

## **Post-transcriptional modulation of Dscam1 enhances axonal growth in development and after injury**

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## Abstract

Injury to the adult central nervous systems (CNS) results in severe long-term disability because damaged CNS connections rarely regenerate. Although several axon regeneration regulators have been proposed, intrinsic regenerative mechanisms remain largely unexplored. Here, we use a *Drosophila* CNS injury model to identify a novel pro-regeneration signaling pathway. We conducted a genetic screen of approximately three hundred candidate genes and identified three strong inducers of axonal growth and regeneration: the Down Syndrome Cell Adhesion Molecule (Dscam1), the de-ubiquitinating enzyme Fat Facets (Faf)/Usp9x and the Jun N-Terminal Kinase (JNK) pathway transcription factor Kayak (Kay)/Fos. Genetic and biochemical analyses link these genes in a common signaling pathway whereby Faf stabilizes Dscam1 protein levels, by acting on the 3'-UTR of its mRNA, and Dscam1 acts upstream of the growth-promoting JNK signal. The mammalian homolog of Faf, Usp9x/FAM, shares both the regenerative and Dscam1 stabilizing activities, suggesting a conserved mechanism.

## Introduction

During CNS development axons grow in a tightly regulated manner to generate an intricate and complex pattern of neuronal connectivity. In most animal species, injury to the adult CNS, either by physical trauma or in the context of neurodegeneration, has devastating long-term consequences- in part because of the inability of mature neurons to regenerate severed axons, in contrast to peripheral nervous system (PNS) neurons. Functional regeneration requires damaged axons to first start re-growing and then to continue to navigate through a strongly inhibitory environment, before they can reach their synaptic partners and establish functional connections. Therefore, both the presence of extrinsic inhibitory factors as well as a lack of intrinsic growth capacity prevent axonal regrowth in the injured CNS (Kaplan et al., 2015). Targeting extrinsic inhibitory factors has so far led to limited regeneration of injured axons (Cafferty et al., 2010; Lee et al., 2010), suggesting that creating a permissive environment is not sufficient to allow regeneration. Therefore, elucidation of the molecular mechanisms underlying the early steps of intrinsic axonal regrowth after CNS damage is critical to fully understand the regeneration process. Even though neural circuits retain a remarkable degree of synaptic plasticity in adulthood, the mature CNS can no longer support the robust axonal growth that was once required to establish neuronal connectivity during development, suggesting that the neuronal intrinsic growth ability is largely lost. Indeed, mammalian CNS axons show a higher regenerative capacity during earlier stages of development, illustrating the importance of intrinsic factors to CNS regenerative failure (Liu et

al., 2011; Shimizu et al., 1990). In PNS neurons, axonal injury results in a regeneration program that shares key molecular features with developmental axon growth (Harel and Strittmatter, 2006; Makwana and Raivich, 2005; Raivich and Makwana, 2007; Yaniv et al., 2012). In particular, the JNK pathway has emerged as a conserved signal for axonal growth and regeneration in the CNS and PNS in mammals, flies and worms (Arthur-Farraj et al., 2012; Ayaz et al., 2008; Li et al., 2012; Nix et al., 2011; Raivich et al., 2004; Raivich and Makwana, 2007). This suggests that conserved developmental axonal growth signaling pathways may be key targets to boost efficient regeneration after injury.

Studies in mice have made unique contributions to our understanding of the molecular basis of axonal regeneration. Nevertheless, the experiments are costly and time-consuming and necessitate a gene-by-gene approach. More recently, simpler genetic model organisms such *C. elegans* and *Drosophila* have proven useful to identify and study novel genes involved in axonal regrowth after injury (Ayaz et al., 2008; Chen et al., 2011; Fang and Bonini, 2012; Fang et al., 2012; Gabel et al., 2008; Kato et al., 2011; Leyssen et al., 2005; Yanik et al., 2004). Interestingly, unlike *C. elegans* neurons and developing *Drosophila* neurons, injured adult *Drosophila* CNS axons fail to regrow after injury, much like their mammalian counterparts (Ayaz et al., 2008). Furthermore, adult *Drosophila* CNS axons show remarkable morphological and genetic hallmarks of mammalian axonal responses to injury, including the formation of retraction bulbs, Wallerian degeneration of the distal fragment, transient upregulation of JNK, and regeneration upon activation of protein kinase A and JNK signaling (Ayaz et al.,



2008; Leyssen et al., 2005; MacDonald et al., 2006). This makes the *Drosophila* adult CNS a particularly powerful model system to systematically search for novel axonal regeneration genes.

Here, we perform a two-step genetic screen of ~300 genes selected by GO term, and identify 13 that promote axonal outgrowth during development in post-mitotic CNS neurons. We then test those genes in an adult *Drosophila* model of CNS injury. Using this approach we identify three robust axonal regeneration regulators, which we find to interact in a novel axonal growth and regeneration signaling pathway. Specifically, the deubiquitinating enzyme Fat facets (Faf) promotes axonal regrowth after injury via the Down syndrome cell adhesion molecule (Dscam1). Faf stabilizes Dscam1 by acting on Dscam1 3'-UTR through the kinase DLK1/Wallenda (Wnd). Faf and Dscam1 act upstream of JNK signaling and its nuclear effector Kayak (Fos). The functional role of Faf in promoting axonal regeneration appears to be conserved in mammals as demonstrated by the ability of the mouse homologue of Faf, Usp9X/FAM to also stabilize Dscam1 and promote axonal regrowth in the injured fly CNS.

## **RESULTS**

### **A genetic screen for axonal growth in development and after injury**

To perform a screen for axonal growth and regeneration (Fig. 1a,b), we selected genes which: 1) are associated with the Gene Ontology (GO) terms neural

development and neurite morphogenesis, 2) have Gal4 inducible transgenes available and 3) represent a diversity of molecular functions, including receptors, protein turnover, transcription factors, and chromatin modifiers (Supplementary data file 1). 307 genes matching these criteria were first tested for their ability to induce developmental axonal over-growth in small Lateral Neurons ventral (sLNv), a small cluster of neurons with a highly stereotyped axonal morphology which can be readily quantified (Helfrich-Forster et al., 2007; Leyssen et al., 2005) and that has been previously used to investigate the molecular mechanisms underlying regeneration in the fly CNS<sup>12</sup> (Fig. 1 a, Fig.2 a and b). The post-mitotic *Pdf-Gal4* driver was used to express GFP together with each of the selected genes, and the length of axonal growth was quantified in comparison with controls. Expression of 13 genes (4.2%) promoted significantly increased axonal growth with no obvious adverse effects on neuronal survival or axonal trajectory (Fig. 2 a-e). In a second selection step (Fig. 1 b), these 13 genes were evaluated in an acute sLNv axonal injury model in *Drosophila* brains explanted and kept in culture (Ayaz et al., 2008; Koch, 2012). Given their superficial location, sLNv axons are easily accessible for injury, and were physically severed using an ultrasonic microchisel. Using the temperature dependency of the UAS/Gal4 system, high expression levels of candidate genes were induced in adult flies starting at 24 h before injury. Axonal regrowth was defined as the growth of novel sprouts from the site of injury within four days. We used three parameters to evaluate axonal regrowth following injury: capacity of regrowth (the percentage of brains that exhibited at least one axonal sprout

grown de novo), total regrowth (defined as the sum of the lengths of novel sprouts), and the maximum projection distance (defined as the distance of the longest novel sprout from the site of injury to their terminus) (Fig. 2 a-d). Of the genes tested, 7 (*drl*, *Dscam1*, *faf*, *kay*, *pdm2*, *pum* and *sens*) showed enhanced regeneration in all three categories (Fig. 2e-h). Kayak is the fly homologue of Fos, a key transcription factor downstream of JNK signaling, confirming that the screen can identify bona fide regeneration genes. Dichaete, D, is an example of a gene that promoted axonal outgrowth during development (Fig 2. a and d), but failed to induce regeneration in most cases (Fig. 3 a and d) and often resulted in short sprouts with poor morphology, making it difficult to measure. Three genes (*dimm*, *dac* and *sqz*) caused axonal phenotypes such as defasciculation, blebbing or fragmentation, and were excluded from further analysis.

### **The ability of Faf to induce axonal regrowth is conserved and depends on its enzymatic activity**

Of the identified 7 genes, 3 in particular (*Dscam1*, *faf* and *kay*) appeared to consistently promote robust growth. We therefore asked whether these genes might be acting together in a novel regeneration pathway linking the cell surface to the nucleus. We began by analyzing the de-ubiquitinating enzyme Faf since it promoted the highest levels of regeneration across all criteria (Fig. 3 a-c and h). First, we confirmed that *faf* is also able to induce axonal overgrowth in other CNS neuronal populations, such as the Dorsal Cluster Neurons (DCNs) (Figure 3 - Figure Supplement 1), suggesting that Faf may be a general CNS axonal growth-

promoting factor. Ubiquitin-dependent protein regulation is critical in regulating many neuronal events, including axonal growth (Ambrozkiewicz and Kawabe, 2015; McCabe et al., 2004). However, the signaling pathways operating downstream of these enzymes are still largely unknown. To test whether the axonal growth induced by Faf was dependent on its deubiquitinase activity, we mutated a critical cysteine 1677 residue in the catalytic protease site to a serine (Chen and Fischer, 2000). In contrast to wild-type Faf, this mutated form of Faf was not able to significantly promote developmental axonal growth (Fig. 4 a,b and d). The mouse homologue of Faf, FAM/Usp9x, which can be active in *Drosophila* in other contexts (Chen et al., 2000; Wood et al., 1997), also induced robust sLNv axonal outgrowth (Fig. 4 c and d). Remarkably, even the yeast homologue of Faf, Ubp2, which only shares homology in the de-ubiquitination domain, induces sLNv axonal outgrowth very similar to Faf (Figure 4 – Figure Supplement 1). More importantly, both FAM and Faf, but not the enzymatic mutant Faf-Ser, induced significant axonal regeneration after injury (Fig. 3 e-h). These data suggest a conserved axonal growth and regeneration activity for Faf as a deubiquitinase enzyme.

### **Faf promotes axon regrowth in a JNK-dependent manner**

Faf has been shown to induce neuromuscular junction growth in *Drosophila* (DiAntonio et al., 2001) in a pathway that requires Wallenda (Wnd), a conserved MAPKK upstream of JNK signaling (Collins et al., 2006). Therefore, we tested whether Faf required Wnd to induce axonal growth. RNA interference knock-

down (RNAi KD) of *wnd* inhibited Faf-mediated axonal outgrowth (Figure 4 – Figure Supplement 2 a,b and i), whereas overexpression of *wnd*, but not a kinase-dead form of it, strongly promoted axonal outgrowth that essentially phenocopied *faf* overexpression (Figure 4 – Figure Supplement 2 a,c and d ). Moreover, overexpression of *wnd* also promoted axonal regrowth after injury (Figure 4 – Figure Supplement 2 g,h and j). Therefore, Wnd likely acts downstream of Faf, to modulate axonal growth and regeneration in response to *faf* overexpression. Similarly, RNAi KD of the *Drosophila* homologue of JNK, *basket* (*bsk*), completely inhibited *faf*-mediated axonal outgrowth (Fig. 4. I and j). Conversely, co-expression of *kay*, the JNK pathway effector we identified as strong promoter of outgrowth in development (Fig. 2 a and c) and after injury (Fig. 3 a,b,c and f) enhanced Faf-mediated axonal outgrowth (Fig. 4 j and k). Together, these data suggest that Wnd and JNK act downstream of Faf to induce axonal outgrowth and regeneration.

### **Faf stabilizes Dscam1 protein levels to promote axonal growth**

How might *faf* activate JNK signaling to induce axonal regeneration? During fly eye development *faf* mediates the internalization of the Notch ligand Delta (Overstreet et al., 2004), and Notch signaling has been proposed to enhance regeneration of developing neurons (Kato et al., 2011), though it has also been shown to act as a repressor of axonal regeneration (El Bejjani and Hammarlund, 2012). To test if *faf* interacted with Delta in the context of sLNv axonal growth, we tested both a RNAi KD as well as a dominant negative (DN) transgene, and

found that loss of Delta function in the sLNvs did not reduce the axonal outgrowth activity of *faf* (data not shown), suggesting an alternative mechanism in the context of axonal growth. Therefore, we reasoned that Faf might interact with different axon growth-promoting effectors.

Mammalian *Dscam1* (Qu et al., 2013) has been shown to be a regulator of JNK signaling. Interestingly, our screen identified *Dscam1* as one of the genes that most strongly and consistently promoted sLNv axonal outgrowth and regeneration (Fig. 2 a, Fig. 3 a-c). This prompted us to investigate the molecular mechanisms underlying the growth and regeneration activity of *Dscam1*. The *Dscam1* gene generates a large number of isoforms by alternative splicing of a plethora of extracellular domains and two transmembrane domains called TM1 and TM2 (Schmucker et al., 2000). We find that different isoforms containing either the TM1 (Fig. 2 a, Fig. 5 a), or the TM2 domain (not shown), and different extracellular domains (see methods) can induce axonal outgrowth, suggesting that induction of axonal growth may be a general property of Dscam1-mediated signaling independent of its isoform specificity. It has previously been reported that isoforms containing TM1 are dendrite specific (Shi et al., 2007). However, we find that upon overexpression these isoforms localize to both cell bodies and axonal terminals (Figure 5 – Figure Supplement 1). Conversely, *Dscam1* knock-down with two different RNAi lines (Watson et al., 2005) resulted in stunted sLNv axonal growth (Fig. 5 b and f). Finally, TM1-containing *Dscam1* isoforms, UAS-*Dscam1* 1.30.30.1 GFP (Fig. 1 j-l) and UAS-*Dscam1* 1.34.31.1. HA induce robust axonal regeneration after injury, with the latter being the strongest line (Fig. 5 c).

Both mammalian and fly Dscam1 are known to interact with p21 activating kinase (Pak) (Li and Guan, 2004; Schmucker et al., 2000) itself an upstream JNK Kinase. We find that inhibition of JNK activity, by using a dominant negative form of Bsk, completely abrogates Dscam1 mediated axonal growth (Fig. 5 d and f). Conversely, the expression of Kay completely reverses the loss of axon growth caused by *Dscam1* RNAi knock-down (Fig. 5 e and f). These data suggest that Dscam1 acts upstream of JNK signaling to induce axonal growth.

The fact that Faf and Dscam1 both promote axonal regeneration after injury and induce strikingly similar JNK-dependent axonal outgrowth phenotypes lead us to hypothesize that Faf and Dscam1 interact in this context. Indeed, we find that Faf-induced axonal outgrowth required Dscam1, as *Dscam1* knock-down almost completely abolished Faf-induced growth (Fig. 6 a and d). Consistent with this, co-overexpression of *faf* and *Dscam1* in the sLNvs induces stronger axonal outgrowth than *faf* overexpression alone (Fig. 6 b and d). Importantly, *Dscam1* knock-down also inhibits FAM/Usp9x mediated axonal outgrowth, indicating a conserved interaction (Fig. 6 c,d).

Faf antagonizes ubiquitination by cleaving the covalent bond between ubiquitin and a substrate protein (Huang et al., 1995), thereby leading to stabilization of proteins targeted for degradation. We asked if Faf might stabilize Dscam1 protein levels. Therefore, we expressed *Dscam1* alone or together with *faf*, *faf-Ser* mutant or mouse *FAM/Usp9x* in *Drosophila* S2 cells. Both Faf and FAM/Usp9x, but not the Faf-Ser mutant lead to a ~30% increase in Dscam1 protein levels

(Fig. 6 e and f), with no change in mRNA levels (Supplementary data file 2). However, we were unable to find evidence for Dscam1 ubiquitination in wild type or proteasome-inhibited S2 cells, nor a change in that status upon overexpression or knock-down of *faf* (data not shown). These data suggest that, at least in this context, Faf does not de-ubiquitinate Dscam1 directly. We then asked whether Faf enhances translation of Dscam1 through a mechanism that is mediated by the 3'UTR of *Dscam1*. To this end, we expressed a *Dscam1*-3'-UTR>GFP reporter in the sLNv alone or together with *faf*. We find that GFP levels are significantly upregulated in sLNv upon Faf overexpression (Fig 6 g,h and i).

Interestingly, Wnd has been shown to be a positive upstream regulator of Dscam1 in neurons. Specifically, using the same *Dscam1*-3'-UTR>GFP reporter construct it has been shown that translation of the reporter protein is enhanced by Wnd (Kim et al., 2013). Therefore, we tested whether Dscam1 is also required for Wnd-induced axonal growth in sLNv neurons and find that Dscam1 RNAi KD significantly decreases Wnd-induced growth (Figure 4 – Figure Supplement 2 e and f). Finally, we asked whether Faf enhances translation of Dscam1 through a mechanism that is mediated by the 3'UTR of *Dscam1*. To this end, we expressed the *Dscam1*-3'-UTR>GFP reporter in the sLNv alone or together with *faf*. We find that GFP levels are significantly upregulated in sLNv upon Faf overexpression (Fig 6 g,h and i).



## Discussion

In contrast to young neurons, injured adult CNS neurons exhibit very limited ability to self-repair, suggesting that the intrinsic regenerative capacity is lost during development. For example, it has been shown that the axon growth rate decreases dramatically with age in post-natal retinal ganglion cells (Goldberg et al., 2002). In addition, pioneer work from Filbin and colleagues demonstrated that developmental loss of the regenerative capacity of neurons in post-natal rats is mediated by a decline in the endogenous levels of neuronal cAMP within a few days after birth (Cai et al., 2001). Consistent with this evidence, transcription factors that regulate developmental axonal growth, such as members of the Kruppel-like family (KLFs), can promote regrowth of adult injured corticospinal tract and optic nerve axons (Blackmore et al., 2012; Moore et al., 2009). Several other intrinsic axonal regulators, including phosphatase and tensin homolog (PTEN), suppressor of cytokine signaling 3 (SOCS3), mTOR, Osteopontin and IGF-1 (Duan et al., 2015; Liu et al., 2010; Sun et al., 2011), have been previously identified in mammalian systems- though mostly on a gene by gene basis. More systematic approaches, such as quantitative proteomic analysis, have been recently employed to identify molecular pathways that are altered in injured retinal ganglion cells, and identified additional intrinsic regulators of regeneration, such as c-Myc (Belin et al., 2015). Taken together, all these studies suggest that manipulation of the intrinsic regenerative ability of mature neurons might be efficient strategies for enhancing the capacity of injured axons to regenerate. It is therefore crucial to discover factors that constitute intrinsic pro-regeneration

signaling pathways in a systematic manner. We have previously shown that the adult *Drosophila* CNS is a suitable model for studying axonal injury and regeneration (Ayaz et al., 2008). By exploiting this model along with the power of *Drosophila* genetic screens, we have uncovered a novel axonal regeneration pathway (Fig. 7) that links the stability of the neuronal cell surface receptor Dscam1, via the de-ubiquitination function of the enzyme Faf, to JNK signal, a major inducer of axonal regeneration in *C. elegans*, *Drosophila* and mouse (Ayaz et al., 2008; Li et al., 2012; Nix et al., 2011; Raivich et al., 2004; Raivich and Makwana, 2007).

E3 ubiquitin ligases are important in many aspects of mammalian brain development and function, by controlling neuritogenesis, modulating axon guidance and pruning, neuronal polarity and synaptic transmission (Ambrozkiwicz and Kawabe, 2015). Not surprisingly, E3 ligase dysfunction and abnormal ubiquitin signaling is implicated in several human brain disorders. In particular, mutations in the mammalian homologue of Faf, FAM/Usp9x, have been associated with X-linked intellectual disability (Homan et al., 2014). Brain specific deletion of FAM/Usp9x results in early postnatal death, and FAM/Usp9x knock-out neurons display reduced axon growth and impaired neuronal migration (Homan et al., 2014; Stegeman et al., 2013).

Ubiquitin-dependent signals that include Faf have also been shown to regulate synaptic development and growth at the *Drosophila* neuromuscular junction (NMJ) (DiAntonio et al., 2001), though its function in the CNS remained elusive.

Both yeast, Ubp2, and mouse homologues of Faf display the ability to induce axonal growth in the CNS, suggesting conservation of this property throughout evolution, similar to what had been shown at the NMJ (DiAntonio et al., 2001)(Kim et al., 2013). Importantly, we show for the first time that both Faf and FAM promote regrowth of injured axons in the adult fly brain.

Ubiquitin signaling is rather complex, and dissecting it downstream players can be challenging. In our screen, overexpression of the neuronal cell adhesion Dscam1 induced robust axonal growth both in development as well as after injury similar to the one induced by Faf, which led us to hypothesize that both genes acted in the same growth-promoting signaling pathway. In *Drosophila*, Dscam1 shows extensive molecular diversity that results from alternative splicing into some 18500 diverse extracellular domains (Schmucker et al., 2000). This isoform diversity has been shown to be critical for neuronal self-recognition and self-avoidance underlying axon growth and dendritic patterning (He et al., 2014; Hughes et al., 2007). Independent of its ectodomain diversity, Dscam1 has also been recently shown to regulate presynaptic arbor growth (Kim et al., 2013).

Our data indicate that post-transcriptional regulation of Dscam1 allows axonal growth after injury (Fig. 6). The Dscam1-stabilizing function of Faf appears to be conserved in mammals, as FAM/Usp9x overexpression also leads to increased levels of Dscam1 protein (Fig. 6 e,f). The placing of Faf upstream of Dscam1 and Wnd in regulating axonal outgrowth suggests that the de-ubiquitination activity of Faf may operate by antagonizing the E3 ubiquitin ligase activity of enzymes such

as Highwire (DiAntonio et al., 2001). Interestingly, Wnd itself is a promoter of mRNA stability and local translation, and is essential for axon regeneration after laser axotomy in adult neurons in *C. elegans* (Byrne et al., 2014; Yan et al., 2009).

Remarkably, a screen for axonal injury response in the mouse identified the differential regulation of transcript-availability and -loading onto ribosomes of CNS development genes as a major feature of abortive regenerative response in the mammalian spinal cord (see accompanying manuscript). Furthermore, overexpression of a post-transcriptional regulator, the cytoplasmic polyadenylation element-binding protein1/Orb, in both *Drosophila* and mouse neurons is sufficient to induce axonal regrowth after injury. Together, our studies, also point to a crucial role of mRNA stability and translational control of neurodevelopmental genes in the design of future therapeutic strategies.

## **METHODS**

### **Candidate gene sample**

We used the Gene Ontology tool (<http://geneontology.org/>) to select candidate genes annotated with the terms 'Neurite Morphogenesis', 'Transcription Factors', 'Receptors', 'Chromatin Modifiers' and 'Ubiquitin Ligases'. We also included an additional set of genes previously implicated in axonal growth and/or involved in actin dynamics (indicated in Supplementary data file 1 by asterisks).

The final candidate gene sample only included genes for which appropriate gain-of-function fly lines were readily available at the stock centers at the time of the study (Supplementary data file 1).

### **Fly stocks and genetics**

*Drosophila melanogaster* stocks were kept on standard cornmeal media. For tissue-specific overexpression of the transgenes, we used the GAL4/UAS system (Brand and Perrimon, 1993). Lines with UAS insertion sites (*i.e.* UAS, EP, EPgy2, XP and Mae-UAS) were received through the Bloomington or Szeged Stock Centres (or from specific laboratories when specified). Loss of function lines (Wnd RNAi GD8365, Dscam1 RNAi KK108835; Dscam1 RNAi (Watson et al., 2005); Bsk RNAi BL35594 and BL36643, and Bsk DN BL6409) were obtained from the Bloomington Stock Centre or from the Vienna *Drosophila* Research Centre (VDRC). UAS-Wnd kinase dead (KD) and UAS-Wnd E flies were a gift from C. Collins. The *PDF-Gal4* line was obtained from P. Taghert. For Faf overexpression in flies, we used the EP3520 line (Szeged Stock Centrum), which was previously reported to induce Faf gain-of-function (DiAntonio et al., 2001). UAS-Faf and UAS-FAM lines were created in house by cloning Faf cDNA and FAM cDNA into a pUAST-attB vector, respectively (Bischof et al., 2007) and injected in an attP2 docking line (BL 8622). A UAS-Faf serine (Faf-Ser) mutant that harbors a cysteine to serine mutation at residue 1677 was also cloned using the same method. UAS-Dscam1 HA-FLAG and UAS-Dscam1 GFP flies, both containing the 1.30.30.1 isoform, were used for the injury experiments. The UAS-

Dscam1 1.30.30.1 GFP flies have been described (Hughes et al., 2007) and the UAS-Dscam1 HA-FLAG flies were created by inserting a HA tag into the intracellular domain (after the 81st bp of exon 22) of isoform 1.34.31.1. The UAS-Dscam1 1.30.30.1 GFP flies have been used for the initial screen in development and after injury, and the UAS-Dscam1 1.30.30.1 HA has been used for additional confirmatory experiments in development and injury, as well as for epistasis experiments. While both lines induce significant axonal growth in development and after injury, UAS-Dscam1 1.30.30.1 HA appears to be the strongest of the two lines. For the genetic screen in development and after injury, pdf-Gal4, UAS-GFP; pdf-Gal4, UAS-2x eGFP/cyo flies were kept as a stock and used to drive expression of the various candidate genes, or crossed to wild-type Canton S (CS) flies. For the genetic epistasis experiments, pdf-Gal4, UAS-GFP; UAS-Dscam1 RNAi and pdf-Gal4, UAS-GFP; Faf EP 3520 flies were maintained as a stock and crossed to overexpression lines to uncover genetic interactions.

### **Developmental outgrowth screen**

To measure axonal outgrowth during development, flies were reared at 25°C and were dissected 2-10 days after eclosion. A minimum of 5 fly brain (10 sLNv projections) per genotype were stained with an anti-GFP antibody (to enhance the GFP signal), visualized under a fluorescent microscope equipped with a GFP filter and scored as 'growth' (when sLNv axonal projections appeared considerably longer than in controls) or 'no-growth' (when the length of sLNv projections was indistinguishable from controls or shorter). All genes were scored

growth promoting genes were confirmed as such in at least one independent experiment, and their growth inducing ability analysed by measuring the axonal sLNv dorsal axonal projections.

### **Whole brain explant culture injury system**

For the axonal regrowth analysis after injury, flies were reared at 18°C, in order to minimize overexpression effects during development, and shifted to 25°C the day before injury to allow optimal transgene expression.

Whole-brain explants on culture plate inserts were prepared and injured as described (Ayaz et al., 2008; Koch, 2012). In brief, Millicell low height culture plate inserts (Milipore) were coated with laminin and poly-lysine (BD Biosciences). Adult female flies were collected 2-10 days after eclosion and placed on ice. Fly brains were quickly and carefully dissected out in a sterile Petri dish containing ice cold Schneider's *Drosophila* Medium (GIBCO). Up to seven brains were placed on the membrane of one culture plate insert and culture medium (10 000 U/ml penicillin, 10 mg/ml streptomycin, 10% Foetal Bovine Serum and 10 µg/ml insulin in Schneider's *Drosophila* Medium) was added. sLNv axonal injury was performed using an ultrasonic microchisel controlled by a powered device (Eppendorf). Culture dishes were kept in a plastic box in a humidified incubator at 25°C

### **Immunohistochemistry**

Freshly dissected brains of adult flies were fixed in 4% formaldehyde and processed for immunohistochemistry as described (Hassan et al., 2000). Cultured brains (four days post-injury) were first fixed by replacing the culture medium in the Petri dish for 30 minutes. Then, 1 ml of fixative was carefully added on top of the filter for 1-2 hours. Brains that detached from the membrane were excluded from further analysis. Immunostaining was performed as for freshly dissected samples. Primary antibodies were rabbit anti-GFP (A-6455, Molecular Probes), rat anti-HA (3F10, Roche); and anti-Pdh (gift from P. Taghert).

### **Imaging and morphological analysis of sLNv axonal projections in development and after injury**

Image J software was used to measure the length of the dorsal axonal projections emanating from the sLNvs. The starting point was set as the point where axons turn medially and start to run parallel to the commissure. Axonal length was measured as a straight line (Computed Distance) from the starting point towards the midline (indicated by an asterisk) and as manual trace using Image J. The maximum computed distance was defined as the distance projected by the longest axonal sprout in a straight line and parallel to the commissure. The Average Length was defined as the average length of the two longest axonal branches traced manually (freehand distance). Imaging was performed on an upright Zeiss Axioscope equipped with a CCD camera, or on a



Zeiss 700 or Nikon AR1 confocal microscope. All measurements were performed using ImageJ.

To analyse the role of the candidate genes in axonal regrowth after injury we imaged cultured brains at two different time points after injury: approximately five hours and four days. Comparison between these two timepoints allowed us to define the location at which the injury took place, in order to define *de novo* growth. Morphometric analysis of axonal regrowth was always performed four days after injury, following fixation and GFP staining of the brains in culture. Capacity of regrowth was defined as the ability of the injured sLNv projection to regrow at least one new axonal sprout. Without the support of the head cuticle, brains will flatten and therefore undergo slight morphological changes during the culture process. To be conservative and account for potential inaccuracies in defining the injury point, only regrown axons with a minimum length of 12  $\mu\text{m}$  were defined as *de novo* growth and taken into account for analysis. To quantify axonal regrowth, newly grown axons were measured in a straight line and manually traced using Image J. In this case, the maximum computed distance was defined as the average of the distance of the two longest axonal sprouts in a straight line in any direction. Maximum growth was defined as the sum of the freehand lengths of all *de novo* grown axons. Images five hours after injury were acquired on an upright Nikon microscope equipped with a Hamamatsu CCD camera ORCA-R2. Imaging 4 days after injury was performed on a Zeiss 700 or a Nikon A1R confocal microscope after GFP immunostaining. See figure legends

for details of individual experiments, including statistical tests used and number of samples tested.

### **Cell culture and western blotting**

*Drosophila* Schneider's (S2) cells were maintained in Sf-900 II SFM medium (Gibco). To achieve transgene overexpression in Schneider's (S2) cells we electroporated a UAS construct in combination with PMT-Gal4, according to previously developed methods (Klueg et al., 2002). For Faf and FAM overexpression, we created a UAS-Faf and a UAS-FAM construct as described in "Fly stocks and Genetics". For Dscam1 overexpression, the UAS-Dscam1 1.30.30.1 GFP construct was used (Hughes et al., 2007).

Cells were electroporated using an Amaxa Nucleofector KitV (Lonza), according to the manufacturer's instructions. Cells were harvested 72-96 hours after copper induction, briefly washed with PBS and pellets frozen until cells were lysed in a 1% NP40 buffer in Tris-HCL. Protein concentration was determined by a modified Lowry assay (Peterson, 1977). Western blotting was performed with a SDS-PAGE Electrophoresis System (Biorad). Briefly, protein samples were diluted in SDS containing sample buffer and 15 µg per sample was loaded onto a 3-8% Tris-Acetate mini gel (Novex, Life Technologies). Samples were blotted using tank transfer to a nitrocellulose membrane (GE Healthcare), blocked with milk and probed with primary antibodies against Dscam1 (1:1000) (Watson et al., 2005) or against Actin (1:5000, ab3280, Abcam), which was used as a protein loading control. Anti-rabbit or anti-mouse horseradish peroxidase conjugated

secondary antibodies (Amersham) were then added, and proteins were detected using enhanced chemiluminescence (ECL Plus, GE Healthcare) on a FUJI LAS imager system (Fuji). Values for Dscam1 were normalized to the values of the loading control (actin) and quantified using the blot analysis function for IMAGE J. Kruskal Wallis test was used to compare the different conditions. Data is shown as mean  $\pm$  SEM and significance was set at  $p \leq 0.05$ .

### **RNA isolation and quantitative PCR**

RNA was extracted with Trizol. 1  $\mu$ g of total RNA was reverse transcribed using the Quantitect RT kit (Qiagen). qPCR using the Taqman Real Time protocol (Applied Biosystems) and probes (See Supplementary data file 2 for info Taqman probes). Data is shown as mean  $\pm$  SEM in Supplementary data file 2.

### **GFP intensity measurements**

Adult brains were dissected and immediately prepared for imaging. Confocal stacks of all sLNv and ILNv cell bodies in each side of the brain were performed. The optimal confocal settings were first adjusted for wild type brains and kept unchanged to allow comparison between genotypes. A maximum projection was created for each brain side and each image was quantified for GFP intensity using the 'Image Analysis' module of Zeiss Zen 2.0 software. All quantifications were done by an investigator blind to experimental conditions. Student T test was used to compare both genotypes. Data is shown as mean  $\pm$  SEM and significance was set at  $p \leq 0.5$ .

## MAIN FIGURE LEGENDS

### **Figure 1: A gain of function screen for axonal growth in development and after injury**

**(A,B)** Schematic illustrating the various outcomes for the development and injury steps of the screen. Both sLNVs (dark brown) and ILNVs (purple) are depicted in (A). Gain of function the candidate genes specifically in PDF neurons appeared to increase growth in 4.2% (n=13) of the cases. Genes that stimulated axonal growth were tested further in an injury paradigm in which the sLNV axonal projection is physically cut and sLNVs were assessed for regrowth four days post-injury (B). Seven genes retained the ability to promote significant regeneration of injured axons.

### **Figure 2: Analysis of axonal outgrowth in the developmental screen**

**(A)** Morphometric analysis (Maximum Computed Distance and Average Length) of sLNV axonal projections where developmental overexpression of candidate genes has been specifically induced in the sLNvs. Axonal outgrowth is measured in  $\mu\text{m}$ . Purple trace in schematic represents measured axonal length.

**(B-E)** Representative images of sLNV axonal arborization in wild type (control) adult flies (B) and in flies where developmental overexpression of *Kayak*, *kay* (C), *Dichaete*, *D* (D) and *Fat Facets*, *faf* (E) has been specifically induced in the sLNvs. Scale bars are 20  $\mu\text{m}$ .

Genotype of flies in (B) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+, in (C) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; EP *Kay*/+, in (D) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-*D*/+, in (E) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-*Faf*/+. Asterisk denotes the brain midline, red arrow denotes the injury point. \*,  $p < 0.05$ ; \*\*\*,  $p < 0.001$ . n.s. indicates no statistical significance. Error bars represent SEM. Dotted insets have been zoomed in to better illustrate the diverse axonal phenotypes obtained. Scale bars are 20  $\mu\text{m}$ .

### **Figure 3: Analysis of axonal regrowth in the regeneration screen**

**(A-D)** Analysis of axonal regrowth four days after injury.

**(A)** Percentage of brains where at least one regenerated axonal sprout is detected (Capacity of regrowth)

**(B,C)** Morphometric analysis (Maximum Computed Distance, (B) and Total Growth (C)) of regenerated sLNv axonal sprouts. Axonal regrowth is measured in  $\mu\text{m}$ .

**(D)** Schematic simplifying how length of the regrown axonal sprouts is assessed. Yellow dot shows the point of injury; red trace represents maximum axonal length; blue trace represents axonal length measured in a straight line.

**(E-H)** Representative images of sLNv axonal regrowth four days after injury in wild type (control) adult flies (E) and in flies where overexpression of *Kayak*, *kay* (F), *Dichaete*, *D* (G) and *Fat Facets*, *faf* (I) has been specifically induced in the sLNvs.

Genotype of flies in (E) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+, in (F) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; EP Kay/+, in (G) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-D/+, in (H) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-Faf/+. Asterisk denotes the brain midline, red arrow denotes the injury point. \*,  $p < 0.05$ ; \*\*\*,  $p < 0.001$ . n.s. indicates no statistical significance. Error bars represent SEM. Dotted insets have been zoomed in to better illustrate the diverse axonal phenotypes obtained. Scale bars are 20  $\mu\text{m}$ .

**Figure 4: Faf and FAM, but not Faf-Ser, promote axonal outgrowth in development and axonal regrowth after injury, and interact with the JNK signaling pathway.**

**(A-C)** Representative images of sLNv axonal arborization in adult flies where developmental overexpression of *Fat facets*, *faf* (A and A'), *Fat-Serine*, *Faf-Ser* (B and B'), and *FAM* (C and C') has been specifically induced in the sLNvs.

**(D)** Morphometric analysis (Maximum Computed Distance and Average Length) of sLNv axonal projections for (A-C). Axonal outgrowth is measured in  $\mu\text{m}$ .

**(E-G)** Representative images of sLNv axonal regrowth four days after injury in flies where overexpression of *faf* (E), *Faf-Ser* (F) and *FAM* (G) has been specifically induced in the sLNvs.

**(H)** Morphometric analysis (Maximum Computed Distance and Total growth) of regenerated sLNv axonal projections in (E-G). Axonal outgrowth is measured in  $\mu\text{m}$ .

**(I,J)** Representative images of sLNv axonal arborization for epistasis experiments between *faf* and *Bsk* (I and I') and *faf* and *kay* (J and J').

**(K)** Morphometric analysis (Average Length) of sLNv axonal projections where developmental overexpression of *faf*; *faf* and *Bsk* RNAi; *kay*; and *faf* and *kay*, has been specifically induced in the sLNvs. Axonal outgrowth is measured in  $\mu\text{m}$ .

Genotype of flies in (A, A' and E) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-Faf/+, in (B, B' and F) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-Faf-Ser/+, in (C, C' and G) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-FAM/+, in (I and I') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-Faf/UAS-*Bsk* RNAi, in (J and J') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; EP-Faf/EP-*kay*. Dotted insets have been zoomed in to better illustrate the diverse axonal phenotypes obtained. Asterisk denotes the brain midline, red arrow denotes the injury point. \*,  $p < 0.05$ ; \*\*\*,  $p < 0.001$ . n.s. indicates no statistical significance. Error bars represent SEM. Dotted insets have been zoomed in to better illustrate the diverse axonal phenotypes obtained. Scale bars are 20  $\mu\text{m}$ .

**Figure 5: *Dscam1* promotes axonal outgrowth in development and axonal regrowth after injury, and interacts with the JNK signaling pathway.**

**(A,B)** Representative images of sLNv axonal arborization in adult flies where developmental overexpression of *Dscam1* (A and A'), and *Dscam1* RNAi (B and B') has been specifically induced in the sLNvs.

**(C)** Representative image of sLNv axonal regrowth four days after injury in flies where overexpression of *Dscam1* has been specifically induced in the sLNvs.

**(D,E)** Representative images of sLNv axonal arborization demonstrating that inhibition of *Bsk* accomplished by overexpression of a dominant negative line of *Bsk* inhibits *Dscam1*- induced outgrowth (D and D'), and that overexpression of *kay* rescues the lack of axonal growth induced by overexpression of *Dscam1*-RNAi (E and E').

**(F)** Morphometric analysis (Average Length) of sLNv axonal projections where developmental overexpression of *Dscam1*; *Dscam1*-RNAi; *Dscam1* and Bsk DN; *Dscam1*-RNAi and *kay*, has been specifically induced in the sLNvs. Axonal outgrowth is measured in  $\mu\text{m}$ .

Genotype of flies in (A,A' and C) is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-*Dscam1*-HA/+, in (B and B') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-*Dscam1*-RNAi/+, in (D and D') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/UAS-Bsk-DN; UAS-*Dscam1*-HA/+, in (E and E') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-*Dscam1*-RNAi /EP-*kay*. Dotted insets have been zoomed in to better illustrate the diverse axonal phenotypes obtained. Asterisk denotes the brain midline, red arrow denotes the injury point. \*,  $p < 0.05$ ; \*\*\*,  $p < 0.001$ . n.s. indicates no statistical significance. Error bars represent SEM. Dotted insets have been zoomed in to better illustrate the diverse axonal phenotypes obtained. Scale bars are 20  $\mu\text{m}$ , with exception of C, which is 30  $\mu\text{m}$ .

### **Figure 6: Faf and Dscam genetically and biochemically interact**

**(A-C)** Representative images of sLNv axonal arborization demonstrating that knock-down of *Dscam1* inhibits Faf- induced outgrowth (A and A'), that co-overexpression of both *faf* and *kay* potentiates axonal growth (B and B') and that knock-down of *Dscam1* inhibits FAM induced outgrowth.

**(D)** Morphometric analysis (Average Length) of sLNv axonal projections where developmental overexpression of *faf*, *FAM*, *Dscam1* and *Dscam1* RNAi has been specifically induced in the sLNvs, uncovering gene interactions. Axonal outgrowth is measured in  $\mu\text{m}$ .

**(E,F)** Western blot and quantification showing increased levels of *Dscam1* protein following S2 electroporation of wild-type Faf and FAM, but not of Faf-Ser in comparison to control (UAS vector).

**(G-I)** GFP fluorescence analysis showing increased levels of 3'UTR-*Dscam1* following overexpression of *faf* (n=28) (H) in comparison to its control (n=24) (G). LNvs GFP average intensities are shown in (I).

Genotype of flies in (A and A') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; / UAS-Faf/UAS-*Dscam1*-RNAi, in (B and B') is PDF-Gal4, UAS-GFP/+;

PDF-Gal4, UAS-2x eGFP/+; / UAS-Faf/UAS-Dscam1-HA, in (C and C') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-FAM/ UAS-Dscam1-RNAi, in (G) is PDF-Gal4, UAS-GFP/+; UAS-3'-UTR-Dscam1-GFP/+; UAS-Faf/+, in (H) is PDF-Gal4, UAS-GFP/+; UAS-3'-UTR-Dscam1-GFP/+; TM6b/+. Dotted insets have been zoomed in to better illustrate the diverse axonal phenotypes obtained. Asterisk denotes the brain midline, \*,  $p < 0.05$ ; \*\*\*,  $p < 0.001$ . n.s. indicates no statistical significance in (D). Error bars represent SEM in (D) and in (I). Scale bars in (A-C) and (G-H) are 30  $\mu\text{m}$ .

## SUPPLEMENTAL FIGURE LEGENDS

### **Supplemental Figure 1: Faf gain of function promotes axonal growth in a distinct neuronal population**

**(A,B)** Overexpression of *faf* specifically in the Dorsal Cluster Neurons (DCNs) results in increased axonal growth (yellow arrows) (B), in comparison to wild-type flies (A).

Genotype of flies in (A and A') is ;UAS-GFP;ato-Gal4 14a, in (B and B') ;UAS-GFP;ato-Gal4 14a/UAS-Faf. Scale bars are 20  $\mu\text{m}$ .

### **Supplemental Figure 2: Ubp2 gain of function also promotes axonal growth**

**(A,B)** Overexpression of the yeast homologue of Faf, *Ubp2*, which shows conservation of the enzymatic domain, also results in increased axonal growth (B), in a similar manner to Faf overexpression (A).

Note that B and B' are the same as in Fig. 4 A and A'

Genotype of flies in (A and A') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-Faf/+; in (B and B') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/UAS-Ubp2. Dotted insets have been zoomed in to better illustrate the diverse axonal phenotypes obtained. Scale bars are 30  $\mu\text{m}$ .

### **Supplemental Figure 3-Wallenda promotes growth in development and after injury and is required for Faf-induced growth.**



**(A,B)** Representative images of sLNv axonal arborization demonstrating that knock-down of *wnd* inhibits Faf-induced outgrowth.

**(C,D)** Representative images of sLNv axonal arborization in adult flies where developmental overexpression of *wnd* (C and C'), but not of a kinase dead form (D and D') in the sLNvs induces axonal growth similar to the one induced by Faf (A and A').

**(E,F)** Representative images of sLNv axonal arborization demonstrating that knock-down of *Dscam1* inhibits Wnd- induced outgrowth (E and E') and results in a phenotype that resembles knock-down of *Dscam1* on its own (F and F').

**(G,H)** Overexpression of *wnd* in the sLNvs (G and G'), but not of Wnd KD (H and H') induces axonal regrowth four days after injury. Five hour after injury timepoints (G and H) have been included to better illustrate the regenerative ability of Wnd, but not of its kinase dead (KD) form. Red arrows point to the place of injury.

**(I)** Morphometric analysis (Average Length) of sLNv axonal projections where developmental overexpression of *faf* and Wnd RNAi has been specifically induced in the sLNvs, uncovering a Faf-Wnd gene interaction. Axonal outgrowth is measured in  $\mu\text{m}$ .

**(J)** Percentage of brains showing at least one regenerated axonal sprout four days after injury (Capacity of regrowth), where overexpression of *wnd* and Wnd KD has been specifically induced in the sLNvs.

Note that A and A' are the same as in Fig. 4 A and A', and F and F' the same as in Fig. 5 B and B'

Genotype of flies in (A and A') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-Faf/+, in (B and B') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-Faf/Wnd RNAi;, in (C, C' and G,G') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/UAS-Wnd E;, in (D,D' and H,H') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-GFP,UAS-Wnd KD, in (E and E') is PDF-Gal4, UAS-GFP/+; /UAS-Wnd E/ Dscam RNAi;, in (F and F') is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/Dscam RNAi;. Dotted insets have been zoomed in to better illustrate the diverse axonal phenotypes obtained. Scale bars are 20  $\mu\text{m}$ .

## **Supplemental Figure 4- Dscam containing the TM1 domain localizes to both axonal projections as well as cell bodies and dendrites of sLNvs.**

**(A,B)** A Dscam form containing the TM1 domain (Dscam1-1.34.31.1 HA) localizes to both dendrites and axonal projections, and to the cell bodies. An antibody against the pigment dispersing factor hormone (PDF) specifically stains PDF neurons (A and B). Dscam expression pattern was visualized using an antibody against HA (A' and B').

Genotype of flies is PDF-Gal4, UAS-GFP/+; PDF-Gal4, UAS-2x eGFP/+; UAS-Dscam1-1.34.31.1.HA. Scale bars are 30  $\mu$ m.

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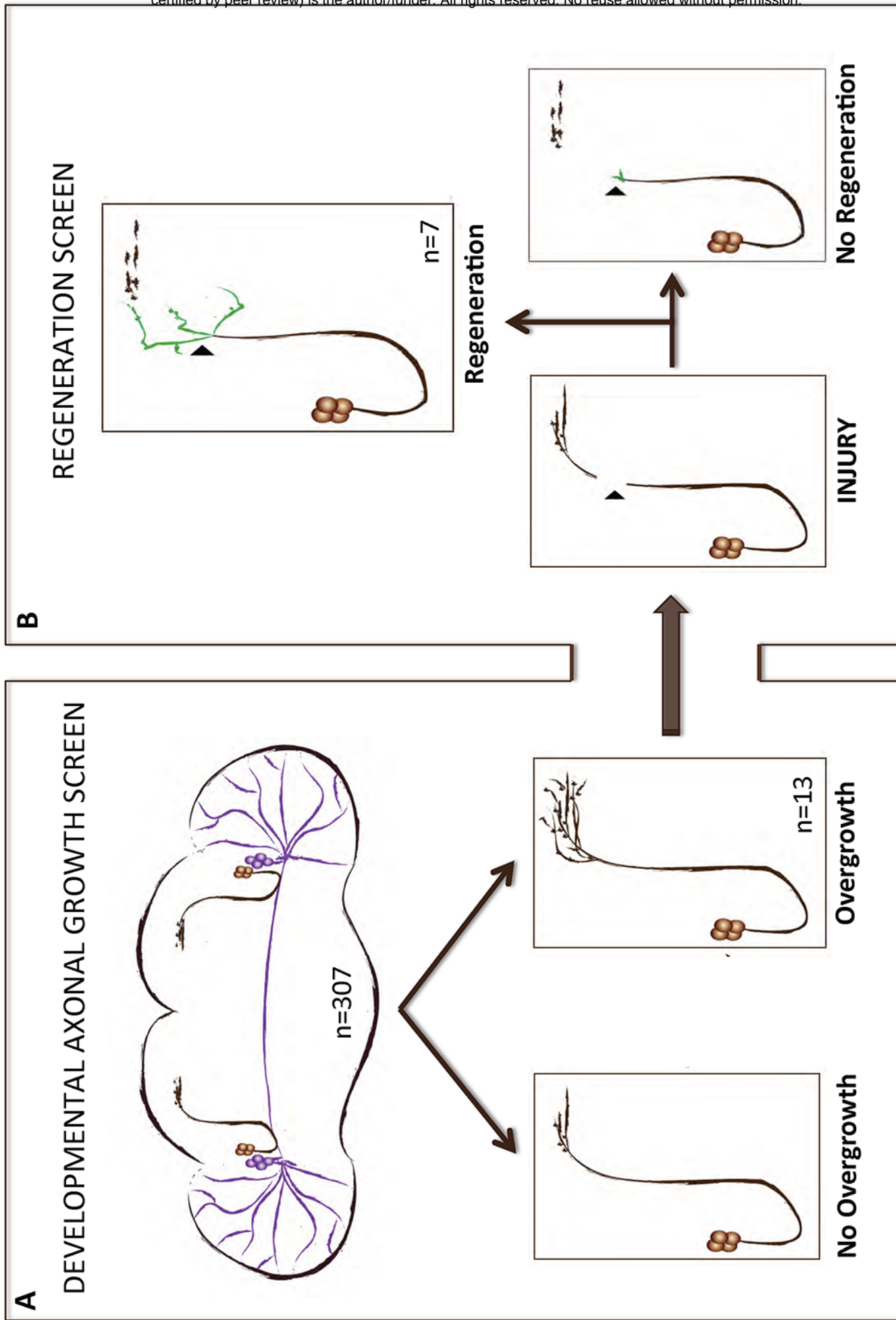


FIGURE 1

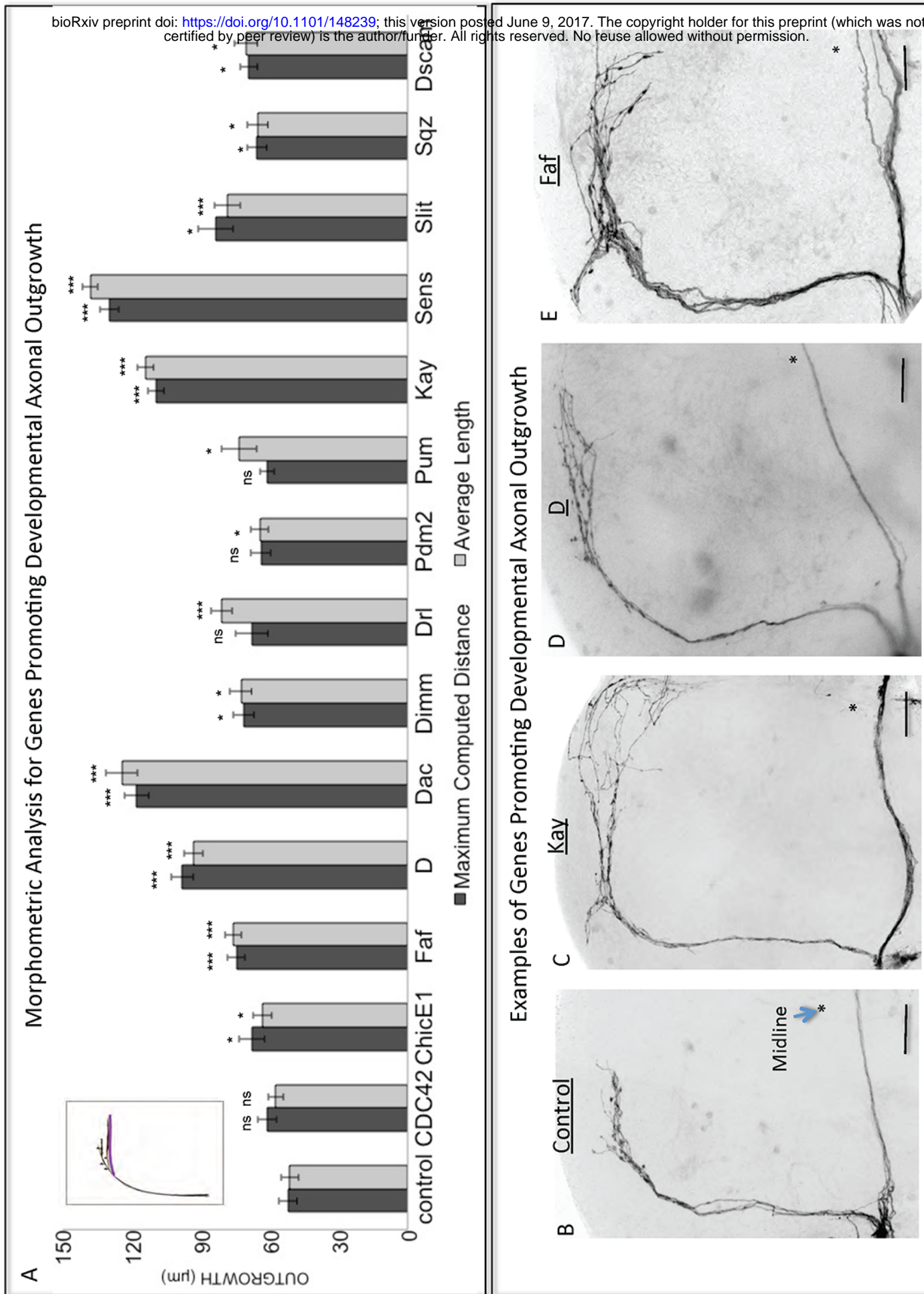
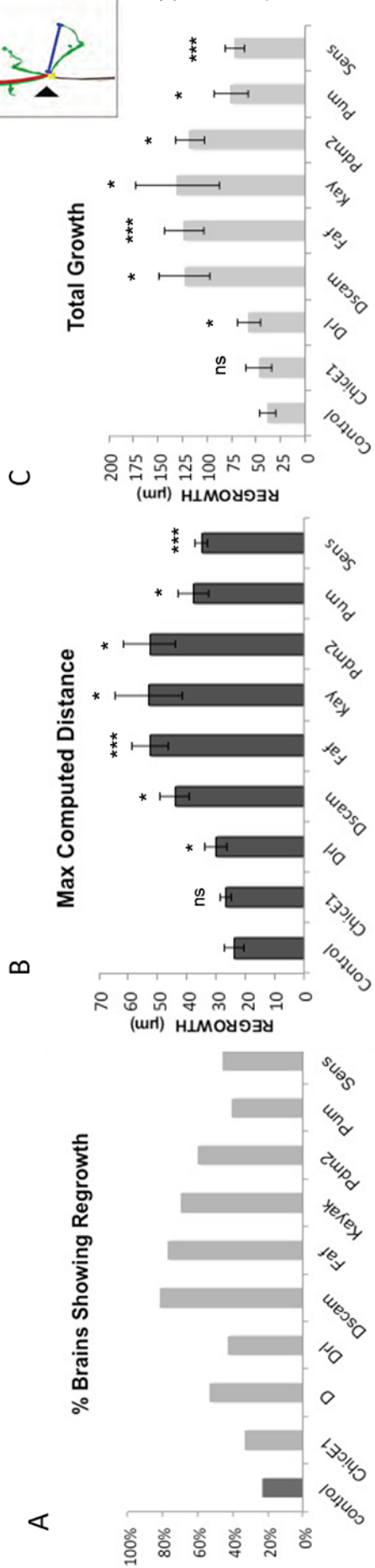


FIGURE 2



### Morphometric Analysis for Genes Promoting Axonal Regrowth After Injury



### Examples of Genes Tested in the Injury Model

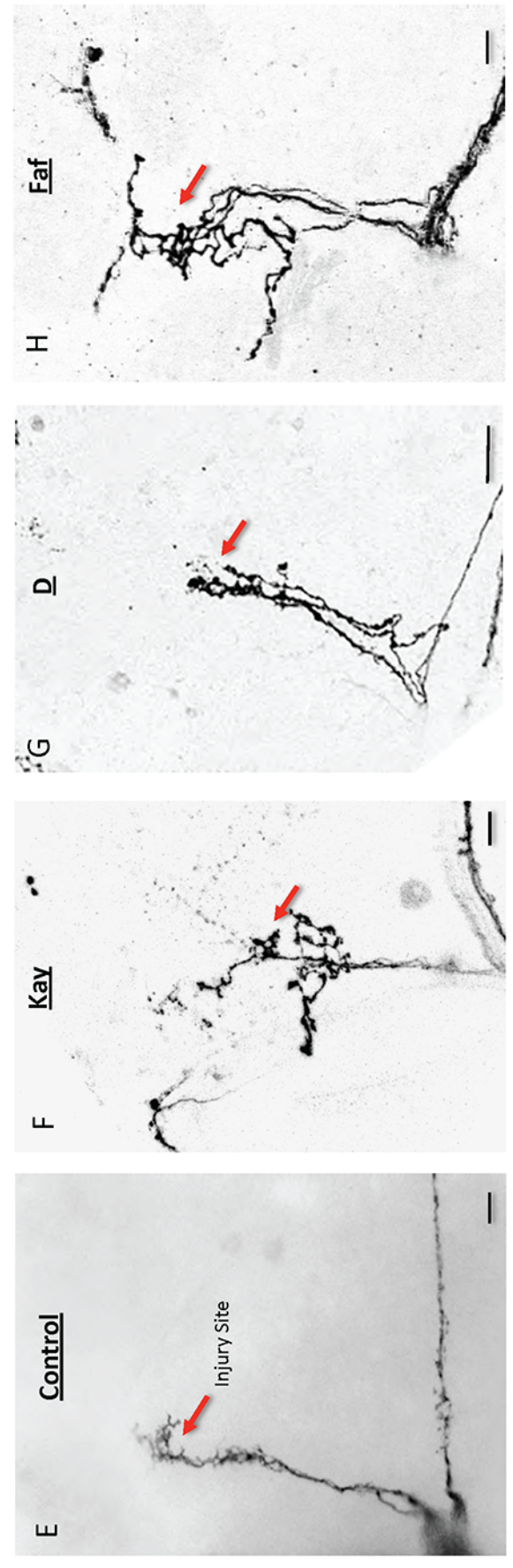


FIGURE 3

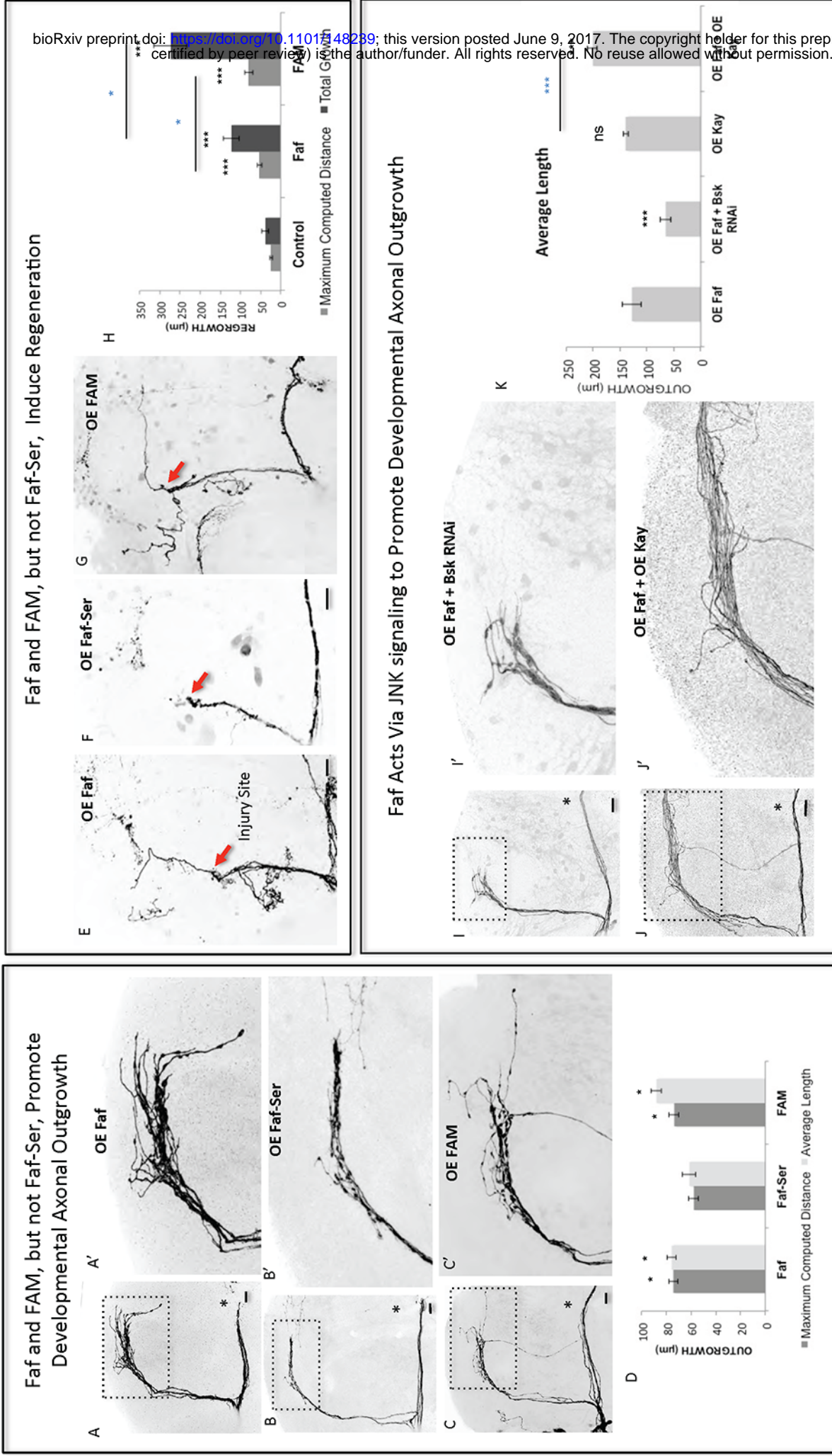


FIGURE 4

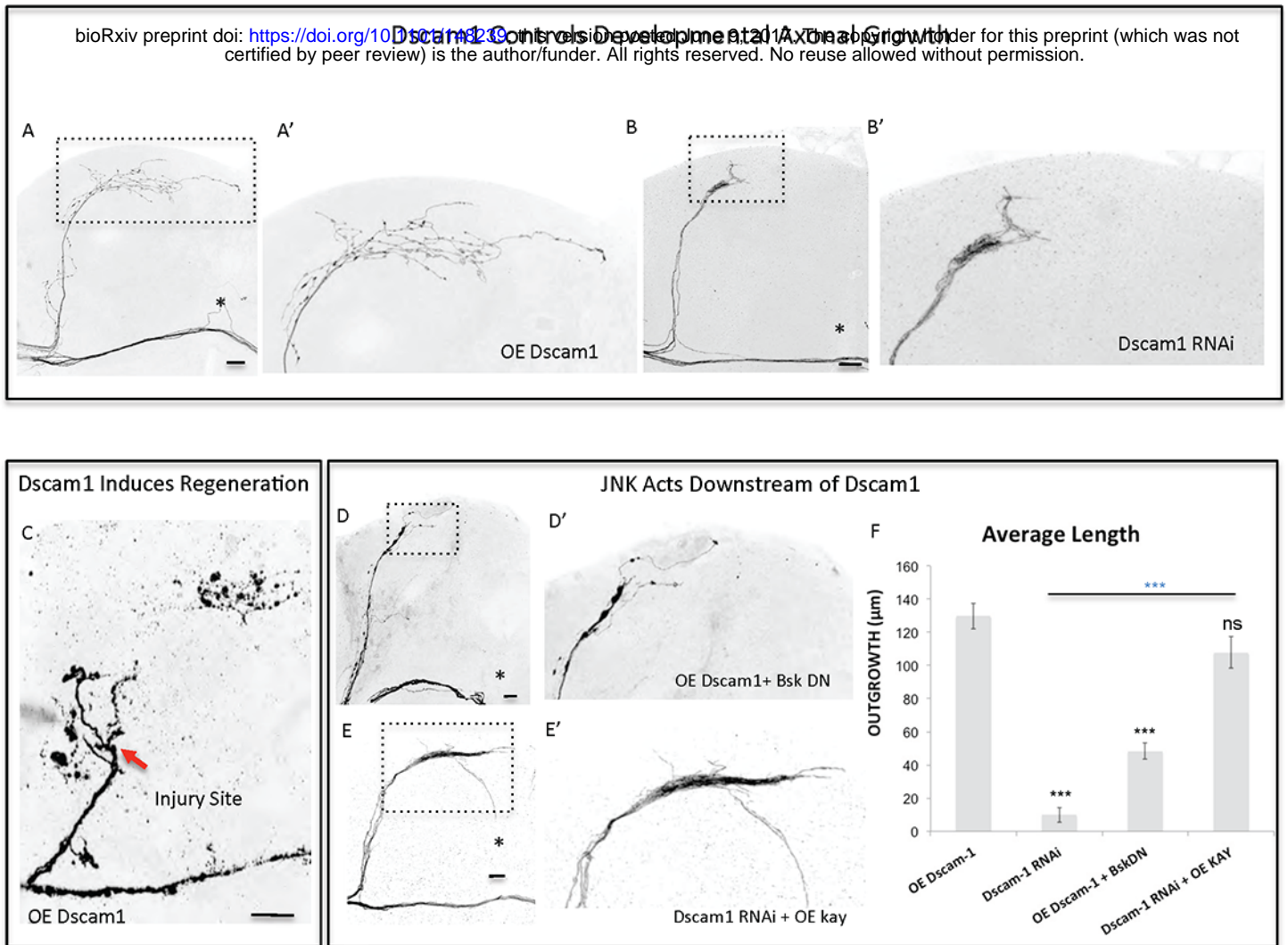
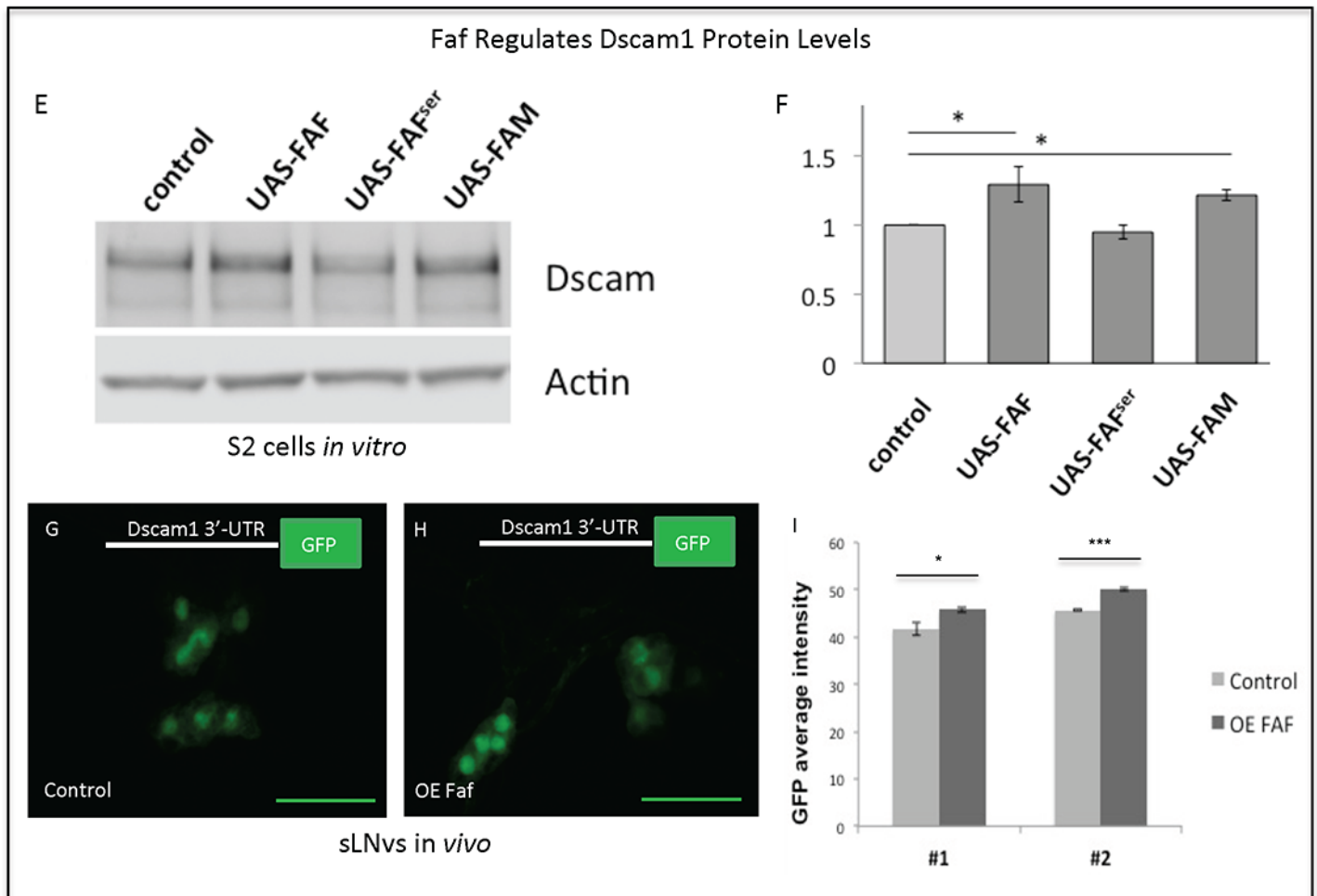
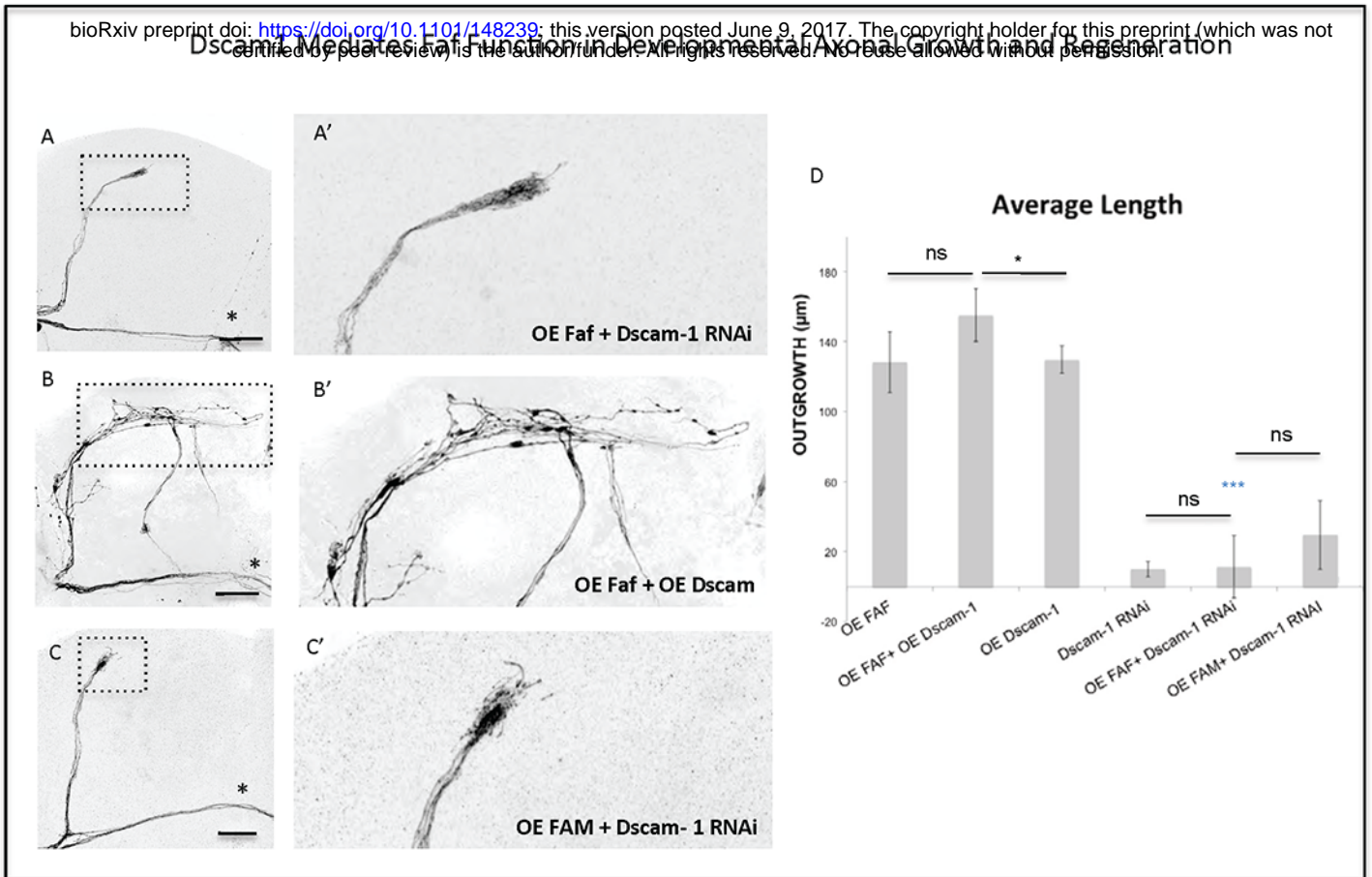


FIGURE 5



**Dscam1 Mediates Faf Function in Developmental Axonal Growth and Regeneration**



**FIGURE 6**

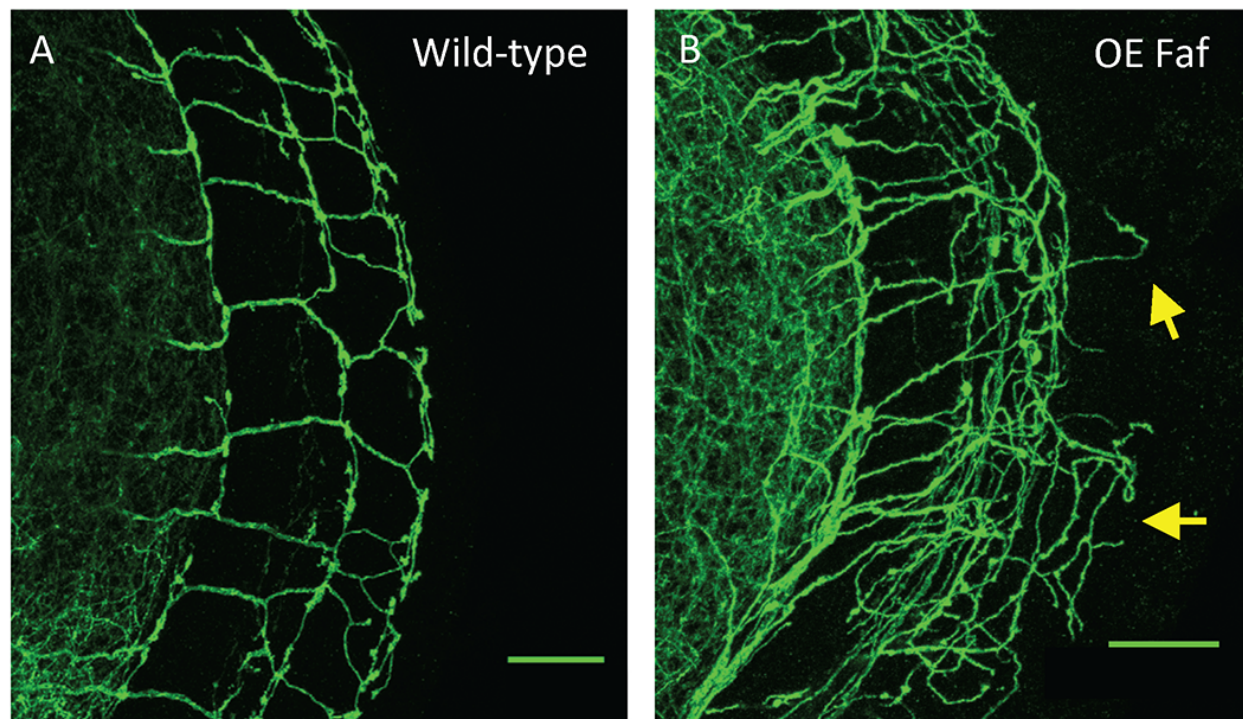


Figure 3 - Figure Supplement 1

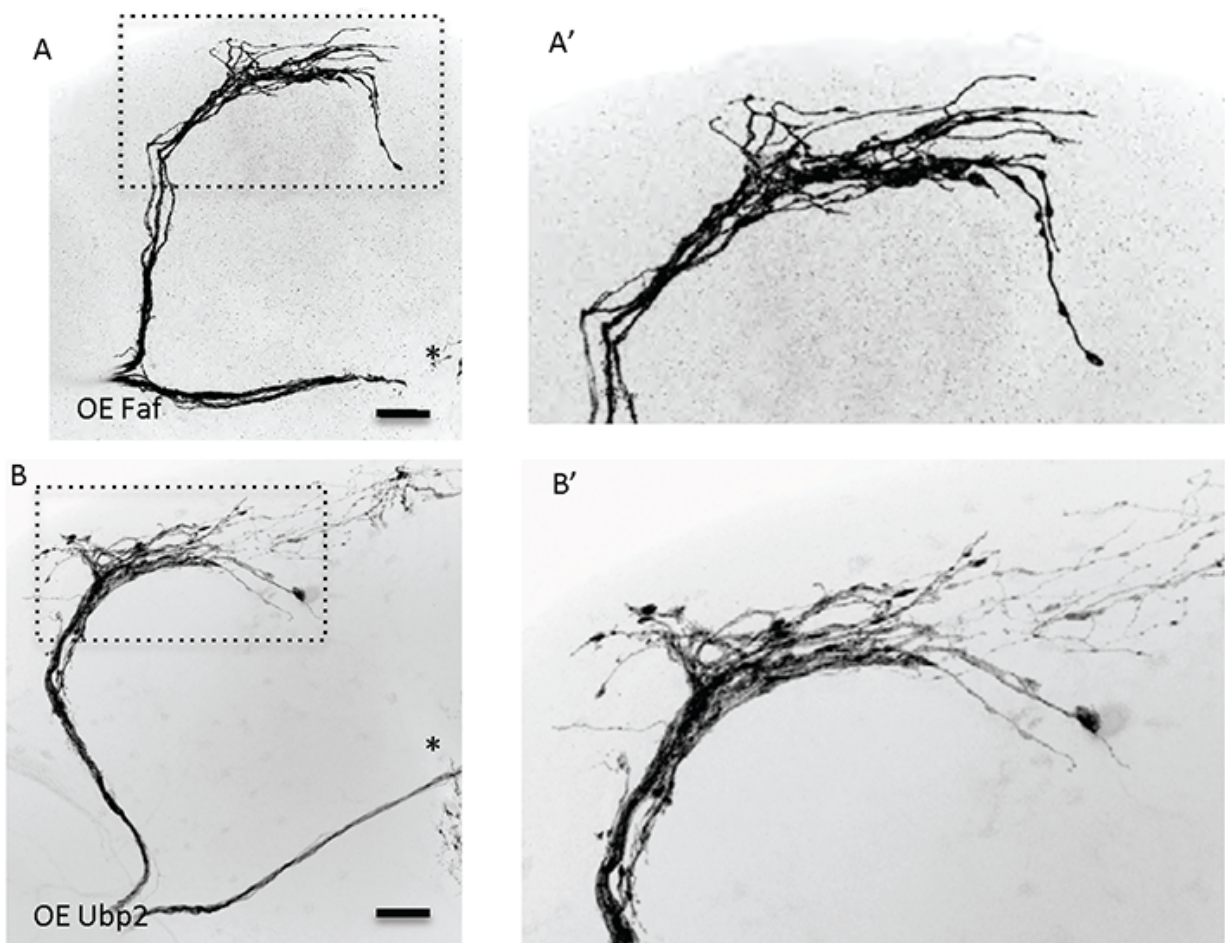


Figure 4 – Figure supplement 1



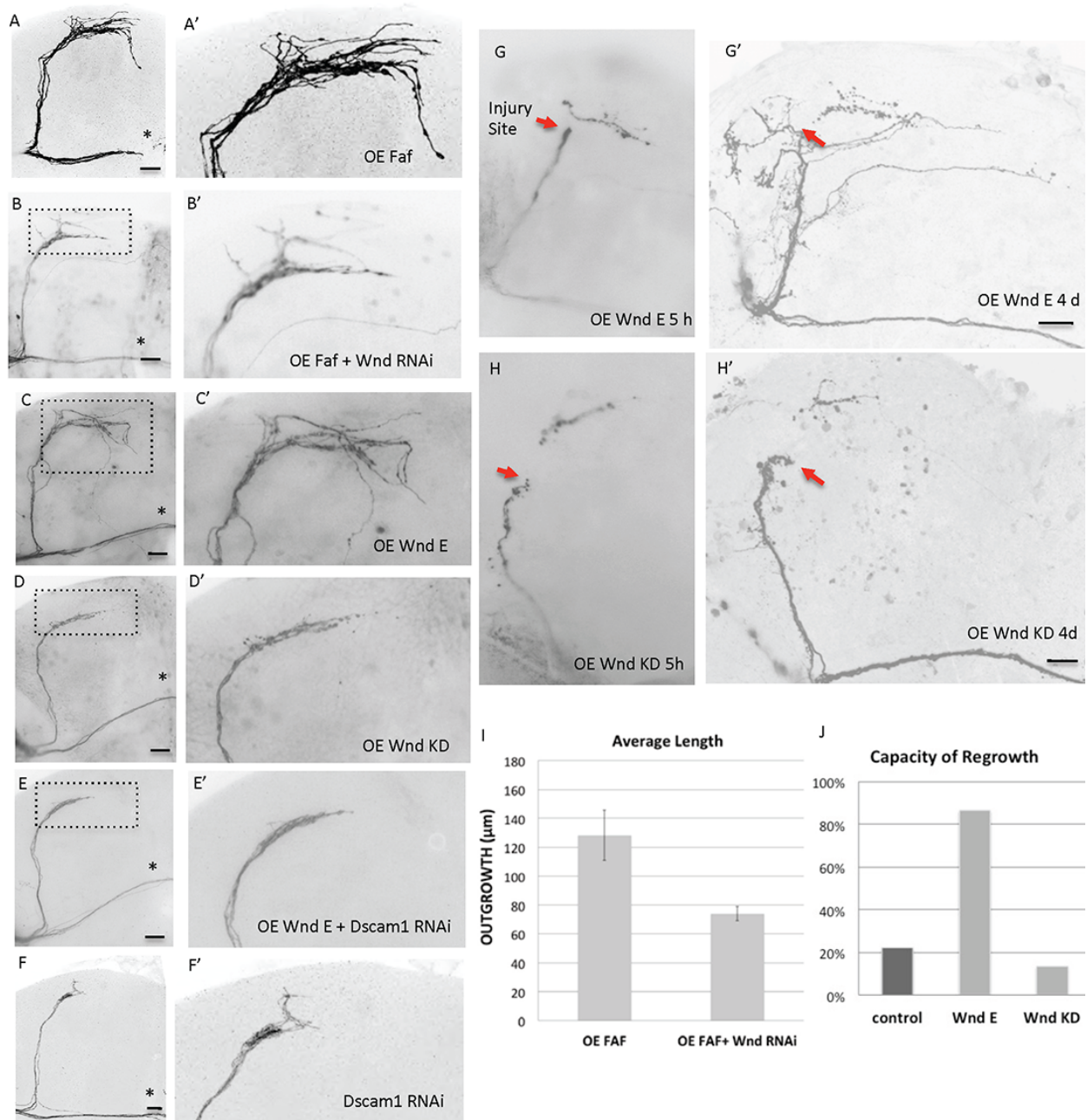


Figure 4 – Figure Supplement 2

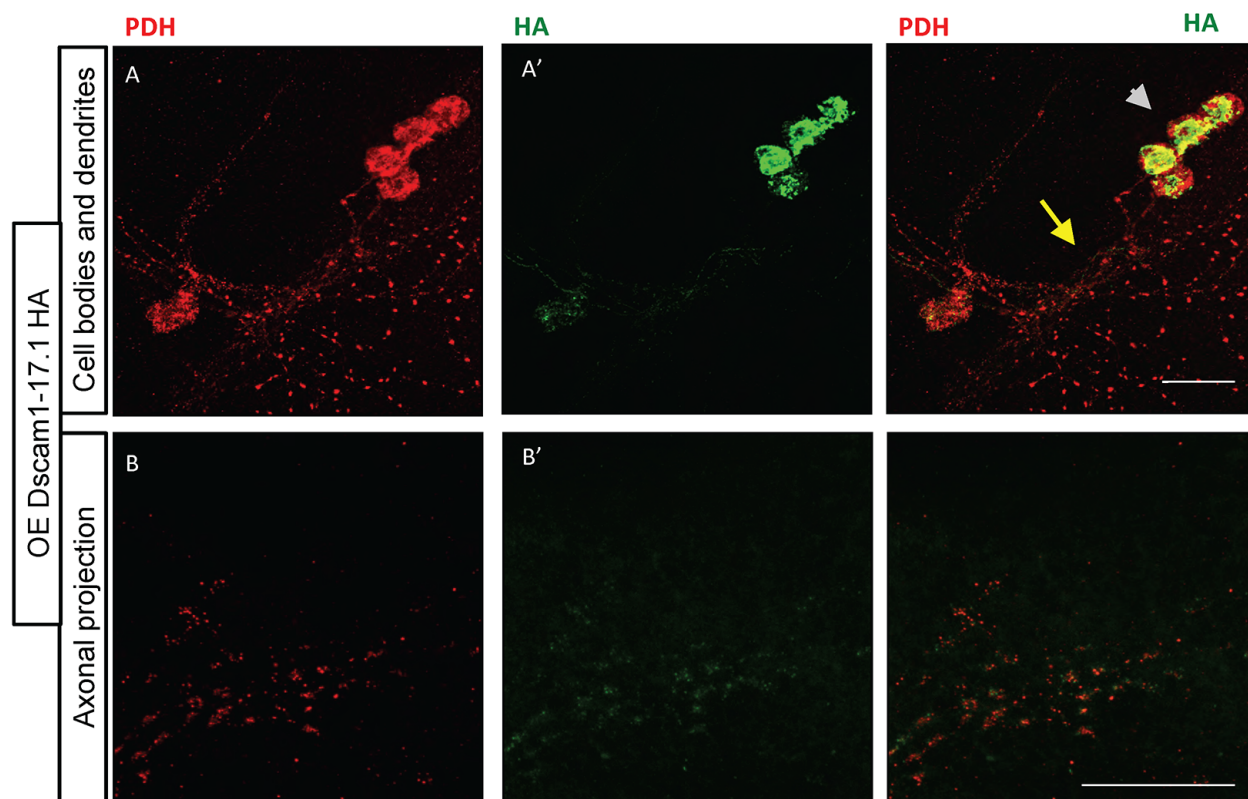


Figure 5 – Figure Supplement 1



<b>GENE NAME</b>	<b>SYMBOL</b>
chromatin modifier	
Argonaute-1	AGO1
ballchen	ball
bonus	bon
brahma	Brm
burgundy	bur
Enhancer of bithorax	e(bx)
Fmr1	Fmr
HDAC6	HDAC6
Histone demethylase	CG15835
Histone demethylase 4B	Kdm4B
Histone methyltransferase 4-20	suv4-20
Imitation SWI	lswi
Invadolysin	Invadolysin
Lamin	lamin
lola like	lolal
moira	mor
Origin recognition complex subunit 6	Orc6
piwi	piwi
Ploy (ADP-ribose) glycohydrolase	Parg
Polycomblike	Pcl
reptin	rept
Rpd3	Rpd3
Sir2	sir2
taranis	Tara

TBP-associated factor 1	Taf1
telomere fusion	tefu
tonalli	tna
Topoisomerase 2	Top2
Tousled-like kinase	tlk

### E3 ubiquitin ligases

archipelago	ago
ariadne	ari1
ariadne-2	ari2
CG2924	CG2924
CG4502	CG4502
CG5823	CG5823
CG8184	CG8184
CG8188	CG8188
COP9 complex homolog subunit	CSN4
courtless	crl
Cullin-2	cul2
Cullin-3	cul3
effete	eff
fat facets	faf
FBX011 ortholog	FBX011
HERC2	Herc2
highwire	hiw
morula	mr
neuralized	neur

partner of paired	ppa
purity of essence	poe
Roc1a	Roc1a
septin interacting protein 3	Sip3
supernumerary limbs	slmb
thread	th
TNF-receptor-associated factor	traf4
Topoisomerase I-interacting prc	Topors
tumor supressor protein 101	TSG101
Ubc-E2H	Ubc-E2H
Ubiquitin activating enzyme 1	Uba1
Ubiquitin conjugating enzyme	UbcD6
Ubiquitin conjugating enzyme 1	UbcD10
Ubiquitin conjugating enzyme 1	Ubc12
Ubiquitin conjugating enzyme 4	UBcD4
vihar	vih
yorkie	yki

## neurite morphogenesis

Abl tyrosine kinase	Abl
abrupt	ab
Actin-related protein 3	Arp3
Adh transcription factor 1	Adf1
anti-silencing factor 1	asf1
argos	aos
armadillo	arm

ATP-dependent chromatin assemblase	Acf1
Axin	Axin
bifocal	bif
Brahma associated protein 55kDa	Bap55
branchless	bnl
Calcium/calmodulin-dependent protein kinase II	cAMKII
cAMP-dependent protein kinase	Pka-R2
capricious	caps
Cdc42	Cdc42
CG1244	MEP-1
chickadee	Chic
Chip	chi
chromosome bows	chb
commissureless	comm
costa	cos
dachshund	dac
Daughters against dpp	Dad
dreadlocks	dock
egghead	egh
Ephrin	Ephrin
failed axon connections	fax
Fasciclin 1	Fas1
Fasciclin 2	FasII
frayed	fray
futsch	futsch
GTPase-activating protein 1	RasGAP1

## Guanine nucleotide exchange factors Gef64c

hedgehog	hh
inscuteable	insc
jelly belly	jeb
Katanin 60	katanin
Kinesin-like protein at 10A	klp10A
kuzbanian	kuz
Laminin A	LanA
LIM-kinase1	LIMK1
Lissencephaly-1	Lis1
longitudinals lacking	lola
Mi-2	Mi-2
multiprotein bridging factor 1	mbf1
nanos	nos
Nedd4	Nedd4
Netrin-B	NetB
Neurotactin	Nrt
non-stop	not
PDZ domain-containing guanine	PDZ-GEF
pole hole	phl
Ptpmed	Ptpmeg
pumilio	pum
pygopus	pygo
Rac1	Rac1
Rac2	Rac2
Ras oncogene at 85D	Ras85D

Rho GTPase activating protein a	RhoGAP93B
Rho GTPase activating protein p	RhoGAPp190
Rho guanine nucleotide exchan	RhoGEF2
rhomboid	rho
sequoia	Seq
Serrate	Ser
shrub	shrb
skittles	Sktl
slit	slit
smooth	sm
specifically Rac1-associated pro	sra-1
spitz	spi
split ends	spen
sprouty	sty, spry
Star	Star
Syndecan	sdc
tramtrack	ttk
trio	trio
Tropomyosin 1	Tm1
twinstar	tsr
unzipped	uzip
WASp	WASp
wingless	wg
Wnt oncogene analog 5	Wnt5

Receptor

## Adipokinetic hormone receptor AkhR

Alk	alk
arrow	arr
breathless	btl
brother of ihog	boi
Cadherin 96Ca	cad96ca
CG12484	CG12484
CG14516	CG14516
CG2061	CG2061
CG31221	CG31221
CG3339	CG3339
CG4313	CG4313
CG7536	CG7536
CG9643	CG9643
croquemort	crq
crumbs	crb
cryptochrome	cry
cueball	cue
Cullin-5	Cul5
dachsous	ds
derailed	drl
distracted	dsd
domeless	dome
doughnut on 2	dnt
Down syndrome cell adhesion n	dscam
draper	drpr

Ecdysone receptor

EcR

embargoed

emb

Epidermal growth factor receptor

Egfr

frazzled

fra

frizzled

fz

frizzled 3

fz3

Furin 2

Fur2

giant fibre A

gfA

heartless

htl

Hormone receptor-like in 38

Hr38

Hormone receptor-like in 39

Hr39

Hormone receptor-like in 46

Hr46

Hr4

Hr4

Insulin degrading metalloproteinase

Idc

KDEL receptor

KdelR

kekkon-1

kek1

LDL receptor-like 1

LRP1

leak

lea

lethal (1) G0232

l(1)G0232

Lipophorin receptor 2

LpR2

methuselah-like 1

mtlh1

methuselah-like 3

mtlh3

moody

moody

mutator 2

mu2



Neurexin IV	Nrx-IV
nicotinic Acetylcholine Receptor	nAcRalpha-30D
nicotinic Acetylcholine Receptor	nAcRalpha-96Aa
NMDA receptor 1	Nmdar 1
Notch	N
off-track	otk
plexin A	plexA
Protein tyrosine phosphatase 10	Ptp10D
Protein tyrosine phosphatase 4E	Ptp4E
Protein tyrosine phosphatase 9A	Ptp99A
proximal to raf	ptr
Resistant to dieldrin	Rdl
Rhodopsin 5	Rh5
rolling pebbles	rols
roundabout	Robo
Ryanodine receptor	RyR
scab	Scb
Sema-1a	Sema-1a
Sema-2a	Sema-2a
Semaphorin-5c	Sema-5c
shifted	shf
shotgun	shg
Sialic acid phosphatase synthase	Sas
Signal recognition particle protein	Srp9
Signal sequence receptor beta	ssRbeta
Tachykinin-like receptor at 99D	Takr99D

thickveins	tkv
Trapped in endoderm 1	Tre1
unc-5	unc-5
Vacuolar H <sup>+</sup> ATPase M8.9 acces	VhaM8.9

## Transcription factor

Alhambra	Alh
Antennapedia	Antp
anterior open	aop
atonal	ato
BarH2	B-H2
bicaudal	bic
bifid	bi
bigmax	bigmax
brinker	brk
broad	br
bunched	bun
cap-n-collar	cnc
CG13624	CG13624
CG30001	CG30001
CG32601	$\beta$ NACtes3
CG32778	CG32778
CG4882	CG4882
CG4914	CG4914
charlatan	chn

## Checkpoint suppressor homologs Ches-1-like

combgap	cg
CTCF	CTCF
CTP synthase	CTPsyn
Cyclic-AMP response element b	CrebB-17A
Cyclin-AMP response element b	CrebA
defective proventriculus	dve
Dichaete	D
dimmed	dimm
DNA replication-related element	Dref
Dorsal interacting protein 3	Dip3
Dorsal-related immunity factor	Dif
doublesex	dsx
E(spl)region transcript m7	HLHm7
E(spl)region transcript mdelta	HLHmDelta
E2F transcription factor	E2f
E2F transcription factor 2	E2f2
engrailed	en
erect wing	ewg
Ets at 97D	Ets97D
forkhead box, sub-group 0	foxo
forkhead domain 3F	fd3F
fruitless	fru
hamlet	ham

Hepatocyte nuclear factor 4	Hnf4
jim	Jim
jing	Jing
jumeau	jumu
kayak	kay
knirps	kni
lilliputian	lilli
Limpet	Lmpt
luna	luna
Meiotic central spindle	Meics
missing oocyte	mio
mitochondrial ribosomal protein mRpL55	mRpL55
Mnt	Mnt
MTA1-like	MTA1-like
Myocyte enhancer factor 2	Mef2
NFAT homologue	NFAT
NK7.1	NK7.1
p53	p53
pebbled	peb
pointed	pnt
POU domain protein 2	pdm2
regular	rgr
Relish	Rel
scalloped	sd
schnurri	shn
scribbler	sbb

senseless	sens
Signal-transducer and activator	stat 92E
similar	sima
Sin3A	sin3A
Six4	six4
squeeze	sqz
TAK1-associated binding protein	Tab2
Ultrabithorax	ubx
vrille	vri
Zinc finger protein 30C	Zf30c

Chromatin Modifier  
E3 ubiquitin ligase  
Receptor  
Transcription factor  
Neurite Morphogene



## **CG number**

CG6671

CG6386

CG5206

CG5942

CG9242

CG32346

CG6203

CG6170

CG15835

CG33182

CG13363

CG8625

CG3953

CG6944

CG5738

CG18740

CG1584

CG6122

CG2864

CG5109

CG9750

CG7471

CG5216

CG6889

CG17603

CG6535

CG7958

CG10223

CG34412



CG15010

CG5659

CG5709

CG2924

CG4502

CG5823

CG8184

CG8188

CG8725

CG4443

CG1512

CG42616

CG7425

CG1945

CG9461

CG11734

CG32592

CG3060

CG11988



CG9952

CG14472

CG16982

CG1937

CG3412

CG12284

CG3048

CG15104

CG9712

CG2257

CG1782

CG2013

CG5788

CG7375

CG8284

CG10682

CG4005



CG4032

CG4807

CG7558

CG15845

CG9383

CG4531

CG11579

CG1966

CG7926

CG1822

CG6546

CG4608

CG18069

CG15862

CG11282

CG12530

CG1244

CG9553

CG3924

CG32435

CG17943

CG1708

CG4952

CG5201

CG3727

CG9659

CG1862

CG4609

CG6588

CG3665

CG7693

CG34387

CG6721

CG32239

CG4637

CG11312

CG30040

CG10229

CG1453

CG7147

CG10236

CG1848

CG8440

CG12052

CG8103

CG4143

CG5637

CG42279

CG10521

CG9704

CG4166

CG9491

CG2845

CG1228

CG9755

CG11518

CG2248

CG8556

CG9375

CG3421

CG32555

CG9635

CG1004

CG32904

CG6127

CG8055

CG9985

CG8355

CG9218

CG4931

CG10334

CG18497

CG1921

CG4385

CG10497

CG1856

CG18214

CG4898

CG4254

CG3533

CG1520

CG4889

CG6407



CG11325

CG8250

CG5912

CG32134

CG32796

CG10244

CG12484

CG14516

CG2061

CG31221

CG3339

CG4313

CG7536

CG9643

CG4280

CG6383

CG3772

CG12086

CG1401

CG17941

CG17348

CG5634

CG14226

CG17559

CG17800

CG2086

CG1765

CG13387

CG10079

CG8581

CG17697

CG16785

CG18734

CG32538

CG7223

CG1864

CG8676

CG33183

CG16902

CG5517

CG5183

CG12283

CG33087

CG5481

CG32697

CG31092

CG4521

CG6530

CG4322

CG1960

CG6827

CG4128

CG5610

CG2902

CG3936

CG8967

CG11081

CG1817

CG6899

CG11516

CG2841

CG10537

CG5279

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CG4700

CG5661

CG3135

CG3722

CG5232

CG8268

CG5474


CG7887

CG14026

CG3171

CG8166

CG8444



CG1070

CG1028

CG3166

CG7508

CG5488

CG3644

CG3578

CG3350

CG9653

CG11491

CG42281

CG17894

CG13624

CG30001

CG32601

CG32778

CG4882

CG4914

CG11798



CG12690

CG8367

CG8591

CG6854

CG6103

CG7450

CG5799

CG5893

CG8667

CG5838

CG12767

CG6794

CG11094

CG8361

CG8328

CG6376

CG1071

CG9015

CG3114

CG6338

CG3143

CG12632

CG14307

CG31753

CG9310

CG11352

CG9397

CG4029

CG33956

CG4717

CG8817

CG42679

CG33473

CG8474

CG7074

CG14283

CG13316

CG2244

CG1429

CG11172

CG8524

CG33336

CG12212

CG17077

CG12287

CG8643

CG11992

CG8544

CG7734

CG5580

CG32120

CG4257

CG7951

CG8815

CG3871

CG5557

CG7417

CG10388

CG14029

CG3998

29

36

80

79

83

307

## **Results qPCR : Dscam mRNA levels upon Faf electroporation**

Taqman:

mRNA levels:	UAS	UAS-Faf	UAS-Fafser	UAS-Fam
EP 23/9	1	0.818	0.56	1.108
EP 17/10	1	1.134	1.21	1.155
EP 25/11	1	0.67	0.862	0.906
mRNA levels: <b>average:</b>	<b>1.000</b>	<b>0.874</b>	<b>0.877</b>	<b>1.056</b>
St.Dev.:				
EP 23/9	0.432	0.331	0.055	0.342
EP 17/10	0.045	0.087	0.117	0.09
EP 25/11	0.032	0.039	0.058	0.062
St.Dev.: <b>average SD:</b>	<b>0.170</b>	<b>0.152</b>	<b>0.077</b>	<b>0.165</b>

<b>UAS-Dscam</b>
53.014
5.034
n.d.
<b>UAS-Dscam</b>
<b>29.024</b>
4.232
0.562
n.d.
<b>UAS-Dscam</b>
<b>2.397</b>

