (PI) for rewarded odor (peppermint), defined as  $t_P/(t_P+t_V)$  \*100 (%), where  $t_P$  is the time spent exploring the peppermint source and  $t_V$  is the time spent exploring the vanilla source. Wilcoxon's (WCX) test was used to compare odor preferences before and after training. For multiple comparisons, Holm's method was used to adjust the significance level.

#### Gene expression analysis

Previously, we generated an antibody against *Gb-oskar*, and showed that this antibody recognizes a band of the correct size in *Gryllus* ovaries, and that the neuroblast-specific signal observed in embryos is abolished following *Gb-oskar* RNAi (Ewen-Campen et al., 2012). However, when we tested this antibody on *Gryllus* brains via Western blot, we observed that this antibody strongly cross-reacted with a ~23-24 kDa protein that is not present in ovaries or embryos. RNAi against *Gb-oskar* in adult brains did not reduce the intensity of this band on

Western blot, and Mass spectrometry of an excised band of 23-24 kDa size from an SDS-PAGE separation of *Gryllus* brain lysate did not identify any peptides corresponding to *Gb-oskar*. Thus, we conclude that, unlike in ovaries and embryos, this antibody recognizes something other than GbOsk protein in *Gryllus* brains. For this reason, we used *in situ* hybridization to detect *Gb-oskar* expression in brains.

For *in situ* hybridization, brains were dissected in ice-cold PBS, and de-sheathed following one hour in 4% paraformaldehyde, followed by an additional overnight fixation at 4°C or 3-4 hours at room temperature. Brains were then stored in methanol overnight at room temperature, due to the surprising observation that storage at -20°C greatly increased background tracheal staining. *Gb-oskar* was detected using a 788 bp probe, following standard protocols (Kainz et al., 2011), with the following modifications to reduce background: a 20 minute Proteinase K treatment followed by a 20-30 minute fix in 0.8% glutaraldehyde and 4% paraformaldehyde. The *Gb-oskar* probe was used at 1.0 ng/μl, and hybridized at 69-70°C.

#### EdU assay

Cell proliferation was assayed using the Click-iT EdU Alexa 488 kit (Life Technologies). Crickets were injected with 15 µl of EdU either into the abdomen or into the head capsule through the median ocellus (both methods successfully labeled dividing neuroblasts), and cell proliferation was detected following manufacturer's instructions. For tissue double-stained by *in situ* hybridization and EdU, the *in situ* hybridization was conducted before the visualization of incorporated EdU.

#### Vibratome sectioning

Gryllus brains were embedded in 4% low-melt agarose, and sectioned at 50-90 μm using a Leica VT1000S vibratome. For antibody staining, brains were sectioned prior to incubation

with the primary antibody. For *in situ* hybridization, brains were sectioned after staining had been completed.

#### **RESULTS**

Gb-oskar RNAi impairs long-term olfactory memory

Crickets have robust olfactory learning capabilities that can be experimentally investigated by training crickets to associate an novel odor with a water reward (Matsumoto and Mizunami, 2000). A behavioral preference for the rewarded odor can be detected one hour after training, and as few as three training sessions are sufficient to form a memory that is that does not significantly decay between 1-7 days (Matsumoto and Mizunami, 2000). In addition, the injection of dsRNA into the hemolymph of adult crickets has been shown to trigger a systemic reduction in target gene levels, and has been used successfully to interfere with the function of genes known to be involved in learning and memory in *Gryllus bimaculatus* (Takahashi et al., 2009). Importantly, because crickets are not injected with dsRNA until adulthood, this approach bypasses any potential developmental requirements for a given gene.

In order to test whether *Gb-oskar* is required in adults for olfactory memory in *Gryllus*, we assayed the olfactory memory of *Gb-oskar* RNAi adult male crickets at one hour and one day after training, compared to control crickets injected with an equal amount of DsRed dsRNA. We used qPCR to quantify RNAi efficiency, and found that *Gb-oskar* levels were reduced to 66.4% of control levels (**Figure 5.1A**). Although this reduction is modest compared to the reductions seen for many maternal or zygotic RNAi experiments (e.g. Ewen-Campen et al., 2013), it is comparable to reported knockdown levels of other genes in the *Gryllus* brain in adult crickets (Takahashi et al., 2009).

In DsRed RNAi control crickets, four training sessions led to a significant short-term preference for the rewarded hour at one hour after training, and this preference was retained after

one day (**Figure 5.1B**). Strikingly, although *Gb-oskar* RNAi crickets formed a short-term preference for the rewarded odor at one hour after training, this memory was lost by one day after training (**Figure 5.1C**). To ensure the specificity of this knockdown, we repeated these experiments using a non-overlapping fragment of *Gb-oskar* dsRNA, and observed a similar result (**Figure 5.1D**). Thus, *Gb-oskar* RNAi impairs long-term, but not short-term, olfactory memory formation. The fact that short-term olfactory learning remains intact in *Gb-oskar* suggests that the effect of *Gb-oskar* RNAi does not globally disrupt such processes as olfaction or locomotion, but is instead specific to long-term memory.

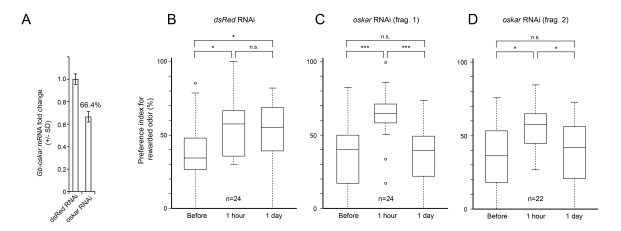


Figure 5.1. *Gb-oskar* RNAi impairs long-term olfactory memory formation in crickets. (A) qPCR validation of *Gb-oskar* RNAi. A modest reduction to 66.4% of control levels is observed. Graph represents *Gb-oskar* fragment #1. (B-D) Effects of *Gb-oskar* RNAi on olfactory learning. Relative preference between the rewarded odor (peppermint) and control odor (vanilla) was tested before training, 1 hour post-training, and 1 day post training for dsRed controls (B), *Gb-oskar* fragment #1 (C) and *Gb-oskar* fragment #2 (D). Boxes represent the  $1^{st}$ - $3^{rd}$  quartiles surrounding the median (middle line). Whiskers extend to extreme values within 1.5x of interquartile range. Wilcoxon's test was used for comparison of preference before and after conditioning. For multiple comparisons, the Holm method was used to adjust the significance level. (\* p < 0.05, \*\*\* p<0.001, n.s. = not significantly different). Behavior experiments and the associated statistical analysis were performed by Ryo Wakuda, Kanta Terao, Yukihisa Matsumoto, and Makoto Mizunami in the Mizunami lab.

Gb-oskar is expressed in adult neuroblasts of the mushroom body

To address the cellular mechanism by which *Gb-oskar* may influence olfactory memory, we examined the expression of *Gb-oskar* in the cricket brain. *Gb-oskar* expression was restricted to a cluster of cells located at the apex of each mushroom body (**Figure 5.2A**), and expression was not detected elsewhere in the brain. These *Gb-oskar*-positive cells matched descriptions of

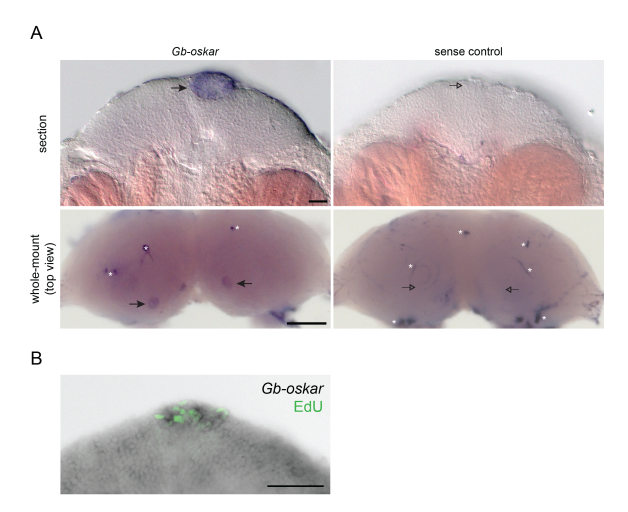


Figure 5.2. *Gb-oskar* is expressed in adult neuroblasts in the mushroom body. (A) *Gb-oskar* in situ hybridization (left panels) and sense controls (right panels), shown in 90  $\mu$ m vibratome section (top row; dorsal is up) and in a top-down view (looking down onto the dorsal surface) in whole-mount preparations (bottom row). *Gb-oskar* is detected in a cluster of cells at the apex of the Kenyon cells (black arrows). (B) The cells that stain for *Gb-oskar* are EdU-positive, confirming that these are mitotically active adult neuroblasts. Crickets were injected with EdU six hours prior to dissection. Scale bars = 50  $\mu$ m in top row of (A) and (B), 200  $\mu$ m in bottom row of (A).

mushroom body neuroblasts, the only proliferative cells in the adult cricket brain, which divide continuously during adult life to produce new Kenyon cells (the cells that make up the mushroom

body) (Cayre et al., 1996). To test whether *Gb-oskar*-positive cells were in fact the neuroblasts, we injected crickets with EdU six hours prior to dissection. Following *in situ* hybridization, we detected EdU specifically in the cells expressing *Gb-oskar* (**Figure 5.2B**), demonstrating that the *Gb-oskar*-expressing cells are indeed the adult neuroblast of the mushroom body.

#### **DISCUSSION**

We have shown that *Gb-oskar* RNAi impairs olfactory learning in the cricket *Gryllus* bimaculatus (**Figure 5.1**), and that *Gb-oskar* is specifically expressed in the neuroblasts of the adult mushroom body (**Figure 5.2**). Intriguingly, proliferation of these mushroom body neuroblasts has previously been linked to olfactory learning in a related cricket species: when these neuroblasts are killed using radiation, olfactory learning is impaired (Scotto-Lomassese et al., 2003). Thus, our results support the hypothesis that *Gb-oskar* is required for neuroblast function (i.e. proliferation, survival, and/or maintenance in an undifferentiated state), and that the role of *Gb-oskar* in olfactory memory is mediated through a cellular role in these neuroblasts. Directly testing this hypothesis will require assaying the effects of *Gb-oskar* RNAi on the proliferation and/or survival of these neuroblasts. Such experiments are currently underway.

The relationship between adult neurogenesis and long-term olfactory memory formation is intriguing, because it is well-established that the formation of long-term memory, including olfactory memory in insects, does not strictly require the birth of new neurons (Heisenberg, 2003; Kandel, 2012). Accordingly, we emphasize that killing adult neuroblasts impairs, but does not abolish, olfactory learning in the cricket (Scotto-Lomassese et al., 2002). In fact, adult neurogenesis does not occur in the mushroom bodies of *Drosophila* (or several other insect species)(reviewed in Cayre et al., 2007), yet olfactory memory is robust in these species (Heisenberg, 2003). Classical experiments in a wide range of animals have demonstrated that

long-term memory involves changes in synaptic strength between existing neurons, regulated via cyclic adenosine monophosphate (cAMP) response element-binding protein (CREB)-dependent de novo protein synthesis (reviewed in Kandel, 2012), including in Gryllus bimaculatus (Matsumoto et al., 2006). In *Drosophila*, long-term olfactory memory requires the  $\sim 2,500$ Kenyon cells of the mushroom body (Heisenberg, 2003), each of which receives synaptic inputs from a small and random subset of the antennal lobe projection neurons (which themselves receive direct synaptic input from the odorant receptor neurons of the antennae) (Caron et al., 2013). A given Kenyon cell thus responds with high selectivity to a small number of odors (or other stimuli), allowing the mushroom body to house an "explicit" representation of a large number of olfactory cues (Caron et al., 2013; Heisenberg, 2003). Specific olfactory stimuli are then associated with learned behavioral responses via specific sets of neurons connecting the mushroom body to other brain regions in a protein synthesis-dependent fashion, to form longterm memories (C.-C. Chen et al., 2012; Pai et al., 2013). Thus, it seems possible that adult-born Kenyon cells in *Gryllus* (and other species which display adult neurogenesis in the mushroom body) are recruited into an existing circuit, and allow for a constantly increasing repertoire of olfactory associations, and it could be this process that is disrupted by Gb-oskar RNAi. It is also interesting to note that of the two mammalian brain regions known to undergo adult neurogenesis, one (the subventricular zone) contributes to the olfactory bulb, and neurogenesis in this region is involved in olfactory memory (Lazarini and Lledo, 2011).

It remains unclear whether *oskar* is involved in long-term memory formation in *Drosophila* (Dubnau et al., 2003). Given that adult *Drosophila* lack the mushroom body neuroblasts seen in *Gryllus* (Cayre et al., 2007), a straight-forward test for a directly comparable *oskar* function is not possible. However, although *Drosophila* mushroom body stem cells are absent in adults, analogous mushroom body neuroblasts remain mitotically active late into pupal development (Ito and Hotta, 1992). Thus, it will be interesting to test whether *oskar* functions in

these mushroom body neuroblasts during larval and/or pupal stages, which would suggest a conserved function.

In addition, there is good reason to hypothesize that *oskar* could function in *Drosophila* olfactory long-term memory independent of a possible role in neuroblasts. Specifically, a recent study of the mushroom body output neurons has suggested that long-term memory involves the activity-dependent de-repression of mRNAs localized to granules containing Pum, Staufen, and Orb (Pai et al., 2013). Given that Oskar is thought to nucleate similar granules containing these proteins in the *Drosophila* oocyte (Breitwieser et al., 1996; Chang et al., 1999), it would be very interesting to test whether Oskar is involved in the formation and/or activity of these granules in the brain. Although I have thus far been unable to detect Oskar protein in the adult brain via antibody staining (see Appendix 2 of this dissertation), direct functional tests are necessary to establish whether or not this is the case. In addition, it should be noted that such a function for *Gb-oskar* in the formation of RNP in neurons is formally possible in *Gryllus*; although *Gb-oskar* is detected at highest levels in the mushroom body neuroblasts, we cannot rule out that it is expressed at lower levels in differentiated Kenyon cells (see Figure 2B).

We have previously shown that *Gb-oskar* functions in embryonic neuroblasts (Ewen-Campen et al., 2012), and have now extended these observations to show that *Gb-oskar* is also present in neuroblasts of the adult brain. It will be interesting to test whether *Gb-oskar* functions in neuroblasts throughout the entirety of nymphal development, or whether its activity is limited to the adult neuroblasts. In addition, future studies could test whether additional "germ line" genes also function in these adult neuroblasts, which would suggest that *oskar* may cooperate with conserved molecular partners in different cellular contexts. Indeed, we have previously shown that *Gb-vasa* is co-expressed with *Gb-oskar* in the embryonic neuroblasts (Ewen-Campen et al., 2012), although *Gb-vasa*'s function in these cells was not addressed.

Both germ cells and neuroblasts are stem cells that give rise to highly specialized, postmitotic daughter cells while they themselves remain proliferative for long periods of time. Thus, the role of *oskar* in both cell types could conceivably be related to stem cell maintenance and/or asymmetric division. Indeed, it has been noted that a variety of highly conserved "germ line genes" including *vasa*, *nanos*, and *piwi* are found in a variety of multipotent cells in diverse animals (Juliano and Wessel, 2010), raising the possibility that such genes are involved in establishing multipotency rather than specifically germ cells *per se*. A broader understanding of the function(s) of *oskar* will thus require additional studies of phylogenetically diverse insects, as well as further detailed biochemical analysis in *Drosophila* germ cells and neurons.

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## DISCUSSION AND OUTLOOK

Divergent mechanisms for specifying a conserved cell type

In this thesis, I provide evidence that a highly conserved cell type – the germ cells – can arise via highly un-conserved developmental mechanisms across species. Specifically, whereas *Drosophila* PGCs acquire their identity via maternal factors localized within the oocyte cytoplasm, my dissertation research demonstrates that such maternal determinants do not specify PGCs in two different basally branching insect species. Instead, my data suggest that PGCs arise later during development in both *Gryllus* and *Oncopeltus*, after cellularization has occurred and the zygotic genome has been activated, and therefore this process likely requires cell signaling. In other words, although PGCs are undoubtedly a homologous cell type across all animals, the mechanisms that specify these cells are diverse.

The observation that homologous structures can develop via divergent mechanisms is certainly not new (reviewed in Scholtz, 2005). Structures as obviously homologous and highly specialized as the wings of insects develop via remarkably different mechanisms across species: in holometabolous insects, wings arise from imaginal discs that survive metamorphosis, whereas imaginal discs (and metamorphosis itself) are entirely lacking in hemimetabolous insects (reviewed in Giorgianni and Patel, 2007). Similarly, while all adult insects (and arthropods) share a segmented body plan, the molecular mechanisms of segmentation during embryogenesis are entirely distinct between species (reviewed in Davis and Patel, 2002). On a broader phylogenetic scale, such cell types as neurons or embryonic mesoderm are each believed to be homologous across Bilateria, based on conserved gene expression patterns and analogy of function, yet both cell types arise via distinct developmental mechanisms even between relatively closely related taxa (reviewed in Roth, 2004; Stollewerk, 2008). Altogether, there is a growing body of modern genomic evidence supporting Karl von Baer's famous 1828 observation that the conserved

"phylotypic stage" of a given phylum (that embryonic stage during which all members of a phylum most closely resemble one another) is arrived at via highly diverse embryonic avenues (Domazet-Lošo and Tautz, 2010; Kalinka et al., 2010).

However, while there are plentiful examples of homologous structures arising via divergent developmental mechanisms, there is much less empirical data on *how*, mechanistically, development can evolve while consistently maintaining a conserved outcome. The data presented in this dissertation suggest that, in the case of PGC specification, gene co-option played an important role. Whereas previous studies proposed that the divergent mechanisms of PGC specification involved the *de novo* evolution of a novel gene (i.e. *oskar*; Lynch et al., 2011), the data presented here instead suggest that this process involved the redeployment of a pre-existing gene into a novel function. Below, I discuss how the modularity of the "germ line genes" may have influenced this evolutionary process.

The roles of gene co-option and modularity in developmental evolution

To understand how PGC specification has evolved, it may be helpful to imagine the molecular machinery of germ cells as a functional module. By module, I mean that the "germ line genes" are characterized by relatively strong interactions between one another, and that their expression in a cell is largely co-regulated by common upstream factors. Indeed, across nearly all animals that have been examined, germ cells express some or all of a conserved suite of genes, including Vasa, Nanos, Pumilio, Tudor genes, and PIWI family genes (reviewed in Ewen-Campen et al., 2010). These proteins often co-localize to the same subcellular structures within germ cells, a granular ribonucleoprotein structure lining the cytoplasmic face of the nuclear membrane, forming a germ cell-specific organelle sometimes referred to as the "nuage" (reviewed in Ewen-Campen et al., 2010; Voronina et al., 2011). Furthermore, there is evidence for physical interactions between PIWI proteins and Tudor proteins (Chen et al., 2009), and it has been

proposed that the ensemble of proteins in the nuage collectively act as a "hub" for post-transcriptional regulation of various classes of RNA as they exit the nucleus, a fundamental aspect of germ cell function (Voronina et al., 2011). Thus, we can imagine a germ line module, comprised of a group of largely conserved proteins with essentially conserved functions and interactions, which collectively perform the cellular activities required of germ cells.

To specify a germ cell, therefore, is a matter of correctly co-expressing the proteins in germ line module. Seen this way, a germ line module is conceptually analogous to a "gene regulatory network," a familiar concept in evolutionary developmental biology (reviewed in Peter and Davidson, 2011). Gene regulatory networks are sets of co-regulated genes that respond to common upstream transcription factors; thus, a single so-called "master regulator" (a highly upstream transcription factor) can simultaneously deploy the expression of hundreds of downstream genes, which collectively effect a cellular phenotype. It is widely accepted that, due to the modular organization of gene regulatory networks, relatively major evolutionary change can be achieved through relatively minor changes in the expression of upstream transcription factors (reviewed in Stern and Orgogozo, 2008). For example, such novel structures as beetle horns and the pigmentation patterns on butterfly wings are believed to have evolved via novel expression domains of transcription factors that in turn deploy largely intact downstream gene regulatory networks (Moczek and Rose, 2009; Monteiro, 2012).

This leads to a simple conceptual model for how PGC specification mechanisms could evolve: while the PGC module itself remains largely intact across species, the upstream signal that assembles these factors changes over time. In the case of insects, my data supports the hypothesis that the PGCs of ancestral insects activated this module in response to secreted cell signals. (Indeed, a recent paper from the Extavour lab has identified BMP signaling as one of the pathways implicated in PGC specification in *Gryllus* [Donoughe et al., 2014]). During the course of insect evolution, likely near the base of Holometabola, the *oskar* gene acquired a novel expression domain in the oocyte, where it was became able to physically assemble the members

of the PGC module in a subcellular region of the oocyte, thus nucleating a germ plasm and establishing a continuity of PGC module protein expression between the generations<sup>1</sup>.

On a broader phylogenetic scale, the fact that germ plasm has independently arisen so many times in widely divergent taxa may be the result of the simple fact that embryos all physically begin as germ cells (gametes) of the previous generation, and are thus likely to express at least some genes of the PGC module during the earliest stages of development. Generating a germ plasm may simply require evolving a mechanism to maintain the expression of these genes into embryogenesis and localize them to a subset of embryonic cells (Extavour, 2007). For insects, my data implies that the evolution of this nucleating factor involved the co-option of *oskar* from a somatic role (perhaps in the nervous system) into a novel role in the germ line of holometabolous insects. As *oskar* and additional nucleating factors (such as *e.g. buckyball* in zebrafish and/or the PGL proteins of *C. elegans*) are characterized on a structural and biochemical level, it will be interesting to compare the common properties of these proteins.

Why has oskar been lost in so many lineages?

Given that *oskar* plays an indispensable role in the PGCs of *Drosophila* and *Nasonia*, why has it been repeatedly lost from so many other insect lineages? One explanation for this seeming paradox has been suggested by Lynch *et al.* (2011). In order for germ plasm to have evolved from an ancestral signaling-based PGC specification mechanism, these two processes must have coexisted within an individual for at least some portion of evolutionary history (an example of the so-called "transition model" *sensu* Extavour, 2007). That is, it is difficult to imagine the simultaneous evolution of germ plasm precisely coincident with the loss of signaling-based PGC

<sup>&</sup>lt;sup>1</sup> It is unknown whether the ability of *oskar* to physically recruit other germ line genes predates its expression in the germ line, or whether this represents an additional evolutionary change in the molecular function of *oskar* and/or it's interaction partners. It will be interesting, in the future, to test whether *oskar* has similar interaction partners in the neuroblasts of *Gryllus*.

specification. In those species where both mechanisms are present, there would be a level of redundancy in the specification of PGCs, which could then allow for evolutionary decay of either one or the other mechanisms in various lineages (and also perhaps for the existence of both mechanisms in some extant lineages, including the crustacean *Parhyale hawaiiensis* [MS Modrell, AL Price, J Havemann, CG Extavour, M Gerberding, and NH Patel, *in revision*], the wasp *Pimpla turionellae* [summarized in Lynch et al., 2011] and the sea urchin *Strongylocentrotus purpuratus* [Yajima and Wessel, 2011], all of which show some evidence for germ plasm yet can regenerate germ cells when PGC precursors are embryonically ablated). This would explain why, in some lineages of holometabolous insects, *oskar*-mediated germ plasm has become the exclusive mode of PGC specification, whereas in others *oskar* appears to have been lost altogether and PGCs are specified via other means (Lynch et al., 2011). We note that this scenario would also require that the functions of *oskar* in the nervous system would also have become dispensable in these lineages, perhaps also through the evolution of compensatory mechanisms.

The idea that redundancy can allow for divergence and eventual loss of an ancestral state, while maintaining the functional output of the system, has been explored both theoretically and experimentally (reviewed in Wagner et al., 2007). For example, an analysis of the *cis*-regulatory elements regulating ribosomal gene expression across the yeast phylogeny revealed that a novel regulatory site arose and eventually replaced an ancestral site in derived taxa, but that both sites remain present in several intermediate taxa (Tanay et al., 2005). Future studies examining the function of *oskar* in a wider variety of intermediate taxa between *Gryllus* and Holometabola will be very interesting to explore these ideas.

### Outlook

Given the fundamental differences we observe in PGC specification between *Drosophila* and

the species studied in this dissertation, many new questions arise. Here, I note three questions that I find particularly interesting for future research:

- 1. What is the mechanism for inductive PGC specification in *Gryllus*? My dissertation suggests that a maternally-supplied germ plasm is absent from *Gryllus* ovaries, but does not suggest possible pathways or mechanisms which could induce PGC fate. An important first step has recently been taken by others in the Extavour lab to show that *dpp* signaling is involved in PGC specification (Donoughe et al., 2014). Interestingly, the mouse orthologs of *dpp* (BMP-family ligands) specify PGC fate in the mouse embryo, raising the possibility that this pathway may in fact have an ancestral role in animal PGC specification. Further work to understand the mechanism of inductive PGC specification in *Gryllus* would be very interesting, as many questions remain. Given the extreme pleiotropy of the *dpp* pathway in embryonic development, how is its role in PGCs accomplished specifically? How is PGC fate restricted to abdominal segments 2-5? Are other pathways involved?
- 2. How are PGCs specified in additional insect species? The immense diversity of insects provides investigators with a wealth of opportunities to study evolution, yet this immense diversity also makes it perpetually difficult to generalize from findings from one species to other insects. Historical descriptions of insect embryology suggest PGCs arise in a wide variety of times and places during development (see Introduction), and modern studies using molecular markers and functional manipulations would greatly expand on these classical descriptions. Specifically, focusing on those hemimetabolous species described to have germ plasm and/or pole cells (see Introduction Figure 1.3) would be very interesting. Studying PGC specification in a wide diversity of insects would greatly improve our understanding of how this process evolves.
- 3. What new experimental tools would be most useful for future studies of germ

cells? Although non-model organisms provide important phylogenetic contrasts with such models as *Drosophila*, there are major experimental limitations to working with such organisms. However, with the recent emergence of (nearly) affordable technologies to sequence and annotate genomes, make stable transgenic lines, and knock-in/knock-out genes using such genome-editing techniques as CRISPR, entire new avenues of experimentation are becoming available. A fluorescent reporter of PGCs in Gryllus, such as a Gb-Piwi or Gb-Vasa reporter, would be an extremely valuable tool, and is being developed by Seth Donoughe and Taro Nakamura in the Extavour lab. Currently, screening for PGC phenotypes is time-consuming, as embryos must be dissected, fixed, stained via antibody staining or *in situ* hybridization, and imaged via confocal microscopy. Being able to visualize PGCs in a faster and easier way would allow future researchers to greatly expand the scope of manipulations they test for effects on PGC formation. In mouse PGC research, such tools as Blimp1 reporter lines have allowed for an explosion in mechanistic studies of inductive PGC specification. In addition, such a tool may allow for a more detailed description of PGC specification, as this process has thus far only been examined in fixed tissue, at various snap-shots of development, as levels of Gb-Piwi and Gb-Vasa become differentially higher in PGCs during stages 5-7 of Gryllus development. Live imaging of PGC formation is likely to be quite technically challenging due to the fact that the embryo is deep within yolk and is undergoing significant physical movements during this time. However, such an approach could yield a far more detailed picture of PGC development than is currently known.

This dissertation has also provided evidence that *oskar*, once thought to be a Dipteranspecific novel gene functioning solely in the germline, in fact evolved quite early in insect

evolution and has an additional function in the nervous system of *Gryllus*. In light of recently published work demonstrating a neural role for *Drosophila oskar*, it seems likely that this neural role may in fact be its ancestral function. However, many questions still remain regarding the enigmatic *oskar* gene.

- 1. Is oskar present in additional insect genomes? As additional insect genomes are sequenced, it will be very interesting to know where oskar is found in the insect tree (and possibly other arthropods). This study is currently being initiated by Tamsin Jones, a graduate student in the Extavour lab. These data will provide far greater resolution to say when oskar first arose, and where it has been lost. Furthermore, this study will identify a number of additional insect species that should be studied in detail, specifically to know whether oskar functions in the nervous system, germline, or both.
- 2. What is the molecular function of *oskar* in the nervous system? Given the ambiguity of the *norka* "allele" of *oskar* (see Appendix 2 of this dissertation), a first priority should be to test for behavioral phenotypes in *bona fide* mutants of *oskar*, to know whether this gene does indeed have a function in olfactory learning and/or other neural roles in *Drosophila*. If so, this would provide a new system in which to probe *oskar* function, which has thus far proved remarkably recalcitrant. Does *oskar* function during neural stem cell divisions in *Drosophila*? Does it function in the formation of ribonucleoprotein granules within specific neurons (which are known to contain such Oskar-associated proteins as Staufen, Pumilio, and Orb; see Chapter 5)? Or, is the neural function of *oskar* in *Gryllus* simply distinct from that of its roles in *Drosophila*?

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## The Molecular Machinery of Germ Line Specification

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#### SUMMARY

Germ cells occupy a unique position in animal reproduction, development, and evolution. In sexually reproducing animals, only they can produce gametes and contribute genetically to subsequent generations. Nonetheless, germ line specification during embryogenesis is conceptually the same as the specification of any somatic cell type; germ cells must activate a specific gene regulatory network in order to differentiate and go through gametogenesis. While many genes with critical roles in the germ line have been characterized with respect to expression pattern and genetic interactions, it is the molecular interactions of the relevant gene products that are ultimately responsible for germ cell differentiation. This review summarizes the current state of knowledge on the molecular functions and biochemical connections between germ line gene products. We find that homologous genes often interact physically with the same conserved molecular partners across the metazoans. We also point out cases of nonhomologous genes from different species whose gene products play analogous biological roles in the germ line. We suggest a preliminary molecular definition of an ancestral "pluripotency module" that could have been modified during metazoan evolution to become specific to the germ line.

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Sexually reproducing animals must ensure that one particularly important cell type is determined: the germ cells

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## INTRODUCTION

Across the plant and animal kingdoms, embryogenesis is that crucial developmental phase during which a single pluripotent cell, the fertilized ovum, must divide and differentiate to produce a plethora of differentiated, unipotent cell types. Sexually reproducing animals must ensure that one particularly important cell type is determined: the germ cells. These cells will be the sole progenitors of eggs and sperm in the sexually mature adult, and as such, their correct specification during embryonic development is critical for reproductive success and species survival. Germ cells and their embryonic origin have fascinated biologists for centuries, resulting in an enormous amount of primary literature on the subject (last comprehensively reviewed by Nieuwkoop and Sutasurya,

1979, 1981). The time and place when germ cells are first observed in embryogenesis, their histological and cytological characteristics, and the results of experimental manipulation of embryos on germ cell formation have been described in great detail for dozens of different species across the metazoa. All studies coincide in their observation of germ cell-specific cytoplasmic inclusions, visible under transmitted light and electron microscopy alike. Molecular studies from the last three decades have shown that this special cytoplasm, often called germ plasm, houses germ cell-specific gene products. Several excellent reviews have examined germ cell formation in specific animals (Saffman and Lasko, 1999; Raz, 2003; Strome, 2005; Hayashi et al., 2007; Saitou, 2009), the genetic mechanisms of specific germ line specification modes (Houston and King, 2000a; Strome and Lehmann, 2007), general molecular

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characteristics of germ cells (Seydoux and Braun, 2006; Cinalli et al., 2008; Nakamura and Seydoux, 2008), or the function of germ cell-specific genes (Raz, 2000; Noce et al., 2001). As we strive to put biological processes into an evolutionary perspective, however, we now need to begin to consider the ancestral histories of not just germ cell-specific genes themselves, but also their molecular interactions and collective functions in the germ plasm. While it is clear that many of these genes are conserved across metazoa, it is less clear to what extent the specific molecular interactions of these mRNAs, proteins, and cellular organelles have changed or remained the same throughout evolution.

The recent molecular revisitation of classical comparative embryology, otherwise known as evolutionary developmental biology or "evo-devo," has clarified a key paradigm that is relevant to the germ cell problem in this context. It is now possible, and moreover useful, to speak of molecular modules comprising gene regulatory networks (GRNs) (see e.g. Davidson et al., 2002). Such modules consist of a group of genes whose genetic interactions, or physical interactions of their gene products, are highly biochemically stable and thus highly conserved. The result of this genetic and molecular interaction stability is that the same batteries of genes, or modules, are found to operate in similar ways both in different organisms, and in different places and/or times during the development of a single organism (discussed in Wagner et al., 2007; Monteiro and Podlaha, 2009). The Notch-Delta pathway, for example, is a ligand/receptor-activated signal transduction pathway that ultimately regulates gene expression at the level of transcription. All members of this pathway are both highly conserved and operate together in all metazoans (reviewed by Kopan and Ilagan, 2009). Over the course of animal development, this module participates in a wide variety of developmental processes, including segmentation, neuroblast specification, and stem cell maintenance (reviewed by Artavanis-Tsakonas et al., 1999; Lai, 2004).

We can therefore ask, in the case of germ line-specific molecules, if it is possible to identify a group of genes that are not only highly conserved, but whose products also display conserved molecular interactions. If so, does this putative "module" also participate, like the Notch pathway, in a variety of developmental decisions, or is it confined to germ line specification? In this review, we will establish a framework for answering these questions by reviewing and summarizing recent data on the molecular functions and interactions of several genes that are critical for germ cell specification. Because examining all known genes involved in the process is beyond the scope of this review, several genes that are conserved in animal genomes, but whose role in germ line specification is either poorly understood or likely to be indirect, are indicated in Table 1, but not discussed further. Instead, we have focused on a subset of genes whose germ line specificity and critical roles in specification are well established. Some of these genes are highly conserved across the Metazoa, while for others, either their presence in the genome or their role in germ line specification, are lineage-specific.

# CONSERVED MOLECULAR COMPONENTS OF GERM LINE SPECIFICATION AND DIFFERENTIATION

#### Vasa

Products of the *vasa* gene family are the most widely used molecular germ cell markers for the Metazoa (discussed in Raz, 2000; Noce et al., 2001; Extavour and Akam, 2003). Vasa proteins are ATP-dependent RNA helicases of the DEAD box class, which was originally identified as a helicase family based on conservation of eight functional domains (Linder et al., 1989). DEAD box helicases are generally involved in RNA metabolism and can mediate both RNA–RNA and RNA–protein interactions (Rocak and Linder, 2004). A significant body of functional data for these helicases exists, based largely on studies of yeast DEAD box proteins (reviewed by Rocak and Linder, 2004). However, much less is known about the specific molecular function of Vasa, the germ cell-specific member of this class.

The vasa (vas) locus was first identified in *Drosophila* in a screen for maternal effect genes involved in anterior–posterior axis formation (Schüpbach and Wieschaus, 1986). *Drosophila* Vasa protein localizes to cytoplasmic granules within pole plasm (Lasko and Ashburner, 1988; Hay et al., 1988a,b), and localization of the mRNA, protein, or both to germ plasm and germ cells at some stage of development is a universal characteristic of the *vasa* gene family (see e.g. Lasko and Ashburner, 1988, 1990; Hay et al., 1988a,b; Fujiwara et al., 1994; Komiya et al., 1994; Ikenishi and Tanaka, 1997; Yoon et al., 1997; Braat et al., 2000; Knaut et al., 2000; Tanaka et al., 2000; Toyooka et al., 2000; Özhan-Kizil et al., 2009).

Localization of vasa gene products to germ plasm is consistent with its loss-of-function phenotypes in *Drosophila*, which are loss of or defective primordial germ cells (PGCs; also called pole cells in Drosophila) (Lasko and Ashburner, 1990), with additional oogenesis defects seen for null mutations (Styhler et al., 1998). Similarly, nematode (Gruidl et al., 1996; Kuznicki et al., 2000; Spike et al., 2008), frog (Ikenishi and Tanaka, 1997), flatworm (Ohashi et al., 2007), crustacean (Özhan-Kizil et al., 2009), tunicate (Sunanaga et al., 2007), and mouse (Tanaka et al., 2000) vasa knockdowns or mutants show germ line defects at various stages of germ cell development, including gametogenesis. In zebrafish, however, morpholino-mediated protein knockdowns of Vasa affect neither germ cell number nor fertility (Braat et al., 2001). While vasa is almost always required for some stage of germ cell development, in no animal has it been shown to be sufficient (see, e.g., Ikenishi and Yamakita, 2003). However, a recent study (Lavial et al., 2009) has shown that experimentally induced vasa expression can reprogram chicken embryonic stem cells and direct them toward a germ cell fate. This suggests that vasa might be able to function as a germ cell determinant for cells that are already pluripotent.

Genetic interactions between *vasa* and other germ line genes have suggested a complex network of positive and negative regulation at multiple levels, including transcription, translation, and post-translational modification. In *Caenorhabditis elegans* and mice, various components of germ

cell-specific cytoplasmic aggregations such as P granules, chromatoid bodies, and nuage lose their localization in vasa mutants (Chuma et al., 2006; Hosokawa et al., 2007; Spike et al., 2008). Vasa's identity as an RNA helicase suggests a role in translational regulation, and indeed, higher levels of some proteins in vasa mutants (Johnstone and Lasko, 2004) and a physical and genetic interaction with a translation initiation factor (Carrera et al., 2000) are both consistent with this hypothesis. However, very few direct molecular interactors have been identified for Vasa to date, and most of them effect or stabilize its localization (but see Carrera et al., 2000; Johnstone and Lasko, 2004). The SOCS-box/SPRYdomain gene gustavus was identified in a yeast two-hybrid screen that used Drosophila Vasa protein as bait (Styhler et al., 2002). Gustavus is a highly conserved protein whose zebrafish homolog localizes to germ plasm (Li et al., 2009b), suggesting an ancient origin for this protein interaction. Drosophila gustavus mutants fail to localize Vasa protein to the germ plasm, and other identified binding partners of Vasa protein also appear to play a role in localization, rather than function, of Vasa. The novel protein Oskar (discussed below) and the ubiquitin-specific protease Fat facets (Liu et al., 2003) interact physically with Vasa, and both are required for Vasa's correct localization to germ plasm. Detailed studies of multiple vasa mutant alleles have shown that the RNA-binding domains of the Vasa protein are not necessary for its localization to the pole plasm, but are necessary for its germ cell function (Liang et al., 1994). While further work will be needed to identify the molecular partners and direct targets of Vasa's regulatory function, it is clear that Vasa co-localizes to the germ plasm together with several other highly conserved germ cell gene products. Those for which the most functional data are available are discussed in the following sections.

## Nanos and Pumilio

Orthologs of Nanos localize to germ cells of nearly all taxa studied (Extavour and Akam, 2003). The specific functions played by Nanos vary, but the phylogenetically widespread expression of these proteins in germ cells suggests that a germ line function of Nanos may have evolved very early in animals (Extavour and Akam, 2003; Extavour, 2007). Pumilio, which has orthologs in organisms as diverse as yeast and plants (Zamore et al., 1997), has been shown to physically interact with Nanos proteins in flies (Sonoda and Wharton, 1999), nematodes (Kraemer et al., 1999), frogs (Nakahata et al., 2001), and humans (Jaruzelska et al., 2003), implying that this interaction is ancestral in bilaterians.

Like vasa, nanos and pumilio were first discovered in Drosophila (Nüsslein-Volhard et al., 1987) where both genes play essential roles in abdominal patterning and germ cell survival (Nüsslein-Volhard et al., 1987; Lehmann and Nüsslein-Volhard, 1991; Wang and Lehmann, 1991; Kobayashi et al., 1996). The molecular functions of Nanos and Pumilio were first investigated in studies of their role in repressing anterior identity in Drosophila embryos (Irish et al., 1989; Tautz and Pfeifle, 1989), and subsequent biochemical studies have suggested mechanisms by which

these two proteins form a complex that binds target RNAs and regulates their translation.

The Nanos protein contains a highly conserved C-terminal domain encoding two CCHC zinc-finger domains that bind RNA with high affinity but low sequence specificity (Curtis et al., 1997). Specificity is provided through complexing with the conserved Puf domain of Pumilio (named for Pum and FBF, its ortholog in C. elegans), which binds specific sequences in the 3'-UTRs of target RNAs (Zamore et al., 1997; Zhang et al., 1997; Sonoda and Wharton, 1999). Structural analyses of Puf domains reveal a "rainbow-shaped molecule" formed of eight tandem helical repeats (Edwards et al., 2001), each of which usually binds a single RNA nucleotide (Wang et al., 2002; Miller et al., 2008; Gupta et al., 2009). Both Nanos and Pumilio proteins thus bind RNA and each other, and can conditionally recruit additional protein factors to regulate target RNAs (see. e.g., Sonoda and Wharton, 2001).

The mechanisms by which the Nanos/Pumilio complex regulate translation likely involve recruitment of the deadenylation machinery to target RNAs. In both flies and worms, binding of Nanos and Pumilio orthologs to target RNAs correlates with translational repression (Wreden et al., 1997; Zhang et al., 1997). In flies, such binding drives RNA deadenylation (Wreden et al., 1997), and Nanos itself has been shown to physically interact with the Ccr4p-Pop2p-Not deadenylase complex member Not4 (Kadyrova et al., 2007). Additionally, Puf proteins in yeast are able to bind Pop2, another member of this deadenylation complex (Olivas and Parker, 2000; Goldstrohm et al., 2006), suggesting that both Nanos and Pumilio have active roles in regulating translation.

Importantly, the ultimate roles played by Nanos and Pumilio orthologs vary in the germ cells of different organisms. In Drosophila, Nanos and Pumilio directly regulate many RNAs in migrating PGCs to repress somatic identity (Kobayashi et al., 1996; Deshpande et al., 1999; Hayashi et al., 2004), halt the cell cycle (Asaoka-Taguchi et al., 1999), and prevent apoptosis (Hayashi et al., 2004; Sato et al., 2007). Similarly, in C. elegans, nos-1 and nos-2 are not necessary for the initial formation of PGCs, but rather for their maintenance and survival during embryogenesis (Subramaniam and Seydoux, 1999). In contrast, the C. elegans NOS-3/FBF complex is not involved in germ cell development, but rather in the sperm-to-oocyte transition in hermaphrodites (Kraemer et al., 1999). In zebrafish (Koprunner et al., 2001) and mice (Tsuda et al., 2003) nanos-related genes are required for PGC survival in both sexes, but specific targets of Nanos are largely unreported in vertebrates.

In both *Drosophila* and *C. elegans*, Nanos is also genetically implicated in the maintenance of a specific "chromatin architecture" that is associated with general transcriptional repression (Schaner et al., 2003). However, it has been pointed out that the cytoplasmic localization of Nanos protein, as well as our knowledge of its molecular function, implies that this function may be indirect (discussed in Seydoux and Braun, 2006).

Finally, there is also evidence for Nanos proteins functioning in the absence of known interactions with Pumilio

proteins. As mentioned above, *C. elegans* NOS-3 physically interacts with the Puf protein FBF, but the two other nematode *nanos* orthologs, *nos-1* and *nos-2*, do not do so in a yeast two-hybrid assay (Kraemer et al., 1999). However, the *C. elegans* genome encodes eight Puf proteins, and knock down of several of these proteins produces PGC defects indistinguishable from those observed in *nos-1* and *nos-2* knock downs (Subramaniam and Seydoux, 1999), suggesting that these two *nanos* orthologs could interact with other Puf proteins. In *Drosophila*, protein expression and detailed mutant analysis suggest that Nanos and Pumilio may have nonoverlapping roles in early oocyte development (Forbes and Lehmann, 1998). Therefore, while most of the studied roles of Nanos involve Pumilio-related proteins, it remains to be seen how the two may function in each other's absence.

#### Tudor

The "grandchildless" phenotype of tudor mutants was first described in Drosophila by Boswell and Mahowald (1985). tud- mutants do not maintain expression of germ granule components Oskar and Vasa (Thomson and Lasko, 2004), form abnormal germ granules, and ultimately fail to produce pole cells (Boswell and Mahowald, 1985; Thomson and Lasko, 2004). Proteins containing the so-called Tudor domains have since been found in organisms ranging from yeast to humans (Ponting, 1997), and Tudor proteins localize to germ granules of flies (Arkov et al., 2006), zebrafish (Mishima et al., 2006; Strasser et al., 2008), and male mice (Chuma et al., 2006; Hosokawa et al., 2007). In flies, Tudor protein also localizes between mitochondria and germ granules, and is required for transferring ribosomal RNAs from the mitochondria (Table 1) to germ granules, an essential process in germ cell specification (Amikura et al.,

Insight into the molecular basis for Tudor function has come from studies of the protein and its interactors. Studies of Tudor domain proteins in humans revealed that these domains interact with methylated arginine and lysine residues of diverse protein partners, including Sm proteins of the spliceosome (Friesen and Dreyfuss, 2000; Brahms et al., 2001). Recent studies suggest that Tudor's interactions with methylated proteins, as well as with proteins of the methylosome itself, may be required for the formation of germ granules. Localization of Drosophila Tudor to germ granules genetically requires the activity of the methylosome components capsuleen (also called dart5, the fly ortholog of human dPRMT5) and Valois (the fly ortholog of human MEP50) (Anne and Mechler, 2005; Gonsalvez et al., 2006). Further, Tudor can bind both Capsuleen and Valois in vitro, and these latter two proteins methylate Sm proteins, with which Tudor also physically interacts (Anne and Mechler, 2005). As Seydoux and Braun (2006) have pointed out, Sm proteins are common components of germ granules from vertebrates to C. elegans, and have been shown to be required for P granule localization and function in C. elegans (Barbee et al., 2002; Barbee and Evans, 2006), suggesting that the role of Tudor in assembling germ granules may involve its association with methylosome components and Sm proteins.

#### A Link Between Tudor and PIWI-Family Proteins

An additional role for Tudor was recently suggested by the finding that PIWI-family proteins in mice, frogs, and flies contain symmetrically methylated arginine residues of the type recognized by Tudor proteins (Kirino et al., 2009; Vagin et al., 2009). In the *Drosophila* ovary, PIWI-family proteins (discussed below) require capsuleen-dependent methylation in order to maintain their own expression and to maintain wild-type levels of piRNAs. In addition, capsuleen mutant ovaries accumulate abnormally high levels of retrotransposons that are normally silenced by PIWI-family proteins (Kirino et al., 2009). In mice, the three PIWI proteins were recently shown to directly interact with the methylosome complex of PRMT5 and WDR77/MEP50. This complex methylates arginine residues of Mili, Miwi and Miwi2, which in turn interact with Tudor domain-containing proteins. Additionally, specific PIWI and Tudor proteins also colocalize to nuage components (Vagin et al., 2009, Wang et al., 2009). Together, these results suggest a previously unrecognized connection between the interacting networks of proteins and RNAs in germ granules.

The PIWI family of proteins (called Piwi, Aubergine, and Ago3 in *Drosophila*; Ziwi and Zili in zebrafish; Miwi, Miwi2, and Mili in mice; and Xiwi, Xili, and Xiwi2 in frogs) were named for a founding member (*P*-element-induced wimpy testis) uncovered in a screen for genes involved in maintaining germ line stem cells in the *Drosophila* ovary (Lin and Spradling, 1997). This protein family has since been intensively studied for its role in silencing retrotransposons in the germ line through interactions with a special class of small RNAs called piRNAs or rasiRNAs (reviewed by Hartig et al., 2007).

Specific roles for PIWI proteins in the specification and/or maintenance of germ cells have been suggested by mutant analyses. Piwi mutant flies have reduced numbers of pole cells (Megosh et al., 2006). Both Piwi and Aubergine localize to germ granules in nurse cells and pole cells, and Piwi physically interacts with Vasa until Piwi translocates to the nucleus, where it remains throughout germ cell migration and gametogenesis (Megosh et al., 2006). In zebrafish. ziwi RNA (Tan et al., 2002) and protein (Houwing et al., 2007) co-localize with Vasa to germ line-specific ribonucleoprotein complexes (RNPs), and ziwi mutants are agametic owing to progressive apoptosis of germ cells (Houwing et al.. 2007). In mice, mutants for miwi, mili, or miwi2 fail to complete spermatogenesis, although these genes are not required for female germ line development (Deng and Lin, 2002; Kuramochi-Miyagawa et al., 2004; Carmell et al., 2007). Recently it has been shown that Miwi and Miwi2 also form a complex with Mvh, the mouse vasa homologue (Vagin et al., 2009).

Importantly, although the PIWI-family proteins are best known for their germ cell function, some members of the related Argonaute family localize not only to germ plasm, but also to somatic RNA processing organelles such as P bodies (see, e.g., Lin et al., 2006). P bodies and germ plasm granules may thus contain organelles with closely related roles in RNA processing in both germ line and soma. While it is clear that the germ line and somatic organelles are

TABLE 1. Additional Genes With Roles in Germ Cell Specification and/or Function

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Gene name	Functional gene product(s)	Molecular/genetic germline role demonstrated	Molecular function	Descriptive summary	References
Maelstrom	Protein	Dm, Mm	Putative protein binding	Nuage localization; piRNA pathway	Costa et al. (2006) Findley et al. (2003), Lim and Kai (2007), Soper et al. (2008), Zhang et al. (2008)
Par-1	Protein	Dm, Mm, Ce	Kinase	Cytoskeletal polarization; asymmetric cell division; Osk/PIE-1 stabilization	Cheeks et al. (2004), Doerflinger et al. (2003, 2006), Guo and Kemphues (1995), Reese et al. (2000), Riechmann et al. (2002), Shulman et al. (2002), Shulman et al. (2000), Tian and Deng (2009), Vaccari and Ephrussi (2002), Zimyanin et al. (2007)
Mitochondrial rRNA	RNA	⊼, ×	Ribosomal component	Polar granule localization; rescues UV-ablated PGCs	Amikura et al. (2001, 2005), Kashikawa et al. (2001), Kloc et al. (2001, 2002), Kobayashi et al. (1993), Kobayashi and Okada (1989)
Germ cell less	Protein	Dn, Mn	Protein binding	Cell cycle and transcriptional regulation; PGC nuclear envelope localization	de la Luna et al. (1999), Jongens et al. (1992), Kimura et al. (1999, 2003), Leatherman et al. (2000), Li et al. (2006), Nii et al. (2001), Scholz et al. (2004)
Dead end	Protein	Mm, Dr, XI, Gg	RNA binding	Protects germline RNAs from miRNA-based degradation; vertebrate-specific	Kedde et al. (2007)
Staufen	Protein	Dm, Mm, XI	dsRNA binding	osk localization; germ plasm component maintenance	Ephrussi et al. (1991), Irion et al. (2006), Ramasamy et al. (2006), St Johnston et al. (1991, 1992), Thomas et al. (2009), Yoon and Mowry (2004)
Dazi	Protein	Dm, Mm, Dr, XI, Ce, Ol, Hs, Am	RNA binding	Germ plasm component	Anderson et al. (2007), Hashimoto et al. (2004), Houston and King (2000b), Houston et al. (1998), Johnson et al. (2001), Karashima et al. (2000), Kosaka et al. (2007), Moore et al. (2003), Reynolds et al. 2005, Saunders et al. (2003), Tung et al. (2006), Venables et al. (2007), Xu et al. (2007)

Am, Ambystoma mexicanum; Dm, Drosophila melanogaster, Mm, Mus musculus; Hm, Homo sapiens; Ol, Oryzias latipes; XI, Xenopus laevis; Ce, Caenorhabditis elegans; Gg, Gallus gallus; Dr, Danio rerio. Due to space limitations, we have focused here on those primary references that focus on the germline role of these gene products.

not identical, there is some overlap in the proteins and RNAs that they contain (Megosh et al., 2006; Gallo et al., 2008; Lykke-Andersen et al., 2008). These observations highlight the molecular similarities underpinning the functional analogies between RNA processing organelles in both germ cells and somatic cells, and are consistent with the hypothesis that RNPs are repressive regulatory organelles with an ancient eukaryotic history, predating the origin of the dedicated metazoan germline (see Eulalio et al., 2007 for a detailed review).

Silencing of transposable elements in the germ line is the most well-established role of PIWI-family proteins. In the above-listed PIWI-family mutations, germ cell failure is correlated with reduced piRNA levels and abnormal accumulation of transposable elements. This function is mediated through the interaction of PIWI-family proteins and additional factors with a special class of small RNAs that provide sequence specificity to a transcript-silencing complex (see Klattenhoff et al., 2007 for additional details). Two exciting studies have recently demonstrated a role for the piRNA pathway in silencing transposable elements in somatic cells of the gonad rather than in the germ cells themselves (Malone et al., 2009; Li et al., 2009a).

Among many other defects, PIWI family mutants also exhibit defects in maintaining the localization of essential germ granule components in *Drosophila*. For example, although *piwi* is not required for the initial expression of Oskar, Vasa, or Nanos, ectopic expression of Piwi protein is

able to recruit these maternal factors and increase their expression levels (Megosh et al., 2006), suggesting that Piwi acts in a positive feedback loop with these factors. Recent studies have also shown that Vasa localization genetically requires *aubergine* and *ago3*, and that these two proteins require one another for their own localization (Li et al., 2009a). Additionally, *aubergine*—mutants fail to properly localize the RNAi pathway members Krimper and Maelstrom (Lim and Kai, 2007). The mechanism by which PIWI-family proteins act to recruit and/or maintain localized expression of other factors to germ granules is unknown.

#### SYSTEM-SPECIFIC MOLECULAR COMPONENTS OF GERM LINE SPECIFICATION Rmps

In contrast to organisms where germ cell determination relies on the inheritance of germ plasm (reviewed by Extavour and Akam, 2003), in the mouse this process requires inductive signals (Tam and Zhou, 1996) (Fig. 1). The first germ cell-inducing signal in mouse embryos was identified only a decade ago as Bmp4, a member of the Bone morphogenetic protein family. Prior to PGC induction in the proximal epiblast, Bmp4 is expressed in the tissue directly adjacent to the epiblast, the extraembryonic ectoderm (ExE). This expression is essential for PGC determination, as Bmp4 mutant mice do not form PGCs (Lawson et al., 1999).

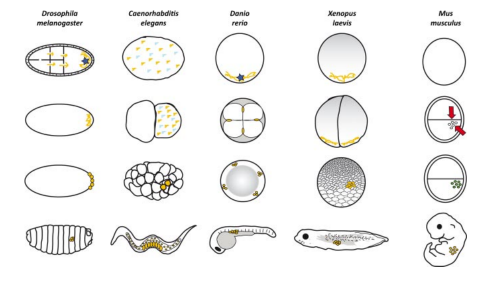


Figure 1. Localization of germ line specification molecules throughout animal development. Developmental progression in time goes from top to bottom. Germ cell specification and the localization of molecules discussed in this review are schematized for the five genetic laboratory organisms that have contributed the most to our current understanding of the molecular mechanisms underlying germ cell specification. In all organisms except for mice (*Mus musculus*), germ cell-specific gene products (yellow), including mRNAs and/or proteins of vasa, nanos, pumilio, piwi, and tudor, are localized to the cytoplasm of germ cells either late in oogenesis or early during embryogenesis. The fruit fly (*Drosophila melanogaster*) protein Oskar and the zebrafish (*Danio rerio*) protein Bucky ball (dark blue star) can autonomously assemble many of these germ plasm components in oocytes and early embryos. The nematode (*Caenorhabditis elegans*) protein PIE-1 (light blue) plays an important role in regulating germ line gene expression (yellow). In mice, somatic signals (red) trigger the expression of Blimp1 (green) in PGCs, followed by the expression of conserved germ line genes (yellow).

Similar to Bmp4, Bmp8b is also expressed in the ExE, and Bmp8b mutants lack or show very reduced numbers of PGCs. However, the effects of Bmp4 and Bmp8b are not additive (Ying et al., 2000). Very recently it was shown that the role of Bmp8 is very different from that of Bmp4. Rather than being directly required for induction of PGCs, Bmp8b signaling from the ExE restrains growth of the anterior visceral endoderm (AVE), which sends still-unidentified signals to the epiblast, thereby inhibiting Bmp4 (Ohinata et al., 2009).

Unlike Bmp4 and Bmp8b, Bmp2 is predominantly expressed in the visceral endoderm (VE). *Bmp2* mutants show a reduced number of PGCs, and Bmp2 and Bmp4 together have an additive effect on PGC development, while Bmp2 and Bmp8b together do not. Thus, Bmp signals from both the VE (Bmp2) and the ExE (Bmp4 and 8b) are required for PGC induction (Ying and Zhao, 2001).

Bmp ligands exert their function by binding and recruiting transmembrane type I and II Bmp serine/threonine kinase receptors on the cell surface. These receptors transmit the Bmp signal by phosphorylating Smad1, Smad5, or Smad8, which enter the nucleus as heterodimers with Smad4 and then serve as transcriptional regulators (Shi and Massagué, 2003). The precise downstream molecules that transduce Bmp signals and result in PGC formation are still largely unknown. There are only reports of three type I Bmp receptors possibly being involved in PGC determination. Alk2. Alk3, and Alk6 (de Sousa Lopes et al., 2004; Ohinata et al., 2009). Several Smads have been shown to be involved in PGC formation. Smad4, Smad1, and Smad5 mutants show reduced numbers or a complete lack of PGCs (Chang and Matzuk, 2001; Tremblay et al., 2001; Hayashi et al., 2002; Chu et al., 2004: Arnold et al., 2006), Furthermore, Smad1 and Smad5 are sufficient for PGC induction in combination with Bmp4 and Alk3 or Alk6 (Ohinata et al., 2009). However, a type II receptor involved in this process, as well as genes that are directly regulated by the Smads during PGC determination, remain to be identified.

Bmp signals have not yet been reported to be involved in germ line specification outside of mice, even though the epigenetic mode of germ cell formation has been hypothesized to be ancestral in metazoans (Extavour and Akam, 2003). More investigation will be necessary to provide functional evidence for or against this hypothesis, to determine whether the Bmp signal is conserved for germ cell induction, or whether different signaling pathways can be used by different animals to induce germ cells.

### Blimp1

The earliest marker of mouse PGCs, Blimp1 (B lymphocyte induced maturation protein 1, also known as Prdm1) was discovered only recently (Ohinata et al., 2005). It was first described in the context of plasma cell differentiation (Turner et al., 1994). The histone methyl transferase Blimp1 contains a PR domain and a proline-rich region at the Nterminus, five C2H2 zinc fingers, and a C-terminal acidic domain (Turner et al., 1994; Tunyaplin et al., 2000). Blimp1 orthologs are found in many bilaterian animals (see e.g. de Souza et al., 1999; Tunyaplin et al., 2000; Hinman and Davidson, 2003; Ng et al., 2006; Arenas-Mena, 2008), but

expression data do not suggest a role in germ cell specification outside of mammals. Among a variety of lethal defects, Blimp1 mutants exhibit only a very small number of PGC founder cells that fail to proliferate or migrate (Ohinata et al., 2005; Vincent et al., 2005; Robertson et al., 2007). These PGC-like cells do not show the wild-type pattern of Hox gene repression (Ohinata et al., 2005), and Blimp1 mutant cells fail to repress other somatic genes (Kurimoto et al., 2008). Consistent with this observation, in other cellular differentiation processes, Blimp1 has been observed to act as a transcriptional repressor that recruits a complex of Groucho-family proteins (Ren et al., 1999) as well as a histone deacetylase (Yurke et al., 2000). The molecular interactions of Blimp1 during germ cell specification are still unknown. Interestingly, Blimp1 co-immunoprecipitates with the mammalian ortholog of Drosophila Capsuleen/dart5 (Prmt5, discussed above), which is required for Tudor localization. This complex of Blimp1 and Prmt5 might play a role in germ line maintenance during PGC migration, as both proteins co-localize to the nucleus during this process. After migration they co-localize in the cytoplasm, which coincides with H2A/H4R3me2s downregulation in PGCs (Ancelin et al., 2006). As mentioned above, fly Capsuleen also binds Tudor and is necessary for its localization to the germ plasm in Drosophila. However, whether mouse Tudor-related proteins bind the Blimp1-Prmt5 in a functional complex during PGC specification remains an open question.

#### Pie-1 and Polar Granule Component

In *C. elegans*, one of the best-understood germ line determinants is the PIE-1 protein. The *C. elegans pie-1* mutant (*p*haryngeal and *i*ntestinal excess) was discovered in a screen as being required for the PGC precursor (Fig. 1) to follow germ line rather than somatic fate (Mello et al., 1992). The PIE-1 protein contains two CCCH zinc fingers (ZF1 and ZF2), which are separated by arginine—serine dipeptide repeats. PIE-1 is expressed maternally, is then asymmetrically distributed to the germ line blastomeres, and continues to be expressed in the germ line throughout development (Mello et al., 1996).

The ZF2 domain of PIE-1 is required for the translation of NOS-2 protein from maternal nos-2 mRNA in the germ line (Tenenhaus et al., 2001). However, PIE-1's principal role is to mediate transcriptional repression in the C. elegans germ line from P2 onwards (Seydoux et al., 1996). mRNA transcription requires phosphorylation of the C-terminal domain (CTD) of RNA polymerase II on Serine 5 (Ser5) for transcriptional initiation, and on Serine 2 (Ser2) for elongation (reviewed in Peterlin and Price, 2006; Saunders et al., 2006; Corden, 2007). The PIE-1 C-terminal region contains a motif that resembles the CTD, but has no phosphorylatable sites. This CTD-like motif, an HLX homology region, and additional C-terminal repeats, are sufficient for transcriptional repression (Batchelder et al., 1999). The PIE-1 CTD-like domain is thought to directly compete with the CTD for binding cyclin T (CycT), thereby inhibiting Ser2 phosphorylation of the CTD by the kinase CDK9 (Cyclindependent kinase 9). CycT and CDK9 together form the

positive transcription elongation factor b (P-TEFb) (Zhang et al., 2003). The CTD-like motif of PIE-1 is essential for inhibiting Ser2 phosphorylation, but does not play a role in Ser5 phosphorylation; the latter activity is mediated by sequences around the CTD-like motif (Ghosh and Seydoux, 2008)

Pgc (polar granule component) accomplishes a similar transcriptional repression function by inhibition of RNA PollI via P-TEFb inhibition in Drosophila pole cells (Nakamura et al., 1996; Martinho et al., 2004). However, the 71-aminoacid Pgc protein does not bear any resemblance to C. elegans PIE-1, even though it interacts with P-TEFb and represses CTD Ser2 phosphorylation, thereby inhibiting its recruitment to transcription sites (Hanyu-Nakamura et al., 2008). Furthermore, pgc is essential only for germ cell migration and not for pole cell specification (Nakamura et al., 1996: Martinho et al., 2004). This is therefore an interesting case of independent evolution of two unrelated proteins that play an analogous role in the same molecular pathway. The PIE-1/Pgc relationship parallels that of two other proteins involved in germ line specification, Oskar and Bucky ball (discussed below).

#### oskar and bucky ball

All genes discussed thus far are either components of germ plasm (vasa, nanos, pumilio, piwi, tudor, pie-1, Blimp1), or molecules that induce the accumulation of germ plasm components (BMPs). In the case of the germ plasm components described above, these genes are necessary but not sufficient for germ cell specification and function. The BMPs and their downstream effectors, in contrast, are both necessary and sufficient for germ cell specification, but are not themselves germ plasm components. There do exist, however, two molecules that are not only germ plasm components, but are also both necessary and sufficient for germ plasm formation. These genes, oskar and bucky ball, are lineage-restricted genes with independent, recent evolutionary histories, whose shared molecular function makes them of special interest in the context of this review.

oskar (osk) mRNA accumulates in the posterior cytoplasm during oogenesis in Drosophila (Ephrussi et al., 1991; Kim-Ha et al., 1991), and its translation is likewise confined to the posterior germ plasm (Kim-Ha et al., 1995). Loss-offunction mutants do not form germ cells (Lehmann and Nüsslein-Volhard, 1986). The sufficiency of osk for germ plasm assembly and germ cell formation was demonstrated in elegant experiments that drove osk expression in ectopic embryonic locations (Ephrussi et al., 1991). This showed that osk gene products can autonomously recruit germ plasm components, resulting in ectopic germ cells that are capable of functional gametogenesis (Ephrussi et al., 1991). In vas or tud mutants, however, ectopic osk does not lead to ectopic PGCs (Ephrussi and Lehmann, 1992), consistent with the hypothesis that the role of Osk is to recruit germ plasm components rather than to induce PGC fate directly.

osk mRNA localizes to the posterior pole during stages 8–10 of oogenesis, via a mechanism that requires Staufen (Table 1), microtubules, and the plus end-directed motor

protein kinesin (Lehmann and Nüsslein-Volhard, 1986; Brendza et al., 2000; Zimyanin et al., 2008). osk translation is confined to the posterior cytoplasm both by positive regulation of localized transcripts and by negative regulation of unlocalized transcripts (Kim-Ha et al., 1995; Wilson et al., 1996; Micklem et al., 2000; Chekulaeva et al., 2006; Klattenhoff et al., 2007; Klattenhoff and Theurkauf, 2008), When osk mRNA is translated, alternative start codons in the osk message result in two isoforms of Osk protein. Short Osk and Long Osk, which have separable roles in germ plasm assembly (Markussen et al., 1995; Rongo et al., 1995; Breitwieser et al., 1996; Vanzo and Ephrussi, 2002). Both Oskar isoforms are phosphorylated by the Par-1 kinase (Table 1) (Riechmann et al., 2002), which is enriched at the posterior in an actin-dependent but microtubuleindependent step during oogenesis (Doerflinger et al., 2006). Par-1-dependent phosphorylation is thought to stabilize Osk protein in the pole plasm (Riechmann et al., 2002), where it recruits Par-1 and thereby participates in a positive feedback loop that reinforces its posterior localization (Shulman et al., 2000; Zimyanin et al., 2007).

oskar's highly upstream position in the *Drosophila* germ cell specification pathway stems from its ability to ectopically induce germ plasm assembly (Ephrussi et al., 1991; Ephrussi and Lehmann, 1992). Accordingly, Short Osk protein has been shown to directly interact with Staufen and Vasa proteins (Breitwieser et al., 1996), and recruits *nanos* mRNA (Ephrussi et al., 1991; Kim-Ha et al., 1995).

The recently described zebrafish gene bucky ball (buc) has biological properties that are remarkably similar to those of oskar. buc transcripts are localized to the vegetal pole during oogenesis, together with other germ cell-specific molecules, and Buc protein is subsequently localized to germ plasm in early cleavage stage embryos (Marlow and Mullins, 2008). Loss-of-function mutations in buc lead to defects in both anterior-posterior patterning (Marlow and Mullins, 2008) and germ cell formation, including a failure of vas, dazl, nos, and buc mRNAs, and other germ plasm organelles, to localize to germ cells (Bontems et al., 2009). Ectopic expression of buc in non-germ line cell lineages of early embryos results in supernumerary germ cells that are derived from the cells containing ectopic buc (Bontems et al., 2009). In summary, buc, like osk, appears to be both necessary and sufficient for germ plasm assembly and germ cell specification.

These two genes share another striking similarity: they are both very recently evolved and do not contain any recognizable functional domains. Both genes encode novel proteins, and while *osk* is restricted to the Diptera (two-winged flies), *buc* is restricted to the vertebrates (Bontems et al., 2009). Despite the presence of germ plasm and pole cells in several other insects, *osk* is not found in nonfly insect genomes (Extavour, unpublished observations). The fact that the Diptera are the insect order furthest removed from the last common insect ancestor suggests that *oskar* may be a recent evolutionary innovation associated with germ cell segregation only in this derived lineage. Similarly, early determination of germ cells is observed in some nonvertebrate deuterostome lineages, but *buc* is not found in their genomes. Despite their evolutionary unrelatedness and

lineage restriction, their biological function is highly similar. This appears to be because their molecular interactors, all of which are conserved across metazoans, are the same in both cases: for example, both Osk and Buc recruit gene products of the vasa and nanos loci to form germ plasm. More biochemical studies on the transcriptional and translational regulation of buc, and on its direct physical molecular interaction partners, will be necessary to determine the extent of the apparent similarity between the biological functions of buc and osk. Given that orthologs of buc exist across vertebrates, it will be fascinating to see whether this gene plays a role in germ cell specification of mice, whose germ cells are specified through inductive signalling rather than the cytoplasmic inheritance of germ plasm. It is conceivable, for example, that interactions between buc and germ line factors are conserved in the germ cells of mice, but that the expression of buc itself is induced through BMP signalling rather than through the localization of maternal buc.

#### **SUMMARY**

We have seen that many genes involved in germ cell specification are conserved across evolution, and expression studies have demonstrated some similarities in their modes of localization to germ cells (Fig. 1). Moreover, these molecules often also interact biochemically in similar ways in phylogenetically distant animals. The Nanos/Pumilio complex, the Tudor domain/PIWI family interaction, and the Tudor/spliceosome component association may therefore represent evolutionarily ancient interactions. In other cases, proteins that are not homologous serve analogous functions in germ cells: PIE-1 in nematodes and Polar granule component in flies both regulate transcriptional elongation by inhibiting RNA polymerase II phosphorylation. Moreover, some of these molecules, and in some cases their molecular roles, are conserved not just in germ cells but are also found in pluripotent cells of many types, and in RNA-processing bodies of somatic cells (see e.g. Lin et al., 2006; Megosh et al., 2006; Gallo et al., 2008; Jud et al., 2008; Lykke-Andersen et al., 2008).

There is still too little biochemical information for us to be able to know how extensive a putative metazoan germ line GRN could be. However, the conserved protein-protein and protein-RNA interactions described above could represent components of an ancestral pluripotency module, which would have likely contained Tudor domain protein, PIWI family members, and a DEAD box helicase. In early multicellular animals where a pluripotent stem cell population produced gametes, those stem cells that entered into gametogenesis would have tailored this module by the addition of unique germ cell genes, such as nanos, vasa, and Aub/ Ago3. With the advent of dedicated germ cells in metazoans, this specialized germ line module would have come under the control of cytoplasmic inheritance or inductive mechanisms that operated exclusively in the germ line, preventing somatic cells (including somatic stem cells) from producing gametes.

One of the predicted consequences of modularity in development is that modules themselves can remain highly

conserved throughout evolution, while their upstream effectors and downstream targets can evolve independently. The germ line specification pathways fulfill this prediction: the robustness of the molecular interactions between the conserved germ line gene products links them together into a module that can be either induced by BMP signals (mouse), assembled autonomously in oocytes (nematode, frog), or possibly even nucleated by a single molecule (fruit fly, zebrafish). In fact, the mechanisms that localize germ line determinants to germ cells appear to be relatively flexible not only on an evolutionary time scale, but also even within developing individuals. A fascinating study has recently demonstrated that several germ line genes become ectopically expressed in somatic tissues of long-lived C. elegans mutants that lack insulin signalling (Curran et al., 2009). Although the restriction of germ line factors such as PIE-1 to the germ cells of *C. elegans* is normally achieved through asymmetric cell divisions, in this case the ectopic expression of PIE-1 is effected at the transcriptional level. Moreover, these somatically-expressed germ line factors appear to serve a crucial function, as knocking down any of these genes drastically reduces the longevity of these mutants. Such apparent flexibility in the spatial and temporal deployment of multiple functional germ line factors further supports the notion that such factors may operate as an interacting module, capable of being induced by a variety of upstream signals. Similarly, the downstream targets of germ line factors have evolved in lineage-specific ways.

Although we have been able to identify a few conserved molecular interaction motifs among germ line specification gene products, we still have far less biochemical data than we do genetic data on these mechanisms. We have knowledge of local interactions between a few pairs of molecules, but still lack information on how the entire suite of genes is linked together biochemically. To improve our definitions of the extent and limits of this modular network, many more biochemical studies, whose results are placed into evolutionary context, are needed.

In addition to molecular studies on the traditional laboratory model organisms that have provided us with the most data thus far, work on the physical interactions of germ line genes in "nonmodel" organisms will also be extremely informative and should be pursued in future. A thorough understanding of the genetic control of germ line development in any organism clearly requires adequate functional tools (Sommer, 2009). However, understanding the extent of the evolutionary conservation of biochemical interactions between germ line molecules is not dependant on a complete knowledge of the developmental genetics of germ line specification. Such studies therefore need not be confined to organisms for which functional genetic analysis tools have been established.

Finally, construction of a GRN for the germ line will require a somewhat different approach to those that have been undertaken thus far. Many of the powerful GRNs that have been constructed to model aspects of somatic differentiation rely largely on transcriptional regulation (see, e.g., Davidson et al., 2002; Loose and Patient, 2004; Koide et al., 2005; Imai et al., 2006). However, most of the molecules for which functional biochemical data are available appear to be

involved in translational regulation and protein–protein interactions, suggesting that post-transcriptional gene regulation is particularly crucial in the germ line (reviewed in Cinalli et al., 2008; see e.g. Merritt et al., 2008). Moreover, it is becoming increasingly apparent that several other recently discovered mechanisms of gene regulation play a critical, albeit not yet well-defined, role in the germ line. These include piRNA-mediated transposon silencing, RNP formation to repress translation, and chromatin architecture-mediated gene regulation. We may therefore need new ways of building GRNs in order to create a framework for understanding the molecular network of the germ line.

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15

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## Appendix B:

# Preliminary experiments regarding a possible role for *oskar* in the *Drosophila* nervous system

#### INTRODUCTION

In Chapter 2 and Chapter 5 of this thesis, I present evidence that *oskar* has a function in the nervous system of the cricket *Gryllus bimaculatus*. Specifically, I show that *Gb-oskar* functions in embryonic neuroblasts (Chapter 2), and also in neuroblasts present in the adult brain (Chapter 5). These data, together with reports of neural function of *oskar* in *Drosophila* (Dubnau et al., 2003; Xu et al., 2013) suggest that a role for *oskar* in the nervous system, but not in the germ line, is conserved between these two species, and may therefore be the ancestral function of *oskar*.

However, the precise roles of *oskar* in the *Drosophila* nervous system are not understood, making it unclear to what extent the specific neural roles are in fact conserved between *Gryllus* and *Drosophila*. Two studies have reported a phenotype for *oskar* in the *Drosophila* nervous system. Xu *et al* (2013) demonstrate that *osk* RNA particles, like *nanos* RNA particles, are motile in larval dendritic arborization (da) neurons, and further show that *osk* mutant larvae display defective motility of *nanos* particles. Furthermore, da-specific *osk* RNAi leads to defects in response to a mechanical stimulation (Xu et al., 2013). However, a direct comparison of this function between *Drosophila* and *Gryllus* is difficult to conceptualize, as hemimetabolous insects such as *Gryllus* do not pass through a larval stage at all.

The other report of a role for *oskar* comes from a forward genetic mutagenesis screen for defects in olfactory learning (Dubnau et al., 2003). However, these *oskar* data are ambiguous, as the putative *oskar* allele (dubbed *norka*), obtained via GAL4 insertional mutagenesis, does not map within the known *osk* locus (Fig 1). In fact, the *norka* insertion is markedly closer to *polychaetoid* (*pyd*), which encodes a cell adhesion molecule with known neural function in

development and in learning (Chen et al., 1996; Eddison et al., 2012; Seppa et al., 2008). In addition, the *norka* allele does not drive expression in the germ line (**Figure A.1**), where *oskar* has a well-characterized role (Ephrussi and Lehmann, 1992; Jenny et al., 2006). Despite these facts, Dubnau *et al* (2003) present northern blot data suggesting that the *osk* transcript is unusually long in *norka* mutants. Clarifying these issues will require additional studies with *bona fide* alleles of *osk*.

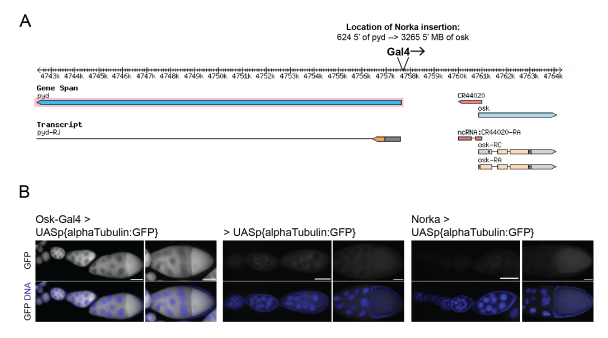


Figure A.1 The *norka* mutant may not be a true *osk* allele. (A) The genomic location of the Gal4 insertion in the *norka* allele. Data from (Dubnau et al., 2003). Note that the insertion is far upstream of *osk*, outside of any known transcription unit, but is quite close to *pyd*, a gene with several known neural functions (see text). (B) The *norka* GAL4 insertion does not drive GFP expression in the germ line, where Oskar is expressed. An Osk-Gal4 line (containing 1,743 bp of genomic sequence upstream of *oskar* fused to Gal4) drives expression throughout oogenesis (left panels), whereas *norka* (right panels) does not drive expression above background levels (compare to the no control, middle panel). Anterior is to the right. Scale bars =  $50 \mu m$ .

In a preliminary attempt to clarify some of these issues, and to test whether there may be a more direct conservation of *oskar* function between *Drosophila* and *Gryllus*, I have undertaken preliminary studies of *oskar* in the *Drosophila* nervous system. In this Appendix, I present data suggesting that Oskar protein is undetectable by antibody staining in the adult brain. I then show

that an Oskar-Gal4 line does not drive expression in neurons of the adult brain. Finally, I present preliminary data suggesting that neither *oskar* RNAi nor *oskar* over-expression in the embryonic neuroblasts disrupt CNS development, in contrast to *Gryllus*. In sum, these initial data do not indicate a function for *oskar* in the embryonic nervous system or in the adult brain, but these studies are far from complete. I suggest several additional experiments that could help extend these data.

#### **METHODS**

Drosophila strains

The following stocks were used:

osk<sup>norka</sup> (Dubnau et al., 2003) - gift of B. di Bivort, Harvard University.

*oskar-Gal4* – a 1,743-bp genomic region upstream of *oskar* fused to Gal4 (Telley et al., 2012) - (Bloomington 44241, 44242)

oskar<sup>54</sup>/oskar<sup>84</sup> – strong osk hypomorphic transheterozygote (Lehmann and Nüsslein-Volhard, 1986; Vanzo and Ephrussi, 2002) (gift of R. Lehmann, NYU)

oskar<sup>A87</sup>/Df(3R)PXT103 – osk null, over a deficiency (Jenny et al., 2006) (gift of E. Gavis, Princeton)

pJFRC7 =  $p{20XUAS-IVS-:mCD8::GFP}$ attP2 – Membrane-GFP reporter for somatic expression (Pfeiffer et al., 2010) (gift of J. Tuthill, R. Wilson Lab, Harvard Medical School)

UASp:GFP::alphaTub (w; p{w[+mC]=UASp-GFPS65C-alphaTub84B) – Cytoplasmic GFP reporter for germ line (Bloomington 7373)

UAS-*osk* (w; p[w+, UAS:osk]) – Line for *osk* over-expression (Zimyanin et al., 2007) (gift of D. St. Johnston, Gurdon Institute)

osk-RNAi – RNAi line for osk knockdown (VDRC 107546)

asense-Gal4 / CyO (Zhu et al., 2006) – Gal4 driver for neuroblasts (gift of T. Lee, Janelia Farms)

OK107 (w;  $P\{w[+mW.hs]=GawB\}ey[OK107]$  – Gal4 driver for the mushroom body. (Gift of W. Tobin, R. Wilson Lab, Harvard Medical School)

Antibody staining

Adult brains were dissected and de-sheathed using fine forceps in ice-cold PBS. Brains were fixed for 60-90 minutes in 4% paraformaldehyde, and were permeabilized with 1% Triton-X in PBS prior to beginning antibody staining following standard protocols. Primary incubation was overnight at room temperature, and secondary incubation was overnight at 4°C. Primary antibodies were *nc82* (1:50; labels neuropil, Iowa Developmental Hybridoma Bank), anti-Eve 28B (1:30, Iowa Developmental Hybridoma Bank), anti-HRP Alexa647 congujate (1:50, gift of S. Kunes), anti-Oskar (1:1500, preabsorbed against *Drosophila* embryos, gift of A. Ephrussi), and anti-GFP Alexa488 conjugate (1:250; Molecular Probes).

#### Embryonic CNS Scoring

Embryos were double-stained for Eve and HRP, then individually mounted on slides, oriented ventral side towards the coverslip. Confocal z-stacks covering the entirety of the embryonic CNS (all visible Eve+ and HRP staining) were captured, and were manually scored for any discernable defects in any of the EL, U/QC, RP2, aCC/pCC, and axonal scaffold. Between five and seven segments were scored per embryo, and the percentage of affected segments for each neuron cluster was used as the single data point for that embryo. Maximum intensity projections were made of representative embryos, but scoring was done on the full stacks to ensure that all Eve+ cells could be identified.

## **RESULTS**

Oskar protein cannot be detected in the adult brain by antibody staining

Dubnau *et al* (2003) report that the *norka* mutation represents an allele of *oskar*. Because the allele was generated via GAL4 insertional mutagenesis, they were able to use a UAS:GFP reporter to reveal expression driven by the *norka* insertion. They show that *norka* drives expression in a variety of neurons in the adult brain, including the mushroom body, the

anatomical substrate associated with olfactory learning and memory (see Figure 3C in Dubnau et al., 2003).

Given the ambiguity of whether the *norka* allele is a true *osk* allele, I wished to test whether Oskar protein could be detected in the mushroom body or other regions of the adult brain via antibody staining. Using an antibody that is specific to Oskar protein in ovaries (**Figure A.2A**), I was unable to detect specific signal in adult brains above background levels observed in two *osk* mutants (**Figure A.2B**). These data are not consistent with the conclusion that the expression driven by the *norka* GAL4 insertion captures wildtype Oskar expression, although it remains a formal possibility that levels of Oskar below the detection limit of antibody staining may function in the adult brain.

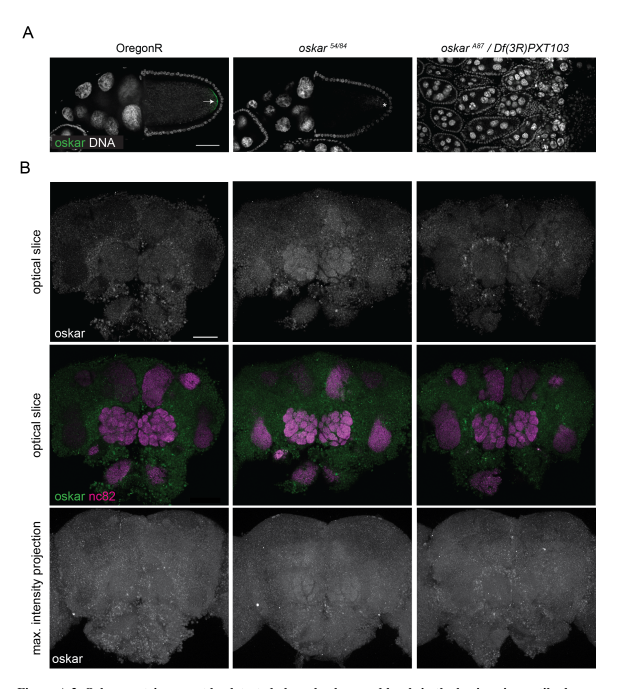


Figure A.2. Oskar protein cannot be detected above background levels in the brain using antibody staining. (A) anti-Oskar antibody is highly specific to Oskar protein in oocytes (left panel), and gives no signal in two strong *oskar* mutant ovaries (middle and right panels). Note that oogenesis arrests at stage 7 in oskar[A87] ovaries, as this is an RNA null (Jenny et al., 2006). Anterior is to the left. (B) In brains, Oskar protein cannot be detected above the background levels present also in oskar mutants. Signal is shown in single optical slices (top row), overlaid with the neuropil maker nc82 (middle row) to see brain structure, and in maximum intensity projection (bottom row). Scale bars = 50  $\mu$ m.

As an additional test for Oskar expression in the *Drosophila* brain, I crossed a published Osk-Gal4 line (made by fusing 1,743 bp of genomic DNA upstream of *oskar* to Gal4)(Telley et al., 2012) to a membrane-bound GFP reporter. A mushroom body driver, *OK107*, was used as a positive control, and a reporter-only fly was used as a negative control (**Figure A.3A**).

Osk-Gal4 did not drive GFP in any neurons of the adult brain (**Figure A.3B**). Surprisingly, GFP was instead observed in the non-neuronal sheath that surrounds the brain, which is typically ripped during dissection to aid in antibody penetration. The biological importance of this expression is unknown. Regardless, these experiments are consistent with the antibody staining (**Figure A.2**) and suggest that Oskar expression is either absent from the neurons of the adult brain, or, if present at low level, is driven by an enhancer outside of the 1,743 bp upstream of the *oskar* locus.

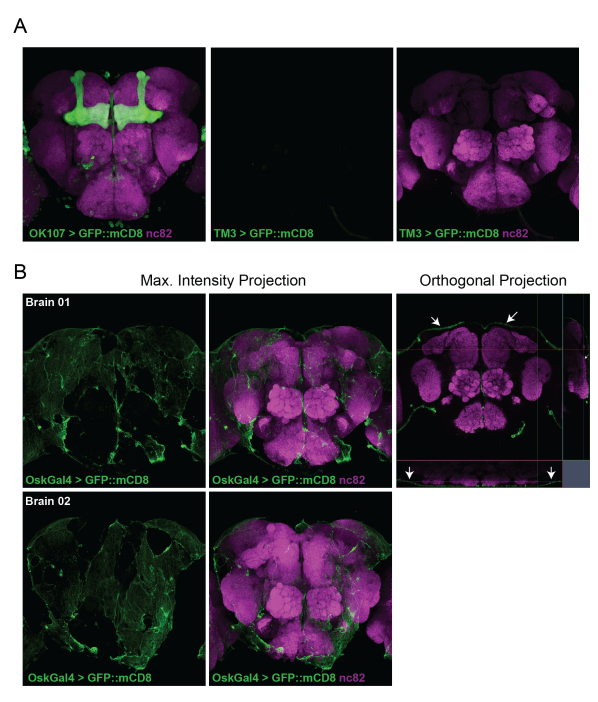


Figure A.3. Osk-Gal4 does not drive expression in neurons of the adult brain. (A) Positive and negative controls for GFP reporter construct. A mushroom body driver, OK107, drives strong, membrane-bound GFP expression in the mushroom body (left panel), and the reporter construct has no leaky expression in the absence of a driver (middle and right panel). (B) Osk-Gal4 drives expression in the sheath surrounding the brain (white arrows). Neurons themselves are not stained, as revealed in an orthogonal projection (top row, right panel). Two separate brains are shown to demonstrate that the staining pattern in the sheath results from mechanical ripping done during dissection to allow antibody penetration into the brain.

Preliminary evidence suggests that oskar RNAi and over-expression in the embryonic neuroblasts does not perturb nervous system development

We have previously shown that *Gb-oskar* is expressed in embryonic neuroblasts of crickets, and that *Gb-oskar* RNAi interferes with the development of the central nervous system (CNS) (Chapter 2; Ewen-Campen et al., 2012). I therefore wished to test for a comparable function of *oskar* in the embryonic neuroblasts of *Drosophila*, as a direct comparison to the cricket.

I drove expression of *osk*-RNAi and UAS-*osk* (an *osk* over-expression construct) using an Asense-GAL4 line (Zhu et al., 2006) which is expressed in neuroblasts and their progeny. I assayed CNS development using two metrics: (1) *even-skipped* antibody staining, which labels a small number of well-characterized neurons present in each segment, and which are often disrupted in mutants that affect neuroblast division (e.g. Dorfman et al., 1991; Ikeshima-Kataoka et al., 1997) and (2) the axonal scaffolding, visualized via anti-HRP staining, which can also reveal neuroblast defects (e.g. Ikeshima-Kataoka et al., 1997).

Neither *osk*-RNAi nor UAS-*osk* led to a detectable phenotype in *even-skipped*-positive cells nor in the morphology of the axonal tracts compared to controls (**Figure A.4**). In contrast to mutants defective for neuroblast division, which disrupt or abolish Eve+ neurons, all Eve+ cells could be readily identified in both *osk* manipulations, and were indistinguishable from wild type in location and appearance (**Figure A.4**). In addition, breaks or defects in the axonal tract, which can result from defects in neuroblast division cells (e.g. Dorfman et al., 1991; Ikeshima-Kataoka et al., 1997), were not observed (**Figure A.4**). Thus, these data did not reveal a function for *oskar* in the embryonic neuroblasts.

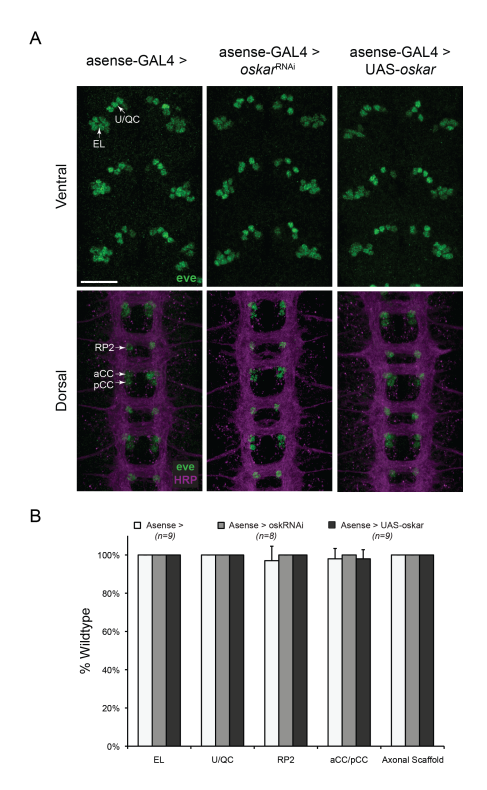


Figure A.4. CNS development is unperturbed in asense: GAL4 > osk-RNAi and asense: GAL4 > UAS:osk (A) Example images of eve-positive cells in the indicated genotypes. Two focal planes are shown, revealing ventral clusters (EL and U/QC) and dorsal motor neurons (RP2, aCC/pCC), as well as the axonal tracts visualized with HRP. (B) Quantification of scoring for each neuron cluster/axonal scaffold in each of the three genomes. Scale bar =  $20 \mu m$ .

#### **DISCUSSION**

I have presented several initial investigations into possible roles for *oskar* in the *Drosophila* nervous system, in order to extend a recent report of *oskar* function in the larval da neurons (Xu et al., 2013). I was unable to detect Oskar protein in the adult brain via antibody staining (**Figure A.2**), and I also showed that an Osk-Gal4 reporter construct does not drive expression in the neurons of the adult brain (**Figure A.3**). These results conflict with the interpretation of the data obtained from the *norka* mutant (**Figure A.1**) (Dubnau et al., 2003), which drives expression in neurons including those of the mushroom body, and are consistent with the hypothesis that the *norka* mutant may not in fact be a *bona fide* allele of *oskar*.

Furthermore, these expression data do not immediately suggest that *oskar* plays a role in the adult brain, although functional studies of additional *oskar* alleles are required to directly test whether or not this is the case. Specifically, it would be interesting to assay for olfactory learning and memory phenotypes in known *oskar* mutants (which are viable, as the gene is a maternal effect gene), and in flies containing UAS:*osk*-RNAi under the control of an adult brain-specific GAL4 driver.

In addition, I have presented preliminary data showing that the embryonic CNS develops normally when *oskar* levels are perturbed in neuroblasts using an asense-Gal4 driver. With the important caveat that the sample sizes were <10 per genotype for these experiments, these data did not reveal a phenotype for *osk*-RNAi or *UAS:oskar*. It will be important to repeat these experiments using additional GAL4 drivers and *oskar*-RNAi constructs, in addition to testing whether additional Dicer can increase RNAi efficacy. The *elav*-GAL4 driver, which drives expression in all embryonic neurons and a small number of glial cells (Berger et al., 2007), may be a good candidate, as are several others including *deadpan*-GAL4 (Lin et al., 2010), *prospero*-GAL4 (Atwood et al., 2007), and *worniu*-GAL4 (Atwood et al., 2007). If, following such

additional experiments, no embryonic CNS phenotype is observed in *Drosophila*, it would suggest that function of *Gb-oskar* in the *Gryllus* embryonic CNS is not conserved.

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