Axonal injury and regeneration in the adult brain of *Drosophila*


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**Abstract**

*Drosophila melanogaster* is a leading genetic model system in nervous system development and disease research. Using the power of fly genetics in traumatic axonal injury research will significantly speed up the characterization of molecular processes that control axonal regeneration in the Central Nervous System (CNS). We developed a versatile and physiologically robust preparation for the long-term culture of the whole *Drosophila* brain. We use this method to develop a novel *Drosophila* model for CNS axonal injury and regeneration. We first show that, similar to mammalian CNS axons, injured adult wild type fly CNS axons fail to regenerate, whereas adult-specific enhancement of Protein Kinase A activity increases the regenerative capacity of lesioned neurons. Combined, these observations suggest conservation of neuronal regeneration mechanisms following injury. We next exploit this model to explore pathways that induce robust regeneration and find that adult-specific activation of JNK signalling is sufficient for *de novo* CNS axonal regeneration after injury, including the growth of new axons past the lesion site and into the normal target area.

**Keywords**

Axon; *Drosophila*; Explant; Injury; Regeneration; Signal transduction

**Introduction**

*Drosophila melanogaster* is a leading genetic model organism in deciphering neuroscience problems like memory formation or odor processing (Hallem and Carlson, 2004; Margulies et al., 2005), as well as modelling neurodegenerative diseases (Shulman et al., 2003; Bilen and Bonini, 2005). Recently, we and others developed methods to study traumatic brain injury in the fly CNS. Similar to their mammalian homologues, fly CNS neurons respond to injury by transient upregulation of the stress response JNK pathway (Leyssen et al., 2005) and by
Wallarian degeneration of their distal axonal stumps (Berdnik et al., 2006; Hoopfer et al., 2006; MacDonald et al., 2006).

The principle problem in CNS injury is the lack of efficient regeneration, which underlies long-term functional impairments. Molecular pathways that modulate axonal regeneration are therefore promising targets in the treatment of injuries. Mammalian studies suggest that stimulating intracellular cAMP signalling helps overcome growth inhibition of injured CNS axons (Cai et al., 2001; Neumann et al., 2002; Qiu et al., 2002).

Thus far, molecular insights into axonal regeneration have been gained from vertebrate models. However, an invertebrate model would lend itself to a faster and larger scale genetic analysis. A model for PNS axon regeneration in *C. elegans* has been described (Yanik et al., 2004) but no paradigms are available yet to study regenerating CNS axons in worms or flies.

A significant impediment to developing a CNS axonal injury model in *Drosophila* is the inaccessibility of the adult fly CNS to reproducible, live, physical manipulation. To solve this problem, we optimized a method to explant, immobilize and culture the *Drosophila* brain (Gibbs and Truman, 1998; Wang et al., 2003; Gu and O'Dowd, 2006), increasing its accessibility to various manipulations. We show that a subpopulation of neurons- the small Lateral Neurons ventral (sLNv)- retain their morphological and functional properties for up to a week in culture. Using a microdissection technique, sLNv axons are reliably severed and their response to injury studied over several days. We find that the distal, severed, axon fragments degenerate in this preparation, as has been shown in *vivo* (Berdnik et al., 2006; Hoopfer et al., 2006; MacDonald et al., 2006). More importantly, as in mammals, very limited and weak regeneration of CNS axons occurs in the wild type fly brain. Instead, injured adult CNS axons appear to form retraction bulbs with small local filopodial sprouts and in only a minority of cases is *de novo* growth observed. Also, as in mammalian systems, the regrowth capacity of injured LNv axons is significantly enhanced by genetic, adult-specific, increase in Protein Kinase A (PKA) activity in the LNv.

Our previous observations that injured fly brains upregulate JNK signalling (Leyssen et al., 2005) and the requirement of c-Jun for PNS regeneration (Raivich et al., 2004) lead us to hypothesize that JNK may play an important role in CNS axonal regeneration as well. We exploited the brain culture system to show that activation of JNK signalling strongly enhances CNS axonal regeneration after injury, including the growth of novel axons re-entering the correct target area.

**Material and Methods**

**Drosophila lines and fly husbandry**

The *PDF-Gal4* line was obtained from P. Taghert, *UAS>CD2,y+ >CD8-GFP* from G. Struhl, *UAS-CD8-GFP, UAS-Bsk* and *UAS-hep* from the Bloomington *Drosophila* stock centre at the University of Iowa, *UAS-PKAc* and *UAS-PKAcA* from D. Kalderon, *tubGAL80ts* from R.L. Davis. The *UAS-PKAc* alleles were checked and verified by sequencing with the forward primer 5’AGGAGTTCGAGGACAAGTGG3’ and the reverse primer 3’CTAATGTTCCTGCTACTGCTA 5’. All flies were raised on standard fly food at room temperature, except where indicated. Flies of the genotype: *PDF-Gal4, UAS-CD8-GFP; UAS-CD8-GFP* were created by recombination. To visualize single sLNv neurons, adult flies of the genotype *hs-FLP; PDF-Gal4, UASLacZ/UAS>CD2,y+ >CD8-GFP* were heat shocked at 37°C for 45 min. They were left to recover at 28°C for at least two days to maximize GFP expression, dissected and examined.
Culturing whole brain explants on culture plate inserts (See supplementary materials for more details)

Millicell low height culture plate inserts (Cat No: PICMORG50, Milipore) were first coated with Laminin (3.3 μg/mL; BD Biosciences Cat No: 354232) and Polylysine (33.3 μg/mL; BD Biosciences Cat No: 354210). Adult female flies were collected within 4 days after eclosion and placed on ice. Before dissection a fly was washed in 70% Ethanol for a few seconds and placed in a sterile Petri dish containing ice cold Schneider’s Drosophila Medium (GIBCO). The brains were quickly and carefully dissected out in this medium (< 3 min/brain), washed in ice-cold dissection medium. For larval brains the same procedure was followed. The eye/antennal disc complex was left attached to the brains. After dissection and washing, up to twelve brains were placed in a drop of medium on the membrane of the culture plate insert. 1.1 ml of culture medium was then added to the Petri dishes that were kept in a plastic box in a humidified incubator at 25°C. The culture medium was refreshed every two days. The culture medium used is modified from Gibbs and Truman (1998) (Gibbs and Truman, 1998) (10 000 U/ml penicillin, 10 mg/ml streptomycin, 10% Foetal Bovine Serum and 10 μg/ml insulin in Schneider’s Drosophila Medium (GIBCO)). For the culture of larval brains 1 μg/ml of Ecdysone was added.

Immunohistochemistry

Freshly dissected brains of adult flies were processed for immunohistochemistry as described (Hassan et al., 2000). Primary antibodies were mouse anti-β-galactosidase (Promega; 1/1000), Rabbit anti-GFP (Molecular Probes; 1/1000); Rabbit anti-PDF (P. Taghert; 1/40000) rat anti-TIM (A. Sehgal 1:1000). Cultured brains were first fixed by replacing the culture medium in the Petri dish with fixation solution (3.7% formaldehyde in PBS) for 30 minutes. Then 1 ml of fixation solution was carefully added on top of the filter for 2 hours. Brains that detached from the membrane were excluded from further analysis. Immunolabeling was performed as for freshly dissected samples.

Imaging and quantifications

Living brains on the culture plate inserts were visualized with a Leica MZ FLIII stereomicroscope. Images were acquired on a Leica DC300F digital camera using Leica IM50 software. Overview images of brains were made at a total magnification of 100X and more detailed images at 200X. For confocal microscope image acquisition, a Leica TCS SL-system was used. Confocal stacks at 0.5μm distances were obtained and projection images were generated with Leica confocal software. Snapshots of the LNv neurons were taken approximately 6 hours after injury from living brains in culture. 550μl of culture medium was put on top of the filter covering the brains in which a 63X water-dip lens was immersed to take the snapshots. Quantification of the newly grown sprouts was performed with ImageJ and the Zeiss KS300 image analysis software. Student’s t-tests were used to calculate the significance of axonal length and distances covered by novel axons. Chi-square tests were used to test significance for the percentage of brains showing regrowth and the percent of axonal regrowth. P-values >0.05 were considered nonsignificant, *p<0.05; ***p<0.001. The contralateral, uninjured sLNv tract was used as a measure of the health of the preparation. Brains in which the uninjured sLNv tract showed signs of degeneration (see Figure S2) were excluded from analysis (~10%). Representative images were used in the Figure panels, which were prepared in Photoshop 7.0.

Electrophysiological recordings of LNv in cultured brains

For each electrophysiological recording experiment, all explants were maintained in 12hr:12hr light-dark cycle. Electrophysiological recordings were made using modification of previously described methods (Sheeba et al., 2008a; Sheeba et al., 2008b) as originally developed by Gu...
and O’Dowd (Gu and O’Dowd, 2006). The composition of the external (bath) solution was: 101 mM NaCl, 1 mM CaCl₂, 4 mM MgCl₂, 3 mM KCl, 5 mM Glucose, 1.25 mM NaH₂PO₄ and 20.7 mM NaHCO₃, 250 mOsm, pH 7.2. The composition of the standard internal solution was: 102 mM K-Gluconate, 0.085 mM CaCl₂, 1.7 mM MgCl₂, 17 mM NaCl, 0.94 mM EGTA, 8.5 mM HEPES, 235 mOsm, pH 7.2. Dissected Drosophila adult fly brains (yw; pdf-Gal4; UAS-dORK-NCl-GFP / +) were cultured for 3 days and immobilized to prevent detachment by the perfusion flow. The insert was visualized using a BX51WI microscope (Olympus, NJ, USA) and all patch-clamp recordings were made using this microscope under a continuous perfusion of the external solution which was bubbled by a gas mixture of 95% O₂ and 5% CO₂. 10 MΩ resistance microelectrodes were fabricated from glass borosilicate capillaries using a PP-83 two-step gravity puller (Narishige, NY, USA). Pipets were directed to neurons close to the explant brain surface using a MP-285 micromanipulator (Sutter Instruments, CA, USA). Cell attached seals of 2–3 GΩ were formed by gentle negative pressure. Whole cell configurations were established by subsequent short negative pressure pulses. Current clamp recordings were made using an Axopatch 200B amplifier (Molecular Devices, CA, USA) connected to a Digidata 1322A acquisition system (Molecular Devices, CA, USA). Experiments were controlled by the pClamp 8.2 Clampex software (Molecular Devices, CA, USA). Whole cell capacitance and series resistance compensations were applied in whole cell configuration. Recordings were lowpass Bessel filtered at 5 kHz. Data analysis was performed using the pClamp 8.2 Clampfit software (Axon Instruments, CA, USA). Traces were lowpass filtered by the 3-point Boxcar method, electrical interference-filtered at 60 Hz, and leak current-subtracted, and corrected for liquid junction potential as described in (Sheeba et al., 2008a).

**Injury of sLNv axonal tracts in cultured brains**

Under visual control at 100X using a Leica MZ FLIII stereomicroscope, the tip of an Eppendorf Microdissector (Cat No: 920 00 401−6, Eppendorf, Germany) was placed on the axonal tract (Figure 3). The Piezo Power MicroDissection device was activated (frequency between 30–60 Hz, and amplitude 75−100%=1.5μm) which cut the axons. The injury was verified at 200X magnification. Axons were only injured on one side of the brain. For cohorts of injury that were compared at different time points, the brains of female sibling flies were dissected, cultured and injured on the same day and fixed and stained at the appropriate time. Experiments were performed at least in duplicate. For the PKAc and JNK injury experiments, flies were raised at 18°C and < 1 week old flies shifted to 28°C two days prior to culturing. Adult brains were dissected, cultured, injured, imaged with a water-dip lens and kept at 25°C for three days and then processed and quantified as double blinds.

**TARGET system**

The efficiency of TubPGAL80⁴⁸ in our experimental setup was tested by raising flies of the genotype PDF-Gal4, UAS-CD8-GFP; Tubulin-Gal80⁴⁸/+ at 18°C and counting the number of sLNv cells that expressed GFP. Adult flies were then shifted to 28°C and the number of sLNv cells expressing GFP counted.

**Results**

**The LNv as a model for CNS axonal injury in Drosophila**

As a first step in the development of a model for CNS axonal tract injury in *Drosophila*, we identified a subpopulation of neurons in the adult brain that 1) forms stereotypic axonal tracts, 2) is easy to reach i.e. superficial in the brain and 3) can be genetically manipulated. The LNv fit all these requirements. These neurons are a component of the neuronal circuit that controls circadian rhythmicity of fly behaviors (Renn et al., 1999; Kaneko and Hall, 2000). There are two subpopulations of LNv (Fig. 1A-E); the small LNv (sLNv) composed of 4–5 neurons with small cell bodies (Fig. 1A, D; arrow) and dorsally projecting axons (Fig. 1A, D; arrowhead).
and the large LNv (ILNv) with large cell bodies (Fig. 1B, E; arrow) and extensive ipsilateral optic lobe neurites (Fig. 1B, E; double pointed arrow) as well as a commissural axon (Fig. 1B; arrowheads) projecting towards the contralateral optic lobe. Importantly, the dorsally projecting axons of the sLNv lie superficially on the posterior side of the brain (Fig. 1F; arrowheads). This makes them easily accessible for visualization and physical manipulation in a brain that is dissected out of the head cuticle (Fig. 1H vs 1G) and oriented with the posterior side up (Fig. 1I, I’). Their cell bodies and the optic lobe innervation are found on the anterior side of the brain (Fig. 1F; arrow, double pointed arrows; Fig. 1H’ arrows, double pointed arrows). Transgene expression specifically in the LNv can be achieved using the Gal4/UAS-system activated by the PDF-Gal4 driver line (Brand and Perrimon, 1993; Kaneko and Hall, 2000) and the LNv axons traced by expressing a membrane-bound fluorescent molecule CD8-GFP in the LNv. This allows easy visualization of the LNv axonal tracts in a living brain using a GFP-filter on a stereomicroscope (Fig. 1H, I).

A long-term, whole-brain explant culture

The LNv axons are, like the rest of the brain, separated from the environment by a hard cuticle (Fig. 1G). In order to overcome this barrier for visualisation and micromanipulation of the axonal tracts, we attempted to remove the cuticle while keeping the brain alive and healthy. Explant cultures of Drosophila brains have already been successfully used to study axon remodelling during metamorphosis (Gibbs and Truman, 1998; Brown et al., 2006) and studying neuronal activity (Wang et al., 2003; Gu and O’Dowd, 2006). It is therefore in principle possible to keep Drosophila brains alive ex vivo. However, induction of reliable axonal injury and subsequent visualization require the explant preparation to be immobilized to make the precise micromanipulation of the brains possible.

The problem of immobilization as well as long term survival of mammalian brain slice preparations has been solved by using culture plate inserts (Stoppini et al., 1991; Belenky et al., 1996; Collin et al., 1997; Weissman et al., 2004). These inserts consist of a porous membrane attached to a plastic ring and are placed in a Petri dish filled with medium (Fig. 1J). When positioned on the culture plate insert, the explant attaches to the membrane and is covered by a small film of medium (Fig. 1J). At this liquid-air interface exchange of nutrients and air takes place, ensuring optimal conditions for the health of the explanted tissue. To obtain healthy, immobilized and easily accessible Drosophila brain preparations, we modified the culture media and conditions described for Drosophila tissues (Gibbs and Truman, 1998) in combination with the physical support of the culture plate insert described for mammalian brain slices.

The Drosophila brain is less than 300μm thick (the thickness of mammalian brain slices). We therefore reasoned that Drosophila whole brain explants could be used. Whole brains were dissected, thus leaving the connectivity pattern within the brain preserved. Fig. 1J shows a schematic of the setup we used for these experiments (for details see Experimental Procedures). Brains from transgenic adult flies, expressing GFP under control of the PDF-Gal4 are carefully and rapidly dissected out of the head cuticle in ice-cold dissection medium. The dissected brains are transferred to the coated culture plate insert (Fig. 1K) and culture medium is added to the Petri dish. When examined with a fluorescence stereomicroscope, the axonal pattern of the posteriorly located sLNv dorsal projections and ILNv commissural axons is easily visualized (Fig. 1L). Explanting and culturing the brains using this new protocol thus simultaneously solves the accessibility, immobilization and visualization problems inherent in whole animal preparations.

Using the medium described in Gibbs and Truman (Gibbs and Truman, 1998) (See materials and methods), a relatively high percentage (~70%, n=18) of the sLNv axon bundles innervating the dorsal brain remained intact (Fig. S1A). To further improve the health of the sLNv axons
we made a few changes to the composition of the medium (Fig. S1A). We found that a medium lacking antibiotics and the fungicide Amphotericin B increases the percentage of intact sLNv axons at day 4 to 94% (n=16). This high percentage of morphologically intact axons was maintained up to 6 days in culture (Fig. S2, supplementary text 1). We therefore used these conditions that enhanced the preservation of sLNv morphology for all further experiments.

To examine the evolution of the gross morphology of cultured brains, we imaged brain explants every day over a period of 20 days (Fig. 1M; n=6). General brain morphology remains remarkably constant over this period of time. The only obvious change is an overall flattening of the preparation after a few days, a phenomenon well known to occur on culture plate inserts (Stoppini et al., 1991). Slight asymmetries in the contact face between the brain and the filter are also accentuated by this process (Fig. 1M; note the difference in optic lobe (ol) size). Importantly, strong GFP expression in LNv cell bodies and axons is maintained (Fig. 1M, last panel), suggesting that the LNv neurons continue to express this protein for >20 days. On a global morphological level the brains therefore support the culture conditions very well for at least three weeks.

**Functional characteristics of the LNv are maintained in cultured brains**

To test if the LNv neurons also retain their functional properties in an explanted brain, we examined three characteristic features of the LN. First, if the LNv retain their fate in culture they should continue to express the LNv specific PDF hormone (Renn et al., 1999). Indeed, a strong PDF immuno-signal is observed in seven day-old cultured brains in cell bodies and axonal bundles of the sLNv (Fig. 2A), suggesting that the hormone is still expressed and transported into axons. Second, if the circadian rhythm controlling-LNv neurons function like in a living fly, they should retain their cyclic molecular clock properties in explants. This clock can be traced by immunostaining against PERIOD (PER) and TIMELESS (TIM) proteins (Hardin, 2005). In flies that have been entrained in 12h:12h Light/Dark (L/D) cycles, high levels of these proteins are normally detected 22 hours after lights ON (ZT22) which corresponds to the end of the night while low levels are detected 10 hours after lights ON (ZT10), close to the end of daytime. To test whether this molecular oscillation is intact in explanted brains, we immunostained three day old explants from entrained flies at two time points corresponding to the entrainment schedule, 10 hours and 22 hours after lights ON (ZT10 and ZT22). We find that the LNv express significantly higher levels of PER and TIM proteins at ZT22 as is the case in vivo (Figure 2B, C). In addition, the relative change in the levels of TIM in explanted brains at the different time points is similar to those reported in vivo (Sehgal et al., 1994; Kaneko and Hall, 2000; Nitabach et al., 2002; Shafer et al., 2002). This means that the LNv not only retain their identity, but also preserve their unique molecular characteristics in explants. In addition, FasII staining after three days in culture shows that other neuronal populations remain healthy in the explant (data not shown). Third, we tested whether neurons in the explant brains retain electrophysiological properties such as the firing of spontaneous action potentials. We attempted to record from large LNv as we have done previously in acute dissected brains (Sheeba et al., 2008a; Sheeba et al., 2008b). However for unknown technical reasons, we were not able to form high-quality giga-ohm pipette seals on LNv. While we cannot rule out the possibility that compromised health of LNv neurons in long term cultures may have contributed to the difficulty of making recordings, we note that these neurons remain morphologically normal and all LNv neurons tested appear to retain clock cycling. Thus, methodological reasons appear to be the more likely cause of the difficulties of making LNv recordings in long term cultures.

To investigate the general health of the cultured brain, unidentified neurons close to the explant brain surface were probed using whole-cell patch clamp recordings (Gu and O'Dowd, 2006; Sheeba et al., 2008a; Sheeba et al., 2008b). The cells were accessible to the patch pipette in

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brains oriented with the anterior side up. As compared to the relative ease of making recordings from acutely dissected whole brains, it was difficult to establish a cell attached configuration and to break into the membrane after establishing the cell attached configuration. In spite of this technical difficulty, whole cell patch clamp recordings in current clamp mode were made successfully from two different unidentified neurons in 3 day old cultured brains. Both cells show healthy spontaneous action potential firing. One of the two recorded neurons exhibited a resting membrane potential of $-53\text{mV}$ and spontaneous firing frequency of $0.5/\text{sec}$ (Figure 2D), the other neuron exhibited a resting membrane potential of $-45\text{mV}$ and spontaneous firing frequency of $0.9/\text{sec}$. These electrophysiological values are similar to those obtained from identified LNv neurons in freshly dissected whole brain preparations (Sheeba et al., 2008a). A third unidentified neuron recorded from a 3 day old explant brain was initially silent, but could be induced to fire action potentials by depolarizing current injection (data not shown).

### Lack of efficient spontaneous regeneration in the Drosophila CNS

Having made the superficial sLNv axonal tracts accessible to micromanipulation, we designed a simple, inexpensive and accurate setup to sever them using a Piezo Power MicroDissection device (Fig. 3A; see supplementary text 1 for details). Briefly, the sharp tip of the microdissector is positioned on the sLNv axonal tract using a micromanipulator. The device is activated, causing an oscillation of the tip and thereby severing of the axons (Fig. 3B-D; arrowhead). The GFP labelled axons are directly visualized throughout the procedure (Fig. 3C, D) and successful injury readily observed. To further confirm the success of the injury procedure, we examined brains shortly after making the lesion using confocal microscopy. We find that in $100\%$ of the cases ($n=18$) the dorsally projecting axonal bundle was fully severed.

We also noted that immediately after the injury, the severed axons retracted away from the lesion site, a process that has been observed previously in the mouse spinal cord (Kerschensteiner et al., 2005). On average, the stable gap size from the retracted site of injury till the beginning of the distal stump was $36.14\mu\text{m}$ with $67\%$ of the samples showing a gap size of $\sim 20\mu\text{m}$ or more ($\text{S.D.} \pm 22.35$; $n=18$).

A key difference between injury to PNS versus CNS axons in mammals is that PNS axons regenerate spontaneously, while CNS axons largely fail to regenerate. To test if, similar to mammals, the fly CNS fails to spontaneously regenerate after injury, we used the explant system to lesion sLNv axonal tracts in cultured brains and examined their morphology at different times after injury. When injured brains are examined within 15 minutes after the lesion, the proximal axonal stump is uniform in width (Fig. 4A-C) and the distal, unconnected axonal ends appear smooth and continuous (Fig. 4D). One day later (Fig. 4E-H) the proximal axons develop bulbous enlargements at their tips (Fig. 4G, arrows) and on these bulbs what appears like filopodial sprouts are formed in $59\%$ of brains (Fig. 4F, arrowhead; $n=17$). The distal axonal ends appear more vesicular and fragmented (Fig. 4H). Over the next 4 days an increase in proximal sprouting is observed. In most cases, the extensive axonal sprouts either remain stalled (Fig. 4J, N, R; $57\%$ [8/14] on day 2; $47\%$ [8/17] on day 3; $34\%$ [5/15] on day 4), or grow laterally for short distances (Fig. 4M, S; full arrowheads; $36\%$ [5/14] on day 2; $35\%$ [6/17] on day 3; $40\%$ [6/15] on day 4). In no case did these sprouts reach or traverse the lesion site. LNv cell body counts show that all these neurons survive (data not shown). These experiments thus provide good evidence that the proximal severed axonal stump is extensively remodelled over a period of several days, but the large majority of axons completely fail to regrow after injury. The distal axonal ends undergo progressive fragmentation and degeneration (Fig. 4D, H, L, P, T). Together, these findings suggest that, as in mammals, injured wild type adult *Drosophila* CNS axons do not regenerate.
Intact axonal outgrowth in larval brain explants

One caveat is that the failure to efficiently regenerate axons may be due to culture conditions, rather than the intrinsic characteristics of the adult fly brain and its neurons. We therefore tested if developing axons can grow in an explanted brain under our culture conditions, taking advantage of the extensive axonal reorganization that takes place during metamorphosis from the larval to an adult CNS. Figure 5A shows a *Drosophila* CNS very shortly before pupariation. The sLNv neurons are present and have already formed their dorsal projections, but the lLNv are not visible at this stage. Larval brains were imaged each day over a period of seven days. We observe that new LNv cell bodies start to express GFP (Fig. 5B; arrows) suggesting that LNv are born and acquire their correct PDF-positive identity in the explant. Next, confocal imaging studies of brains at 24h versus 5 days in culture were performed. At 24h in culture only the four sLNv cell bodies (Fig. 5C, arrows) and their dorsal projections (Fig. 5C, arrowhead) are apparent. However, after five days in culture we find axons projecting towards the brain centre and the optic lobes in 100% of the cases (n=13; Fig. 5D, white arrows and arrowheads). Finally, to visualize the morphology of the growth cones on newly formed axons in brain explants, we imaged the tips of the axonal bundles that project towards the contralateral side of the brain at high resolution. We find that these axons form extensive growth cones with a large number of small sprouts (Fig. 5E, arrowheads). When growing larval axons were injured, outgrowth was still observed after five days in culture (data not shown). However, since uninjured axons also grow during this period, it is very difficult to ascertain whether the growth observed is regenerative or developmental. Nonetheless, these data clearly show that the culture conditions used are not per se growth inhibitory.

These results indicate that larval LNv axons can grow in explanted brains and that this growth correlates with the formation of elaborate growth cones. This morphology contrasts with the dystrophic growth cones of injured adult brain axons (Fig. 4) indicating that, rather than being inhibited by the culture conditions per se, injured *Drosophila* adult brain neurons, like their mammalian counterparts, largely fail to regenerate.

The role of PKAc in stimulating growth after injury is conserved

Mammalian CNS axon injury studies show that elevating cAMP levels can enhance regrowth after injury (Neumann et al., 2002; Qiu et al., 2002). These studies also suggest that a major effector of the cAMP signalling pathway for regrowth is PKA since cAMP-induced regrowth can be blocked by PKA inhibitors (Qiu et al., 2002). Together these studies imply a crucial role for PKA in the regenerative capacity of injured CNS neurons. If axonal injury in the explanted *Drosophila* brain is to serve as a genetic paradigm for the study of traumatic axonal injuries in the CNS, modulation of the regenerative response like switching the damaged neurons to a stronger growth state should be possible. To test if PKAc activity could modulate the regenerative capacity of cut CNS axons in our injury paradigm, we used the binary Gal4-UAS system to overexpress different UAS-PKAc alleles specifically in the LNv neurons. Transgenes encoding PKAc<sup>5.9</sup>, a wild type, fully active, Flag tagged, PKAc allele (henceforth PKAc<sub>wt</sub>; (Kiger et al., 1999) and PKAc<sup>A75</sup>, a mutant allele with at least 1000 fold less catalytic activity (Kiger and O'Shea, 2001) than PKAc<sub>wt</sub> were used to genetically manipulate PKAc activity. Both transgenes were expressed in the LNv’s together with CD8-GFP using the pdf-Gal4 driver. To overcome the developmental effects of PKAc overexpression and allow delivery of PKAc activity specifically in adult neurons prior to inducing the injury onwards, we turned to the TARGET system (McGuire et al., 2003). The TARGET system uses a ubiquitously expressed, temperature-sensitive Gal80 (tubP-GAL80<sup>ts</sup>) to repress Gal4 activity at the permissive temperature. Using this approach we largely suppressed developmental expression and thus developmental effects of increased PKAc activity (data not shown).
To assess the capacity of PKA to induce axonal regeneration, we imaged each injured brain approximately 6 hours (Fig. 6A) and 3 days (Fig. 6B) after injury. By comparing the images from the same axon stump six hours and three days after the injury, the novel axons were identified (Fig. 6C, yellow circles; Fig. 6D pink lines) and their response quantified based on the following criteria (for further details on the definition of these criteria see Supplementary text 2). First, we calculated the penetrance of the regenerative response at the level of individual brains, by determining the percentage of brains showing regrowth after injury. Specifically, we counted the number of brains sprouting at least one new axon after injury. Second, we calculated the penetrance of the regenerative response at the level of individual axons. Specifically, we counted the number of newly generated axons and divided this by the total number of injured axons, thus determining the propensity of single axons to regenerate. Third, we calculated the extent of axonal regrowth, by measuring the length of each newly grown axon extension by freehand tracing (Fig. 6D, pink lines). In addition, we computed the direct distance these new axons cover as a straight line from the site of injury to the tip of the new axonal sprout (Fig. 6E, blue lines) to test if regenerated axons can bridge the lesion gap and grow towards the original target area. In wild type brains this gap was measured to be 36 μm.

Using these three criteria, we are able to objectively quantify the regenerative response at the level of individual brains as well as at the level of single axons.

At day 3 after injury the percentage of brains showing regrowth after injury in control samples is 35% (n=20) (Fig. 6J). Expression of the catalytically mutant version PKA<sup>A75</sup> significantly decreases the percentage of brains sprouting new axons from the site of injury to (27.2%, n=11, p=0.0402). Conversely, significantly more brains expressing PKA<sup>wt</sup> sprouted new axons from the injury site (61.9%, n=21, p=0.0078; Fig. 6J and Fig. 7A-C). The regenerated axons transport PDF showing they are functionally connected to the original bundle (Fig. S3 A-C). These results suggest that PKA activity can increase the percentage of brains showing regrowth after injury.

Next we asked if, in addition to increasing the percentage of brains showing regrowth after injury, PKA has also an effect on the percentage of axons showing regrowth as well as on the length and the distance the new axons cover. In control brains, the percentage of axonal regrowth is 13.7% [11/80], the length of a newly extended axon is 24.8μm and the average distance in a straight line a new axon can bridge is 18.3μm (Fig. 6 K and L). Expression of PKA<sup>wt</sup> significantly increases the percentage of axonal regrowth (33.3% [28/84], p = 0.001), but not the average length of the new axons (21.7μm; n= 21, p=0.62), or the distance they are able to cover (18μm; n=21, p=0.96) (Fig. 6 K and L).

In conclusion, PKAc<sup>wt</sup> enhances both the percentage of brains showing regrowth as well as the percentage of regenerating axons, but it did not have an effect on the extent of regrowth in terms of the length and distance the new sprouts cover. Because the PKAc<sup>wt</sup> allele is subject to negative regulation by the endogenous regulatory subunit of PKA, a constitutive-active PKAc allele, PKA<sup>mC<sup>*</sup></sup> (Li et al., 1995), was also tested in the injury model. As expected, PKA<sup>mC<sup>*</sup></sup> also significantly increased both the percentage of brains and axons showing regrowth after injury (64.2%, n=14, p=0.019 and 17.8% [10/56], p=0.0053 respectively; Fig. 6J and 6K). Furthermore, LNv expressing activated PKAc did grow significantly longer axons than controls (45.8μm; p=0.023; Fig. 6L). Interestingly, the average distance these new axons had moved from the point of injury was not significantly different from wild type (30μm; p=0.17; Fig. 7D, Fig. 6L).

Taken together, these data suggest that at a certain threshold PKAc activity can enhance the potential of damaged axons to regrow. Constitutive activation of PKAc can increase the length of newly generated axons, but not the distance they cover. These results show that our injury model is amenable to modifying the regenerative response of damaged neurons. In addition,
these results are in agreement with the observations in mammalian CNS regeneration studies and strongly suggest conservation of injury, regrowth and regeneration processes. Importantly, despite the robust growth induced by constitutively active PKA\textsuperscript{mC*}, the average distance covered by these axons (30\textmu m) is still short of bridging the gap (36\textmu m) between the lesion site and the normal target area.

Activated JNK signalling induces regeneration back into the original target area

Our analyses so far suggest that the brain explant system is a robust, quantifiable and genetically amenable system for the study of conserved axonal injury and regeneration processes. The data also clearly highlight the need for the discovery of pathways that might induce better regeneration. We hypothesized that a good candidate may be the JNK pathway. First, JNK is a general stress response pathway (Leppa and Bohmann, 1999). Second, we have shown that JNK signalling is necessary and sufficient for axonal development in the fly CNS (Srahna et al., 2006) and that it is transiently induced by brain injury in Drosophila (Leyssen et al., 2005). Third, c-Jun is necessary for spontaneous regeneration in the mouse PNS (Raivich et al., 2004). Despite these hints in flies and mammals, a role for JNK activity in CNS regeneration has not been investigated.

To test a role for JNK in CNS axonal regeneration, we inactivated JNK signalling solely in the adult LNv neurons by overexpressing a dominant negative allele of the Drosophila JNK Basket (Bsk\textsuperscript{DN}) (Boutros et al., 1998; Reed et al., 2001; Srahna et al., 2006), and examined axonal regeneration using the three main criteria described above. Adult-specific inhibition of JNK signalling in injured LNv does not significantly alter the percentage of brains or single axons showing regrowth (43.7%; n=16, \(p=0.26\) and 10.9% [7/64], \(p=0.059\) respectively; Fig. 6J and K, Fig. 8A). In contrast JNK inhibition shows a strong tendency to reduce the length of the newly extended axons from 24.8\textmu m in control neurons to 11.2\textmu m (\(p=0.053\)). As a result the average gap bridged by these axons is reduced from 18.3\textmu m to 10.2\textmu m (\(p=0.18\); Fig. 6L). These data suggest that JNK signalling is not necessary for injured neurons to initiate axonal regrowth, but may rather regulate the extent of this regrowth.

Next, we tested if activation of JNK signalling might positively influence axonal regeneration after injury. To this end we expressed a constitutively active form of the Drosophila JNKK Hemipterous (Hep\textsuperscript{CA}) (Adachi-Yamada et al., 1999) specifically in injured adult LNv. As in previous experiments, we imaged the severed LNv axons at approximately six hours and three days after injury. We observe that JNK activation significantly enhances all aspects of axonal regrowth after injury. First, the percentage of brains showing regrowth expressing Hep\textsuperscript{CA} was 86.6\% (n=15, \(p=0.001\)) in comparison to 35\% of control brains (Fig. 6J). Second, the percentage of axonal regrowth increased 3.5 fold to 48.3\% in comparison to the control brains (13.7\% \(p=0.001\); Fig. 6L). Third, the average length of the new axons is increased approximately 3 fold to 76.7\textmu m (\(p=0.001\); Fig. 6L). In addition, these new axons cover an average distance of 49\textmu m (\(p=0.01\)), almost 2.5 fold the length observed in control brains, and well beyond the \(\sim 36\textmu m\) necessary to bridge the injury gap (Fig. 6L, Fig. 8B). Although the average distance covered by novel axons in control brains is only 18.3\textmu m –almost half of the average gap size induced by the lesion– occasionally (1/20) there is a brain that can grow an axon able to cover a distance longer than 36\mu m. Therefore, the number of axons growing towards and into the target area can be precisely quantified (Fig. 6 G-I, supplemental text 2,

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Fig. S3 D and E). We find that 46.6% [7/15] of HepCA expressing brains grow at least one axon towards the target area (Fig. 8 B-D). Since activation of JNK signalling results in axons that are of sufficient length to potentially bridge the injury gap, we finally quantified the number of brains in which novel axons re-enter the denervated target area (Fig. 8D bottom panel, shaded pink). We do not find any examples of this in genetic controls. In contrast 1/3 of all HepCA expressing brains show novel axons re-entering the target area ([5/15], p=0.02; Fig. 8C-E).

Discussion

Why a Drosophila model for traumatic axonal injury?

*Drosophila melanogaster* is a leading genetic model organism not only in basic neuroscience but also in biomedical research. Using the power of this genetic model organism, difficult problems like the molecular mechanisms behind human neurodegenerative diseases are being successfully addressed (Shulman et al., 2003; Bier, 2005; Bilen and Bonini, 2005). These advances have depended largely on the development of good *Drosophila* models for these human diseases (Feany and Bender, 2000). Very recently the first models for axonal severing in the CNS of the fly have been developed and have unveiled a marked conservation in the molecular mechanisms underlying neuronal responses to injury in flies and mammals (Leyssen and Hassan, 2007). While these valuable paradigms hold promise to significantly enhance our insights into axonal degeneration in the future, they do not allow the evaluation of axonal regeneration. Increased insight into the process of regeneration is crucial from a therapeutic point of view that aims at stimulating axons to reconnect to their postsynaptic target. In this work we describe a protocol that allows, in addition to a versatile series of manipulations, the precise and reliable severing of sLNv axons in explanted adult *Drosophila* brains. This technique provides, for the first time, the opportunity to study axonal regeneration in the *Drosophila* CNS and use genetic screens to identify factors that promote axonal regeneration.

The responses of injured axons in the Drosophila brain

The regenerative responses of injured axons in *Drosophila* brain explants show remarkable similarities to those seen in mammals. First, the distal, severed ends of the axons became fragmented in a manner that is reminiscent of Wallerian degeneration in mammals, as was described also in living *Drosophila* (Hooper et al., 2006; MacDonald et al., 2006). In both cases unconnected axons were rapidly split into small vesicles, which then disappeared. A major advantage of the explant model is that it also allows the study of the proximal, injured axon. One day after injury, the proximal axonal stump forms a bulbous structure that shows morphological similarities to the dystrophic retraction bulbs seen in injured mammalian CNS axons (Silver and Miller, 2004). This structure is extensively remodelled and develops small, and occasionally longer, thin axonal sprouts. Similar morphological changes have been recently described in severed, GFP-marked axons of the mouse spinal cord (Kerschensteiner et al., 2005).

The lack of efficient regeneration of injured *Drosophila* axons in brain explants is not due to the experimental setup. First, sLNv neurons in the preparation appear healthy both morphologically and functionally. Second, axons in explanted larval brains do form elaborate growth cones and grow large distances over the course of a few days. Third, the proximal axonal stumps are clearly dynamic and capable of sending out axonal sprouts. We therefore suggest that the lack of efficient sLNv axonal regeneration is an intrinsic feature of the adult fly brain. Indeed we show that the adult specific, cell autonomous manipulation of the catalytic activity of PKA can stimulate a generally dormant regrowth capacity. A significantly larger proportion of lesioned brains become capable of extending new axons at the cut tip. This is very similar to the mammalian cAMP based enhancement of regrowth (Cai et al., 2001; Neumann et al., 2002; Qiu et al., 2002). In both cases regrowth is achieved by increasing PKA
catalytic activity, either reducing the negative regulation of PKAc through adding of cAMP in the mammalian studies, or overexpressing a catalytic mutant version of PKA that is free from endogenous inhibition. In summary, the data presented herein support the concept that the fly CNS is closer in its response to injury to the mammalian CNS than the mammalian PNS. The mammalian PNS regenerates spontaneously, while, in contrast, both the fly and mammalian CNS fail to regenerate except when additional signals are activated. In the case of PKA, the same signal enhances regeneration in both models. This, in addition to several other details such as the anatomy of the retracting injured axons and the appearance of a lesion gap support the conservation of key injury response mechanisms between the two CNS systems.

JNK signalling regulates axonal regeneration

In mammalian models for CNS regeneration, several modes of neuronal repair are proposed. These neuronal recoveries can either be accomplished directly by the transected axons itself, or by sprouts from unlesioned axons in the neighbourhood into the denervated target area (Raivich and Makwana, 2007). However, what is defined as true axonal regeneration is the re-entry of the cut axons themselves onto the denervated target area. Axonal regeneration encompasses several types of neuronal responses that can be grossly divided into the following steps: 1) regrowth of the cut axons, 2) guidance of the newly growing axons towards the normal area of innervation and finally 3) re-innervation of the denervated target area.

Despite the promising results of PKA manipulation, a constitutively active form of PKA is not sufficient to produce axons that can extend into the original target area. We therefore tested the hypothesis that JNK signalling might support better regeneration. Our data suggest that JNK is likely to be a key pathway in CNS axonal regeneration after injury. Inhibiting JNK activity results in a high tendency to inhibit the length of the newly grown axons compared with control brains, whereas its constitutive activation results in a massive increase in the penetrance and extent of regeneration, with one third of all brains showing regenerated axons re-entering the target area. It is surprising that activation of only a single pathway is sufficient to enhance the three major anatomical criteria of regeneration. This may be explained by at least two alternative models. It may be that LNv neurons, and presumably other CNS neurons as well, retain their proper guidance and targeting signals well into adult life and as such genetic manipulations stimulating the frequent growth of lengthy axons can result in axons reaching the correct target area. Alternatively, JNK activation may act as a reprogramming signal allowing neurons to reactivate their specific developmental pathways. Although we do not exclude a role for JNK in axonal navigation our previous work (Leyssen et al., 2005; Srahna et al., 2006) shows that JNK can stimulate different cell types to grow longer axons, without influencing their guidance properties. In this context it may seem surprising that the induction of JNK signalling observed upon axonal injury (Leyssen et al., 2005) is not sufficient to induce regeneration. However, it is in fact likely that the transient upregulation of JNK does play a role in stimulating growth after injury, because when injured axons express dominant negative forms of JNK, they show a strongly reduced regrowth tendency compared to wild type brains (Fig. 6), although this reduction remained just below statistical significance. The reason why activated JNK signaling has such a dramatic influence on regeneration is two fold. First, an activated form of the protein is no longer subject to endogenous negative regulation. Second, this form is expressed as a UAS-transgene which means that its expression is permanent and not transient, in contrast to what we observe under wild type injury conditions.

Future applications of the injury paradigm and of Drosophila whole brain culture

Since the system described here is amenable to cell-type specific and adult-specific manipulation of gene activity, future genetic screens will be able to identify new molecules involved in CNS axonal regeneration. A selected set of several hundreds of genes could easily be screened for enhancing the regenerative response of damaged neurons using this novel injury
paradigm. Alternatively, genome-wide primary reverse genetic screens for axon outgrowth can be performed using the LNv to enrich for a set of genes involved in the induction of axonal outgrowth. These can then be taken further to test for their capacity of inducing regeneration after injury. These kinds of approaches will result in a selection of potential regeneration genes and as such result in a very valuable contribution to the studies of axonal regeneration. Such molecules should be prime candidates for further testing in mammalian systems. In addition, the brain culture system makes a living *Drosophila* brain accessible for various micromanipulations. We show that cultured brains retain overall axonal integrity, expression of immunohistochemical markers, molecular characteristics, like circadian oscillation of protein expression, and electrophysiological activity for several days in the explant culture. Many other applications for this explant and culture technology can be imagined: direct visualization of dynamic processes like vesicular transport (e.g. by expression of Synaptobrevin-GFP); activity of neurons in networks (Miesenbock and Kevrekidis, 2005) in a living brain; addition of pharmacological compounds to the culture medium to interfere with pathways or molecules of interest; transplantation of transgenic cell populations from one brain to the other; study of cell migration in living *Drosophila* brain tissue.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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**References**


Figure 1. Description of the LNV neurons and *Drosophila* brain explant culture on organotypic culture plate inserts

A, B, D and E are projection images of 1μm confocal stacks through adult *Drosophila* brains of the indicated genotype. Panels A and B are compiled from separate images of the brain halves of the same brain. (A) A single cell sLNV FLP-out clone shows the cell body (arrow) and dorsally projecting axon (arrowhead) are marked with GFP (green), all LNV are marked by lacZ (red). Scale bar = 80μm. (B) A single cell lLNV FLP-out clone shows the cell body (arrow), commissural axon (arrowhead) and optic lobe innervation (double pointed arrows) are marked with GFP (green), all LNV are marked by lacZ (red). (C) Schematic of an adult *Drosophila* brain indicating morphology and localization of the sLNV (green) and the lLNV.
(D) Higher magnification of the GFP pattern in the single sLNv neuron shown in A. Scale bar = 40μm. (E) Higher magnification of the GFP pattern in the single lLNv neuron shown in B. Arrows indicate the cell bodies, arrowheads the axons. Scale bar = 80μm. (F) Lateral view on a three dimensional reconstruction of an adult *Drosophila* brain. The sLNv dorsal projections and the commissural axons (white arrowheads) are on the posterior side of the brain (P). The LNv cell bodies (arrow) and the optic lobe innervation (double pointed arrow) lie anteriorly (A). Genotype of the flies is: *PDF-Gal4; UAS-CD8-GFP*. (G) Intact adult *Drosophila* head (H, I) Dissected adult *Drosophila* brain seen in brightfield (H, I) or GFP filter (H’, I’) of a stereo microscope (100X magnification). (H, H’) Anterior view. Arrows point to the LNv cell bodies; double pointed arrows indicate optic lobe innervation by the lLNv. (I, I’) Posterior view. Red arrowheads indicate the sLNv dorsal projections, white arrowheads the lLNv commissural axons. Genotype of the flies is *PDF-Gal4, UAS-CD8-GFP; UAS-CD8-GFP*. (J) Schematic of the culture setup. *Drosophila* brains are positioned on the membrane (grey) of a low height organotypic culture plate insert (blue) that is placed in a Petri dish containing culture medium (yellow). The brain is separated from the air by a thin film of medium (inset). (K) Overview image of several adult *Drosophila* brains positioned on the membrane of a culture plate insert (L) Higher magnification image of one brain with bright field (top) or GFP-filter (bottom; pseudocolored), showing the posterior LNv axonal tracts. (M) Gross morphology of a cultured brain over the course of 20 days in culture remains intact. A GFP signal can still be detected after 20 days (ol=optic lobe). All images are taken with a stereomicroscope. The blue panels in C and D are a merge between a brightfield image pseudocolored in blue and the GFP signal in green. The genotype of the flies is *PDF-Gal4, UAS-CD8-GFP; UAS-CD8-GFP*. 
Figure 2. The functional characteristics of LNv are maintained in cultured brains

Brains were maintained in a 12hr:12hr Light/Dark cycle. (A) Immunostaining for PDH (red) shows the presence of PDH in the sLNv dorsal projection after 7 days in culture (B) Immunolabeling of small (arrow) and large (line) LNv in a 3-day old cultured brain. Time points ZT10 and ZT22 correspond to 10 and 22 hours after lights ON. A significantly reduced level of TIM (red) is seen in the brains after 10 hours of light (ZT 10) while cells show higher levels and nuclear localization during the late hours of the night (ZT 22) both in the large and small LNv. (C) Quantification of TIM levels in brain explants (t-test for ILNv ZT10 versus ZT22, t = 1.72, p = 0.024; t-test for sLNv ZT10 versus ZT22: t=1.75, p = 0.001, n=5 brains for each comparison). The genotype of the fly in (A) is PDF-Gal4, UAS-CD8-GFP; UAS-CD8-GFP; in (B) yw; pdf-Gal4; dORK-NC1 / +. (D) Spontaneous action potentials of unidentified surface neuron in a 3 day old cultured brain. Large amplitude spontaneously firing action potentials and after-hyperpolarizations are observed in whole cell patch current clamp recordings day old cultured brain. Large amplitude spontaneously firing action potentials and after-hyperpolarizations are observed in whole cell patch current clamp recordings.
Figure 3. Injury to the sLNv axonal bundle

(A) Schematic showing the setup for the injury of sLNv axonal tracts in explanted brains on a culture plate insert. The axonal GFP pattern is observed using a stereomicroscope equipped with a GFP filter (blue ray). The micromanipulator (grey) tip is positioned on an axonal tract using a micromanipulator. The schematic is not drawn to scale. (B) Schematic of sLNv before (top) and after (bottom) injury. (cb= cell body; D= Distal part of the axon; P= Proximal part of the axon; arrowhead=injury) (C) Fluorescent signal in sLNv of the same brain before (top) and after (bottom) injury. Arrowhead indicates the disruption in GFP signal. (D) Brightfield (left) and GFP (right) images during (top) and after (bottom) the injury procedure. The genotype of the fly is PDF-Gal4, UAS-CD8-GFP; UAS-CD8-GFP. Images are made with a stereo microscope at 200X (C) or 100X (D).
Figure 4. Response of sLNv axons to injury

(A-D) 15 minutes after the lesion all axons are clearly severed (B,C). The distal axon ends appear uniform and continuous (D). (E-H) 1 day after the lesion the proximal stumps form bulbous ends (G, arrows) from which axonal spikes can grow (F, arrowhead). The distal axon ends appear more vesicular (H). (I-L) 2 days after the lesion axonal sprouts are formed (I-K, full arrowheads). The distal axonal ends appear vesicular (L). (M-P) 3 days after the lesion axonal sprouts are present on the proximal stump. In some cases, axons grow away from the target area (M, full arrowheads) and degenerated fragments of axons can be observed (empty arrowheads). In other cases axons remain stalled before the lesion site (N). The distal axonal fragments appear vesicular (P, empty arrowhead). (Q-T) 4 days after the lesion axonal sprouts
of different lengths (full arrowheads) grow laterally (Q, S; full arrowheads), or towards the injury site (Q, full arrowheads). The distal axons are fragmented (T). (*) lesion site. The genotype of the flies is PDF-Gal4, UAS-CD8-GFP; UAS-CD8-GFP. All panels are projections of confocal 0.5μm stacks. Scale bar of overview pictures = 40μm, of proximal stumps = 8μm and of distal stubs = 20μm.
Figure 5. Axonal outgrowth and growth cone formation in explanted larval brains

(A) Morphology of the third instar larval CNS (left panel) (ad: antennal disc; ed: eye disc; ol: future optic lobe; vc: ventral cord; red arrows: nerves). The GFP signal in the brain lobes (right panel) corresponds to the sLNv neurons. (B) Morphological changes in the GFP pattern over time include the appearance of new GFP positive cell bodies (arrows). (C) Detail of the GFP expression pattern after 24h in culture showing the sLNv cell bodies (arrows) and the dorsally projecting axons. Scale bar = 20μm. (D) Detail of the GFP expression pattern after 5 days in culture showing dorsally projecting axons of the sLNv (red arrowhead). New axonal tracts project towards the contralateral brain half (white arrowheads), the future optic lobes (white arrows) and the central brain (red arrows). Scale bar = 40μm. (E) Detail of the growth cones. Note the presence of filopodia at the axonal end (arrowheads). Scale bar = 8μm. A and B are stereo microscope images. The blue panel in A is a merge between a brightfield image pseudocolored in blue and the GFP signal in green. C, D and E are projection images of 0.5μm confocal stacks. The genotype of the flies is PDF-Gal4, UAS-CD8-GFP; UASCD8-GFP.
Figure 6. Quantification of novel axonal outgrowth after injury

(A) Injured axons were imaged approximately 6 hours after injury to verify the injury and capture the morphology of the cut proximal stump. Scale bar = 40 μm. (B) The same brain was fixed and processed for imaging 3 days after injury and the proximal stump imaged again at the same magnification as A. Scale bar = 40 μm. (C) By comparing the images from the same axon stump approximately six hours and three days after the injury, the novel axons were identified and their point of origin from the cut axonal stump determined (yellow circles). Based on this determination of novel growth, the percentage of brains showing regrowth was calculated by counting the number of brains that sprouted at least one novel axon from the site of injury. Scale bar = 8 μm. (D) To measure the extent of regrowth, the length of a novel axon is traced by free-hand tracing (pink lines) which is a criterion to assess how long the novel axons can grow. (E) In addition to the length of the new axons, the distance these novel axons can cover in a straight line is also measured (blue line) which serves as a criterion to measure how far the novel axons can reach. (F) The size of the gap induced by axonal lesion was measured in μm by drawing a line connecting the proximal stump and closest distal stub (n=18). (G) Measurement of the direction of normal target area to a reference line running with the axonal shaft. The results of F and G were calculated as averages (n=18). (H) The direction of the novel axons was measured by first drawing a reference line running parallel with the axonal...
shaft and then (I) measuring the angle of the novel axon to the reference line. Scale bar in H = 40μm and in I is 8μm. (J) Graph showing the percentage of brains regrowing at least one axon with different genetic manipulations. The frequency is calculated as the percentage of brains on the total number of brains regrowing at least one novel axon after injury. *p<0.05; ***p<0.001. (K) Graph showing the percentage of axonal regrowth. This is calculated as the number of newly grown axons on the total number of cut axons. To calculate the total number of cut axons, the number of examined brains was multiplied by four (the average number of sLNv in one brain hemisphere) *p<0.05; ***p<0.001. (L) A comparative graph showing the extent of regrowth under different genetic manipulations. The black bars represent the average length of the new axons, the grey bars represent the average computed distance of the new axons. For statistical analysis, values of the different genotypes were compared with the values for WT brains. *p<0.05; ***p<0.001.
Figure 7. 18°C - 28°C injury experiments of PKA alleles with tubPGAL80ts

(A, A', A'') Injury on control brain. (A') Snapshot taken from a control brain ~6 hours after injury. (A'') The same brain was fixed at day 3 and imaged again. (A) A high magnification of the proximal stump. Scale bar = A', A'' is 40μm and A= 8μm. (B, B', B'') Overexpression of PKAc.A75. (B') Snapshot ~6 hours after injury. (B'') The same brain imaged at day 3. (B) High magnification of proximal stump. Scale bar in B', B'' = 40μm and B= 8μm. (C, C', C'') Overexpression of PKAc.wt has a strong positive effect on the penetrance of regrowth, but not the extent of axonal growth. (C') Snapshot of brain ~6hours after injury. (C'') Snapshot of the same brain 3 days after injury. (C'') High magnification of proximal stump. Scale bar = C', C'' is 40μm and C= 8μm. (D, D', D'') Overexpression of PKAmc* induces significantly more regrowth. Also the length of the newly grown axons is significantly bigger, but not the average distance they bridge. (D') Snapshot of brain ~6hours after injury. (D'') Snapshot of same brain 3 days after injury. (D) High magnification of proximal stump. Scale bar = D', D'' is 40μm and D= 8μm.

Genotype of (A, A', A'') is PDF-Gal4, UAS-CD8-GFP; tubPGAL80ts/+; (B, B', B'') is PDF-Gal4, UAS-CD8-GFP; tubPGAL80ts/ UAS-PKAc.A75, (C, C', C'') is PDF-Gal4, UAS-CD8-GFP; tubPGAL80ts/+; UAS-PKAc5.9F/+, (D, D', D'') is PDF-Gal4, UASCD8-GFP; tubPGAL80ts/ UAS-PKAmC*
Figure 8. 18°C - 28°C injury experiments on JNK signalling with tubPGAL80ts

(A, A’, A”) Injury on brains expressing a BskDN allele in the LNv (A’ and A”) A snapshot of the brain ∼6hours after injury and 3 days after injury and (A) a higher magnification of the injured proximal stump 3 days after injury. Scale bar in A’, A” = 40μm and A= 8μm (B, B’, B” and C’, C”) Injury on brains expressing a HepCA allele. (B’, C’ and B”, C”) Snapshot ∼6 hours after injury, 3 days after injury respectively and (B, C) higher magnifications showing massive growth. Scale bar = B’, B” is 40μm and B= 8μm (C-D) Quantifying the extent of growth observed in brains expressing HepCA in comparison to control brains. (C) Percentage of brains regrowing at least one novel axon longer than 36μm (dark blue bar); the percentage of brains growing at least one novel axon towards the target area (light blue bar); the percentage of brains growing at least one novel axon into the target area (pink bar). (D) Schematic drawing of the different quantifications described above in C. All values are significant (*P<0.05, ***P<0.001). (E, E’, E”) Example of HepCA expressing neurons showing regrowth back into the target area. Scale bar in E’, E” = 40μm and E= 8μm

Genotype of (A, A’, A”) is PDF-Gal4, UAS-CD8-GFP; tubPGAL80ts/BskDN, (B, B’, B”, E, E’, E”) is PDF-Gal4, UAS-CD8-GFP; tubPGAL80ts/+; UAS-hepCA

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