



Genetic characterization of candidate ecdysteroid kinases in Drosophila melanogaster

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Ecdysteroids are major hormones in insects and control molting, growth, reproduction, physiology, and behavior. The biosynthesis of ecdysteroids such as 20-hydroxyecdysone (20E) from dietary sterols is well characterized, but ecdysteroid catabolism is poorly understood. Ecdysteroid kinases (EcKs) mediate the reversible phosphorylation of ecdysteroids, which has been implicated in ecdysteroid recycling during embryogenesis and reproduction in various insects. However, to date, only 2 EcK-encoding genes have been identified, in the silkworm Bombyx mori and the mosquito Anopheles gambiae. Previously, we identified 2 ecdysteroid kinase-like (EcKL) genes— Wallflower (Wall) and Pinkman (pkm)—in the model fruit fly Drosophila melanogaster that are orthologs of the ecdysteroid 22-kinase gene BmEc22K. Here, using gene knockdown, knockout, and misexpression, we explore Wall and pkm's possible functions and genetically test the hypothesis that they encode EcKs. Wall and pkm null mutants are viable and fertile, suggesting that they are not essential for development or reproduction, whereas phenotypes arising from RNAi and somatic CRISPR appear to derive from off-target effects or other artifacts. However, misexpression of Wall results in dramatic phenotypes, including developmental arrest, and defects in trachea, cuticle, and pigmentation. Wall misexpression fails to phenocopy irreversible ecdysteroid catabolism through misexpression of Cyp18a1, suggesting that Wall does not directly inactivate 20E. Additionally, Wall misexpression phenotypes are not attenuated in Cyp18a1 mutants, strongly suggesting that Wall is not an ecdysteroid 26-kinase. We hypothesize that the substrate of Wall in this misexpression experiment and possibly generally is an unknown, atypical ecdysteroid that plays essential roles in Drosophila development, and may highlight aspects of insect endocrinology that are as-yet uncharacterized. We also provide preliminary evidence that CG5644 encodes an ecdysteroid 22-kinase conserved across Diptera.

Keywords: ecdysone; 20-hydroxyecdysone; 3-dehydroecdysone; trachea; UAS/GAL4; RNAi; CRISPR; steroid

Introduction

Ecdysteroids are polyhydroxylated steroids that control numerous aspects of insect biology, most notably molting, growth, and reproduction (Riddiford 1993). Twenty-hydroxyecdysone (20E) is responsible for most of the functions attributed to ecdysteroids, although its precursor ecdysone (E) may also have specific developmental roles (Beckstead et al. 2007; Ono 2014). The nuclear receptor heterodimer EcR/USP and the G protein-coupled receptor DopEcR are thought to modulate most genomic and non-genomic responses to ecdysteroids (Baker et al. 2000; Wang et al. 2000; Ishimoto et al. 2013), although the nuclear receptor DHR38 may also be involved in ecdysteroid signaling (Baker et al. 2003). The biosynthesis of ecdysteroids from dietary sterols involves the highly conserved "Halloween"? enzymes (Rewitz et al. 2006, 2007) and the tight transcriptional regulation of Halloween genes is essential for controlling insect growth and development (Yamanaka et al. 2013; Kannangara et al. 2021). However, relatively little is known about ecdysteroid inactivation through catabolism and how this is regulated.

Ecdysteroid inactivation can modulate ecdysteroid signaling by reducing levels of active hormone and/or producing inactive "storage"? forms that can be recycled into active hormones. This can play important roles in reproduction, where maternal- or matecontributed storage hormones can influence early development or reproductive physiology, respectively (Sonobe and Ito 2009; Crocker and Hunter 2018; Peng et al. 2022). It can also function in development, by shaping the pulse-like titers of 20E required for developmental transitions (Rewitz et al. 2010). Ecdysteroid recycling might also be responsible for the mid-pupal pulse of 20E in some brachyceran dipterans (Ohtaki 1981; Scanlan et al. 2023), although this does not appear to be the case in Drosophila melanogaster (Zhang et al. 2024).

Ecdysteroid catabolic reactions can be broadly classified into modifications and conjugations. Modifications involve changes to various hydroxyl groups on the ecdysteroid nucleus. 26-Hydroxylation occurs via Cyp18a1 (Guittard et al. 2011) and can result in ecdysteroids that may be functional in some insects but not others (Kaplanis et al. 1973; Baker et al. 2000). 26-Hydroxylated ecdysteroids can be further oxidized by Cyp18a1 to 26-ecdysonoic acids, which is an irreversible process and widespread across taxa (Fig. 1; Rewitz et al. 2010; Li et al. 2014). 3-Oxidation and 3-epimerization also occur through ecdysteroid 3-oxidases and 3-alpha-reductases (Takeuchi et al. 2001; Sun et al. 2012), although the former modification can be reversed through the action of 3-beta-reductases (Sakurai et al. 1989; Chen et al. 1996). Conjugations involve the

ecdysonoic acid 26-hydroxyecdysone (26E) E 2-phosphate ecdysone (E) 26E 26-phosphate E 22-phosphate

Fig. 1. Diagram of the four types of ecdysteroid phosphorylation reactions and their corresponding ecdysteroid kinase enzymes, and the hydroxylation and carboxylation reactions of Cyp18a1, based around the ecdysone nucleus. Key carbon positions are shown with small blue numbers on the ecdysone molecule.

addition of a bulky polar or nonpolar moiety that is thought to broadly limit the binding affinity to ecdysteroid receptors (Baker et al. 2000; Makka et al. 2002). Conjugated moieties include glucose (Thompson et al. 1987), fatty acids (Hoffmann et al. 1985), and acetate (Modde et al. 1984), although the most common is phosphate.

E 3-phosphate

Phosphorylation reactions can occur at 4 potentially hydroxylated carbons on the ecdysteroid nucleus—C-2, C-3, C-22, and C-26 (Fig. 1)—but the prevalence of each seems to vary broadly across both substrates and taxa (reviewed in Lafont et al. 2012). Some ecdysteroid-phosphates can be hydrolyzed in vivo to active ecdysteroids through the action of phosphatases (Isaac, Sweeney, et al. 1983; Yamada and Sonobe 2003; Davies et al. 2007) but others may be non-hydrolyzable, and therefore terminal, catabolites (Lagueux et al. 1984; Sonobe and Yamada 2004).

The most well-defined function for ecdysteroid-phosphates in insect biology is ovary-embryo recycling: phosphate conjugates are formed in, or transported to, the oocyte and hydrolyzed to active hormones after fertilization. Bombyx mori (Lepidoptera: Bombycidae) recycle 22-phosphates of 20E and some precursors in its eggs (Sonobe and Ito 2009), while the related moth Manduca sexta (Lepidoptera: Sphingidae) recycles 26-hydroxy-E 26-phosphate (Feldlaufer et al. 1988). Two locust species, Schistocerca gregaria and Locusta migratoria (Orthoptera: Acrididae), both recycle ecdysteroid 22-phosphates (Isaac, Rose, et al. 1983; Isaac, Sweeney, et al. 1983; Lagueux et al. 1984). Hydrolysis of ecdysteroid-phosphates seems to control embryonic diapause in B. mori (Yamada et al. 2005; Matsushima et al. 2019) and may do the same in locusts (Tawfik et al. 2002). Various ecdysteroid-phosphates are also produced during later life stages of lepidopterans and orthopterans but their functions are unclear (Gibson et al. 1984; Modde et al. 1984; Beydon et al. 1987; Lozano et al. 1989).

In Diptera, ecdysteroid phosphorylation saw comparatively limited study until recently. In the mosquito Anopheles gambiae (Diptera: Culicidae), 3-dehydro-20E 22-phosphate is a male sexual gift that is dephosphorylated in the female reproductive tract to prevent remating (Peng et al. 2022). 20E is also converted to a 22-phosphate in virgin females of this species to modulate the hormonal response to blood-feeding (Peng et al. 2022). In Musca domestica (Diptera: Muscidae), ecdysteroid 2-phosphorylation can be induced by a synthetic ecdysteroid agonist but its physiological function remains unclear (Williams et al. 2002). In D. melanogaster, 4 ecdysteroidphosphates have been identified: 3-dehydro-E 2-phosphate and 3-epi-20E 3-phosphate in larvae (Sommé-Martin et al. 1988a; Hilton 2004); E 22-phosphate in adult ovaries (Grau et al. 1995; Pis et al. 1995); and 20,26-dihydroxy-E 26-phosphate in the S2 cell line, due to the necessity of a phosphatase pretreatment when characterizing Cyp18a1 (Guittard et al. 2011). Possible roles for these metabolites in development or reproduction have yet to be identified.

Ecdysteroid-phosphate conjugates are produced by ecdysteroid kinases (EcKs); reverse genetic analyses of EcK genes could be a powerful method to understand the functions of ecdysteroidphosphates in insect biology. To date, only 2 EcKs have been biochemically and genetically characterized: BmEc22K, which is responsible for ecdysteroid 22-phosphate production in the B. mori ovary (Sonobe et al. 2006); and AgEcK2, which produces 20E 22-phosphate in A. gambiae virgin females (Peng et al. 2022). Both BmEc22K and AgEcK2 belong to the ecdysteroid kinase-like (EcKL) gene family. We recently performed a comprehensive phylogenomic analysis of the EcKL family in 140 insect genomes, classifying them into 13 subfamilies. BmEc22K and AgEcK2 belong to the A and D subfamilies, respectively, making other genes in these clades strong a priori candidates for encoding EcKs (Scanlan and Robin 2024). D. melanogaster contains 2 subfamily A genes: CG13813 which we name Wallflower (Wall) in this study—and Pinkman (pkm), previously named CG1561. Both genes have been lost 2-3 times in Diptera but are otherwise conserved as single-copy orthologs in most species (Scanlan and Robin 2024), and they are more closely related to each other than they are to any lepidopteran subfamily A EcKL (Supplementary Fig. 1). Wall and pkm share 23 and 28% identity, and 42 and 47% identity, to BmEc22K along their alignable amino acid sequence, respectively (Supplementary Fig. 2).

pkm is strongly expressed in the adult eye, with some expression in the pupal CNS and fat body (Xu et al. 2004; Graveley et al. 2011; Leader et al. 2018). The Pkm protein has an N-terminal disordered domain, atypical of most EcKLs, and may physically interact with the small heat shock proteins Hsp23 and Hsp26 and an uncharacterized carbonyl reductase, CG11200 (Guruharsha et al. 2011). It has been proposed that Pkm modulates synapse formation through interactions with Hsp23 and Hsp26 (Santana et al. 2020). pkm also appears to be transcriptionally induced in adult oenocytes during starvation, under the control of insulin signaling (Chatterjee et al. 2014); given this, we hypothesize *pkm* has a role in the metabolic response to starvation.

Wall is primarily expressed in the midgut of embryos, larvae, and adult flies (Weiszmann et al. 2009; Leader et al. 2018). Despite a lack of focused study, multiple transcriptional data suggest that Wall has an ecdysteroid-related function. Specifically, Wall has: (1) enriched expression in the 3rd-instar larval ring gland (RG), which contains the ecdysteroidogenic prothoracic gland (PG) cells (Ou et al. 2016); (2) temporal co-expression with Cyp18a1 (Graveley et al. 2011; Supplementary Fig. 3), especially during the mid-embryonic pulse of ecdysteroids (Maróy et al. 1988); (3) strong induction by exogenous 20E in nearly all surveyed cell lines (Gauhar et al. 2009; Stoiber et al. 2016); and (4) positive regulation by the primary 20E response gene DHR3 (Ruaud et al. 2010; Mazina et al. 2015). Given that Wall is strongly expressed in S2 cells during 20E exposure and is co-expressed with Cyp18a1 during embryogenesis, we hypothesize that Wall could encode the ecdysteroid 26-kinase proposed by Guittard et al. (2011).

Here, we genetically characterize the EcKL genes pkm and Wall through germline and somatic CRISPR knockouts, RNAi knockdown, and UAS-GAL4 misexpression. We also test 2 specific hypotheses: (1) that Wall encodes an ecdysteroid 26-kinase, through epistasis experiments with Cyp18a1; and (2) that pkm is involved in the starvation response in adult flies. We aim to genetically identify EcKs in D. melanogaster, in order to shed light on their functions in this species and across insects more broadly.

Materials and methods

Fly lines and husbandry

The following fly lines (BL, Bloomington line) were obtained from the Bloomington Drosophila Stock Center (BDSC): w*;; Sb¹/TM3 actGFP Ser¹ (BL4534), Tango11¹/CyO actGFP (BL36320), FM7i actGFP/C(1)DX $y^1 f^1$ (BL4559), FM7j (BL6418), $y^1 w^{67c23}$;; P{EPgy2}EY20330 (BL23106, called UAS-Wall^{EY} herein), w*; nos-GAL4; UAS-Cas9.P2 (BL67083), and elav-GAL4; UAS-Cas9.P2/CyO (BL67073). The following UAS-dsRNA (and control genotype) lines were obtained from the Vienna Drosophila Resource Centre (VDRC): VL60100 ("KK control"?), w^{1118} (VL60000; "GD control"?), KK^{Wall} (VL104249), GD^{Wall} (VL45409), KK^{pkm} (VL106503), GD^{pkm1} (VL32634), and GD^{pkm2} (VL32635). UAS-Cyp18a1 (Guittard et al. 2011), UAS-Dcr2/CyO; tub-GAL4/TM6B, and some GAL4 driver stocks (Supplementary Table 2) were a kind gift of Philip Batterham (The University of Melbourne). w*; tub-GAL80^{ts}; TM2/TM6B and y¹ w* P{lacW}Fas2G0032 P{neoFRT}19A/ FM7c; P{ey-FLP.N}5 were a kind gift of Michael Murray (The University of Melbourne). The strong loss-of-function Cyp18a1¹ allele was described in Rewitz et al. (2010); a Cyp18a1¹/FM7i actGFP line was a kind gift of Michael O'Connor (University of Minnesota). For GAL4 driver lines used in this study, see Supplementary Table 1; other fly lines were made by routine crossing (Supplementary Methods).

For routine stock maintenance, flies were kept on yeastcommeal-molasses media ("lab media"?; http://bdsc.indiana.edu/ information/recipes/molassesfood.html) at 18°C, 21°C, or 25°C in plastic vials sealed with cotton stoppers.

Generation of UAS-ORF lines

Cloning of ORFs into the pUASTattB vector (Bischof et al. 2007) and D. melanogaster transformation was as described in Scanlan et al. (2022), with the following modifications. The Wall ORF was isolated by PCR from the DGRC cDNA clones IP11764 and GH09153, using the primer pair CG13813_EagIF and CG13813_KpnIR (Supplementary Table 2). The 2 ORFs of pkm—the full 635 aa ORF ("pkmF"?) and a truncated 430 aa ORF without the N-terminal intrinsically disordered region ("pkm"?)—were synthesized by Integrated DNA Technologies, with Eagl and KpnI restriction sites (plus 6 additional nucleotides to allow for efficient digestion) at the N- and C-termini, respectively. Expected amplicon sizes from recombinant plasmids in colony PCR for Wall, pkm^F , and pkm^T were 1,521 bp, 2,136 bp, and 1,521 bp, respectively (using primers pUASTattB_3F and pUASTattB_5R; Supplementary Table 2). All correctly assembled plasmids, as verified by Sanger sequencing, were sent to TheBestGene Inc. (USA) for microinjection and incorporation into the D. melanogaster genome at the attP40 site on chromosome 2 (BL25709).

Generation of mutants by CRISPR-Cas9

The recombinant pCFD6 plasmids pCFD6-CG13813 pCFD6-CG1561—each of which express, under the control of a UAS promoter, 4 gRNAs that target either the CG13813/Wall locus (Fig. 2a) or the CG1561/pkm locus (Fig. 2b)—were designed in silico using Benchling (http://benchling.com), and cloned as described in Scanlan et al. (2022), with the following modifications: the primer pairs for the amplification of the inserts for pCFD6-CG13813 were pCFD6_CG13813_1F/R, pCFD6_CG13813_2F/R, and pCFD6_CG13813_ 3F/R; and the primer pairs for the amplification of the inserts for pCFD6-CG1561 were pCFD6_CG1561_1F/R, pCFD6_CG1561_2F/R, and pCFD6_CG1561_3F/R (Supplementary Table 2). Correctly assembled plasmids, as verified by Sanger sequencing, were sent to The Best Gene Inc. for microinjection and incorporation into the D. melanogaster genome at the attP40 site on chromosome 2 (BL25709), to produce the homozygous fly lines UAS-Wall^{pCFD6} and UAS-pkm^{pCFD6}.

Mutagenesis of the Wall locus on chromosome 3L followed protocols previously described in Scanlan et al. (2022), using the crossing scheme in Supplementary Fig. S4A. Mutagenesis of the pkm locus on the X chromosome was done using the crossing scheme

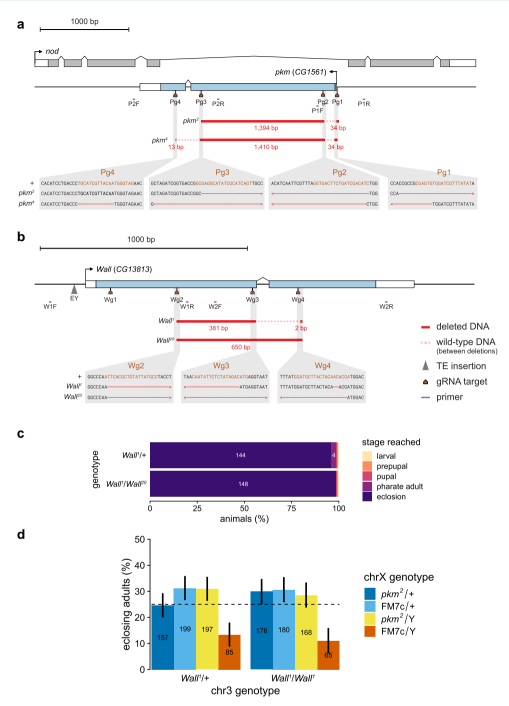


Fig. 2. CRISPR-Cas9 mutagenesis of pkm and Wall. a) The Pinkman/pkm (CG1561) locus on the X chromosome of Drosophila melanogaster, including the pkm gene (blue coding regions) and the nod gene (gray coding regions). Mapped onto the locus are the locations of gRNA target sites (Pg1-4; brown), genotyping primer-binding sites (purple), and 2 CRISPR-Cas9-induced deletion alleles (pkm² and pkm⁴; deletion sizes in red text). P1F, \(\D037_1F; P1R, \(\D037_1F; P2F, \) AD37_2F; P2R, AD37_2R. b) The Wallflower/Wall (CG13813) locus on chromosome 3L of Drosophila melanogaster. Mapped onto the locus are the locations of gRNA target sites (Wg1-4; brown), genotyping primer-binding sites (purple), the EYgp2 element EY20330 (gray triangle), and 2 CRISPR-Cas9-induced deletion alleles (Wall and Wall of the total and Wall of the total and Wall of the total of the Wall and Wall of the Wall of t AD38_2F; W2R, AD38_2R. Wild-type DNA present in-between deletions is indicated with pale dashed lines. Gray boxes (bottom) are sequence-level detail of the deletions with respect to the wild-type genetic background (+). gRNA target sites are highlighted in brown, and deleted bases are red dashes; > and < symbols indicate that the deletion continues out of the frame of the highlighted sequence. c and d) Developmental viability of Wall and pkm single and double mutant animals. c) Larval-to-adult viability of offspring from crosses between Wall¹ females and wild-type (+) males (top) or Wall²⁰ males (bottom). Numbers on the bars are the number of individuals in each lethal phase category (for numbers greater than 3). d) Eclosing adult genotypic ratios of the offspring from crosses between $pkm^2/FM7c_{;;}$ Wall¹ females and Wall¹ (left) or w^{1118} (right) males. Bars are colored by their chrX genotype, as determined by their sex and the presence or absence of the FM7c balancer chromosome; pkm2 hemizygotes are in yellow (single mutants on left, double mutants on right). The dashed line indicates the expected 1:1:1:1 genotypic ratio if all genotypes per cross are equally developmentally viable. Numbers on the bars are the number of adults of each genotype. Error bars are 97.5% confidence intervals (95% CI adjusted for 2 tests) for the proportion of each genotype within each cross.

in Supplementary Fig. S4B. Single founder female flies during pkm mutagenesis were allowed to lay viable larvae before undergoing DNA extraction.

Wall PCR genotyping used the ΔD38_1F/1R and ΔD38_2F/2R primer pairs, which produce amplicons of size 698 and 845 bp, respectively, from wild-type chromosomes, and $\Delta D38_1F/2R$, which produce 1,689 bp amplicons from wild-type chromosomes and amplicons between ~700 and 1,500 bp in the case of deletions between any 2 Wall gRNA pairs. pkm PCR genotyping used the ΔD37_1F/1R and ΔD37_2F/2R primer pairs, which produce amplicons of size 693 and 898 bp, respectively, from wild-type chromosomes, and Δ D37_2F/1R, which produce 2,757 bp amplicons from wild-type chromosomes and amplicons between ~900 and 2,600 bp in the case of deletions between any 2 pkm gRNA pairs. PCR—2 min initial denaturation (95°C), then 2 min denaturation (95°C), 45 s annealing (55°C) and 1.5 min extension (72°C) for 32 cycles, and then a 5 min final extension (72°C)—was carried out with GoTaq Green Master Mix (Promega). Deletion loci from mutant flies were amplified using the genotyping primers above with 4 GoTaq Green PCR reactions per line, which were combined before gel-purification to allow for the detection of early-cycle polymerase-derived errors by close inspection of the sequencing chromatogram output. Gel-purified amplicons were sequenced using the appropriate genotyping primers at AGRF and compared to the D. melanogaster r6 reference genome at FlyBase (Gramates et al. 2022; Öztürk-Çolak et al. 2024).

Germline knockout, somatic knockout, RNAi knockdown, and misexpression developmental viability assays

Egg-to-adult and larval-to-adult viability assays were conducted as described in Scanlan et al. (2022), but at 25°C or 29°C depending on the experiment. The "MultinomCI"? function in the DescTools package in R was used to calculate confidence intervals for multinomial proportions.

Misexpression phenotyping was conducted by crossing 3-5 UAS-responder females to GAL4 males, letting them lay in lab media vials for 24 h, with a minimum of 3 replicate vials per genotype; vials were left to develop at 25°C and were checked every 24 h for developmental arrest phenotypes. Arrest phenotypes were fully penetrant unless otherwise noted.

Delayed-onset misexpression assay

Three groups of 10 act-GAL4/CyO actGFP virgin females were crossed to tub-GAL80ts; UAS-WallEY males and allowed to mate in lab media vials at 18°C for 24 h, then transferred to 3 fresh vials at 18°C (the GAL80^{ts} restrictive temperature) and allowed to lay for 24 h. This transfer-and-lay process was repeated every 24 h (with fresh females added after 9 days) until 19 sets of 3 vials containing offspring were produced, and then all the vials were shifted to 29°C (the GAL80^{ts} permissive temperature) to complete development—each set of vials contained animals that started expressing $Wall^{EY}$ at a different developmental stage. Eclosing adults were genotyped based on the presence or absence of the phenotypic markers on the CyO actGFP chromosome. Adult genotype counts from each set of vials were analyzed by the "binom.test"? function in R (with a Bonferroni correction for multiple tests) to determine if genotypic ratios were significantly different from the Mendelian expectation (1:1).

Food avoidance assay

tub-GAL80ts; da-GAL4 virgin females were crossed to w1118 or $\mathsf{UAS}\text{-}\mathsf{Wall}^{\mathsf{EY}}$ males and laid eggs on juice plates (Supplementary Methods) at 25°C for 8 h. First-instar larvae were transferred to lab media vials (20 larvae per vial, 8 vials per genotype) and kept at 29°C. Two and 3 days post-hatching, the number of larvae located in the food substrate (mouthparts hidden in substrate; "digging"?), on top of the food substrate (mouthparts out of substrate; "on food"?), or on the sides of the vials (body completely off substrate; "side of vial"?) was scored for each genotype. Fisher's exact test was used to determine if there were significant differences in larval position between genotypes and/or timepoints, using the "fisher.test"? function in R.

Misexpression epistasis assays

Six Cyp18a1¹/FM7i actGFP; UAS-Wall^{pU} virgin females or +; UAS-Wall^{pU} virgin females were crossed to da-GAL4 or Mef2-GAL4 males and allowed to lay in lab media vials for 24 h, with 10 vials per genotype. Offspring were left to develop at 25°C for 14 days, and the number of individuals that reached pupariation, pupation (or pupariation/pupation for da-GAL4 crosses), pharate adult differentiation, and eclosion were scored. Fisher's exact test was used to determine if there were significant differences between the developmental outcomes of offspring of each maternal genotype, using the "fisher.test"? function in R.

Starvation assays

Starvation assays were conducted at 21-22°C using the "wet starvation"? method developed by Storelli et al. (2019). Adult flies were collected as virgins in the first 12 h post-eclosion, separated by sex and kept in lab media vials for 2-3 days. Empty vials were halffilled with deionized water and cellulose acetate stoppers (Flystuff, 49-102) were pushed to the bottom and saturated. Excess water was thoroughly removed and 9-10 flies were placed into each vial (6 vials per sex-genotype combination), which were closed with an additional dry stopper. Flies were moved to fresh vials twice a week. Survival was scored every 24 h until all flies were dead. Male and female flies were analyzed separately due to known sex differences in starvation resistance (Jang and Lee 2015). Survival analysis was conducted with the survival (v2.38) and survminer (v0.4.7) packages in R, using a log-rank test to compare survival curves; pairwise P-values were adjusted for multiple tests using the Benjamini-Hochberg procedure.

Additional misexpression assays

UAS-ORF responder lines for 13 EcKLs were obtained from FlyORF at the University of Zurich (Bischof et al. 2013) or the DPiM transgenic fly resource at the Bangalore Fly Resource Center (Guruharsha et al. 2012); FlyORF lines contain 3xHA-tagged ORFs, while DPiM lines contain FLAG-HA-tagged ORFs. FlyORF EcKL lines used were F002982 (CG31300), F002821 (CG10560), F002832 (CHKov2; "CHKov21"?), and F002521 (CG9259). DPiM lines used were 817 (CG10562), 854 (CHKov2; "CHKov22"?), 1332 (JhI-26), 2262 (CG14314), 2439 (CG5644), 2866 (CG10514), 3380 (CG6830), 3774 (CG31102), 3915 (CG31087), and 3916 (CG31288). w^{1118} (VDRC stock 60000) was used as a wildtype control in the absence of true matched genetic backgrounds. tub-GAL4/TM3 actGFP Ser1 and phm-GAL4/CyO actGFP (Guittard et al. 2011) were a kind gift of Philip Batterham (The University of Melbourne).

UAS-ORF responder (and w¹¹¹⁸) males were crossed to tub-GAL4/ TM3 actGFP Ser1 females, which were allowed to lay on lab media food, and the offspring were left to develop at 25°C for 14 days. Adult offspring were collected after eclosion and scored for the presence or absence of the TM3 actGFP Ser^1 balancer chromosome (and the TM6B $Antp^{Hu}$ Tb^1 chromosome in the case of DPiM line 3774). UAS-CG5644 males were also crossed to phm-GAL4/CyO actGFP females; adult offspring were scored for the presence or absence of the CyO actGFP chromosome. The "binom.test"? function in R was used to test deviations from expected Mendelian ratios (1:1 for all crosses except those involving DPiM line 3774, which was 1:3)—significant deviations against misexpression genotypes were considered evidence for developmental lethality due to misexpression of the EcKL ORF in question.

Results

Pkm null mutants have no obvious developmental phenotypes, in contrast to somatic knockout and some RNAi knockdown animals

We explored the developmental essentiality of pkm through 3 gene disruption methods: germline CRISPR, to generate heritable null alleles; somatic CRISPR, to induce gene knockout in somatic tissues; and transgenic RNAi, to knock down mRNA levels. Through germline CRISPR mutagenesis, 18 putative deletions at the pkm locus were generated from 24 founder females. Two pkm composite deletion alleles were kept as homozygous-viable stocks and molecularly characterized: pkm2 and pkm4. pkm2 comprises 2 deletions, one of 34 bp overlapping the transcription start site and another of 1,394 bp that induces a frameshift in exon 1, while pkm⁴ comprises 3 deletions, one of 34 bp that removes the transcription start site, one of 1,410 bp in-frame in exon 1, and another of 13 bp that induces a frameshift in exon 2 (Fig. 2a). Both alleles delete the ATP-binding Brenner's motif (Supplementary Fig. 2), induce frameshifts, and delete significant portions of the coding sequence, and are therefore likely strong loss-of-function (null) alleles.

The developmental essentiality of pkm was tested with egg-to-adult viability assays involving null mutants. In crosses between $pkm^2/FM7c$ females and +/Y males, while FM7c hemizygotes appeared much less viable than other genotypes, there were no clear differences between the proportions of other genotypes, showing that pkm^2/Y hemizygotes are developmentally viable (Fig. 2d).

Somatic knockout of pkm using the ubiquitous da-GAL4 driver resulted in a substantial number of adults falling into the food media, being unable to remove themselves and "drowning"?; these adults were unable to be easily counted. Tipping vials upside-down before eclosion revealed that knockout adults eclosed in a 1:1 ratio with balancer adults, indicating they had no detectable pre-adult lethality (Supplementary Fig. S5B), but had an uncoordinated limb phenotype, effectively resulting in immobility (Supplementary Fig. S5C), a phenotype not seen in pkm null mutants. This immobility phenotype was also observed when UAS-pkmpCFD6 and UAS-Cas9 were driven with the (nonspecific; see Berger et al. 2007; Casas-Tintó et al. 2017) neuronal driver elav-GAL4, although this was not quantified.

RNAi knockdown experiments using the strong ubiquitous tub-GAL4 driver were previously performed at 25°C with KK library UAS-dsRNA lines (Scanlan et al. 2020), but we added to this experiment by also using independently generated GD lines that target different parts of each gene's transcript (Dietzl et al. 2007). We also performed knockdown at a higher temperature (29°C), which increases GAL4 activity and should result in stronger knockdown (Fortier and Belote 2000; Duffy 2002), as well as at 29°C with the addition of a UAS-Dcr2 transgene, which can further increase knockdown efficiency (Dietzl et al. 2007). For pkm, none of the 3 RNAi constructs produced detectable developmental arrest at 25°C (Supplementary Fig. S6A), but KKpkm produced partially penetrant lethality before the adult stage at 29°C (Supplementary Fig. S6B). KKpkm, GDpkm1, and GDpkm2 all had significant effects on developmental viability compared to controls (all P $< 2.2 \times$ 10⁻¹⁶, Fisher's exact test) at 29°C when paired with the UAS-Dcr2 transgene, with KK^{pkm} resulting in pupal and pharate adult lethality,

and GD^{pkm1} and GD^{pkm2} producing a mix of pharate adult lethality and a "drowning"? phenotype whereby adults eclosed from the puparium but fell to the bottom of the vial and failed to escape from the semi-liquid food media (Supplementary Fig. S6D).

Wall null mutants and somatic knockout animals have no obvious developmental phenotypes, in contrast to most RNAi knockdown animals

We also explored the developmental essentiality of Wall through germline CRISPR, somatic CRISPR, and transgenic RNAi. Through germline CRISPR mutagenesis, 22 putative deletions at the Wall locus were detected by PCR from 66 founder males. Two Wall deletion alleles were kept as homozygous-viable stocks and molecularly characterized: Wall¹ and Wall²⁰. Wall¹ comprises 2 deletions in the coding region of the genes—a 381 bp in-frame deletion in exon 1 (between sgRNAs Wg2 and Wg3), and a 2 bp frameshift deletion in exon 2 (at sgRNA Wg4). Wall²⁰ comprises a single 605 bp frameshift deletion that spans exons 1 and 2 (Fig. 2b). Both alleles are predicted to significantly disrupt the function of the encoded protein, due to the deleted coding sequence (including Brenner's motif; Supplementary Fig. 2) and frameshifts, and are therefore likely also null alleles.

The developmental essentiality of Wall was tested with larval-to-adult viability assays involving null mutants. There were no significant differences between the developmental outcomes of Wall¹/+ and Wall¹/Wall²⁰ heterozygotes (P = 0.152, Fisher's exact test; Fig. 2c), with nearly all individuals successfully eclosing as viable adults. To test for possible developmental redundancy between Wall and pkm, we also conducted egg-to-adult viability assays with crosses that produced double mutant animals and compared the results to those that produced pkm single mutants (see above). There was no significant difference in the genotypic ratios between Wall¹/Wall¹ and Wall¹/+ crosses (P = 0.168, Fisher's exact test; Fig. 2d), demonstrating that double mutants (homozygous for Wall¹ and hemizygous for pkm²) do not suffer any developmental viability loss compared to other genotypes.

We also noted that all generated Wall and pkm mutant stocks could be kept for multiple generations as viable homozygous stocks without obvious issues, suggesting that neither gene is required for fertility, although this was not quantified.

Somatic knockout of Wall produced no detectable loss of egg-to-adult viability compared to non-gRNA-expressing animals (Supplementary Fig. S5A), consistent with the viability of germline null mutants. In contrast, most RNAi knockdowns of Wall affected developmental progression. The KKWall RNAi construct (but not GDWall) resulted in detectable developmental arrest before the adult stage at either 25°C (Supplementary Fig. S6A) or 29°C (Supplementary Fig. S6B). Both KKWall and GDWall had significant effects on developmental viability compared to controls (both $P < 2.2 \times 10^{-16}$, Fisher's exact test) at 29°C when paired with the UAS-Dcr2 transgene, with KK^{Wall} resulting in mostly larval lethality, and GDWall resulting in both pharate adult lethality and the "drowning"? phenotype seen with GD^{pkm1} and GD^{pkm2} knockdowns (Supplementary Fig. S6D).

RNAi-induced developmental arrest may be due to off-target effects, not genetic compensation

We considered our knockdown results to be suspicious, given that no developmental phenotypes were detected for Wall and pkm null mutant animals (Fig. 2c and d). However, discrepancies between null mutant and RNAi knockdown animals can be due "genetic compensation"?, wherein gene knockout triggers a compensatory mechanism (such as the up-regulation of genes with similar functions) to restore a wild-type phenotype, while RNAi knockdown fails to trigger compensation, leading to a mutant phenotype (El-Brolosy and Stainier 2017; El-Brolosy et al. 2019). To test the hypothesis that the phenotypic discrepancy between knockdown animals and null mutant animals was due to genetic compensation in mutants, we performed pkm knockdown in a pkm² null mutant background, with the expectation that if genetic compensation was occurring in mutants, knockdown would fail to produce a developmental arrest phenotype, while if the knockdown phenotype was due to off-target effects, the presence of the pkm^2 allele would have no effect on the phenotype. Females homozygous for the KK^{pkm} construct—and homozygous for either the pkm² allele (1 of 2 such independently generated lines) or a wild-type allele (pkm⁺)—were crossed to tub-GAL4/ TM3, actGFP, Ser¹ males, and adult genotypic ratios of the offspring were scored, with half of the knockdown individuals (males) expected to be pkm⁺ or pkm² hemizygotes. The proportion of adult knockdown individuals between knockdown crosses with pkm+ and pkm² was not significantly different (Supplementary Fig. S6C), with a pkm² (line 1) vs pkm⁺ odds ratio of 1.67 (97.5% CI: 0.94, 3.0) and a pkm² (line 2) vs pkm⁺ odds ratio of 1.02 (97.5% CI: 0.54, 1.91). This demonstrates that the presence or absence of pkm loss-of-function alleles does not affect the $tub > KK^{pkm}$ knockdown phenotype, strongly suggesting that these phenotypes are due to offtarget effects and not genetic compensation.

A summary of all pkm and Wall knockout and knockdown phenotypes, for easy comparison, is presented in Supplementary Table 3.

Ubiquitous misexpression of Wall, but not pkm, causes developmental arrest

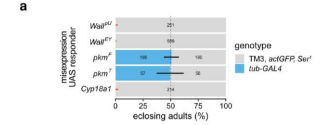
Without obvious requirements for Wall or pkm during development, we sought to test the hypothesis that these genes encode EcKs by misexpression, by driving UAS-ORF constructs (and an EPgy2 insertion upstream of the native Wall locus that contains a UAS and promoter region—UAS-Wall^{EY}) with the strong ubiquitous driver tub-GAL4 at 25°C to explore if ectopic expression arrested developmental progression. As a positive control for ecdysteroid inactivation, we used a UAS-Cyp18a1 responder (Guittard et al. 2011) to misexpress the ecdysteroid 26-hydroxylase/carboxylase Cyp18a1, which phenocopies mutations in the ecdysteroid biosynthetic pathway (Rewitz et al. 2010; Guittard et al. 2011).

Ubiquitous misexpression of Cyp18a1 with tub-GAL4 completely arrested development, as expected, as did ubiquitous misexpression of Wall with either the UAS-Wall^{EY} line or the UAS-Wall^{PU} line (Fig. 3a). Examination of offspring vials under a fluorescent microscope (to detect the presence of the TM3 actGFP Ser1 balancer chromosome) suggested that these misexpression genotypes failed to complete embryogenesis and hatch as larvae. Misexpression of either of the pkm ORF transgenes failed to significantly alter adult genotypic ratios from Mendelian expectations (Fig. 3a), suggesting that ubiquitous expression of either pkm ORF does not affect developmental progression.

Misexpression of Wall using UAS-Wall^{EY} and the strong, ubiquitous act-GAL4 driver at 25°C also caused complete developmental arrest (n = 158 adults, $P < 2.2 \times 10^{-16}$, exact binomial test), although not all animals died as embryos, with some larvae hatching but dying before pupariation, with various morphological phenotypes, such as spiracle and rectal pad pigmentation, and tracheal and denticle belt defects (Fig. 3b-i).

Ubiquitous misexpression of Wall causes developmental arrest until the middle of metamorphosis

While ubiquitous misexpression of Wall was embryonic lethal and occasionally larval lethal, it was unclear if misexpression affected later developmental stages. To investigate this, a delayed-onset



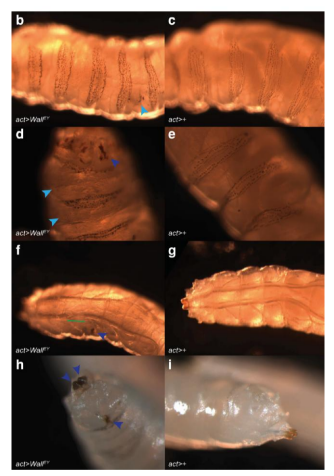


Fig. 3. Ubiquitous misexpression of Wall, pkm, and Cyp18a1. a) Egg-to-adult viability of strong, ubiquitous Wall, pkm, and Cyp18a1 misexpression, estimated from the adult genotypic ratios of offspring from crosses between tub-GAL4/TM3 actGFP Ser1 females (genotypes in bold) and UAS-responder males at 25°C. UAS-pkm^F expresses the full 635 aa pkm ORF, while UAS-pkm^T expresses a truncated 430 aa ORF that does not include the pkm N-terminal disordered region. The dashed line indicates the expected 1:1 genotypic ratio if both genotypes per cross are equally developmentally viable; error bars are 99% confidence intervals (95% CI adjusted for 5 tests) for the proportion of GAL4-containing heterozygotes; black and red bars indicate nonsignificant or significant deviations, respectively, from expected genotypic ratios after correction for multiple tests. Numbers on the bars are the number of adults of each genotype (for numbers greater than 0). b-i) Phenotypes in larvae misexpressing UAS-Wall^{EY} with the act-GAL4 driver (act > Wall^{EY}; left) compared with wild-type, non-misexpressing larvae (act>+; right), at 25°C. b-e) Denticle belt disorganization, including ectopic denticles in between belts (light blue arrowheads) and rectal pad melanization (dark blue arrowheads). (f and g) Fluid-filled section of a tracheal dorsal trunk (green line segment), along with a melanized section of trachea (dark blue arrowhead). (h and i) Melanized posterior spiracles and rectal pad (dark blue arrowheads).

misexpression experiment was conducted using the temperaturesensitive tub-GAL80ts construct, which represses GAL4 activity at 18°C and permits GAL4 activity at 29°C, and the act-GAL4 driver,

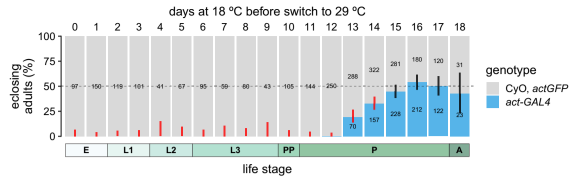


Fig. 4. Delayed onset of ubiquitous misexpression of Wall arrests development up until mid-metamorphosis. Top: egg-to-adult viability of Wall misexpression with the ubiquitous driver act-GAL4, in concert with the UAS-Wall^{EY} element and the temperature-sensitive tub-GAL80^{ts} construct, which inhibits GAL4 at 18°C but not at 29°C; animals were moved from 18°C to 29°C at 1 of 19 timepoints (top), which began Wall misexpression. Viability is $estimated from the adult genotypic ratios of offspring from crosses between \textit{act-GAL4/CyO} \textit{actGFP} females and \textit{tub-GAL80}^{ts}; UAS-Wall^{EY} males. The \textit{act-GAL4/CyO} \textit{actGFP} females and \textit{tub-GAL80}^{ts}; UAS-Wall^{EY} males and$ dashed line indicates the expected 1:1 genotypic ratio if both genotypes per cross are equally developmentally viable; error bars are 99.7% confidence intervals (95% CI adjusted for 19 tests) for the proportion of act-GAL4-containing heterozygotes; black and red bars indicate nonsignificant or significant deviations, respectively, from expected genotypic ratios after correction for multiple tests. Numbers on the bars are the number of adults of each genotype (for numbers greater than 10). Bottom: approximate developmental life stages of D. melanogaster at 18°C, showing Wall-sensitive (red) and Wall-insensitive (black) developmental periods; dashed line indicates region of uncertainty. E, embryo; L1, 1st-instar larva; L3, 2nd-instar larva; L3, 3rd-instar larva; PP, prepupa; P, pupa; A, adult.

which strongly expresses GAL4 in a ubiquitous pattern (tub-GAL4 was not used because pilot experiments suggested that tub-GAL80ts was unable to fully repress its activity). Misexpression of Wall caused significant reductions in egg-to-adult viability when delayed 0-14 days into development, but did not significantly affect viability from 15–18 days (at 18° C adult emerge at \sim 18 days), suggesting that animals become insensitive to Wall misexpression around the middle of metamorphosis (Fig. 4).

Tissue-specific misexpression of Wall causes developmental arrest distinct from Cyp18a1

The developmental arrest phenotype observed upon ubiquitous misexpression of Wall is consistent with an EcK function of its encoded protein. To further test this hypothesis, we conducted tissuespecific misexpression crosses to explore if the patterns of developmental arrest produced by Cyp18a1 misexpression matched that of Wall misexpression; a close match would be good evidence that Wall and Cyp18a1 act on the same or similar substrates.

Misexpression of Wall was conducted with 2 ubiquitous GAL4 drivers (tub-GAL4, used previously, and da-GAL4, a weaker driver) and 14 tissue-specific GAL4 drivers (Supplementary Table 1). The positive control UAS-Cyp18a1 construct produced developmental arrest when misexpressed with all 16 drivers (Fig. 5; Supplementary Table 4). The UAS-Wall^{EY} construct produced developmental arrest when misexpressed with 10 drivers, while the UAS-Wall^{pU} construct produced very similar results, but with no phenotype with the ppl-GAL4 driver and generally caused developmental arrest at later life stages (Fig. 5; Supplementary Table 4). Misexpression of UAS-Wall pU with btl-GAL4 and cg-GAL4 also resulted in a small number of adult escapers (Fig. 5b; Supplementary Table 4). These differences between UAS-Wall transgenes are likely due to differences in expression strength: the EPgy2 element contains 14 copies of UAS in its artificial enhancer region (Bellen et al. 2004), while pUASTattB contains only 5 (Bischof et al. 2007). However, the broad consistency of the results from both an independent transgene construct and a native locus misexpression construct means these phenotypes can be confidently linked to the ectopic expression of the Wall ORF.

Misexpressing UAS-WallEY with the HR-GAL4 driver resulted in a dramatic phenotype: an extended wandering period during the 3rd larval instar, then defects during pupariation and pupation wherein the puparium was thin and elongated and anterior adult tissue differentiation appeared to occur without head eversion, resulting in dark thoracic (likely from wings) and red eye pigmentation developing internally (Supplementary Fig. S7A-C). This phenotype suggests that Wall misexpression with this driver can block epidermal and cuticular development during metamorphosis, but curiously still allows for imaginal discs eversion to take place. The HR-GAL4 driver expresses GAL4 along the whole length of the midgut, as well as in the Malpighian tubules and the fat body (Chung et al. 2007); however, the lack of phenotypes with the enterocyte-specific mex1-GAL4 driver (Phillips and Thomas 2006) suggests that Wall misexpression in these midgut cells does not cause developmental defects. Additionally, given that the fat body drivers cg-GAL4 and ppl-GAL4 failed to phenocopy HR-GAL4, the crucial tissues might be non-enterocyte midgut cells and/or the Malpighian tubules. Alternatively, HR-GAL4 may drive at a low (yet sufficient) level in another tissue to produce this phenotype, such as the muscles.

Conspicuously, while the phm-GAL4 driver, which expresses GAL4 in the prothoracic cells of the ring gland (PG)—the ecdysteroidogenic tissue in larvae (Redfern 1983)—as well as some other tissues such as the wing disc (Casas-Tintó et al. 2017)—caused embryonic lethality when misexpressing Cyp18a1 as expected, it produced no developmental arrest when misexpressing either Wall construct (Fig. 5b). This result appears inconsistent with Wall encoding a kinase that can act on ecdysteroids present in the PG, although it could be explained by the action of an ecdysteroid-phosphate phosphatase in these cells.

Misexpression of Wall with the c204-GAL4 driver caused larval arrest with the UAS-Wall^{EY} construct and arrest during metamorphosis with the UAS-Wall^{pU} construct (Fig. 5). The expression pattern of this driver is not well characterized—while it is known to be expressed in the adult ovary (Manseau et al. 1997), it may be expressed in other tissues and at other life stages, given it causes embryonic arrest with UAS-Cyp18a1 and the aforementioned arrest with UAS-Wall constructs. Curiously, elav-GAL4 misexpression of Wall caused developmental arrest, while nSyb-GAL4 misexpression did not, suggesting that misexpression in neurons is not the cause of elav-GAL4 arrest; the elav-GAL4 driver, while often considered specific to neurons, expresses in other tissues, such as glia, trachea and the imaginal discs (Berger et al. 2007; Casas-Tintó et al. 2017), 1 or more of which is likely the cause of the developmental arrest observed here.



Fig. 5. Tissue-specific misexpression of Wall and Cyp18a1, using the UAS-Wall^{EY}, UAS-Wall^{PU}, and UAS-Cyp18a1 constructs, at 25°C. a) Heat map of qualitative developmental arrest phenotypes for Wall and Cyp18a1 misexpression using 2 ubiquitous GAL4 drivers and 14 tissue-specific GAL4 drivers. E, embryo; L, larva; PP, prepupa; P, pupa; PA, pharate adult; NA, no arrest (successful eclosion). When arrest occurred at more than 1 stage, 2 stages are indicated and separated with a slash. For phenotypic descriptions, see Supplementary Table 4. b) Egg-to-adult viability of Wall and Cyp18a1 misexpression, estimated from the adult genotypic ratios of offspring from crosses between GAL4/balancer females (genotypes in bold) and UAS-responder males. The dashed line indicates the expected 1:1 genotypic ratio if both genotypes per cross are equally developmentally viable; error bars are 98.3% confidence intervals (95% CI adjusted for 3 tests) for the proportion of GAL4-containing heterozygotes; black and red bars indicate nonsignificant or significant deviations, respectively, from expected genotypic ratios after correction for multiple tests. Numbers on the bars are the number of adults of each genotype (for numbers greater than 0).

Overall, these data demonstrate that Wall misexpression in the trachea, muscle, epidermis, or fat body is sufficient to cause developmental arrest, while misexpression in the PG, imaginal discs, salivary gland, neurons, or oenocytes does not cause any obvious phenotype. In addition, combined with the results of the ubiquitous delayed-onset misexpression experiment (Fig. 4), these data suggest that the pharate adult lethality observed through misexpression with some drivers (cg-GAL4 and btl-GAL4) may be due to misexpression of Wall during or before early metamorphosis, not during the pharate adult stage itself.

Larval tracheal misexpression of Wall causes tracheal defects and food aversion

Misexpression of Wall with the trachea-specific btl-GAL4 driver caused developmental arrest at the larval or pharate adult stages, depending on the UAS-responder construct and the temperature (Fig. 5, Supplementary Table 4). We noticed that 1st- or 2nd-instar $btl > Wall^{EY}$ larvae raised at 25°C abandoned the food media to crawl up the sides of the vial and died before pupariation, a phenotype that was shared with ubiquitously misexpressing $da > Wall^{EY}$ larvae that were escaped from embryonic lethality using the tub-GAL80^{ts}

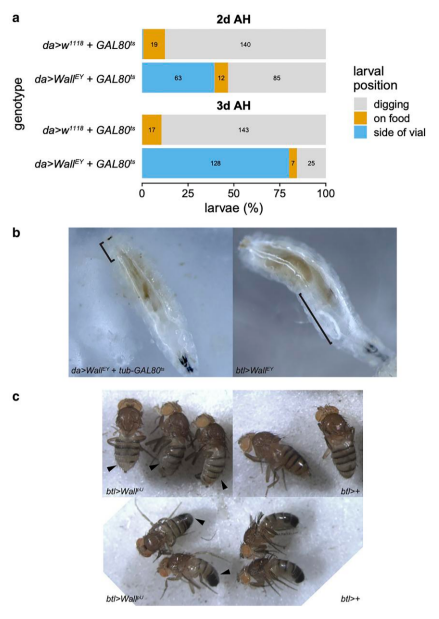


Fig. 6. Misexpression of Wall affects tracheal integrity and adult pigmentation. a) Food avoidance assay comparing da > w¹¹¹⁸ + GAL80^{ts} and da > Wall^{FY} + GAL80^{ts} larvae 2 days (top) and 3 days (bottom) after hatching (AH). Numbers on the bars are the number of larvae in each position (for numbers greater than 5). b) Photos of larvae misexpressing Wall ubiquitously (da > Wall^{EY} + tub-GAL80^{ts}, left) or specially in the trachea (btl > Wall^{EY}, right) retrieved from food media at 25°C, before food aversion occurred. Black segments indicate sections of tracheal dorsal trunks that are filled with liquid. c) Adult flies (females, top; males, bottom) misexpressing Wall in the trachea (btl > Wall^{pU}, left) or with the btl-GAL4 driver alone (btl>+, right). Arrowheads indicate dorsal abdomen pigmentation defects (areas of lightened pigment compared to the rest of the abdomen). Wings have been removed to expose the dorsal abdomen.

construct and then moved to 29°C, as well as the act > Wall^{EY} larvae described previously. Quantification of this food-aversion phenotype showed $da > Wall^{EY}$ larvae were positioned differently within vials compared to non-misexpressing $da > w^{1118}$ controls, at both 2 days and 3 days after hatching (both $P < 2.2 \times 10^{-16}$, Fisher's exact test; Fig. 6a). There was also a significant difference in positions between the 2 timepoints for misexpressing larvae (P = 9.7×10^{-14} , Fisher's exact test), while there was no significant difference between non-misexpressing larvae (P = 0.73, Fisher's exact test), consistent with progression of the misexpression phenotype over time (Fig. 6a). Food aversion is a well-established consequence of hypoxia in both larvae and adult D. melanogaster (Vigne and Frelin 2010; Wang et al. 2015; Wingen et al. 2017)—consistent with this, close examination of $btl > Wall^{EY}$ and $da > Wall^{EY}$ larvae revealed defects in

tracheal filling (similar to those seen in act > Wall^{EY} and c204 > Wall^{EY} larvae; Fig. 3f, Supplementary Table 4), with parts of the dorsal trunks often filled with liquid (Fig. 6b), which can cause severe reductions in gas exchange (Parvy et al. 2012). Taken together, these data suggest that Wall misexpression in trachea causes hypoxia through defects in either tracheal development, maintenance, or integrity. The food-aversion phenotype is also the basis for the gene name Wallflower (Wall), due to the tendency of the larvae to position themselves on the walls of the vial, rather than in the food with other larvae, an allusion to the "wallflower"? metaphor for social behavior.

Misexpression of Wall with the btl driver also appears to affect tissues other than the trachea, as we noticed a pigmentation defect on the dorsal abdomen of both male and female $btl > Wall^{pU}$ adult escapers (Fig. 6c).

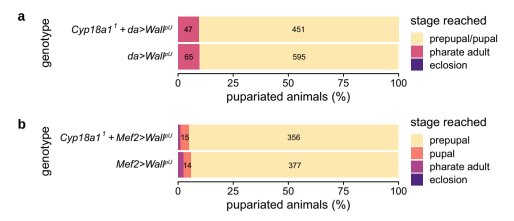


Fig. 7. Tests of hypostasis between Wall^{PU} misexpression and Cyp18a1, using a) the moderate, ubiquitous driver da-GAL4 and b) the muscle-specific driver Mef2-GAI4. Genotypes are pooled offspring genotypes from crosses with Cyp18a1¹/FM7i actGFP chrX mothers (25% of offspring are Cyp18a1¹ hemizygotes) or wild-type chrX mothers (100% of offspring have wild-type Cyp18a1). The number of individuals was not quantified before the prepupal stage, so proportions are given as the percentage of pupariated animals per genotype. Numbers on the bars are the number of individuals in each lethal phase category (for numbers greater than 10).

Wall misexpression phenotypes are not hypostatic to Cyp18a1

We hypothesized that Wall encodes an ecdysteroid 26-kinase, an enzyme that can only act on the products of the ecdysteroid 26-hydroxylase Cyp18a1 (Fig. 1). If this hypothesis is true, Wall misexpression should be hypostatic to the wild-type function of Cyp18a1—i.e. misexpression of an ecdysteroid 26-kinase (Wall) should not cause developmental defects when the source of 26-hydroxyecdysteroids (Cyp18a1) is not functional. Data indicating hypostasis would be strong genetic evidence of a biochemical interaction between the 2 genes and would support the hypothesis that Wall encodes an ecdysteroid 26-kinase. In contrast to Cyp18a1¹ animals, which largely reach the pharate adult stage but fail to eclose (Rewitz et al. 2010), most da > Wall^{pU} animals at 25°C arrest development before the differentiation of adult structures (prepupal/pupal arrest; Supplementary Table 4), while most $Mef2 > Wall^{pU}$ animals arrest development before pupation (prepupal arrest; Supplementary Table 4). We reasoned that if Wall misexpression phenotypes were hypostatic to the function of Cyp18a1, Cyp18a1¹ hemizygotes (25% of the offspring of crosses between Cyp18a11/ FM7i, actGFP; UAS-Wall pU females and GAL4 males) should reach the pharate adult stage instead of earlier developmental arrest due to Wall misexpression, increasing the proportion of pharate adult-lethal animals compared to crosses without the Cyp18a1¹

There were no significant differences in developmental outcomes between Cyp18a1¹, Mef2 > Wall^{pU}, and Mef2 > Wall^{pU} offspring (Fig. 7a; P = 0.336, Fisher's exact test), nor between Cyp18a1¹, da > $Wall^{pU}$, and $da > Wall^{pU}$ offspring (Fig. 7b; P = 0.841, Fisher's exact test). These data indicate that misexpression of Wall in Cyp18a1¹ hemizygotes produces the same developmental arrest phenotypes as misexpression of Wall in a wild-type background, strongly suggesting that Wall is not hypostatic to Cyp18a1, and that Wall is unlikely to encode an ecdysteroid 26-kinase.

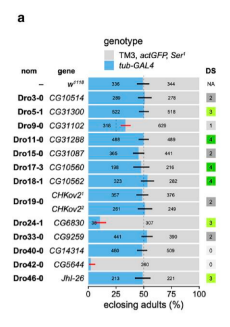
Pkm mutants are not sensitive to starvation

pkm was found by Chatterjee et al. (2014) to be up-regulated in adult female oenocytes upon starvation, which led to the hypothesis that pkm plays a role in the metabolic response to starvation, and that pkm loss-of-function mutants are more susceptible to starvation. Wet-starvation experiments (Storelli et al. 2019) were conducted with virgin male and virgin female flies, the offspring from 3 different crosses: pkm⁴ females and w¹¹¹⁸ males (yielding pkm⁴/pkm⁺ females and pkm⁴/Y males); pkm⁴ females and pkm² males (yielding pkm^4/pkm^2 females and pkm^4/Y males); and w^{1118} females and pkm2 males (yielding pkm2/pkm+ females and pkm+/ Y males). For heterozygous female genotypes, median survival in days were 9 (95% CI: 8, 9), 8 (95% CI: 8, 8), and 9 (95% CI: 8, 9) for pkm⁴/pkm², pkm⁴/pkm⁺, and pkm²/pkm⁺, respectively, with no significant differences between any pairs of genotypes across entire survival curves (all P > 0.6, log-rank test; Supplementary Fig. S8A). For hemizygous male genotypes, median survival in days were 7 (95% CI: 6, 7), 8 (95% CI: 7, 9), and 6 (95% CI: 6, 7) for pkm⁴/Y (pkm² father), pkm⁴/Y (w¹¹¹⁸ father), and pkm⁺/Y (pkm² father), respectively, with the only significant difference between the pkm^4/Y (pkm^2 father) and pkm^4/Y (w^{1118} father) across entire survival curves genotypes (P = 0.0037, log-rank test; all other P >0.067; Supplementary Fig. S8B). These data do not support the hypothesis that pkm is involved in starvation resistance.

Misexpression of CG5644, another strong EcK candidate gene, also causes developmental arrest

As other EcKLs in the D. melanogaster genome may also encode EcKs, we screened an additional 13 EcKLs for misexpression phenotypes using preexisting UAS-ORF fly lines and the tub-GAL4 driver. Six of these genes—CG31300, CG31288, CG10560, CG10562, CG6830, and JhI-26—were previously considered candidate detoxification genes, while the 7 others—CG10514, CG31102, CG31087, CHKov2, CG9259, CG14314, and CG5644—were not (Scanlan et al.

Only misexpression of CG6830 and CG5644 caused significant developmental arrest before the adult stage (Fig. 8a). The UAS-CG5644 line was further crossed to the phm-GAL4 driver at 25°C, resulting in 66 misexpression adults surviving compared to 115 balancer adults (proportion = 0.365, 95% CI: 0.295-0.439, P = 0.0003, binomial test), suggesting that PG misexpression of CG5644 can also disrupt development. These data are consistent with the hypothesis that CG5644 encodes an ecdysteroid 22-kinase, previously proposed based on 1:1 orthology to AqEcK2 in A. gambiae (Peng et al. 2022; Scanlan and Robin 2024). The lack of developmental arrest seen in tub > CG31300 and tub > JhI-26 individuals is consistent with previously published experimental data (Liu et al. 2014; Scanlan et al. 2022).



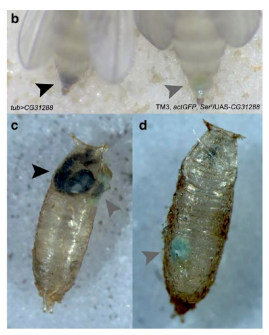


Fig. 8. Misexpression of 13 additional EcKL genes to find candidate EcKs. a) Egg-to-adult viability of the misexpression of 13 EcKLs (and 1 control genotype, w¹¹¹⁸), estimated from the adult genotypic ratios of offspring from crosses between tub-GAL4/TM3, actGFP, Ser¹ females and UAS-responder males at 25°C. The dashed line indicates the expected 1:1 genotypic ratio (or the 1:3 genotypic ratio for CG31102) if both genotypes per cross are equally developmentally viable; error bars are 99.7% confidence intervals (95% CI adjusted for 15 tests) for the proportion of GAI4-containing heterozygotes; black and red bars indicate nonsignificant or significant deviations, respectively, from expected genotypic ratios after correction for multiple tests. Numbers on the bars are the number of adults of each genotype (for numbers greater than 6). nom, Drosophila EcKL nomenclature (Scanlan et al. 2020); DS, detoxification score, where 3 or 4 is considered a candidate detoxification gene (Scanlan et al. 2020); NA, not applicable for w¹¹¹⁸. b) Meconium of freshly eclosed female flies, visible within the posterior tip of the abdomen before excretion. Black arrowhead, dark meconium of a tub > CG31288 fly; gray arrowhead, wild-type (green) meconium of a TM3 actGFP Ser1/UAS-CG31288 fly. c) Meconium deposited on the empty puparium of a fly from the cross between tub-GAL4/TM3 actGFP Ser1 and UAS-CG31288. Black arrowhead, dark meconium of a tub > CG31288 fly; gray arrowhead, wild-type (green) meconium of a TM3, actGFP Ser¹/UAS-CG31288 fly. d) Yellow-green meconium deposited on the empty puparium of a wild-type w¹¹¹⁸ fly (gray arrowhead).

While misexpression of CG31288 did not appear to affect developmental progression, we noticed a visible phenotype for tub > CG31288 adult flies: their meconium was a dark gray, almost black color, compared to the yellow-green of non-misexpressing flies (Fig. 8b-d). This color change was apparent both within the posterior abdomen of recently eclosed flies (Fig. 8b) and when the meconium had been excreted and left on empty puparia (Fig. 8c and d).

Discussion

Here we have performed the first detailed genetic characterization of candidate ecdysteroid kinase (EcK) genes in insects, targeting Wall and pkm, orthologs of the B. mori ecdysteroid 22-kinase gene BmEc22K. Despite phenotypic discrepancies between gene disruption methods, we ultimately found no clear evidence that these genes encode EcKs that have essential functions in development or reproduction. Tissue-specific misexpression was also used to detect possible EcK activity, which produced striking developmental disruption phenotypes for Wall but not for pkm.

The phenotypic differences we observed between our null allele, RNAi knockdown, and somatic CRISPR knockout approaches were unexpected. Broadly, strong RNAi knockdown conditions resulted in developmental arrest for both genes, across independent dsRNA constructs, but homozygous mutant animals (including double mutants) showed no obvious developmental phenotypes and had similar viability to controls (see Supplementary Table 3 for a summary). One explanation is genetic compensation in null mutants that is not trigged during RNAi knockdown (El-Brolosy and Stainier 2017), but knocking down pkm in a pkm mutant background did not rescue developmental arrest, inconsistent with this hypothesis. The more

likely explanation then is that the RNAi knockdown phenotypes were due to off-target effects, which have been well documented in Drosophila (Seinen et al. 2011). For Wall, no phenotypes were detected for both null alleles and somatic CRISPR knockout, however for pkm, somatic CRISPR knockout produced a highly penetrant motor coordination defect not seen in null mutant animals. Explanations for this are unclear, but might be linked to CRISPR-Cas9 off-target cleavage or unintended effects on the splicing or transcriptional regulation of nod, the fourth intron of which contains pkm.

Null alleles are generally considered the gold standard for characterization of gene function. Animals homozygous or hemizygous for null alleles of Wall, pkm, or both genes appeared indistinguishable from wild-type animals respect to both developmental progression and reproduction, strongly suggesting that these 2 closely related EcKLs do not play important roles in these processes, even redundantly with each other. This is in contrast with BmEc22K, another subfamily A EcKL that encodes an ecdysteroid 22-kinase involved in ovary-embryo ecdysteroid recycling (Sonobe et al. 2006), and indicates that this function is not conserved in dipteran subfamily A genes. It is possible Wall and/or pkm null mutants have subtle or environmentally specific phenotypes we were unable to detect in our experiments, although the lack of expression of either gene in reproductive tissues (Leader et al. 2018) makes this unlikely for reproductive functions. The lack of strict conservation of the EcKL clades containing pkm and Wall (referred to as Dip9 and Dip10 in Scanlan and Robin 2024) is consistent with nonessential functions of these genes in Drosophila and other dipteran taxa.

Neither null mutants nor misexpression animals for pkm displayed any obvious phenotypes, offering no clue as to the gene's native function or enzymatic activity. We hypothesized that pkm is involved in the starvation response in adult flies, based on transcriptomic data from Chatterjee et al. (2014), but experiments with null mutants did not show a difference in wet starvation survival compared to control genotypes, for either sex, offering no support for the hypothesis. Using RNAi knockdown with the same dsRNA lines used here, Santana et al. (2020) suggested that the Pkm protein controls mRNA and protein abundances of Hsp23 and Hsp26, leading to modulation of synaptogenesis and neural response to stress. However, our results cast doubt on the interpretability of RNAi knockdown of pkm and suggest that off-target effects could be responsible for these phenotypes. Confirmation of neural functions of pkm could be done with null alleles, which we show here to be viable and therefore could allow the study of subtle neurological phenotypes at any life stage.

Like pkm, Wall null mutants had no obvious phenotypes. However, misexpression of Wall disrupted developmental progression with both ubiquitous and tissue-specific drivers, suggesting that the Wall protein can act on 1 or more developmentally important small molecules. Tissues sensitive to Wall misexpression are the trachea, fat body, epidermis, and muscle, while insensitive tissues are the PG, neurons, oenocytes, imaginal discs, and salivary glands. Ubiquitous misexpression of Wall caused developmental arrest at all life stages, except past the midpoint of metamorphosis, after pupal-adult apolysis has occurred (Bainbridge and Bownes 1981). These data suggest that the substrate(s) of Wall could have essential signaling functions in trachea, fat body, epidermis, and muscle across D. melanogaster development. However, we believe it is unlikely these substrates are E and/or 20E, as: (1) embryonic Wall misexpression did not phenocopy the "Halloween"? phenotype like Cyp18a1 (Rewitz et al. 2010); (2) Wall phenotypes were specific to certain tissues, while Cyp18a1 misexpression reliably caused developmental arrest in tissues known to rely on 20E signaling; and (3) Wall misexpression in the PG, the larval source of E, did not produce developmental arrest, unlike Cyp18a1. Curiously, however, a recent paper reports that the dorsal internal oblique muscles (DIOMs) are the source of the mid-pupal pulse of 20E (Zhang et al. 2024). Metamorphic defects seen upon pan-muscular Wall misexpression could be consistent with interfering with ecdysteroid signaling in DIOMs, although these generally occurred before the middle of metamorphosis (Fig. 5a).

If Wall can indeed phosphorylate E and/or 20E, the tissue-specific phenotypes of Wall misexpression may have alternative explanations. One such explanation could be the presence of tissue-specific phosphatases that hydrolyze ecdysteroid-phosphates back to active hormones. Another explanation could be tissue-specific posttranscriptional or post-translational regulation of Wall that reduces the level of active enzyme and dampens or eliminates the effects of misexpression in certain cell types.

An ecdysteroid 26-kinase is known to be active in the S2 cell line (Guittard et al. 2011), which we hypothesized could be encoded by Wall due to transcriptional similarities to the ecdysteroid 26-hydroxylase/carboxylase gene Cyp18a1. However, this hypothesis was not supported by epistasis experiments, which found that the severity of Wall misexpression phenotypes was not reduced in a Cyp18a1 mutant background, suggesting that Wall-induced developmental defects are to not due to phosphorylation of 26-hydroxylated ecdysteroids produced by Cyp18a1. Alternative candidates for encoding this ecdysteroid 26-kinase are CG2004, CG31975, and JhI-26, which are all basally expressed in S2 cells to appreciable degrees (Supplementary Fig. 9).

Plausible, alternative ecdysteroid substrates for Wall are 3-dehydroecdysteroids and 3-epiecdysteroids, which are produced in the fat body and epidermis of larvae (Sommé-Martin et al. 1988a, 1988b) and may have signaling functions in D. melanogaster (Sommé-Martin et al. 1990; Baker et al. 2003). The physiological functions of these modified ecdysteroids are still unclear in D. melanogaster, but intriguingly, 3-dehydro-20E was recently found to modulate female reproductive physiology in the mosquito A. gambiae (Peng et al. 2022). Additionally, Cavigliasso et al. (2024) recently suggested that fezzik (fiz) encodes an ecdysteroid oxidase that adaptively modulates larval growth rate under malnutrition in D. melanogaster. If oxidized ecdysteroids control larval development, they would be good candidate substrates for Wall. Other possible substrates could be produced by Cyp301a1, Cyp303a1, and Spidey, developmentally important yet biochemically uncharacterized enzymes with various links to ecdysteroid metabolism (Sztal et al. 2012; Chiang et al. 2016; Wu et al. 2019).

The phenotypes associated with Wall misexpression might not necessarily reflect the gene's wild-type function, as the enzyme may be acting promiscuously on substrates that it rarely or never encounters in non-transgenic animals. However, if they are attributable to native substrates, these misexpression phenotypes might still be exposing novel hormonal pathways that underpin important developmental processes. Such native substrates would clearly be involved in tracheal integrity, cuticle and epidermal development, and functions of the fat body and muscles during metamorphosis. These are all known to be regulated by ecdysteroid signaling (Zachary and Hoffmann 1980; Bond et al. 2010; Chanut-Delalande et al. 2014; Buhler et al. 2018), but links to signaling genes and proteins that have been associated with more than 1 of the developmental defects seen in Wall-misexpressing animals could offer mechanistic clues. These include: the scavenger receptor Dsp, which is involved in both tracheal integrity and adult pigmentation (Dembeck et al. 2015; Wingen et al. 2017); Megf8, which is involved in tracheal integrity and denticle belt development (Lloyd et al. 2018); and Svb, which has roles in tracheal and denticle belt development (Chanut-Delalande et al. 2006, 2014). Intriguingly, Svb also acts in the adult midgut to coordinate stem cell behavior under the control of EcR (Al Hayek et al. 2021), which is a major site of native Wall expression (Leader et al. 2018). Ultimately, biochemical characterization of the purified Wall protein is needed to fully understand its signaling interactions upon misexpression.

The results of this study leave as open questions the identities of the ecdysteroid 2-, 3-, 22-, and 26-kinases in D. melanogaster. However, data from our misexpression experiments are consistent with CG5644 encoding an EcK, especially the developmental arrest seen upon PG-specific misexpression of this gene. Curiously, developmental arrest upon PG misexpression was only partially penetrant, raising the possibility that an ecdysteroid-phosphate phosphatase is present in this tissue. Our previous phylogenetic analyses of the EcKL gene family determined that CG5644 is a 1:1 ortholog of AgEcK2, which encodes an ecdysteroid 22-kinase in A. gambiae (Peng et al. 2022; Scanlan and Robin 2024). While CG5644 is predominantly expressed in the adult nervous system (Leader et al. 2018), it also appears expressed in the migratory border cells in the ovary (Borghese et al. 2006), which might make it responsible for the presence of E 22-phosphate in this tissue (Pis et al. 1995; Grau et al. 1995). Reverse genetics experiments like those conducted here for Wall and pkm, as well as biochemical characterization, might uncover the functions of both CG5644 and ecdysteroid 22-phosphates in D. melanogaster development, reproduction, and physiology.

Overall, we here provide the first in-depth functional genetic characterization of candidate EcK genes in insects, focusing on the BmEc22K orthologs Wall and pkm in D. melanogaster. While neither gene is essential for reproduction and development, multiple lines of evidence are consistent with Wall being able to inactivate 1 or more signaling molecules essential for development. Whether these molecules are ecdysteroids or other hormones is currently unclear. This work is suggestive of unknown complexity in tissuespecific hormonal signaling in D. melanogaster and paves the way for further characterization of the likely ecdysteroid 22-kinase gene CG5644. This work also serves to highlight the possible phenotypic inconsistencies that may arise between functional genetic methodologies and reinforces the importance of multiple, complementary gene disruption approaches when characterizing genes of unknown function.

Data availability

Plasmids are available upon request. Fly strains that are not available in stock centers are available upon request. The authors affirm that all data necessary for confirming the conclusions of the article are present within the article, figures, tables, and supplementary material.

Supplemental material available at G3 online.

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Conflicts of interest

The author(s) declare no conflicts of interest.

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