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Understanding the role of dMob4 in postembryonic Neural Stem Cell mitotic reactivation

by

ELEANOR GONZAGA

A thesis submitted to the University of Plymouth in partial fulfilment for the degree of

DOCTOR OF PHILOSOPHY

Peninsula Medical School

July 2020

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Signed..... EGONZAGA

Date.....16.07.2020

For Nan

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ABSTRACT

UNDERSTANDING THE ROLE OF DMOB4 IN POSTEMBRYONIC NEURAL STEM CELL MITOTIC REACTIVATION

Eleanor Gonzaga

Brain homeostasis supports cognition and neural plasticity, the brains' ability to adapt to its environment throughout life. These processes rely on replenishment from stem cells in the brain, termed Neural Stem Cells (NSCs). In the adult brain, most NSCs exist in a quiescent state and must activate to generate new neural progeny. Understanding how NSCs balance quiescence and activation is crucial for the development of brain repair therapies. Drosophila NSCs, termed neuroblasts (NBs), are a model to study reactivation from quiescence. The Hippo and Insulin pathways were reported to promote NB quiescence and reactivation, respectively. However, the underlying mechanisms are not fully understood. This thesis describes my investigation of Monopolar spindle-one-binder 4 (Mob4), a member of the highly conserved Striatin-Interacting Phosphatase and Kinases (STRIPAK) complex, identified by our laboratory in a transcriptome analysis comparing guiescent and reactivating NBs. I show that Mob4 is upregulated in reactivating NBs, and its loss prevents NB reactivation. Mob4 overexpression accelerates reactivation, which can be mimicked by human MOB4 expression, suggesting a conserved function. I provide evidence that Mob4 acts primarily cell-autonomously in pNBs and demonstrate that inactivation or activation of Hippo and Insulin signalling, respectively, in mob4 pNBs can restore reactivation. Finally, I show that Mob4 cooperates with another STRIPAK member, Cka, to promote pNB reactivation, whereas PP2A phosphatase with its regulatory subunit Widerborst maintains NB quiescence. My results, together with data from our group, lead to a model whereby Mob4 and other STRIPAK members assist the coordination of Insulin and Hippo pathways to promote NB reactivation. Given the evolutionary conservation of the molecules investigated, my findings may also be relevant to mammalian NSCs and other stem cells.

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LIST OF ABBREVIATIONS

A.A	Amino Acid
ALH	After Larval Hatching
ana	anachronism
aNSC	Adult Neural Stem Cell
Antp	Antennapedia
APKC	Atypical Protein Kinase C
ASCL1	Achaete-Scute Homolog 1
Ase	Asense
Ban	Bantam
Baz	Bazooka
BBB	Blood-Brain Barrier
b-HLH	Basic Helix-Loop-Helix
BMP	Bone Morphogenetic Protein
Вр	Base Pair
Brat	Brain Tumour
Brdu	Bromodeoxvuridine
BV	Blood Vessels
Cas	Castor
CB	Central Brain
CCM3	Cerebral Cavernous Malformations 3
CD8-GFP	Membrane-Bound Green Fluorescent Protein
ChIP	Chromatin immunoprecipitation
Chro	Chromator
Cka	Connector Of Kinase To AP-1
CLK	CLOCK
CNS	Central Nervous System
CNTR	Control
Co-IP	Co-immunoprecipitation
CRL4	Cullin4-RING
CSF	Cerebral Spinal Fluid
Cul4	Cullin4
CYC	CYCLE
DCX+	Double Cortin Positive
DDB1	Damaged DNA-Binding Protein 1
DG	Dentate Gyrus
DILP	Drosophila Insulin-Like Peptides
DII1	Delta-Like 1
DNA	Deoxyribonucleic acid
Dpn	Deadpan
East	Enhanced Adult Sensory Threshold
ECL	Enhanced chemiluminescence
Edu	5-Ethynyl-2'-Deoxyuridine
ERK	Extracellular Signal-Regulated Kinases
Eve	Even-skipped
EZ	Ependymal Zone

FACS	Fluorescence-Activated Cell Sorting
FB	Fat Body
FDS	Fat Body-Derived Signal
FGFR10P2	Fibroblast Growth Factor Oncogene Partner 2
FMRP	Fragile X Protein
Foxo	Forkhead Box O3
FUCCI	Fluorescent Ubiguitination-Based Cell Cycle Indicator
GABA	Gamma-Aminobutvric Acid
GCK	Germinal Centre Kinase
GCL	Granule Cell Laver
GCL	Granule Cell Laver
GFAP	Glial Fibrillary Acidic Protein
GFP	Green Fluorescent Protein
GMC	Ganglion Mother Cell
GO	Gene Ontology
Grh	Grainvhead
Н	Hour
Hb	Hunchback
Hes1	Enhancer Of Split 1
Hno	Hinno
HRP	Horse Radish Peroxidase
Hsp83/Hsp90	Heat Shock Protein 83
IGE	Insulin-Like Growth Factor
II P	Insulin-Like Pentides
INP	Intermediate Neural Progenitor
InR	Insulin-Like Recentor
Insc	Inscuteable
IPC.	Intermediate Progenitor Cell
INK	C-Jun N-Terminal Kinases
Kr	Krunnel
Mahi	Mahiong
ΜΔΡΚ	Mitogen-Activated Protein Kinase
mbNB	Mushroom Body Nbs
Mfae8	Milk Fat Globule-Epidermal Growth Factor
MI	Molecular Laver
Mob4	Monopolar Spindle-One-Binder 4
mRNA	Messenger RNA
Msn	Misshapen
Mst1/2	Mammalian Hippo Kinase
Mts	Microtubule Star
Mud	Mushroom Body Defect
NR	Nutrition Restricting
NR	Neuroblasts
NDR	Nuclear Dhf2-Related
NEP	Neuroenithelial Progenitors
NEIX	Nuclear Factor One Family Member
NPD	Non-Radial-Precureore
INIAI	

Non-Significant
Neural Stem Cells
Neurotrophin-3
Olfactory Bulb
Optic Lobe
Phosphorylated-AKT
Phosphate Buffered Saline
Proliferating Cell Nuclear Antigen
Polymerase Chain Reaction
Pou Domain 2
Phospho-Histone H3
Pleckstrin Homology Domain-GFP Fusion Protein
Pleckstrin Homology-Like
Phosphatidylinositol 3-Kinase
Partner Of Inscuteable
Postembryonic Neuroblasts
Pointed P1
Partner Of Numb
Protein Phosphatase 2A
Complex Polycomb Repressor Complex 2
Prospero
Protein S
Quiescent Adult Neural Stem Cells
Reversed Polarity
Repressor Element 1-Silencing Transcription
Radial Glial Cell
Radial Glia-Like
Ras Homologue Enriched In Brain
Rostral Migratory Stream
Ribonucleic Acid
RING Of Cullin
Real-Time Quantitative Polymerase Chain Reaction
Salvador
Sodium dodecyl sulphate
Standard Error of the Mean
S/G2 Nuclear Autoantigen
Subgranular Zone
Suppressor Of IKBKE 1
Sirtuin 1
Slimfast
Sarcolemmal Membrane-Associated Protein
Sex Determining Region Y-Box 2
STRN-Interacting Protein 1/2
Striatin-Interacting Phosphatase And Kinases
Striatin
Subventricular Zone
Transient Amplifying Cells

Tara	Taranis
TBS	Tris Buffered Saline
TBST	Tris buffered saline containing Tween-20
Tcs1/2	Tuberous Sclerosis Complex 1 And 2
TEMED	Tetramethylethylenediamine
TLX	Orphan-Nuclear Receptor
TOR	Target Of Rapamycin
Trol	Terribly Reduced Optic Lobes
Tws	Twins
UAS	Upstream Activation Sequence
VCAM1	Vascular Cell Adhesion Molecule-1
VEGF	Vascular Endothelial Growth Factor
VNC	Ventral Nerve Cord
VS	Ventricular Space
Wdb	Widerborst
wdb-DN	Dominant Negative widerborst
Wrd	Well Rounded
WT	Wild-Type
Wts	Warts
Yki	Yorkie

CHAPTER 1

INTRODUCTION

1.1. Neural stem cells, the progenitors of the central nervous system

Stem cells are multipotent, self-renewing cells capable of making differentiated progeny, providing a homeostatic function to the tissue in which they reside (Li and Clevers, 2010; Cheung and Rando, 2013). Neural Stem Cells (NSCs) are the progenitor cells of the central nervous system (CNS), defined by their ability to self-renew and generate further committed progeny, such as neurons and glia (astrocytes and oligodendrocytes). It is the NSCs which are responsible for generating the enormous cell number and diversity of neural tissue which animals rely on for CNS functions including learn- ing, memory formation and plasticity (Blau *et al.*, 2001; Sahay *et al.*, 2011; Spalding *et al.*, 2013; Eichenbaum, 2004). Thus, it is imperative that the correct type and number of these cells are generated at the correct time points (Cheung and Rando, 2013). Understanding the mechanisms controlling NSCs which are affected by various physiological and environmental factors could be the leading avenue for the development of new therapies against a variety of brain disorders.

The process by which NSCs generate new neural cells is called neurogenesis (Altman, 1962). NSCs exist during embryonic development and adulthood. Embryonic NSCs are multipotent and generate multiple subtypes of neural cells. The earliest NSCs are called neuroepithelial progenitors (NEPs) and, in vertebrates, these reside in the neural tube, which arises from the neuroectoderm during embryogenesis. NEPs give rise to NSCs in a spatially and temporally controlled manner (Temple, 2001; Alvarez-Buylla *et al.*, 2009). NEPs initially undergo symmetric division to expand the NSC pool, but later switch to asymmetric divisions to self-renew and give rise to a post-mitotic neuron or an intermediate progenitor cell (IPC) (Temple, 2001; Mori *et al.*, 2005; Kriegstein and Alvarez-Buylla, 2009; Anthony *et al.*, 2004). During this time, the embryonic NSCs acquire astrocyte-like properties and are termed radial glial cells (RGCs), based on their radial processes and morphological resemblance to astrocytes (Mori *et al.*, 2005). IPCs, located in the subventricular zone (SVZ), eventually generate a pair of post-mitotic neurons via symmetrical division (Noctor *et al.*, 2004). The

neurons generated migrate vast distances in the developing cortex using RGCs as contacts or scaffolds (Kriegstein and Alvarez-Buylla, 2009). RGCs and IPCs become a significant source of neural cells; RGCs expand the cortex tangentially, and IPCs fill the cortical layers radially to create the functional CNS. The CNS generated during embryogenesis was thought to be complete and therefore last an adult mammalians' lifetime, incapable of generating new neural tissue for regeneration or repair (Ramón y Cajal, 1909). However, this was overturned by a growing body of evidence showing multipotent NSCs within distinct niches of the adult mammalian CNS (Reynolds and Weiss, 1992; Richards *et al.*, 1992; Gage and Fisher, 1995; Eriksson *et al.*, 1998; Gould *et al.*, 1999a; Gould *et al.*, 1999b; Fuentealba *et al.*, 2012; Fuentealba *et al.*, 2015).

1.2. Adult NSCs and neurogenesis in mammals

Adult neurogenesis is the formation, maturation and integration of new neurons into the existing circuitry. It was previously believed that the mammalian brain lacked adult neurogenesis and was, therefore, incapable of regeneration or repair (Ramón y Cajal, 1909). However, in the 1960s, Joseph Altman discovered the first evidence of adult neurogenesis - the source of adult-born olfactory neurons, in adult rat and guinea pig brains using 3H-thymidine autoradiography (Altman, 1962; Altman, 1963; Altman, 1969). In the 1990s, adult neurogenesis was demonstrated *in-vitro* using mammalian NSCs (Reynolds and Weiss, 1992) and in the adult rat brain (Gage and Fisher, 1995). The excitement was propagated by evidence of adult neurogenesis obtained in the human brain using bromodeoxyuridine (BrdU) and specific neuronal markers such as NeuN (Eriksson *et al.*, 1998). One year later, Elizabeth Gould showed that neurogenesis contributes to the adult macaque neocortex (Gould *et al.*, 1999). Due to the growing body of evidence, neurogenesis in humans is now generally acknowledged (Abdissa, 2020).

Unlike embryonic NSCs, which are the highly proliferative, adult NSCs (aNSCs) have been shown in mammalian models to be mostly quiescent and restricted to only a few neurogenic regions. Most notably are the subgranular zone (SGZ) of the hippocampal dentate gyrus (DG), and the subventricular zone (SVZ) lining the lateral ventricles (**Figure 1.A**). Both regions contain an environment or 'niche' which provides a tightly

controlled microenvironment to regulate the residing aNSCs. Two non-canonical sites of adult neurogenesis have also been reported in the striatum and the hypothalamus (**Figure 1.A**).



Figure 1.1. Mammalian neurogenic regions and NSC lineages. A. A schematic representation of a mouse brain with the neurogenic regions outlined. Neurogenesis in the subgranular zone (SGZ) of the hippocampus produces mature granule neurons for the dentate gyrus (DG). Neurogenesis in the subventricular zone (SVZ) produces neuroblasts which migrate through the rostral migratory stream (RMS) to the olfactory bulb (OB). The SGZ and the SVZ are known as classical neurogenesis regions. Two non-canonical sites of neurogenesis are the hypothalamus and the Striatum. **B.** The SGZ NSC lineages and **C.** the SVZ NSC lineages and common markers used to distinguish them. Molecular layer (ML); Granule cell layer (GCL); Radial glial cell (RGL); Glial fibrillary acidic protein (GFAP); Achaete-scute homolog 1 (ASCL1); Sex determining region y-box 2 (Sox-2); Neuronal Differentiation (NeuroD); Doublecortin (DCX); Prospero homeobox 1 (Prox1); Intermediate progenitor cell (IPC); Cerebral spinal fluid (CSF); Ependymal Zone (EZ); blood vessels (BV).

1.2.1. aNSCs in the hippocampus

In mammalian adult brain models such as in mice and rats, aNSCs in the SGZ of the hippocampus generate mature granule cells of the DG (**Figure 1.1. A**), an area critical for learning and memory (Song *et al.*, 2012a). The DG of the hippocampus has three distinct layers; the molecular layer, the granule layer and the SGZ (**Figure 1.1. B**).

Three types of aNSCs exist in the dentate gyrus: self-renewing 'Type 1 NSCs'; 'Type 2' amplification cells; and migratory 'Type 3 cells/neuroblasts' which give rise to granule cells/neurons. In the DG, aNSCs follow a characterised, multistep lineage development and are identifiable by a subset of temporally expressed markers (summarised in Figure 1.1. B). 'Type 1'/Radial glia-like (RGL) NSCs resemble RGCs of the embryonic brain and reside at the border of the granulate layer. These cells usually exist in a quiescent (G0) state and rarely divide, favouring asymmetric division if they are activated (Kemperman, 2011). They extend an elongated radial process through the granule cell layer (GCL) to the molecular layer (ML) and have a cilium in contact with blood vessels and cerebrospinal fluid (Kemperman, 2004). Two populations of Type 1 NSCs exist in the hippocampus. Both populations express Sex determining region y-box 2 (Sox-2), but one is a quiescent RGL population expressing Nestin and Glial fibrillary acidic protein (GFAP); and another population, of non-radial glia-like cells, are more proliferative (active) and lack radial-glia-like markers (Suh et al., 2007). RGL NSCs give rise to two 'Type 2' intermediate progenitor cell (IPC) subgroups: Type 2a/non-radial- precursors (NRPs) express Achaete-scute homolog 1 (ASCL1); Type 2b/intermediate progenitor cells (IPCs) do not express ASCL1 and are further committed. After a limited round of division, Type 2b IPCs exit the cell cycle and terminally differentiate into astrocytes and neurons (Ming and Song, 2011; Bonaguidi et al., 2011). Both Type 2 populations are relatively active, have limited divisions, demonstrate short processes and differentially express migration and neuronal markers (Kempermann et al., 2004; Encinas et al., 2011). Type 3 cells/neuroblasts tangentially migrate through the SGZ, lack previous lineage markers (Nestin, Sox2, GFAP) and express Neuronal Differentiation 1 (NeuroD) and Prospero Homeobox 1 (Prox1) to give rise to immature granule cells/neurons. The immature granule cells/neurons migrate radially into the GCL, mature and differentiate into dentate granular cells/neurons. After 7-8 weeks, they integrate into the hippocampal circuitry (Göritz and Frisén, 2012; Jessberger and Kempermann, 2003) and contribute to hippocampus-associated learning/memory (Kempermann et al., 2015). It has been estimated that the young rat hippocampus produces approximately 9000 new cells every day (Cameron and McKay, 2001), with ~30% of these cells (~800) thought to survive past four weeks and develop dendritic and axonal structures required for proper functioning (Snyder and Cameron et al., 2009; Sailor et al., 2017; Sun and Song et al., 2013). Defects in hippocampal neurogenesis are linked to brain-related

disorders, for example, depression, impaired spatial learning and memory, pattern separation and mood regulation (Clelland *et al.*, 2009; Hill *et al.*, 2015; Lee *et al.*, 2013; Anacker, 2014; Anacker and Hen, 2017; Apple *et al.*, 2017; Oppenheim, 2019).

1.2.2. aNSCs in the subventricular zone

The subventricular zone (SVZ) is located next to the lateral ventricles and is the largest germinal region in the adult brain (Obernier and Alvarez-Buylla, 2019; Lim and Alvarez-Buylla, 1999; Doetsch *et al.*, 1999). The SVZ is a thin layer of proliferating cells located along the ependymal cell layer/zone (EZ), which separates the ventricular space (VS) from the SVZ (**Figure 1.1. C**); at the edge of the striatum (**Figure 1.1. C**). SVZ aNSCs arise from striatal radial glia and function to continually give rise to oligodendrocytes, astrocytes and olfactory bulb (OB, in rodents) or striatal interneurons (in humans) (Lim and Alvarez-Buylla, 2016; Obernier and Alvarez-Buylla, 2019; Ming and Song, 2016). Overall, the SVZ contains four cell types: Type B/RGL cells (NSCs); Type C/transient amplifying cells (TACs); Type A cells/neuroblasts, and multiciliated ependymal cells, which produce and circulate cerebral spinal fluid (CSF), essential for the niche (**Figure 1.1. C**).

Type B/RGL NSCs have a radial/astrocyte-like morphology, extend an apical process with a primary cilium through the EZ and into the VS, and a basal process which terminates on the blood vessels (BVs) (Mirzadeh *et al.*, 2008). Type B/RGL NSCs of the SVZ express stem cells markers such as GFAP, Nestin and Sox2 (Liu *et al.*, 2006; Zhang and Jiao, 2015). Type B/RGL NSCs are not described as quiescent but do divide slowly to produce highly proliferative type C/TACs (Doetsch *et al.*, 1999; Zhang and Jiao, 2015), also known as the progenitors of the SVZ. The Type C/TACs lack GFAP expression but do express Nestin, Sox2 and ASCL1 (Kempermann, 2011; Zhang and Jiao, 2015). After multiple rounds of divisions, TACs become Double cortin positive (DCX+) type A cells/neuroblasts. These neuroblasts form a chain and migrate radially along the rostral migratory stream (RMS) to the olfactory bulb (OB), and differentiate into dopamine and GABA-producing interneurons that integrate into the OB circuitry (Ming and Song, 2011; Zhu *et al.*, 2018). The OB receives an estimated 30,000 new neuroblasts every day (Lois and Alvarez, 1994), with ~40% of these cells (12,000) surviving past four weeks and effectively integrating into the

circuitry (Mouret *et al.*, 2008). SVZ neurogenesis is linked to olfactory function and learning (Lledo *et al.*, 2006; Bragado *et al.*, 2019), and decreased SVZ neurogenesis causes impaired odour discrimination (Valley *et al.*, 2009; Sahay *et al.*, 2011). SVZ proliferation and differentiation are altered in response to stroke/ ischemia, whereby SVS aNSCs and neuroblasts are redirected to the injury site and can directly or indirectly generate the appropriate neural cells towards attempt repair (Arvidsson *et al.*, 2002; Zhang *et al.*, 2004; Zhang *et al.*, 2008b; Faiz *et al.*, 2015). Reduced SVZ neurogenesis is linked to neurodegenerative diseases in rodents (Curtis *et al.*, 2007b; Rodríguez and Verkhratsky, 2011; Winner and Winkler, 2015; Scopa *et al.*, 2019; Sung *et al.*, 2020).

1.2.3. Non-canonical adult NSC niches: striatum and hypothalamus

The striatum is located in the subcortical basal ganglia of the forebrain and is a significant input area for the basal ganglia and critical to the reward system (Figure **1.1. A**). Local astrocytes in the mouse striatum can produce new neurons in response to stroke (Magnusson et al., 2014). Striatal adult neurogenesis has been observed in adult humans (Bergmann et al., 2012; Ernst et al., 2014; Wang, C. et al., 2014; Wang et al., 2011), where it continually gives rise to interneurons (Ernst et al., 2014); and neuroblasts in the SVZ are observed to migrate to the striatum in humans (Ernst et al., 2014). The striatum is associated with motor function, cognitive flexibility, cognitive function and, more recently, schizophrenia pathology (Simpson et al., 2010; Ernst and Frisén, 2015). Impaired striatal neurogenesis is observed in patients with Parkinson's disease (Ernst et al., 2014); the striatal cell death found is thought to induce SVZ growth to aid migration via the caudate nucleus (Curtis et al., 2007a). However, as decreased neuronal production has been observed in Parkinson's disease patients, it suggests that the increased neuronal migration is insufficient to replenish the lost neurons (Ernst et al., 2014). Also, the striatum is associated with Huntington's disease, whereby striatal adult-born neurons are selectively depleted 2014). Although there are currently fewer studies on striatal (Ernst *et al.*, neurogenesis compared to other neurogenic regions, there is growing evidence for its importance.

Most recently, adult neurogenesis has also been observed in the hypothalamus, a small area and the organisms' homeostatic regulator (**Figure 1.1. A**). It was first shown

that cells from the hypothalamic VZ could generate neurospheres *in-vitro* (Weiss *et al.*, 1996). Later it was reported that the tanycytes (hypothalamic RGCs) function as neural precursors (Robins *et al.*, 2013; Xu *et al.*, 2005; Haan *et al.*, 2013; Lee *et al.*, 2012). Tanycytes resemble embryonic neural progenitors and are exposed to CSF (Bolborea and Dale, 2013; Rodríguez *et al.*, 2005). They extend one or two apical cilia into the VZ and a long basal process into the parenchyma or pial surface, depending on location. Much like the NSCs in the SVZ and SGZ, tanycytes are reportedly influenced by diet and also regulate weight gain (Niwa *et al.*, 2016; Lee *et al.*, 2014; Whalley, 2012); hypothalamic adult neurogenesis helps regulate body weight and energy balance by replacing degenerated neurons (Pierce and Xu, 2010; McNay *et al.*, 2012). Overall levels of hypothalamic adult-born neurons are low compared to the other neurogenic niches (Lee and Blackshaw, 2014), which adds difficulty in exploring the functional significance of hypothalamus neurogenesis.

It is becoming clear that adult neurogenesis in areas of the mammalian brain, canonical and non-canonical, offer advantages; and its deregulation is associated with cognitive disadvantages. How exactly adult neurogenesis is achieved, the pathways and regulators involved, and therapeutic advantages are significant questions in stem cell biology.

1.3. Quiescence and activation of aNSCs

Mammalian aNSCs interconvert between actively proliferating and temporary quiescent states. The transition is regulated by a variety of extrinsic and intrinsic cues, which we are only beginning to understand (Basak *et al.*, 2012; Basak *et al.*, 2014; Basak *et al.*, 2018; Costa *et al.*, 2011; Giachino *et al.*, 2014; Ziebell *et al.*, 2018; Velthoven and Rando, 2019; Morales and Mira, 2019; Mohammad *et al.*, 2019). Below, I describe some of the key literature contributing to our current knowledge of aNSC quiescence and activation.

1.3.1 aNSCs are actively maintained in cellular quiescence but can reactivate in response to appropriate signals

Quiescence is a highly conserved, reversible cell cycle arrested state. Most of our knowledge on cell quiescence comes from *Saccharomyces cerevisiae*, *Schizosaccharomyces pombe*, and *Escherichia coli* (Werner-Washburne *et al.*, 1993; Lillie

and Pringle, 1980). Stem cell quiescence is conserved in mammals across various tissues such as skin, gut and brain and all share commonalities: decreased transcription of cell-cycle progression genes increased expression of genes involved in suppressing apoptosis, differentiation and senescence, a decreased cell size, stress resistance, condensed chromosomes and elevated autophagy (Cho et al., 2019; Velthoven and Rando, 2019). The quiescent state was first observed *in-vitro* in 1951 and later observed in unicellular and multicellular organisms (Dubrovsky, 2003; Velthoven and Rando, 2019; Howard and Pelc, 1986). In mammals, during G1, before committing to mitosis, somatic cells may withdraw from replication and enter quiescence (Pardee, 1974). Quiescent aNSCs (qaNSCs) only rarely divide, making them challenging to label using lineage-tracing methods labelling dividing cells only. Quiescence allows increased longevity, tolerance to metabolic stress, and ultimately acts to preserve the stem cell pool, maintaining tissue homeostasis. Previously, quiescent NSCs were thought to exhibit little activity; however, it is now regarded as an actively 'restrained' state. These 'poised' quiescent somatic cells are ready for activation in response to appropriate stimuli to regenerate tissue. Cell-extrinsic stimuli such as nutrition, exercise, stress, and cell-intrinsic epigenetic, transcriptional and post-transcriptional mechanisms regulate quiescence (Cheung and Rando, 2013).

The heterogeneity and dynamics of aNSCs populations have made qaNSCs challenging to study. Quiescence was previously assumed based on low RNA levels, poor label-retaining ability and lack of proliferation markers such as Proliferating cell nuclear antigen (PCNA), Ki67 and phospho-histone H3 (PH3). The last decade has provided technology to study qaNSCs such as *in-vivo* imaging, clonal lineage tracing, fluorescence-activated cell sorting (FACS), whereby subpopulations of aNSCs can be purified and subjected to a microarray, and single-cell RNA sequencing (Ming and Song, 2014; Bonaguidi *et al.*, 2011; Shin *et al.*, 2015; Marcy, G. and Raineteau, 2019). The advancement has allowed a single-cell transcriptomic analysis of qaNSCs and activated aNSCs cells at population and cell level. Multiple studies show that aNSCs (RGLs) are actively maintained in quiescence (Codega *et al.*, 2014; Llorens-Bobadilla *et al.*, 2015; Dulken *et al.*, 2017; Velthoven and Rando, 2019). Mammalian qaNSCs can be identified by specific markers such as GFAP, Sox2, and Nestin. They appear smaller in size with a more radial morphology and favour glycolytic and lipid metabolism compared to active NSCs (Alvarez-Buylla *et al.*, 2001; Llorens-Bobadilla

et al., 2015; Shin *et al.,* 2015; Renault *et al.,* 2009). Interestingly, most aNSCs exist primarily in a quiescent state (Codega *et al.,* 2014; Llorens-Bobadilla *et al.,* 2015).

aNSC quiescence is described to enhance stress resistance, defends against deleterious gene mutations and exhaustion to secure the NSC pool (Urbán and Guillemot, 2014); it is therefore tightly regulated by extracellular and intracellular signals. The neurogenic niche communicates the needs of the organism to the qaNSCs, including via the vasculature and CSF, which are in contact with the NSC basal and apical processes, respectively. Extracellular receptor activation can lead to the activation of intrinsic factors such as signal transduction pathways, transcription pathways, metabolism and epigenetic modifications. These extrinsic/intrinsic mechanisms crosstalk and feedback on each other to modulate aNSC cellular activity within the neurogenic niche (Otsuki and Brand, 2017; Faigle and Song, 2013; Ming and Song, 2015; Tian *et al.*, 2018; Chaker *et al.*, 2016; Cavallucci *et al.*, 2016). Many studies describe the effects of molecules and signalling pathway that enhance neurogenesis and NSC proliferation in general., Below, I provide examples of molecules and pathways implicated in aNSC quiescence, specifically.

Notch signalling regulates tissue homeostasis and cell-fate determination throughout development. Direct cell-cell contact is required as the transmembrane Notch receptor is activated by binding to its neighbouring cell via Delta or Jagged ligands (Ables *et al.*, 2011). Notch regulates aNSCs by stimulating cell-cycle exit and decreasing the neural progenitor pool (Hitoshi *et al.*, 2002; Imayoshi *et al.*, 2010; Ehm *et al.*, 2010). Notch signalling has also been shown to maintain aNSCs in quiescence by repressing genes associated with the cell cycle and inhibiting differentiation (Ottone *et al.*, 2014; Engler *et al.*, 2018) and via the Delta-like 1 (DII1) ligand as a homeostatic feedback mechanism (Kawaguchi *et al.*, 2013). Recently, a downstream target of the Notch pathway, *Id4*, was shown to maintain quiescence of hippocampal NSCs (Zhang *et al.*, 2019). Cell-cell contact is required for signalling pathways such as Notch, and cell adhesion molecules that help position NSCs, such as vascular cell adhesion molecule-1 (VCAM1) and the laminin receptor $\alpha 6\beta$ 1-integrin; support NSC quiescence (Kokovay *et al.*, 2012; Shen *et al.*, 2008).

Bone morphogenetic protein (BMPs) growth factors/BMP signalling has been shown

to regulate mammalian neurogenesis (Wang, R. *et al.*, 2014; Lim *et al.*, 2000; Bonaguidi *et al.*, 2008; Mira *et al.*, 2010; Ming and Song, 2015; Mira *et al.*, 2016; Joppé *et al.*, 2015; Morell *et al.*, 2015). BMP restricts proliferation; BMP inactivation was shown to promote aNSC proliferation but later caused decreased proliferation and production of new neurons (Mira *et al.*, 2016). However, this study partially contradicts other studies (Bonaguidi *et al.*, 2008; Gobeske *et al.*, 2009; Bond *et al.*, 2012) which did not find any subsequent loss in NSC proliferation or new neuron production. Bonaguidi *et al.* showed that overexpressing the BMP antagonist, Noggin, gave rise to increased numbers of proliferative GFAP+ NSCs *in-vitro*, and in the mouse SGZ (Bonaguidi *et al.*, 2008). Interestingly, Noggin is secreted by ependymal cells and aids NSC activation (Lim *et al.*, 2000). Also, the anti-proliferative effects of BMP signalling are dominant over the pro-proliferative signalling molecule, EGF, *in-vitro* and *in-vivo* (Joppé *et al.*, 2015). Further studies are therefore needed to elucidate the function of BMP signalling on aNSCs.

Several transcription factors promote aNSC quiescence, for example, the Nuclear factor one (NFI) family member NFIX is expressed during aNSC quiescence, and activation, but is upregulated in quiescent aNSCs compared to activated aNSCs; NFIX directly activates quiescence-promoting genes, and loss of NFIX leads to a loss of aNSC quiescence (Martynoga *et al.*, 2013).

The Repressor Element 1-Silencing Transcription factor (REST) regulates unique aspects of neural development embryonically and postembryonically (Ballas *et al.*, 2005; Mukherjee *et al.*, 2016; Gao *et al.*, 2011; Gao *et al.*, 2012). REST has been shown to bind and control the expression of different gene targets in a developmental and cell-context dependent fashion (Ballas *et al.*, 2005; Mukherjee *et al.*, 2016; Gao *et al.*, 2005; Mukherjee *et al.*, 2016; Gao *et al.*, 2011). REST is highly expressed in embryonic pluripotent stem cells (ES cells), and its expression decreases via post-translational degradation as these cells develop into NSCs. REST is completely transcriptionally repressed in mature neurons (Ballas *et al.*, 2005). In the transition from pluripotent ES to NSCs, the expression of REST is thought to be just enough to maintain neuronal gene chromatin inactive, whereas as progenitors neural cells differentiate into neurons, REST together with its co-repressors dissociate from their DNA target binding sites, allowing activation of neuronal genes (Ballas *et al.*, 2005). Postembryonically, REST is required to

maintain aNSC quiescence, and to prevent neurogenesis over time (Mukherjee *et al.*, 2016; Gao *et al.*, 2011). Mukherjee *et al.* found that conditional deletion of REST in adult mouse hippocampal quiescent aNSCs results in their premature mitotic activation. Furthermore, its deletion from adult neural progenitors (transient amplifying cells, TAPs) led to increased TAP proliferation as well as increased numbers of resulting immature and mature neurons. The authors concluded that REST deletion from TAPs activated the neuronal differentiation program (Mukherjee *et al.*, 2016). The same laboratory team used genome-wide chromatin immunoprecipitation sequencing (ChIP-seq) and RNA-sequencing profiling to identify REST binding sites and targets, and found, not surprisingly, that many were neuronal genes (Mukherjee *et al.*, 2016).

Furthermore, the transcription factor Forkhead Box O3 (FoxO) promotes aNSC quiescence; loss of FoxO leads to an initial rise in aNSC proliferation but an overall depletion of the stem cell pool (Paik *et al.*, 2009; Renault *et al.*, 2009; Santo and Paik, 2018). FoxO is inhibited by AKT, the downstream component of the conserved Insulin signalling/Insulin-like growth factor (IGF) pathway (Tzivion *et al.*, 2011). Also, within this pathway, upstream of AKT is the phosphatase and tensin homolog (Pten) which is also required for NSC quiescence (Bonaguidi *et al.*, 2011). The tumour suppressor, Pten, maintains NSC quiescence by suppressing self-renewal and regulating the G0-G1 transition cell cycle entry; Pten depletion causes NSC activation from quiescence and depletion of the NSC pool (Bonaguidi *et al.*, 2011). In addition to Pten, other cell cycle regulators include the cyclin-dependant kinase inhibitors p27 and p57 which are expressed in non-dividing aNSCs and inhibit the cell cycle from maintaining the NSC pool; depletion of which leads to loss of NSC quiescence and increased neurogenesis (Andreu *et al.*, 2015; Furutachi *et al.*, 2013).

Some neurotransmitters are also known to promote aNSC quiescence. Gammaaminobutyric acid (GABA) and its precursor glutamate create a negative feedback mechanism (Liu *et al.*, 2005) and promote quiescence by preventing aNSC-cycle progression (Fernando *et al.*, 2011); decreasing GABA signalling causes aNSCs to exit quiescence (Song *et al.*, 2012b). Pineda and colleagues found that using a GABA antagonist caused activation of quiescent NSCs (Pineda *et al.*, 2013). Vascular-derived growth factors also maintain NSC quiescence. The endothelial-

derived neurotrophin- 3 (NT-3) mediates quiescence; NT-3 mutant mice showed higher aNSCs proliferation and increased number of neural cells (Delgado *et al.,* 2014). Recently, Zhou *et al.* showed that Milk fat globule-epidermal growth factor (Mfge8), otherwise known as lactadherin, promotes quiescence via mTOR signalling and its deletion causes NSC depletion and decreases neurogenesis (Zhou *et al.,* 2018).

Lastly, the secreted glycoprotein Protein S (Pros1) maintains quiescence by mediating Notch expression and activity; loss of Pros1 in aNSCs leads to increased proliferation (Zelentsova-Levytskyi *et al.*, 2017). Interestingly, in *Pros1* knockout mice, the increase in aNSC proliferation leads to an enriched aNSC pool and does not lead to aNSC pool depletion up to one year of age (Zelentsova-Levytskyi *et al.*, 2016).

The mechanisms governing aNSC quiescence are still being elucidated. **Figure 1.2** summarises some of the known involved factors and molecular mechanisms mentioned above. Variable states (or levels) of quiescence have also been identified, suggesting there is a 'gradient' of quiescence from 'dormant' to 'primed' (Llorens-Bobadilla *et al.*, 2015). This has also been observed in other adult stem cell systems (Rodgers *et al.*, 2014; Malam *et al.*, 2014; Zismanov *et al.*, 2016). aNSCs may also differ in their ability to return to quiescence after initially activating (Urbán *et al.*, 2016). In this study, a 'resting' population and a 'dormant' population of qNSCs was described (Urbán *et al.*, 2016). The majority of the stem cells are 'dormant' and thought to have never divided, whereas the resting population are fewer in number and have previously activated but returned to quiescence; the latter pool is essential for adult neurogenesis (Urbán *et al.*, 2016).

There is increasing evidence of heterogenic populations of quiescent aNSCs. Contrary to the long-held belief that quiescent stem cells are arrested in the G_0 phase of the cell cycle, emerging evidence suggests NSCs can be arrested in G_2 and that the latter are more readily reactivated than NSCs in G_0 (Otsuki and Brand, 2018). Deregulation of quiescence or activation from quiescence can lead to impaired homeostasis. How precisely the NSCs are maintained in quiescence, as well as how they can be

activated are essential questions in stem cell biology with implications for therapeutic strategies, which remain to be fully elucidated.

1.3.2. Mitotic activation of quiescent aNSCs

aNSCs exit from quiescence and entry into an active proliferative state requires tight coordination between the extracellular signals from the neurogenic niche and intrinsic factors. For example, one of the vascular endothelial growth factor (VEGF) receptors are expressed by aNSCs; VEGF receptor activation causes activation from quiescence (Han *et al.*, 2015). Significant evidence shows a variety of external stimuli can modulate mammalian adult neurogenesis, and specifically the activation of aNSCs. For example, aNSCs were shown to become activated upon brain injury (Wang *et al.*, 2016; Chang *et al.*, 2016; Llorens-Bobadilla *et al.*, 2015), ischemia (Nakatomi *et al.*, 2002; Nemirovich-Danchenko and Khodanovich *et al.*, 2019), and medial temporal lobe epileptic seizures (Parent *et al.*, 1997). Although a great deal of literature exists surrounding the regulation of mammalian NSC proliferation and other stages of neurogenesis (Recently reviewed by Obernier and Alvarez-Buylla, 2019; Morales and Mira, 2019; Petrik and Encinas, 2019), there is less understanding about the process of NSC activation from quiescence specifically. The few factors that have been identified to regulate this process are described below.

One of the identified aNSC activators is the basic helix-loop-helix (bHLH) transcription factor, Achaete-scute-like 1 (ASCL1), also known as Mash1. ASCL1 is a mammalian proneural gene (Guillemot and Hassan, 2017; Guillemot, 2007) which has been associated with NSC activation in multiple studies (Llorens-Bobadilla *et al.*, 2015; Dulken *et al.*, 2017; Basak *et al.*, 2018). ASCL1 is required throughout mouse embryogenesis to promote neuronal fate and progenitor proliferation (Castro *et al.*, 2011; Bertrand *et al.*, 2002), and has been observed in the developing human neocortex suggesting a conserved function (Hansen *et al.*, 2010). Postembryonically, ASCL1 is famously required for NSC activation from quiescence and neurogenesis but is also expressed in quiescent NSCs, although at low levels (Anderson *et al.*, 2014). ASCL1 loss blocks quiescent NSC activation, the production of new neurons and causes a depletion of the stem cell pool (Anderson *et al.*, 2014). ASCL1 can also promote NSC cell-cycling by activating pro-proliferative genes such as Cyclin D1 (Castro *et al.*, 2011; Urbán *et al.*, 2016).

ASCL1 is regulated by Notch signalling; Notch inactivation causes ASCL1 upregulation (Anderson *et al.,* 2014). Hairy and enhancer of split 1 (Hes1), a transcriptional repressor and downstream target of the Notch pathway, inhibits ASCL1 in embryonic NSCs (Shimojo *et al.,* 2008; Imayoshi *et al.,* 2013) and adult mammalian NSCs (Sueda *et al.,* 2019). Sueda and colleagues showed that Hes1 expression oscillates in both quiescent and activated NSCs; however, more so during quiescence to subsequently periodically suppress ASCL1 expression (Sueda *et al.,* 2019). Hes1 levels were higher in quiescente and oscillating ASCL1 expression controlled NSC activation, whereas continued ASCL1 expression controlled differentiation (Sueda *et al.,* 2019). Inactivating Hes1 lead to more ASCL1 expression and subsequent NSC activation from quiescence but also decreased overall neurogenesis (Sueda *et al.,* 2019).

ASCL1 is also controlled via post-translational modification by the ubiquitin ligase Huwe1 that stimulates ASCL1 degradation, causing more NSC activation from quiescence (Urbán *et al.*, 2016). Inactivating Huwe1 increases ASCL1 stability and causes the overproliferation of NSCs, which are unable to return to quiescence (Urbán *et al.*, 2016). In embryonic NSCs, both AKT (Oishi *et al.*, 2009) and cyclindependant kinases (Ali *et al.*, 2014) promote or inhibit ASCL1 function, respectively. Given the role of Insulin signalling in promoting adult neurogenesis, it would be interesting to investigate its activity upon ASCL1 and NSCs activation.

Another intrinsic regulator of NSC activation is the Orphan-Nuclear receptor (TLX), also known as Nr2e1 and Tailless, belongs to a family of intracellular transcription factors involved in embryonic patterning (Monaghan *et al.*, 1995). TLX is part of a superfamily of nuclear hormone receptors which recruit corepressors to cause transcriptional repression of target genes (Niu *et al.*, 2011; Wang and Xiong, 2016; Wang *et al.*, 2013). Although TLX is expressed during embryonic development, after initially declining, TLX expression later increases to peak within the adult mouse brain (Monaghan *et al.*, 1995). Postembryonically, TLX is expressed highly in the stem cells of neurogenic niches and maintains adult NSCs self-renewal capacity and its overexpression or deletion results in enhanced neurogenesis or reduced NSC

proliferation, respectively (Murai et al., 2014; Shi et al., 2004; Li et al., 2012; Sobhan et al., 2017; Zhang et al., 2008a). TLX represses transcription of some of its target genes, for example, Pten and p21 (Niu et al., 2011; Sun *et al.*, 2007), alternatively, it activates proneural gene transcription of ASCL1 (Elmi et al., 2010). TLX has been shown to regulate the Wnt signalling pathway (Qu et al., 2010) and be regulated by MicroRNA's such as microRNA-9 (Zhao et al., 2009; Sun et al., 2011). MicroRNA's are highly conserved short strands of non-coding RNA, which interfere with mRNAs via complementary base pairing that can lead to mRNA destabilisation (Fabian *et al.*, 2010). TLX was previously shown to be upregulated in activated NSCs, and NSCs in *tlx* mutant mice have cell-cycle entry and proliferation defects (Obernier *et al.*, 2011). Most recently, TLX was shown to negatively regulate transcription of the Notch effector, Hes1, in the mouse brain (Luque-Molina et al., 2019). Elevated Hes1 levels and Notch signalling in *tlx* mutants lead to disrupted quiescence and the downregulation of ASCL1/Mash1 (Luque-Molina et al., 2019).



Figure 1.2: Summary of factors known to regulate mammalian adult NSCs transit between quiescence and activation states. Notch signalling maintains quiescence by repressing genes associated with the cell cycle and inhibiting differentiation (Sueda et al., 2019; Ottone et al., 2014; Engler et al., 2018; Kawaguchi et al., 2013; Zhang et al., 2019). Vascular cell adhesion molecule-1 (VCAM1) and the laminin receptor $\alpha 6\beta$ 1-integrin help position aNSCs and support quiescence (Kokovay et al., 2012; Shen et al., 2008). Bone morphogenetic protein (BMP) is thought to support quiescence, as Noggin, a BMP inhibitor secreted by ependymal cells, aids NSC activation (Lim et al., 2000; Martynoga et al., 2013). Nuclear factor one (NFI) family member (NFIX) stimulates quiescencepromoting genes (Martynoga et al., 2013). The transcription factor repressor element 1-silencing transcription (REST) inhibits proneural genes to support quiescence (Ballas et al., 2005; Mukherjee et al., 2016; Gao et al., 2011). The Forkhead Box O3 (FoxO); however, the mechanisms are not fully understood transcription factors (Paik et al., 2009; Renault et al., 2009; Santo and Paik, 2018). The phosphatase and tensin homolog (Pten) promotes quiescence by inhibiting self-renewal and the G0-G1 transition (Bonaguidi et al., 2011). The neurotransmitter Gamma-aminobutyric acid (GABA) prevent NSC cell-cycle progression (Song et al., 2012; Pineda et al., 2013). Cell-cycle regulators such as the cyclin-dependant kinase inhibitors p27 and p57 (Andreu et al., 2015; Furutachi et al., 2013) and growth factors received from the niche such as the endothelial-derived neurotrophin- 3 (NT-3) (Delgado et al., 2014), and the Milk fat globule-epidermal growth factor (Mfge8) (Zhou et al., 2018) and promote NSC quiescence. Molecules regulating the activation of quiescent adult NSCs is far less known, the Vascular endothelial growth factor (VEGF) receptors (VEGF3) expressed on NSCs cause activation upon receptor activation (Han et al., 2016). The basic helix-loop-helix (bHLH) transcription factor, Achaete-scute-like 1 (ASCL1) promotes activation from quiescence by promoting pro cell-cycle and proliferation genes and is regulated by Notch signalling and post-translationally by the ubiquitin ligase Huwe1 (Anderson et al., 2014; Castro et al., 2011; Urbán et al., 2016; Llorens-Bobadilla et al., 2015; Dulken et al., 2017; Basak et al., 2018). The Orphan-Nuclear receptor (TLX) promotes activation of quiescent NSCs by promoting ASCL1, NSC self-renewal, and repressing quiescence-promoting genes such as pTen and p21 (Luque-Molina et al., 2019; Obernier et al., 2011; Yu et al., 1994; Monaghan et al., 1995; Wang et al., 2006; Zhang et al., 2006; Sun et al., 2007; Yokoyama et al., 2008).

1.4. Drosophila melanogaster, a model organism in neurobiology

The fruit fly *Drosophila melanogaster* is a model organism that has been extensively used in neurobiology studies, including postembryonic neurogenesis (Ramon-Cañellas *et al.*, 2019; Doe, 2017; Harding and White, 2017; Sullivan, 2019). The benefits of *Drosophila* as a model include a short life cycle of approximately 11 days at 25°C, cost-effectiveness compared to other models such as mice, highly fecund and, most importantly, *Drosophila* shares over 70% disease gene sequences with humans (Reiter *et al.*, 2001).

Moreover, there is a vast genetic toolkit available for easier gene manipulation, and, significantly for my work, one can study NSCs at a single-cell resolution in vivo (Schnorrenberg *et al.*, 2016; Jeibmann and Paulus, 2009; Pandey and Nichols, 2011). One of the key gene manipulation techniques used with *Drosophila* is the UAS-GAL4 system, a genetic tool allowing targeted gene expression (Brand and Perrimon 1993; Elliott and Brand, 2008). Since its establishment, the UAS/GAL4 system has been widely used, modified and expanded to allow for more specific temporal and spatial control of targeted gene expression. For example, it can be used for a variety of loss and gain of function assays, employed simultaneously with other independent expression systems such as LexA/LexAop allowing additional manipulation, and depending on the system's reporters, it can inform on a variety of parameters including visualisation of cells and their organelles, cell activity status and behaviour (Caygill and Brand, 2016).

In its simplest form, the UAS/GAL4 system is based on a driver and a responder fly line. A transgenic fly line will contain a desired promoter gene region fused to the yeast transcriptional activator GAL4 - the 'driver line'. The latter must be crossed with a transgenic fly line containing a gene region of interest fused to an Upstream Activation Sequence (UAS), to which the GAL4 protein specifically binds, and is known as the 'responder line'. GAL4 only activates transcription when bound to its UAS promoter sequence. For example, an NSC-GAL4 'driver line' will only drive the expression of transgenes under the control of an Upstream Activation Sequence (UAS) promoter, such as UAS-Green Fluorescent Protein (GFP). Thus, when the transgenic driver and responder flies are crossed, their progeny will express the gene sequence of interest

(GFP) in the tissue pattern directed by the desired promoter (NSCs). In my studies, I took advantage of *Drosophila* as a model and extensively used the UAS/GAL4 system in the assays performed.

1.1.1. Drosophila embryonic NBs generate the functional larval CNS

NSCs in Drosophila are termed neuroblasts (NBs), the progenitors to the CNS neurons and glia. Drosophila NBs divide asymmetrically to self-renew and generate further committed progeny. Like mammalian neurogenesis, Drosophila neurogenesis is controlled by intrinsic programs and extrinsic stimuli (Doe, 2017; Harding and White, 2017). Yet, the Drosophila CNS offers a simpler model system to investigate conserved mechanisms. Neurogenesis in Drosophila occurs embryonically producing the larval CNS and ~ 10% of the adult fly CNS, and postembryonically producing the majority of the adult fly CNS (Prokop and Technau, 1991; Truman and Bate, 1988; Doe, 2017; Harding and White, 2017). The two neurogenic waves are separated by a period of cellular quiescence (Figure 1.3.A) (White and Kankel, 1978; Truman and Bate, 1988; Maurange et al., 2008).

Depending on their location, NBs are referred to as being thoracic, abdominal, central brain (CB) and optic lobe (OL). During embryogenesis, Drosophila NBs delaminate from the neuroectodermal cells (Bossing et al., 1996; Egger et al., 2007). OL NBs are generated only postembry onically (Doe, 2017; Harding and White, 2017). Delamination and NB identity are specified by Notch-mediated lateral inhibition, members of the Sox family (SoxN and Dicheate) and the Snail family (wornio, escargot and snail) and subsequent expression of proneural genes (Ashraf, 2001; Harding and White, 2017). Neural diversity is achieved by spatial patterning and temporal patterning (Miyares and Lee, 2019; Doe, 2017; Harding and White, 2017). Spatial determination along the anterior-posterior axis is established by Hox genes: abdominal-A and ultrabithorax are differentially expressed between NBs to generate diverse neurons or glia (Prokop and Technau, 1994). The temporal transcription factors Hunchback (Hb), Kruppel (Kr), Pou domain 2 (Pdm), Castor (Cas), and Grainyhead (Grh) are sequentially expressed in NBs and their progeny to give rise to distinct cell types (Kambadur et al., 1998; Doe, 2017; Harding and White, 2017).

The CNS is generated by around 1200 NBs found in the embryo (Bossing *et al.*, 1996; Schmidt et al., 1997; Urbach et al., 2003; Wheeler et al., 2009; Birkholz et al., 2015). To expand the embryonic CNS, each newly born embryonic NB begin limited rounds of asymmetric divisions, self-renewing and generating a smaller terminally differentiating ganglion mother cell (GMC). The NBs shrink with each division. Type 0 NBs do not produce a GMC (see below section 1.4.2.) and are an exception (Karcavich and Doe, 2005; Ulvklo et al., 2012; Baumgardt et al., 2014). The GMCs produce neurons required by the larval CNS. Each NB generates a stereotyped family of neural progeny to the CNS (Bossing et al., 1996). The post-mitotic neurons generated display a three-layered structure whereby Hb is detected in the deepest layer and Cas in the superficial layer, reminiscent of the developing mammalian cortex (Grieg et al., 2013). Grh is the last factor expressed and persists into larval NBs, stimulating cell cycle progression and cell survival (Cenci and Gould, 2005). The NBs were reported to exit the cell cycle at around stage 16 of embryogenesis (Figure 1.3.B) either by programmed cell death (Maurange and Gould, 2005; Ulvklo et al., 2012) or by entering cellular quiescence (Truman and Bate, 1988; Chell and Brand, 2010; Sousa-Nunes et al., 2011). Recently, two types of NB quiescence, G2 and G0, were reported (Otsuki and Brand, 2018). Furthermore, at embryonic stage 17, thoracic NBs display both G2 and G0 quiescence, whereas abdominal NBs enter G0 quiescence or apoptose (Harding and White, 2019).



Figure 1.3. *Drosophila* neurogenesis and proliferation patterns of NBs. A. *Drosophila* neurogenesis occurs embryonically to generate the larval CNS, and postembryonically throughout the larval stages. Embryonic neuroblast (NB) size decreases at each division throughout embryonic neurogenesis, and a temporal transcriptional program determines if embryonic NBs enter apoptosis (Figure 1.3. A, 'A') or enter quiescence (Figure 1.3. A, 'Q'); except for the mushroom body NBs (B, orange). A nutritional stimulus stimulates quiescent NBs to enlarge and re-enter the cell cycle to proliferate and generate the adult fly CNS. **B.** *Drosophila* larval brain depicting the postembryonic NBs (pNB) populations and their locations. **C.** The proliferation patterns of postembryonic NBs (pNB) and their lineages. Most pNBs follow the Type I pattern of division, present in the brain lobes (BL), ventral nerve cord (VNC) and Optic lobes (OL). A small population follow the Type II pattern of division and are only located in the central brain region. Type 0 is utilised by NBs in the outer proliferative centre of the OL. Hunchback (Hb), Kruppel (Kr), Pou domain 2 (Pdm), Castor (Cas), Grainyhead (Grh); Quiescent (Q); Apoptosis (A); Reactivated (R); Ganglion Mother Cell (GMC); Intermediate Neural Progenitor (INP). Schemes of larval brain depicted by arrows: A, anterior; P, posterior; D, dorsal; V, ventral., (Adapted from Hakes and Brand, 2019; Doe, 2017; Harding and White, 2017).

1.4.2. Proliferation patterns and lineages of Drosophila NBs

Three types of *Drosophila* NBs are identifiable based on their mode of division: Type 0, Type I and Type II (**Figure 1.3.C**). Type I NBs are prominent within the embryo, however, unlike embryonic NBs, type I postembryonic NBs (pNBs) maintain their original size after divisions. Embryonic NBs have been shown to switch from Type I divisions to Type 0 divisions (Doe, 2008; Baumgardt *et al.*, 2009; Baumgardt *et al.*, 2014; Karcavich and Doe, 2005; Monedero Cobeta *et al.*, 2017. Type 0 NB division produces a smaller terminally differentiating cell to generate a single neuron. Type 0 divisions occur during embryogenesis and postembryonically in the optic lobes (Baumgardt *et al.*, 2014; Bertet *et al.*, 2014; Mora *et al.*, 2018). Interestingly, the optic lobe NBs switch from Type 0 to Type I divisions which, like VNC pNBs, is under the control of a distinct temporal transcriptional program (Bertet *et al.*, 2014).

The two main types of *Drosophila* pNBs are type I and type II. Both types I and II pNBs divide asymmetrically and unequally segregate cell fate determinants but differ in their lineage pattern development (Homem and Knoblich, 2012; Doe, 2017; Harding and White, 2017). Type I is the most common mode of division; Type I NBs generate a terminally differentiating GMC, giving rise to a pair of sibling neural cells (neurons or glia). Type I pNBs are the most abundant, found throughout the larval brain lobes (mammalian hemispheres equivalent) and VNC (mammalian spinal cord; **Figure 1.3.B, C**). A subset of NBs located in the BLs called the mushroom body NBs (mbNBs) also follow the Type I division pattern (**Figure 1.3.B, C**). Type I pNBs are frequently distinguished by the self-renewal markers Deadpan (Dpn) and Asense (Ase), as well as Inscuteable (Insc). Their GMCs express nuclear Prospero (Pros), a pro-neural homeobox protein, which inhibits cell cycle genes (Choksi *et al.,* 2006; Homem and Knoblich, 2012; Doe, 2017; Harding and White, 2017). The GMC progeny, a pair of neurons or glia, can be distinguished by the neuronal marker Elav or the glia marker Reverse polarity (Repo), respectively.

Type II pNBs generate an intermediate neural progenitor (INP) which, upon maturation, undergoes limited rounds of division to self-renew and generate a GMC, giving rise to a pair of sibling neural cells (Homem and Knoblich 2012; Harding and White, 2017). There are less Type II pNBs, around ~8 per BL compared to ~90 for
Type I pNBs. Type II NBs are restricted to the central brain (CB; **Figure 1.3.B, C**) and generate higher numbers of neural progeny due to the amplifying INPs. Type II pNBs in the CB originate from the embryo whereby they specifically acquire expression of the transcription factor, Pointed P1 (Pnt), mediated by EGF signalling (Walsh and Doe, 2017; Álvarez and Diaz-Benjumea, 2018). Pnt is absent in Type I pNBs and is therefore used to distinguish the Type II pNBs and their lineages. In addition, unlike the Type I pNBs, Type II pNBs do not express Ase but do express the transcription factor/ self-renewal marker Dpn. Newborn immature INPs are smaller and do not express the transcription factors Ase, Dpn or Pros until they have 'matured' after 4-6 hours; at which point the mature INP will re-express those markers in that sequence (Bayraktar *et al.,* 2010). In addition, mature INPs express the transcription factor, Earmuff, which the Type I and II pNBs do not (Weng *et al.,* 2010). The mature INPs divide asymmetrically to generate a terminally differentiating GMC. INPs have less proliferative potential than pNBs but serve as an amplification step, like mammalian TACs (Bello *et al.,* 2008; Boone & Doe, 2008; Bowman *et al.,* 2008).

1.4.3. Asymmetric division in *Drosophila* NBs

Asymmetric division produces two cells with distinct fates (recently reviewed by Loyer and Januschke, 2020). Asymmetric division in Drosophila NBs is controlled by cellintrinsic cues and requires subcellular localisation and segregation of specific cell fate determinants. Precise spindle orientation along the apical-basal axis is required to separate the apical cortex, retained by the NB, from the basal cortex, transmitted to the progeny. The process is mediated by the apical Par, Insc, atypical protein kinase C (aPKC) complexes (Sousa-Nunes and Somers, 2013; Yu et al., 2006). The apicalbasal polarity is inherited during embryogenesis from the neuroectoderm; the apical Par complex (Par3/Bazooka, Par6 and aPKC) is retained by the NB (Petronczki and Knoblich, 2001; Wodarz et al., 2005; Wodarz et al., 2000; Wodarz et al., 1999; Bello et al., 2006; Betschinger et al., 2006; Ikeshima- Kataoka et al., 1997; Knoblich et al., 1995; Lee et al., 2006; Rhyu et al., 1994). The Par complex recruits the adaptor protein Insc, which subsequently recruits the Partner of inscuteable (Pins), Mushroom body defect (Mud), and the Drosophila Gai to form a Pins/Mud/ Gai complex. The apical complexes undergo a phosphorylation cascade to direct cell fate determinants Brain tumour (Brat), Numb and Pros to the basal cortex of the dividing NB via the scaffolding proteins Miranda and Partner of Numb (Pon). These cell fate determinants

segregate into the progeny (GMC), inhibit self-renewal and promote differentiation (Doe *et al.*, 1991; Rhyu *et al.*, 1994; Hirata *et al.*, 1995; Knoblich *et al.*, 1995; Bello *et al.*, 2006; Betschinger *et al.*, 2006; Lee *et al.*, 2006; Doe, 2017; Harding and White, 2017). Miranda sequesters and prevents Pros activity in the NB, however within the GMC Miranda is degraded, and Pros can enter the nucleus to stimulate differentiation (Ikeshima-Kataoka *et al.*, 1997; Hirata *et al.*, 1995). Interestingly, cell cycle gene repression has been observed by Prox1, the mammalian ortholog of Pros (Foskolou *et al.*, 2013; Dyer, 2003). Incorrect spindle orientation and failure in proper unequal segregation of cell fate determinants can lead to excess progeny with stem-cell characteristics and even tumour formation (Choksi *et al.*, 2006; Shen *et al.*, 1997; Bello *et al.*, 2006; Betschinger *et al.*, 2000). Also, loss of Insc function leads to Type I progeny with INP-like characteristics (An *et al.*, 2017).

1.4.4. Drosophila NBs enter quiescence at the end of embryogenesis

At the end of embryogenesis, after the embryonic lineages are generated, NBs switch from Type I to Type 0 divisions and many *Drosophila* NBs enter cellular quiescence (Tsuji *et al.*, 2008; Lai and Doe, 2014; Baumgardt *et al.*, 2014; Karlsson *et al.*, 2010; Monedero Cobeta *et al.*, 2017). These quiescent NBs are poised to mitotically reactivate postembryonically ~24 hours during the first instar larval stage to produce the neurons and glia required by the adult fly CNS. The only exceptions are the four mbNBs and one lateral NB which do not enter quiescence and will continually proliferate throughout development (Ito and Hotta, 1992).

As previously mentioned, quiescence is historically described as a G0 arrested state. However, recently *Drosophila* NB quiescence was shown to contain a mix of G2 and G0 quiescent pNBs (Otsuki and Brand, 2018) using cell cycle markers and the Fluorescent Ubiquitination-based Cell Cycle Indicator (FUCCI) system, which is a set of fluorescent probes enabling visualisation of the cell cycle (Sakaue-Sawano *et al.,* 2008). Here, the authors demonstrate ~75% of pNBs express Dpn, cyclin A, cyclin B and are arrested in G2. The remaining ~25% of pNBs express Dpn only and are arrested in G0 (Otsuki and Brand, 2018). Furthermore, the pNBs in G2 are mitotically reactivated faster than their G0 counterparts. The pseudokinase, *tribbles*, known to inhibit cell cycle progression by degrading Cdc25/String, regulates G2 quiescence by inhibiting the AKT pathway (Otsuki and Brand, 2018). Tribbles expression in NBs begins at embryonic stage 15 and continues to mark quiescence until nutritional cues stimulate NB reactivation; it is not expressed in the mbNBs that do not undergo quiescence (Otsuki and Brand, 2018). It is the first gene identified that distinguishes quiescent pNBs in *Drosophila* (Otsuki and Brand, 2018). Tribbles expression continues under starved conditions, and expressing constitutively active AKT, but not Pi3K, in *tribble* mutants could fully rescue the defects observed in *tribble* mutant pNBs (Otsuki and Brand, 2018). In addition, they showed that tribble activity is normally inhibited by AKT upon the nutritionally-dependant secretion of *Drosophila* Insulin-like peptides (dilps) from the niche glia (Otsuki and Brand, 2018).

NB entry into quiescence is determined by numerous factors, including the Hox genes Antennapedia (Antp) and Abdominal-A (Abd-A). At the end of embryogenesis, Antp and Abd-A determine NB cell cycle exit and whether the NB will enter quiescence or apoptosis (Bello *et al.*, 2003; Tsuji *et al.*, 2008; Arya *et al.*, 2015; Khandelwal *et al.*, 2017; Otsuki and Brand, 2018; Harding and White, 2019). Interestingly, the Polycomb Repressor Complex 2 (PRC2 complex) is known to suppress Hox gene expression, counteracting their known anti-proliferating action and regulation of programmed cell death in the early *Drosophila* larval brain (Curt *et al.*, 2019; Yaghmaeian Salmani *et al.*, 2018; Bahrampour *et al.*, 2019). PRC2 mutants display anterior expansion of Hox gene expression, resulting in reduced NB proliferation in the brain (Curt *et al.*, 2019; Yaghmaeian Salmani *et al.*, 2018). It would be interesting to examine if PRC2 may also be involved in Hox gene regulation of NB entry into quiescence.

The temporal transcription factor cascade (**Figure 1.3.A**) determines NB fate; although the majority of NBs enter programmed cell death, and a smaller subset of Cas+ NBs enter quiescence (Tsuji *et al*, 2008; Miyares and Lee, 2019; Doe, 2017; Harding and White, 2018). It was recently found that a small subset of NBs belonging to the G2-quiescence category are terminated through an unknown mechanism (Harding and White, 2018).

Downstream of the temporal transcription cascade, Nab and Squeeze (also transcription factors) were found to be required for entry into quiescence (Tsuji *et al.,* 2008). The homeodomain transcription factor Pros, usually associated with

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neuronal differentiation was found to regulate NB quiescence and act as a 'binary switch' between NB and differentiated progeny (Choksi *et al.,* 2006; Lai and Doe, 2014). Lai and Doe showed that different Pros levels determine various NB fates (Lai and Doe, 2014). A transient pulse of low-level Pros in the NB nucleus induces quiescence, but how this is achieved or the regulation by upstream temporal transcription factors is unknown (Lai and Doe, 2014).

1.4.5. Maintenance of Drosophila NB quiescence

NB quiescence must be properly maintained to ensure the appropriate NB lineages are only generated at the correct developmental time. Maintenance of quiescence is regulated by a variety of factors, including intrinsic and extrinsic. Glia cells surrounding the pNBs secrete a glycoprotein, anachronism (Ana), to promote quiescence in pNBs and prevent premature cell-cycle re-entry (Ebens *et al.*, 1993). The *Drosophila* Perlecan, Terribly reduced optic lobes (Trol), promotes cell-cycle progression by either inhibiting *ana* activity or functioning downstream of *ana* (Datta 1995; Voigt *et al.*, 2002). Fragile X protein (FMRP), which codes for the inherited mental retardation Fragile X syndrome, is reported to function first in pNBs and then in glia to maintain pNB quiescence and is regulated by Insulin signalling (Callan *et al.*, 2010; Callan *et al.*, 2011; Monyak *et al.*, 2016).

Maintenance of pNB quiescence is further regulated by Hippo signalling (Ding *et al.*, 2016; Poon *et al.*, 2016). Hippo signalling is a growth inhibition pathway and highly conserved in *Drosophila* and vertebrates (Harvey and Tapon, 2007; Halder and Johnson, 2011; Lee *et al.*, 2018). Active canonical Hippo signalling activates a phosphorylation cascade whereby the Hippo kinase activates the Warts kinase, which phosphorylates Yorkie (Yap/Taz in mammals) via a 14-3-3 binding site; sequestering the transcriptional co-activator Yorkie in the cytoplasm causes its deactivation (Oh and Irvine, 2008). Inactive Hippo signalling allows Yorkie translocation to the nucleus where it activates the transcriptional program, promoting growth and proliferation (Saucedo and Edgar, 2007). Ding *et al.*, found that the Hippo pathway maintains pNB quiescence; the core kinases, Hippo and Warts, are phosphorylated (activated) in quiescent pNBs (Ding *et al.*, 2016). Also, quiescent pNBs and non-quiescent pNBs (the MBNBs)

can be identified by cytoplasmic-Yorkie (inactive), and nuclear-Yorkie (active), respectively. Loss of Hippo/Warts or upstream regulators caused premature pNB enlargement and mitotic reactivation from quiescence (Ding *et al.*, 2016). This process is mediated by the upstream glial transmembrane proteins, Echinoid and Crumbs, which lead to inhibition of Yorkie activity but are downregulated by nutrition to enable mitotic reactivation (Ding *et al.*, 2016; Poon *et al.*, 2016). Premature loss of Crumbs or Echinoid in NBs or glia resulted in earlier reactivation shown by NB enlargement at 4h ALH (Ding *et al.*, 2016). The Hippo pathway was also shown to regulate larval brain size and cell cycle speeds (Poon *et al.*, 2016). Here, altering Hippo pathway components caused increased NB size, precocious NB proliferation, faster cell cycle speeds, and clone size; leading to larval brain overgrowth (Poon *et al.*, 2016).

In vertebrates, deactivation of YAP stimulates proliferation of stem cells (Tremblay and Camargo, 2012) and the Hippo pathway is implicated in stem cells within the liver (Zhou *et al.*, 2009); the skin (Schlegelmilch *et al.*, 2011; Zhang *et al.*, 2011); and the intestines (Barry *et al.*, 2013; Zhou *et al.*, 2011; Staley *et al.*, 2010).

1.4.6. Drosophila NBs mitotically reactivate to generate the adult fly CNS

The exit from NB quiescence, termed reactivation, requires the coordination of multiple signalling pathways, in which the pNB niche glia cells has been shown to be crucial. Anti-proliferative Hox genes regulating patterning and quiescence must be inactivated (Prokop *et al.*, 1998; Tsuji *et al.*, 2008; Harding and White, 2019; Curt *et al.*, 2019); which is controlled by brain transcription factors and the PRC2 complex (Curt *et al.*, 2019; Monedero Cobeta *et al.*, 2017; Yaghmaeian Salmani et al., 2018). Reactivation of pNBs occurs in a 'wave' along the anterior to posterior (AP) axis (Doe, 1992; Skeath and Carroll, 1992; Skeath, 1992, Skeath, 1999; Tsuji et al., 2008; Kemelman et al., 2012; Rosenberg et al., 2009), beginning in the BL NBs first, and propagating through the VNC; the abdominal NBs are the last to reactivate. Anterior propagation, governed by Hox genes, is a requirement for proper CNS development and appears to be conserved in mammals (Monedero Cobeta *et al.*, 2017; Philippidou and Dasen, 2013; Yaghmaeian Salmani *et al.*, 2018; Metzis *et al.*, 2018).

Upon reactivation, pNBs grossly enlarge, enter S-phase and begin proliferating to generate the adult CNS. This provides a unique opportunity to study pNB mitotic reactivation using a simpler model. Reactivated pNBs can be initially distinguished

from their quiescent counterparts by their larger size; an increase from ~4-5 µm (quiescent) up to ~10-15 µm (Chell and Brand, 2010). Most pNBs reactivate within 24 hours of larval hatching and reactivation is initiated by a nutritional stimulus following larval feeding (Figure 1.4). Larval feeding increases the amount of circulating amino acids which is sensed by the fat body (FB, the Drosophila liver equivalent, Figure **1.4**). This hepatic/adipose-like tissue was found to release a FB-derived signal (FDS) which enables pNB reactivation (Britton and Edgar 1998). Larval CNSs cultured alone or co-cultured with a gut or CNS from fed larvae failed to reactivate the NBs. Only when quiescent larval CNSs were co-cultured with a FB, were the NBs able to reactivate (Britton and Edgar, 1998). Further work revealed that the FB senses nutrition via the cationic amino-acid transporter Slimfast (slif) (Colombani, Raisin et al., 2003; Géminard et al., 2009), which activates the Target of Rapamycin (TOR) signalling pathway (Sousa-Nunes et al., 2011). Upon TOR activation, the FDS is released and sensed by glia cells surrounding the NBs. The glia cells release Insulinlike peptides (ILPs, totalling 7) to which the single NB Insulin-like Receptor (InR) responds, activating the phosphatidylinositol 3-kinase (PI3K) and TOR pathway within the NB. As a consequence, NBs characteristically enlarge and re-enter the cell cycle (Truman and Bate 1988; Chell and Brand 2010; Sousa-Nunes et al., 2011).

Of the 7 ILPs, ILP6 was found to be the most influenced by nutrition and a potent activator of NBs' reactivation (Sousa-Nunes *et al.*, 2011). Global inactivation of Tor, inhibition of slif within the FB, or FB specific activation of TOR inhibitors Tuberous sclerosis complex 1 and 2 (Tcs1/2) reduced the number of NBs exiting quiescence. Furthermore, inhibition of PI3K signalling also inhibits neuroblast reactivation. Conversely, overexpression of the TOR pathway activator Ras homologue enriched in brain (RHEB) and of PI3K pathway activators specifically in the NBs caused premature pNB reactivation (Chell and Brand, 2010; Sousa-Nunes *et al.*, 2011).

More recently, the highly conserved heat shock protein 83 (Hsp83/ Hsp90 in mammals) and its co-chaperon Cdc37 were found to function upstream of the InR/PI3K/AKT cascade to promote pNB reactivation; Hsp83 was shown to directly bind to the InR (Huang and Wang, 2018) (**Figure 1.4**). Insulin signalling and insulin-like growth factors (IGF) are highly conserved between invertebrates and mammals, regulating metabolism and growth. In mammals, dietary restriction or a high-fat diet causes increased or decreased levels of Sirtuin 1 (SIRT1), respectively (Cohen *et al.,*

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2004; Chen *et al.*, 2008; Spéder and Brand, 2011). SIRT1 correlates to the nutritional status of the animal and modulates systemic IGF-1 (Cohen *et al.*, 2004; Cohen *et al.*, 2009; Cohen *et al.*, 2010). Like *Drosophila* glia cells, mammalian astrocytes have also been shown to release local insulin/IGF-1 (Garcia-Estrada *et al.*, 1992; Shetty *et al.*, 2005), however, unlike in *Drosophila*, a direct link between these glia and NSC reactivation has not yet been shown.

As described in section **1.3.2**, nutritional dependant neurogenesis has also been observed in rodents (Lee *et al.*, 2000; Poulose *et al.*, 2017; Shohayeb *et al.*, 2018; Norman *et al.*, 2008; Stranahan *et al.*, 2008; Alvarez *et al.*, 2009), and associated to cognitive decline in human patients (Spinelli *et al.*, 2019; Kodl and Seaquist 2008). In the *Drosophila* larval CNS, Spéder and Brand (2014) discovered a cell-autonomous role of the gap junction proteins within the glia forming the Blood-Brain Barrier (BBB, **Figure 1.4**). They found that the nutrient dependant ILP secretions essential for pNB reactivation are coordinated by gap junction proteins (Innexin 1/2) across the BBB and conditions that reduce/block Ca2+ oscillations impair pNB reactivation (Spéder and Brand 2014). Similar mechanisms are also seen in vertebrates, whereby the pancreatic beta cells release insulin in response to Ca2+ (MacDonald and Rorsman 2006).

The glia niche performs additional functions for pNB reactivation. Cortex glia has been shown to progressively remodel their membranes around each NB and its lineage in response to local and systemic signals (Spéder and Brand, 2018). The cortex glia undergoes three developmental steps: membrane expansion, encasing, and extension, creating cortex glia 'chambers' around pNBs (Spéder and Brand, 2018). A nutritional signal first induces expansion of the glia membranes via Pi3K/Akt, and the reactivating NBs signal for the remodelling process; the new glia niche structure is critical for the new neurons to survive and inhibiting pNB reactivation disrupts chamber formation. In addition, knocking down BBB glia gap junctions, or Insulin signalling in the cortex glia or pNBs, or the spindle matrix complex component, *east*, in pNBs, all impairs the chamber formation (Spéder and Brand, 2018). Similarly, mammalian astrocytes, and the local neurogenic niche, support neurogenesis (Gengatharan *et al.*, 2016; Goldman and Chen, 2011; Tavazoie *et al.*, 2008; Ottone *et al.*, 2014).

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The *Drosophila* Perlecan homolog, Terribly reduced optic lobes (Trol), functions noncell-autonomously to promote pNB reactivation, by antagonising Ana activity in niche glia (Ebens *et al.*, 1993; Datta, 1995; Voigt *et al.*, 2002). Trol also performs structural and function roles in FGF-2 signalling and Hedgehog signalling to stimulate pNB reactivation at the G1-S-phase transition (Park *et al.*, 2003). Thought to function similar to the mammalian Perlecan, Trol is proposed to function in the *Drosophila* extracellular matrix, modulating extrinsic signalling factors such as Ana, to stimulate pNB reactivation (Voigt *et al.*, 2002) (**Figure 1.4**). Furthermore, Trol-dependant action is enhanced by a transacting signal produced by *even-skipped* (Eve); a regulator of embryonic neurogenesis (Park *et al.*, 2001; Akam, 1987).

Also downstream of nutritional signals and InR/PI3K/AKT signalling, the spindle matrix protein complex, comprised of Chromator (Chro), Megator, Skeletor and Enhanced adult sensory threshold (East), stimulate pNB reactivation (**Figure 1.4**). Chro is critical for Insulin signalling detection by pNBs and maintaining pNB proliferation (Li *et al.*, 2017). Grainyhead (Grh) prevents pNB nuclear Pros to maintain proliferation in the VNC (Maurange *et al.*, 2008). Interestingly, Chro was also found to repress *pros* expression and stimulate *grh* expression, which further represses *pros*, to promote pNB reactivation (Li *et al.*, 2017). Chro and East have been found to modulate gene expression via chromatin modification (Rhee *et al.*, 2014). In pNBs, the spindle matrix complex prevents re-entry into cellular quiescence by targeting Grh and *pros* (Li *et al.*, 2017).

An additional level of cell-autonomous regulation exists via *taranis* (*tara*), the mammalian SERTAD orthologue. Tara promotes the E2F1 pathway in pNBs to promote pNB reactivation; *tara* is expressed in mitotically active type I pNBs and absent in quiescent pNBs (Manansala *et al.*, 2013). More recently, a conserved ubiquitin ligase complex was shown to be essential for pNB reactivation (Ly *et al.*, 2019). Knocking down the complex components with mutations or by RNAi in pNBs impaired reactivation (Ly *et al.*, 2019). Here, the conserved ubiquitin ligase family, Cullin4-RING (CRL4) complex consisting of the adaptor damaged DNA-binding protein 1 (DDB1), the scaffold Cullin4 (Cul4), and the catalytic RING of Cullin (ROC), together with its substrate receptor Mahjong (Mahj; CRL4^{Mahj}) form a protein complex with Warts, causing warts ubiquitination and inhibition in S2 cells (Ly *et al.*, 2019).

Simultaneously expressing Warts-RNAi could suppress the reactivation defects observed upon Mahj-RNAi in pNBs (Ly *et al.*, 2019). Interestingly, rat Cul4B loss in neural progenitors leads to G2/M phase cell cycle arrest, and mutations of the human Cul4B gene is often associated with mental impairments (Tarpey *et al.*, 2007; Zou *et al.*, 2007; Liu *et al.*, 2012).



Figure 1.4. A simplified summary of main regulators of Drosophila pNB reactivation. Drosophila pNB reactivation is stimulated by larval feeding. Circulating amino acids are sensed by the Slimfast amino acid transporter (Slif) (Colombani et al., 2003; Géminard et al., 2009), on the fat body (FB). This activates Target of Rapamycin (TOR) in the FB (Sousa-Nunes et al., 2011), stimulating the release of a fat body derived signal (FDS) (Britton and Edgar, 1998), which is sensed by the glia surrounding the NBs. The glia release Insulin-like peptides (ILPs). This release also requires coordinated Ca2+ oscillations across the gap junctions of the Blood-Brain Barrier (BBB) glia (Spéder and Brand 2014). Local ILP secretion is sensed by the NBs' InR, activating PI3K and TOR pathway within the NB (Chell and Brand 2010; Sousa-Nunes; Yee et al., 2011). In contrast, the Hippo pathway functions to maintain NBs in quiescence, and therefore must be switched off for reactivation to occur (Ding et al., 2016; Poon et al., 2016). Phosphorylation and activation or Hippo and Warts leads to the inactivation of Yorkie, which is sequestered in the cytoplasm and degraded. Upstream regulators are the transmembrane proteins, Echinoid and Crumbs which stimulate the pathway until a nutritional stimulus inhibits their action. Inactivation of the Hippo pathway allows Yorkie to translocate to the nucleus and activate growth targets. Spindle matrix proteins Chromator (chro), Megator and Enhanced adult sensory threshold (East) (Li et al., 2017; Song et al., 2017), FMRP, Tribbles, and Ana (Ebens et al., 1993; Datta, 1995; Voigt et al., 2002) inhibit pNB reactivation whereas Heat shock protein 83 (Hsp83) (Huang and Wang, 2018), Taranis (Manansala et al., 2013) and Trol (Ebens et al., 1993; Datta, 1995; Voigt et al., 2002) promote pNB reactivation.

1.4.7. Drosophila Mob4, a potential new player in NB reactivation

To enhance our knowledge of postembryonic neurogenesis and the process of pNB reactivation, a single-cell transcriptome microarray screen was performed in our laboratory comparing Drosophila quiescent versus reactivating pNBs (Gill-Renado et al., 2019). The single cells were obtained directly from live transgenic Drosophila larval brains, in which pNBs were visualised by membrane-tagged Green Fluorescent Protein (GFP) driven by the grainy-head (grh) Gal4 driver (Ding et al., 2016; Chell and Brand, 2010). pNBs were removed from the second and third thoracic segments of the VNCs to minimize potential differences from spatial positioning and avoiding retrieving a mix of type I and II NSCs, as the latter are not present in VNCs. The pNBs were identified by their size as quiescent pNBs are much smaller, measuring 4-5 µm (maximum cell diameter), than reactivating (enlarging) pNBs (Chell and Brand, 2010; Ding et al., 2016). The pNBs were harvested at 17h after larval hatching (ALH) as, at this particular age, both quiescent and reactivating pNBs can be distinguished and simultaneously harvested. Only reactivating (enlarged) pNBs not in division and without any progeny were harvested. cDNA was obtained from each cell using an established single-cell transcriptome protocol (Liu and Bossing, 2016; Bossing et al., 2012).

Expression of the NB markers *dpn* and *ase* was confirmed via real-time quantitative PCR (RT-qPCRs) in both quiescent and reactivating cell samples harvested, with levels being higher in the later, as expected. Single pNB samples were compared in pairs (quiescent versus reactivating; n=3 pairs) on whole-genome microarrays (FlyChip, Cambridge). The screen analysis revealed 145 transcripts up-regulated and 51 transcripts down-regulated (p<0.05) in reactivating compared to quiescent NBs. 90% of identified targets have human orthologues, and most (66%) show high orthology conservation levels. The data was independently validated for a subset of candidates, and differential expression for all targets tested (18 in total) was confirmed (Gil-Ranedo *et al.*, 2019).

Among the candidates identified as potential novel regulators of postembryonic pNB reactivation were genes encoding for several members of the highly conserved Striatin Interacting Phosphatase and Kinases complex (STRIPAK) protein complex (Shi *et al.*, 2016; Ribeiro *et al.*, 2010; Zheng *et al.*, 2017): MOB kinase activator 4, also known as Monopolar spindle-one binder (Mob4), the only *Drosophila* Striatin Connector of kinase to AP-1(Cka) and Microtubule Star (Mts), which is the catalytic subunit of Protein Phosphatase 2A (PP2A). *mob4 and cka* were found upregulated, whereas *mts* was downregulated. Interestingly the STRIPAK complex, through its PP2A phosphatase, has been shown to be is an inhibitor of the Hippo pathway (Goudreault *et al.*, 2009; Ribeiro *et al.*, 2010), a known regulator of pNB quiescence (Ding *et al.*, 2016; Poon *et al.*, 2016). Yet, no member of STRIPAK had been reported to be involved in the process of pNB reactivation.

The above mentioned data and literature, the availability of cellular and genetic tools (e.g. transgenic *Drosophila* lines and antibodies) and preliminary observations in our laboratory (Dr C. Barros) suggesting that the STRIPAK member Mob4 may be involved in pNB reactivation, formed the basis for my PhD project focused primarily on investigating this candidate gene and its mode of action.

In addition, to be a STRIPAK member, Mob4 belongs to the Mob family of noncatalytic, kinase interacting proteins highly conserved from yeast to humans (Luca and Winey 1998; Lai *et al.*, 2005; Mrkobrada *et al.*, 2006; Wei *et al.*, 2007). The two founding members of the Mob protein family, first identified in *Saccharomyces* *cerevisiae* (Mob1 and Mob2), are involved in regulation and activation of Nuclear dbf2-related (NDR) kinases, which are cell cycle regulators (Komarnitsky *et al.*, 1998; Luca and Winey 1998; Lee *et al.*, 2001; Mah *et al.*, 2001; Wei *et al.*, 2007; Lai *et al.*, 2005; Mrkobrada *et al.*, 2006). Mob4, however, is the most divergent member of its family (Schulte *et al.*, 2010). On the other hand, RNA interference knocking down Mob4 in *Drosophila* embryonic cells (Schneider 2 cells, S2) showed a role in mitotic spindle assembly, specifically in fibre spindle focusing, and phenotypes similar to centrosomal protein mutants were reported (Trammell *et al.*, 2008). Furthermore, *invivo* analysis of *Drosophila mob4* null mutants showed abnormal synaptic development with a supernumerary bouton phenotype, disrupted axonal transport, disorganised microtubules and larval lethality (Schulte *et al.*, 2010). Interestingly, the lethality observed was rescued by ubiquitous expression of the human orthologue of Mob4, MOB4 (also called Phocein), which is over 80% identical at the amino acid level, highlighting also functional evolutionary conservation among the molecules (Schulte *et al.*, 2010) (see also Discussion).

1.5. Hypothesis and objectives of proposed PhD study

The hypothesis underlying my PhD project is that the candidate gene *mob4*, identified in a transcriptome screen performed in our laboratory as upregulated in reactivating compared to quiescent *Drosophila* postembryonic NSCs, is required for NSC mitotic reactivation. The overall aim is to characterize the expression and function of Mob4 towards identifying its potential mechanism of action regulating NSC mitotic reactivation.

The specific objectives of my PhD project are:

- 1. Characterise Mob4 expression in quiescent and reactivated postembryonic NBs
- 2. Perform functional analysis of Mob4 in *Drosophila* pNBs during mitotic reactivation.
- 3. Identify a potential mechanism of Mob4 action on pNB reactivation.

CHAPTER 2

RESULTS

2.1 Mob4 is upregulated in reactivated versus quiescent pNBs

mob4 transcript was found upregulated in reactivating versus guiescent pNBs in a single-cell transcriptome analysis performed in our laboratory, and the result was confirmed by RT-qPCR (see Introduction). Thus, my first step was to analyse Mob4 protein expression in the larval CNS, specifically in pNBs upon reactivation. The UAS-GAL4 system (Brand and Perrimon, 1993; Duffy, 2002) was used to visualise the larval pNBs. Membrane-bound GFP (CD8-GFP) was expressed using grainyhead-GAL4 (grh-Gal4), specifically highlighting ~one-third of all pNBs in the larval CNS (Chell and Brand, 2010). *elav-Gal⁸⁰* was simultaneously employed to repress any potential mild GFP expression in neurons driven by the *grh* promotor (Bray *et al.*, 1989), which are of similar size to quiescent pNBs. Larval brains (grh-Gal4, UCD8-GFP; elav-Gal⁸⁰) were dissected and immunostained first for Mob4 (red) and Green Fluorescent Protein (GFP, green) at 22h ALH. Consistent with previous observations of *mob4* mRNA and protein localisation in the Drosophila CNS (Schulte et al., 2010), Mob4 was observed ubiquitously in the brain lobes (BLs, Figure 2.1. A-A') and the ventral nerve cords (VNCs, **Figure 2.1. B-B**'), including within the cytoplasm of pNBs (white arrowheads) and their lineages (Figure 2.1. A-B'). Larval brains of the same genotype were immunostained with Mob4 (red, also shown separately in grey), GFP (green), and the NB marker Deadpan (Dpn, blue) to assess Mob4 protein levels in reactivated versus quiescent pNBs, at 18h ALH. Mob4 protein levels in reactivated versus quiescent pNBs were compared by measuring the Mob4 antibody signal intensity (average pixel intensity per cell area; see Methods). pNBs were determined to be quiescent (Figure **2.10.** E) or reactivating (Figure 2.10. F) by their size, i.e. quiescent cells: 4-5 µm (maximum diameter), reactivating cells: enlarged (approximately 7 µm) (Chell and Brand, 2010; Ding et al., 2016; Truman and Bate, 1988). The results show that Mob4 protein expression is significantly higher (p<0.01) in reactivating pNBs (**Figure 2.10**. E, F) compared to quiescent pNBs (Figure 2.10.G), in line with the microarray and RT- qPCR observations of our group.

Glia cells are vital transducers of the Insulin signal to reactivate pNBs (Chell and

Brand, 2010; Sousa-Nunes *et al.*, 2011). Since Mob4 is present ubiquitously in the brain, 24h ALH wild-type (WT) larvae were also immunostained for Mob4 (red, also shown in grey), Dpn (green) and Reversed polarity (Repo, a nuclear pan-glia marker, blue) antibodies to investigate expression in Glia. Mob4 seems to localise to the cytoplasm of glia (yellow arrowheads (**Figure 2.1. C-D'**) as inferred by absence from the nucleus, but its expression appears weaker compared to neighbouring pNBs (white arrows, **Figure 2.1.D'**).

In conclusion, the above assays show that Mob4 is abundant throughout the larval CNS and significantly upregulated in reactivating versus quiescent pNBs.



Figure 2.1. Mob4 protein is upregulated upon pNB mitotic reactivation: (A, A') *grh-Gal4, UCD8-GFP; elav-Gal⁸⁰* larval brain lobe (BL) and (B-B') Ventral Nerve Cord (VNC) at 22 hours after larval hatching (ALH) immunostained for Mob4 and Green Fluorescent Protein (GFP). Mob4 channel also shown in monochrome. (C-D') Wild-type (WT) VNCs were immunostained for Mob4, Deadpan (Dpn, NB marker) and Repo (glia nuclear marker) at 24h ALH. Mob4 channel also shown in monochrome. Mob4 is expressed ubiquitously throughout the BLs and VNCs and is found mainly in the cells' cytoplasm, including in pNBs (white arrows and dashed lines) and their progeny (dashed lines only) and glia (yellow arrowheads). (E-F) To quantify Mob4 expression levels in quiescent and reactivating pNBs, we used the same *Drosophila* line shown in monochrome. Mob4 signal intensity in small quiescent pNBs (E) is significantly less than in enlarged reactivating pNBs (50 cells, 8 brains). Values are normalised average intensities +/- s.e.m. Wilcoxon rank-sum test ** *p*<0.01. All images are single focal planes, anterior up. Scale bar: 10 μ m.

2.2 pNBs in mob4 mutants fail to mitotically reactivate

Confident that Mob4 is upregulated in pNBs upon their reactivation, pNBs in *mob4* null mutants were next investigated. Previously in another lab, *mob4*^{EYAL3} null mutants (herein *mob4* mutants) were generated via a P transposable element excision screen (Schulte et al., 2010). The P-element localised 33 base pairs upstream of the mob4 start codon. In the *mob4*^{EYAL3} line, the excision resulted in a 357 base pair deletion including the sequence encoding the initiator methionine. Western blots on larval brain lysates made from these mutants show no Mob4 protein, which would be expected at ~25kDa weight (Schulte *et al.*, 2010). Homozygous *mob4*^{Δ L3} null mutants are larval lethal, but ~10% of larvae can survive to the third instar larval stage (Schulte et al., 2010). To evaluate pNB reactivation in *mob4*^{ΔL3} mutants, pNBs' cell size (maximum) diameter) and mitotic index were quantified and compared to Wild-type (WT) controls, using Dpn, phospho-HistoneH3 (Ph3) labelling mitosis and Discs-large marking cell membrane antibodies. At 1h ALH, there are no significant differences in pNB size or divisions (Figure 2.11). The proliferating NBs detected and scored correspond to the four mushroom body NBs (mbNBs, white asterisk) and one ventrolateral NB, which do not enter quiescence (Figure 2.11. A, B; Ito, K. & Hotta, 1992). These NBs remain proliferative from embryonic throughout larval stages development and are included in the proliferation scores. Overall at 1h ALH, the pNBs in mob4^{AL3} mutants are quiescent similar to those in WT controls.



Figure 2.2. pNBs are quiescent in both mob4 mutants and controls immediately after larval hatching: (A, D) Wild-type (WT, control) and (B, E) mob4^{EYAL3} (herein mob4 ^{AL3}) mutant larval brain lobes (BLs) and ventral nerve cords (VNCs) at 1h after larval hatching (ALH), immunostained for Deadpan (Dpn), Discs large (Dlg) and Phospho-Histone-H3 (Ph3). (A-F) All pNBs in mutants and controls (arrowheads) are quiescent (small size and not mitotic), except mushroom body pNBs (white asterisks). (C) Quantification of pNB diameters in BLs and VNCs shows that pNBs in mob4^{ΔL3} mutants are not statistically different compared to controls at 1h ALH (Control BL: median 4.5 µm, average 5.19 μm n⁼³⁵⁹/10 BLs, 5 brains; mutant BL: median 4.5 μm, average 5 μm n^{= 441}/10 BLs, 7 brains; Control VNC: median 3.6 µm, average 3.64 µm n⁼³²¹/7 VNCs; mutant VNC: median 3.7 µm, average 3.73 µm $n^{=171}/5$ VNCs). (F) Quantification of pNB divisions in BLs and VNCs in *mob4*^{Δ L3} mutants also shows no statistical differences compared to controls at 1h ALH (Control BL: median 2, average 1.75 Dpn/Ph3+ cells n⁼²⁴/24 BLs, 12 brains; mutant BL: median 2, average 1.65 Dpn/Ph3+ cells n⁼²⁰/20 BLs, 12 brains). There are no pNB divisions in the VNCs at this age. In this figure and throughout this thesis, the box and whisker plots show the data outliers (symbols above/below the boxplot); the minimum value (excluding outliers, lower whisker); the lower 25% quartile (box start); the average (dashed line); the median (solid line); upper 25% quartile (box end); the maximum value (excluding outliers, upper whisker). p=Wilcoxon rank-sum test. n.s non-significant p>0.05. All images are single focal planes, anterior up. Scale bar: 10 µm.

By 4h ALH, *mob4*^{Δ L3} mutant pNB diameters in BLs and VNCs are smaller compared to WT controls (**Figure 2.3. C**). There were no differences in pNB divisions, and the proliferating cells observed correspond to the mbNBs, and ventrolateral NB described earlier (**Figure 2.3. F**).



Figure 2.3. pNBs in *mob4* mutants do not begin to enlarge in contrast to control pNBs: (A, D) Wild-type (WT, control) and (B, E) *mob4*^{ΔL3} mutant larval brain lobes (BLs) and ventral nerve cords (VNCs) at 4h after larval hatching (ALH), immunostained for **Deadpan (Dpn)**, **Discs large (Dlg)** and **Phospho-Histone-H3 (Ph3)**. (C) Quantification of pNB diameters in BLs and VNCs shows that pNBs in *mob4*^{ΔL3} mutants are statistically smaller compared to controls at 4h ALH (Control BL: median 5.3 µm, average 5.5 µm n⁼¹¹²¹/14 BLs, 7 brains; *mob4*^{ΔL3} mutant BL: median 4.4 µm, average 4.8 µm n^{= 552}/10 BLs, 5 brains; Control VNC: median 4.4 µm, average 4.4 µm n⁼⁷⁰⁰/6 VNCs; *mob4*^{ΔL3} mutant VNC: median 3.8 µm, average 3.8 µm n⁼³³¹/5 VNCs). (F) Quantification of pNB divisions in BLs and VNCs in *mob4*^{ΔL3} mutants show no statistical difference compared to controls at 4h ALH (Control: median 1, average 1.3 Dpn/Ph3+ cells n⁼²⁴ 12 brains; mutant: median 0, average 0.7 Dpn/Ph3+ cells n^{= 24} 12 brains). The only pNBs dividing at this stage are mushroom body pNBs (mbNBs, white asterisks). There are no pNB divisions in the VNCs at this age. *p*=Wilcoxon rank-sum test. *** *p*<0.001. n.s *p*>0.05. Arrowheads indicate pNBs. All images are single focal planes, anterior up. Scale bar: 10 µm.

At 24h ALH, the end of the first instar larval stage, the majority of pNBs have exited quiescence (Truman and Bate, 1988; Chell and Brand, 2010). As expected, the pNBs in the WT controls incorporate 5-ethynyl-2'-deoxyuridine (EdU) indicating entry into S-phase (DNA synthesis) of the cell cycle (**Figure 2.4. A, D**). However, pNBs in $mob4^{\Delta L3}$ mutants do not incorporate EdU, except the mbNBs (**Figure 2.4. B, E**). The $mob4^{\Delta L3}$ mutant BLs and VNCs are also dramatically smaller compared to the WT controls (**Figure 2.5. A, D** versus **Figure 2.5. B, E**).



Figure 2.4. *mob4* pNBs do not enter S-phase: (A, D) Wild-type (WT, control) and (B, E) $mob4^{\Delta L3}$ mutant larval brain lobes (BLs) and ventral nerve cords (VNCs) at 24h after larval hatching (ALH), labelled with 5-ethynyl-2'-deoxyuridine (EdU). (C) Normalised EdU voxel quantification in either BLs or VNCs shows that pNBs in $mob4^{\Delta L3}$ mutants do not incorporate EdU in contrast to controls, indicating no entry into S-phase of the cell cycle, with the exception of mushroom body NBs (Control BL: average $1\pm0.2 \text{ n}^{=8}/8$ brains; $mob4^{\Delta L3}$ mutant BL: average $0.03\pm0.01 \text{ n}^{=8}/8$ brains; Control VNC: average $1\pm0.15 \text{ n}^{=7}/7$ brains; $mob4^{\Delta L3}$ mutant BL: average $0 \text{ n}^{=8}/8$ brains). Normalised average scores +/- s.e.m. are shown. *p*=Wilcoxon rank-sum test. *** *p*<0.001. White dashed lines mark BLs and VNCs. All images are Z-stack projections. Anterior up. Scale bar: 10 µm.

In addition to failing to enter S-phase, pNB size (white arrows) and mitotic index (yellow arrows) are dramatically smaller in the $mob4^{\Delta L3}$ mutants compared to WT controls at 24h ALH (**Figure 2.5.C, F**). At 48h ALH, pNBs in the $mob4^{\Delta L3}$ mutants are still unable to re-enter division (**Figure 2.5.G-I**). Interestingly, a tiny increase in the average size of pNBs in $mob4^{\Delta L3}$ mutants is observed at 4h ALH (**Figure 2.3.C**), and 24h ALH (**Figure 2.5.C**) compared to 1h ALH (**Figure 2.3C**), suggesting that the mutant pNBs attempt to enlarge but are unable to succeed (see Discussion).



Figure 2.5. mob4 mutant pNBs fail to mitotically reactivate: (A, D, G, J) Wild-type (WT, control) and (B. E. H. K) $mob4^{\Delta L3}$ mutant larval brain lobes (BLs) and ventral nerve cords (VNCs) immunostained for Deadpan (Dpn), Discs large (Dlg) and Phospho-Histone-H3 (Ph3) at 24h ALH (A, B, D, E) and 48h ALH (G, H, J, K). (C) Quantification of pNB diameters in BLs and VNCs shows that pNBs in $mob4^{\Delta L3}$ mutants are smaller compared to controls at 24h After Larval Hatching (ALH) (Control BL: median 9.7 μm, average 9.5 μm n⁼⁷⁹⁹/10 BLs, 7 brains; *mob4*^{ΔL3} mutant BL: median 6.2 μm, average 6.5 μm n⁼ 395 /18 BLs, 9 brains; Control VNC: median 8.3 µm, average 8.3 µm n⁼⁵⁸⁴/5 VNCs; *mob4* $^{\Delta L3}$ mutant VNC: median 4.7 μ m, average 4.9 μ m n⁼⁸⁸/8 VNCs). (F) Quantification of pNB divisions in BLs and VNCs in $mob4^{\Delta L3}$ mutants shows no dividing pNBs compared to controls at 24h ALH, except mbNBs (Control BLs: median 25.5, average 26.8 Dpn/Ph3+ cells $n^{=24}$ 12 brains; *mob4* $^{\Delta L3}$ mutant BLs: median 2, average 1.7 Dpn/Ph3+ cells n^{= 24} 12 brains; Control VNCs: median 19, average 26.3 Dpn/Ph3+ cells n⁼¹²/12 VNCs; mob4 $\Delta L3$ mutant VNCs: median 0, average 0 Dpn/Ph3+ cells n⁼¹²/12 VNCs). (G-I) At 48h ALH, pNBs in mob4 mutants remain not reactivated with no pNB divisions seen in BLs (except mbNBs) or VNCs, in contrast to the controls (I; Control BLs: median 19, average 21.2 Dpn/Ph3+ cells $n^{=11}/11$ brains; *mob4*^{Δ L3} mutant BLs: median 1, average 1 Dpn/Ph3+ cells $n^{=11}/11$ brains; Control VNCs: median 30, average 31.3 Dpn/Ph3+ cells $n^{=11}/11$ VNCs; $mob4^{\Delta L3}$ mutant VNCs: median 0, average 0 Dpn/Ph3+ cells $n^{=11}/11$ VNCs). p=Wilcoxon rank-sum test. *** p<0.001. White arrowheads indicate pNBs; yellow arrowheads indicate Ph3+ pNBs. White asterisks label Ph3+ mbNBs. All images are single focal planes, anterior up. Scale bar: 10 µm.

Upon reactivation, type I pNBs divide asymmetrically to self-renew and generate GMCs which express the cell fate determinant *prospero* (*pros*) in their nuclei (reviewed in Jan and Jan, 1998). WT and *mob4*^{Δ L3} mutant CNSs were immunostained for Grainyhead (Grh, labelling NBs and GMCs, red), Pros (green) and Ph3 (blue) at 24h ALH. As expected, in WT, most of the pNBs in VNCs (type I) are reactivated and have generated GMCs (yellow arrows, **Figure 2.6. A**); contrasting with *mob4*^{Δ L3} mutant pNBs (white arrows, **Figure 2.6. B**). Together, the above results demonstrate that pNBs in *mob4*^{Δ L3} mutants fail to mitotically reactivate and generate progeny.



Figure 2.6. *mob4* pNBs fail to generate neural progeny: (A) Wild-type (WT, control) and (B) $mob4^{\Delta L3}$ mutant ventral nerve cords (VNCs) at 24h after larval hatching (ALH), labelled with Grainyhead (Grh), Prospero (Pros) and Phospho-Histone-H3 (Ph3) antibodies. (A) WT pNBs (Grh+, no nuclear Pros) have mitotically reactivated to generate Ganglion Mother Cells (GMCs, Grh/nuclear Pros+). (B) *mob4* mutant pNBs remain small, do not divide (no Ph3 signal) and no GMCs can be observed. White arrowheads indicate pNBs. Yellow arrowhead labels a GMC. All images are single focal planes, anterior up. Scale bar: 10 µm.

2.3 Mob4 functions cell-autonomously in pNB reactivation

Since Mob4 expression in the CNS is not restricted to pNBs, an investigation was carried out to discern if Mob4 acts in a cell-autonomous way in pNBs. The *mob4*^{ΔL3} mutant strain was recombined with an *inscuteable*-Gal4 (*insc*-Gal4) line (see Materials and methods). This fly line allowed for any UAS constructs to be expressed in pNBs in a *mob4*^{ΔL3} mutant background. To validate the efficacy of this fly line, Mob4 antibody (red) was applied to 18h ALH *mob4*^{ΔL3} mutants with ectopic Mob4 expression in the pNBs (*mob4*^{ΔL3}, *insc*-Gal4; UAS-*mob4*; herein *mob4*^{ΔL3}, NB>*mob4*; Figure 2.7. C, F), control brains (*insc*-Gal4 herein CNTRL, Figure 2.7. A, D) and *mob4* mutants recombined with insc-GAL4 (*mob4*^{ΔL3}, *insc*-Gal4) (Figure 2.7. B, E). The brains were also immunostained for Dpn (green) and Dlg (blue). As anticipated, Mob4 protein is ubiquitous in control (CNTRL) brains including pNBs (white arrows, Figure 2.7. A, D) and *mob4* mutants is absent in the *mob4*^{ΔL3}, *insc*-Gal4 mutant brains (white arrows, Figure 2.7. B, E). However, upon *mob4* ectopic expression in *mob4*^{ΔL3} mutant pNBs, Mob4 protein (red) can be readily seen in pNBs and their progeny (white arrows, Figure 2.7. C, F).



Figure 2.7. Validation of *Drosophila* line allowing targeted Mob4 ectopic expression in *mob4* **mutant pNBs:** Brain lobes (A-C) and ventral nerve cords (D-F) immunostained for Mob4, Deadpan (Dpn) and Discs large (Dlg) at 18h ALH. (A, D) Mob4 is expressed throughout the larval brain lobes and ventral nerve cords of *Insc-Gal4* brains (control, CNTRL) and is absent in *mob4*^{$\Delta L3$} mutants recombined with *Insc-Gal4* (*mob4* $^{\Delta L3}$, *insc-Gal4*) (B, E). However, Mob4 protein is expressed in the pNBs and their progeny of these mutants when *mob4* is ectopically driven by *insc-Gal4*. White arrowheads indicate pNBs. All images are single focal planes, anterior up. Scale bar: 10 µm.

Once validated, the *mob4*^{Δ L3}, NB>*mob4* flies were used for Mob4 rescue experiments (**Figure 2.8**.). First, it was determined if reintroducing Mob4 in pNBs of *mob4* mutants would rescue the observed pNB reactivation defects. Three fly strains were used for this experiment: *insc-Gal4* controls (CNTRLS, **Figure 2.8. A, D**); *mob4*^{Δ L3},*insc-Gal4* mutants (**Figure 2.8. B, E**) and *mob4*^{Δ L3},*insc-Gal4*; UAS-*mob4* (*mob4*^{Δ L3},NB>*mob4* Rescue, **Figure 2.8. C, F**). The brains were immunostained for Dpn (red), Dlg (green) and Ph3 (blue) to quantify pNB size and mitotic index. Predictably, at 18h ALH, many CNTRL pNBs are enlarged (white arrows, **Figure 2.8. A, D**) and proliferate (yellow arrows, **Figure 2.8. B, E**). Remarkably, ectopic expression of Mob4 in pNBs of

 $mob4^{\Delta L3}$ mutants can rescue pNB enlargement and re-entry into mitosis (yellow arrows, **Figure G, H**) to control values. The $mob4^{\Delta L3}$ mutant larval brains were small, whereas the pNB>mob4 rescue larval brains were more akin to the controls. These results demonstrate that re-introduction of Mob4 in pNBs is sufficient to rescue pNB reactivation defects in mob4 mutant brains.





Figure 2.8. Ectopic expression of Mob4 in pNBs of mob4 mutants rescues mitotic reactivation. (A, D) Insc-Gal4 (Control, CNTRL), (B, E) mob4^{ΔL3}, insc-Gal4 (mob4 mt) and (C, F) dmob4^{ΔL3}, insc-Gal4; UAS-mob4 (mob4^{ΔL3}, NB>mob4 Rescue) brain lobes (BLs) and ventral nerve cords (VNCs) immunostained for Deadpan (Dpn), Discs large (Dlg) and Phospho-Histone-H3 (Ph3), at 18h ALH. pNBs of mob4^{ΔL3} mutants have impaired enlargement and cannot re-enter division compared to controls. However, these defects can be rescued by ectopically expressing Mob4 in pNBs using the NB Gal4 driver inscuteable-Gal4 (insc-Gal4). (G) Quantification of pNB sizes (BL pNB size - Control: median 7.8 µm, average 7.7 µm n⁼²¹⁹/3 BLs, 3 brains; *mob4* mutant: median 4.9 µm, average 5 µm n⁼ ⁵²⁰/7 BLs, 7 brains; *mob4*^{Δ L3}, NB>*mob4* Rescue: median 7.8 µm, average 7.8 µm n^{= 669}/10 BLs, 5 brains. VNC pNB size – Control: median 6.9 µm, average 6.9 µm n⁼²⁵⁶/3 VNCs; *mob4* mutant: median 3.6 µm, average 3.6 μ m n⁼³⁷⁷/4 VNCs; *mob4^{ΔL3}*, NB>*mob4* Rescue: median 6.4 μ m, average 6.6 μ m n⁼⁶⁴⁹/7 VNCs). (H) Quantification of pNB divisions (BL pNB divisions - Control: median 7.5, average 9.4 Dpn/Ph3+ cells n⁼¹²/12 brains; *mob4* mutant: median 1, average 1.2 Dpn/Ph3+ cells n^{= 10}/10 brains; mob4^{ΔL3},NB>mob4 Rescue: median 8.5, average 10.4 Dpn/Ph3+ cells n^{= 10}/10 brains. VNC pNB divisions Control: median 7, average 7.2 Dpn/Ph3+ cells n=6/6 VNCs; mob4 mutant: median 0, average 0 Dpn/Ph3+ cells n⁼¹⁰/10 VNCs; *mob4*^{ΔL3}, NB>*mob4* Rescue: median 4, average 8.1 Dpn/Ph3+ cells $n^{=10}/10$ VNCs). *p*=Wilcoxon rank-sum test. n.s. *p*>0.05, * *p*<0.05, *** *p*<0.001. White arrowheads indicate pNBs; yellow arrowheads indicate Ph3+ pNBs. All images are single focal planes, anterior up. Scale bar: 10 µm.

Second, since the glial niche performs vital roles in pNB quiescence and reactivation (Chell and Brand, 2010; Sousa-Nunes et al., 2011; Ding et al., 2016), the next step was to perform rescue experiments by expressing Mob4 in the glia of $mob4^{\Delta L3}$ mutant brains. The *mob4*^{ΔL3} mutant allele was combined with *repo-Gal4* to allow ectopic Mob4 expression in CNS glia in a *mob4*^{Δ L3} mutant background (*mob4*^{Δ L3} mt, Glia>*mob4* rescue). At 18h ALH, the overall brain size of the *mob4* mutants expressing Mob4 in glia is comparable to the *mob4*^{Δ L3} mutants; both are much smaller than the controls Figure A-H). The pNBs in *mob4*^{∆L3} mutant brains (white arrows, Figure 2.9. D, E) and in the *mob4*^{AL3} mutants expressing Mob4 in glia (white arrows, **Figure 2.9. G, H**) are markedly smaller compared to the controls (white arrows, Figure 2.9. A, B). Quantifications confirmed that forcing Mob4 expression in the glia of *mob4*^{ΔL3} mutants cannot rescue pNB size (Figure 2.9. C, F) or mitotic activity defects (Figure 2.9. I, L) to the control values. However, a minimal yet significant increase is observed in both size and divisions when comparing brain lobe or VNC pNB sizes, as well as brain lobe pNB divisions, in the *mob4*^{Δ L3} mt, Glia>*mob4* expressing brains with *mob4*^{Δ L3} mutants (Figure 2.9. C, F, I, L). Overexpressing mob4 in glia in a WT background using repo-Gal4; UAS-mob4 (Glia>mob4) caused a tiny but significant VNC pNB size increase compared to the controls, at 18h ALH (**Figure 2.9. F**; see Discussion). However, analysis of the size of brain lobe pNBs (**Figure 2.9. C**), or pNB divisions in either brain lobes or VNCs show no significant differences (**Figure 2.9. I, L**) compared to controls.



Figure 2.9. Mob4 ectopic expression in glia is unable to rescue pNB reactivation in mob4 mutants and its overexpression in control brains causes a minor premature enlargement only in VNC pNBs: (A, B) repo-Gal4 (control, CNTRL), (D, E) mob4^{ΔL3} mutant (mob4^{ΔL3} mt), (G, H) mob4^{ΔL3}: repo-Gal4; UAS-mob4 (mob4^{ΔL3} mt, Glia>mob4 Rescue), (J, K) repo-Gal4; UAS-mob4 (Glia > mob4). Larval brain lobes (BLs) and ventral nerve cords (VNCs) are immunostained for Deadpan (Dpn), Discs large (DIg) and Phospho-Histone-H3 (Ph3), at 18 h After Larval Hatching (ALH). (C, F) Quantification of pNB diameters in BLs (C) and VNCs (F) shows that driving *mob4* in glia of *mob4*^{ΔL3} mutants does not rescue the pNB enlargement defects to control values. However, there is a small increase in pNB size when compared to mob4 mutant values. Also, mob4 overexpression in glia in control background does not affect the size of pNBs in BLs but leads to a small increase in VNC pNBs. (C) pNB size quantifications in BLs (Control: median 7.8 μm, average 7.9 μm n¹⁸⁸/3 BLs, 2 brains; *mob4*^{ΔL3}mt: median 5.1 µm, average 5.2 µm n⁼³⁰⁷ /5 BLs, 4 brains; $mob4^{\Delta L3}$ mt, Glia> mob4 Rescue: median 5.4 µm, average 5.5 μ m n^{= 310}/4 BLs, 3 brains; Glia> *mob4*: median 7.6 μ m, average 7.7 μ m n^{= 278}/4 BLs, 2 brains). (F) pNB size quantifications in VNCs (Control: median 6.3 µm, average 6.4 µm n⁼²⁷⁸/3 VNCs; *mob4*^{Δ L3} mt: median 3.9 µm, average 4 µm n⁼²⁰⁶/4 VNCs; *mob4*^{Δ L3} mt, Glia> *mob4* Rescue: median 2.8 μ m, average 4.3 μ m n⁼²⁵³/5 VNCs; Glia> *mob4*: median 6.5 μ m, average 6.7 μ m n⁼²³³/3 VNCs). (I, L) Quantification of pNB divisions (Dpn/Ph3+ cells) in BLs (I) and VNCs (L) shows that ectopic expression of mob4 in glia of mob4 mutants cannot rescue the mitotic defects to control values. However, there is a small increase in dividing pNBs in the BLs when compared to mob4 mutant values. mob4 overexpression in glia in control background does not affect pNB divisions compared to controls. (I) BL pNB division quantifications (Control: median 12, average 12.3 Dpn/Ph3+ cells n⁼²⁸ 28 BLs, 14 brains; *mob4*^{$\Delta L3$} mt: median 1, average 1.2 Dpn/Ph3+ cells n⁼³⁴ 17 brains; *mob4*^{$\Delta L3$} mt, Glia> *mob4*: median 3, average 3.7 Dpn/Ph3+ cells n⁼⁵⁷ 26 brains; Glia> mob4: median 11, average 11.8 Dpn/Ph3+ cells n⁼⁶⁶ 66 BLs, 33 brains). (L) VNC pNB division quantifications (Control: median 7, average 8.4 Dpn/Ph3+ cells n⁼¹⁴/14 VNCs; mob4^{Δ L3} mt: median 0, average 0.4 Dpn/Ph3+ cells n⁼¹⁷/17 VNCs; mob4^{Δ L3} mt, Glia> mob4: median 0, average 0.4 Dpn/Ph3+ cells n⁼²⁹/29 VNCs; Glia> mob4: median 8, average 8.8 Dpn/Ph3+ cells $n^{=33}/33$ VNCs). p=Wilcoxon rank-sum test. n.s (non-significant) p>0.05 * p<0.05, ***p<0.001. White arrowheads indicate pNBs; yellow arrowheads indicate Ph3+ pNBs. The white asterisk indicates mbNBs. All images are single focal planes, anterior up. Scale bar: 10 µm.

Finally, RNA interference was used to knockdown *mob4* expression, specifically in pNBs using *insc-Gal4* (**Figure 2.10**.). Membrane-tagged GFP was simultaneously expressed (UAS-CD8-GFP, *UCD8-GFP*). Larval brains with *mob4* knock-down in the pNBs (UAS-*dicer*; *insc-Gal4, UCD8-GFP*; UAS-*mob4*-RNAi) were dissected and immunostained for Dpn (red), GFP (green) and Ph3 (blue), and compared to the controls (UAS-*dicer*; *insc-Gal4, UCD8-GFP*, CNTRL), at 18h ALH. *mob4*-RNAi specifically in pNBs RNAi (NB>*mob4*-RNAi, **Figure 2.10. B, E**) leads to a small but significant reduction in pNB enlargement (**Figure 2.10. C**) compared to the controls

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(Figure 2.10. A, D). NB>*mob4*-RNAi pNBs also showed significantly reduced mitotic reactivation compared to the controls (yellow arrows, Figure 2.10. F). Together, the above results indicate that Mob4 acts primarily cell-autonomously in pNBs promoting reactivation.



Figure 2.10. Mob4 inhibition via RNAi delays pNBs mitotic reactivation: (A, D) UAS-dicer; insc-Gal4, UCD8-GFP; UAS-mob4-RNAi (NB>mob4-RNAi) brain lobes (BLs) and ventral nerve cords (VNCs) immunostained for **Deadpan** (Dpn), green fluorescent protein (GFP) and Phospho-Histone-H3 (Ph3), at 18h After Larval Hatching (ALH). (C) Quantification of pNB diameters in BLs and VNCs shows that pNBs with mob4-RNAi are smaller compared to controls (Control BL: median 6.8 µm, average 7 µm n⁼⁶⁸⁴/9 BLs, 9 brains; mob4-RNAi BL: median 6.3 µm, average 6.6 µm n^{= 614}/8 BLs, 8 brains; Control VNC: median 5.8 µm, average 6 µm n⁼⁴⁴¹/5 VNCs; mob4-RNAi VNC: median 5 µm, average 5.1 µm n⁼⁶⁸⁶/5 VNCs). (F) pNB divisions in BLs and VNCs in pNBs with mob4-RNAi are also statistically less compared to controls (Control BL: median 5, average 7.9 Dpn/Ph3+ cells n⁼²⁸ 14 brains; Control VNC: median 3, average 4.3 Dpn/Ph3+ cells n⁼²³/23 VNCs; mob4-RNAi VNC: median 1, average 1.8 Dpn/Ph3+ cells n⁼¹⁴/14 VNCs). *p*=Wilcoxon rank-sum test. *** *p*<0.001. Yellow arrowheads indicate Ph3+ pNBs. All images are single focal planes, anterior up. Scale bar: 10 µm.

2.4 Mob4 or human MOB4 (Phocein) overexpression leads to premature pNB reactivation

To strengthen the results indicating that Mob4 is required for pNB reactivation, the next step was to analyse the potential effects of Mob4 overexpression in pNBs. Mob4 was overexpressed using *insc-Gal4* during a period of when pNB are still mitotically quiescent (6h ALH) and when pNB reactivation is ongoing (18h ALH). Larval brains with Mob4 overexpressing pNBs (insc-Gal4, UCD8-GFP; UAS-mob4, NB>mob4) were dissected and immunostained for Dpn (red), GFP (green) and Ph3 (blue), and compared to controls (*insc*-Gal4, UCD8-GFP, CNTRL) (Figure 2.11). At 6h ALH, pNBs are of small size and only the mbNBs/vNB are mitotically active (white asterisk, Figure 2.11. A, B). In contrast, pNBs in NB>mob4 larval brains (white arrows, Figure 2.11. B, E) showed a significant increase in pNB size (Figure 2.11. C) compared to the controls (white arrows, Figure 2.11. A, D). However, there were no significant differences in pNB divisions between the two groups at this time point (Figure 2.11. F). At 18h ALH, overexpression of Mob4 (NB>*mob4*; Figure 2.11. H, K) did not cause a further increase in pNB diameters (Figure 2.11. I) compared to the controls (Figure 2.11. G, J), yet more pNBs in division were detected compared to the controls (yellow arrows, Figure 2.11. L). These experiments demonstrate that Mob4 overexpression leads to premature pNB reactivation.



Figure 2.11. pNBs overexpressing Mob4 show premature enlargement and entry into division: (A, D, G, J) insc-Gal4. UCD8-GFP (control; CNTRL) and (B, E, H, K) mob4 overexpression specifically in pNBs (insc-Gal4, UCD8-GFP; UAS-mob4) larval brain lobes (BLs) and ventral nerve cords (VNCs) immunostained for Deadpan (Dpn), green fluorescent protein (GFP) and Phospho-Histone-H3 (Ph3) and at 6h and 18h after larval hatching (ALH). (C) Quantification of pNB diameters in BLs and VNCs shows that pNBs with Mob4 overexpression (NB>mob4) enlarge prematurely compared to controls at 6h ALH (Control BL: median 4.6 µm, average 4.9 µm n⁼⁶⁶⁴/10 BLs, 5 brains; NB>*mob4* BL: median 5.3 μm, average 5.6 μm n^{= 478}/10 BLs, 5 brains; Control VNC: median 4.1 μm, average 4.1 μm n⁼⁶¹³/6 VNCs; NB>*mob4* VNC: median 4.6 μm, average 4.7 μm n⁼⁶⁵⁷/6 VNCs). (F) pNB divisions in BLs show no statistical difference between mob4 overexpression compared to controls at 6h ALH (Control BL: median 1, average 1.2 Dpn/Ph3+ cells n⁼²⁴ 12 brains; NB>mob4 BL: median 1.5, average 1.8 Dpn/Ph3+ cells n^{= 22} 11 brains). pNB division corresponds to only mbNBs. No Dpn/Ph3+ cells were observed in the VNCs of either group at this age. (I) At 18h ALH, pNB diameters in BLs and VNCs show no statistical differences (Control BL: median 7.1 µm, average 7.3 µm n⁼¹⁰⁵²/14 BLs, 7 brains; NB>mob4 BL: median 7.2 μm, average 7.3 μm n^{= 1004}/14 BLs, 9 brains; Control VNC: median 5.9 μm, average 6.1 μ m n⁼⁶⁵⁷/6 VNCs; NB>*mob4* VNC: median 6 μ m, average 6.1 μ m n⁼⁶³⁰/6 VNCs). (L) At 18h ALH overexpression of *mob4* in pNBs shows increased numbers of dividing pNBs compared to controls in both BLs and VNCs (Control BL: median 10, average 10.6 Dpn/Ph3+ cells n⁼²⁴ 12 brains; NB>mob4 BL: median 12, averaged 12.9 Dpn/Ph3+ cells n⁼²⁰ 10 brains; Control VNC: median 4, average 4.9 Dpn/Ph3+ cells n⁼¹⁰ 10 brains; NB>mob4 VNC: median 10, average 10.7 Dpn/Ph3+ cells n⁼¹⁵ 15 brains). p=Wilcoxon rank-sum test. *** p<0.001, p>0.05 ns (non-significant). White arrowheads indicate pNBs; yellow arrowheads indicate Ph3+ pNBs. Mushroom body pNBs (white asterisks). All images are single focal planes, anterior up. Scale bar: 10 µm.

Mob4 shares 80% amino acid homology with its human ortholog, MOB4 (also known as Phocein), and can rescue the lethality of *mob4* mutants (Schulte *et al.*, 2010). To assess whether Mob4 action in pNB reactivation may be evolutionary conserved, *phocein* was overexpressed in pNBs at 6h ALH and 18h ALH (**Figure 2.12**.). Larval brains with *phocein* overexpressing pNBs (*insc*-Gal4, *UCD8-GFP*; UAS-*phocein*, NB> *phocein*) were dissected and immunostained for Dpn (red), GFP (green) and Ph3 (blue), and compared to the controls (*insc*-Gal4, *UCD8-GFP*, CNTRL) (**Figure 2.12**.). At 6h ALH the pNBs are small with only the mbNBs/vNB enlarged and mitotically active (white asterisk, **Figure 2.12. A, B**). pNBs in NB>*phocein* larval brains (white arrows, **Figure 2.12. B, E**) showed a significant increase in pNB size and a small increase in divisions (**Figure 2.12. C, F**) compared to the controls (**Figure 2.12. H, K**) also displayed significant increases in pNB diameters and divisions (yellow arrows, **Figure 2.12. L**)

compared to the controls (yellow arrows, **Figure 2.12. G**, **J**). The results indicate that Phocein overexpression causes precocious pNB reactivation, similar to mob4 overexpression, which suggests a conserved function.



Figure 2.12. Overexpressing the human ortholog of mob4, phocein, results in premature pNB reactivation: (A, D, G, J) Insc-Gal4, UCD8-GFP (controls; CNTRL) and (B, E, H, K) Human MOB4 (Phocein) overexpression specifically in pNBs (Insc-Gal4, UCD8-GFP; UAS-phocein, NB>phocein) larval brain lobes (BLs) and ventral nerve cords (VNCs) immunostained for Deadpan (Dpn), green fluorescent protein (GFP) and Phospho-Histone-H3 (Ph3) at 6h and 18h after larval hatching (ALH). (C, F) Quantification of pNB diameters in BLs and VNCs shows that pNBs with *phocein* overexpression enlarge and start dividing prematurely compared to controls at 6h ALH. (C) pNB diameters quantifications at 6h ALH (Control BL: median 4.7 µm, average 5.1 µm n⁴⁶⁶/32 BLs, 16 brains; NB>*phocein* BL: median 5.7 µm, average 6.2 µm n^{= 508}/6 BLs, 3 brains; Control VNC: median 3.9 µm, average 4 μ m n⁼⁶⁸²/5 VNCs; NB>*phocein* VNC: median 5 μ m, average 5 μ m n⁼⁶⁵⁴/5 VNCs). (F) pNB divisions at 6h ALH (Control BL: median 1, average 1.5 Dpn/Ph3+ cells n⁼³² 16 brains; NB>*phocein* BL: median 2, average 2.5 Dpn/Ph3+ cells n^{= 28} 14 brains; Control VNC: median 0, average 3.2 Dpn/Ph3+ cells n⁼³² 12 VNCs; *phocein* VNC: median 2, average 2.5 Dpn/Ph3+ cells n^{= 28} 14 VNCs). (I, L) At 18h ALH pNBs with *phocein* overexpression also enlarge and mitotically reactivate prematurely compared to controls. (I) pNB diameters quantifications (Control BL: median 7.4 µm, average 7.6 µm n⁵²¹/7 BLs, 5 brains; NB>*phocein* BL: median 8.5 µm, average 8.6 µm n^{= 533}/7 BLs, 5 brains; Control VNC: median 6.8 µm, average 6.8 µm n⁼⁶⁹⁷/6 VNCs; NB>*phocein* VNC: median 7.3 µm, average 7.4 µm n⁼⁷¹⁰/6 VNC). (L) pNB divisions quantifications (Control BL: median 8.5, average 9.1 Dpn/Ph3+ cells $n^{-26}/13$ brains; NB>*phocein* BL: median 13, average 12.9 Dpn/Ph3+ cells n^{= 40}/20 brains; Control VNC: median 9.5, average 9 Dpn/Ph3+ cells n=13/13 brains; NB>phocein VNC: median 15.5, average 15.2 Dpn/Ph3+ cells n^{= 20}/ 20 brains). p=Wilcoxon rank-sum test. **p<0.01 ***p<0.001. White arrowheads indicate pNBs; yellow arrowheads indicate Ph3+ pNBs. Mushroom body pNBs (white asterisks). All images are single focal planes, anterior up. Scale bar: 10 µm.

Given that *mob4* overexpression in pNBs leads to premature reactivation, the next step was to determine if it also leads to overproliferation of pNBs later in development. Using the same fly strains for the *mob4* overexpression experiments, larval brains with *mob4* overexpressing pNBs (*insc*-Gal4, *UCD8-GFP*; UAS-*mob4*, NB>*mob4*) were dissected and immunostained for Dpn (red), GFP (green) and Ph3 (blue), and compared to the controls (*insc*-Gal4, *UCD8-GFP*, CNTRL) (**Figure 2.13**) at 48h and 94h ALH. pNBs in NB>*mob4* VNCs only (white arrows, **Figure 2.13. B, E**) showed a significant increase in pNB divisions (**Figure 2.13. C**) compared to the controls (white arrows, **Figure 2.13. A, D**) at 48h ALH, a time when pNB reactivation still occurs but is about to cease (White and Kankel, 1978; Truman and Bate, 1988; Maurange *et al.*, 2008). However, there were no significant differences in pNB divisions in NB>mob4 brain lobes or VNCs (white arrows, **Figure 2.13. H, K**) compared to the controls (white

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arrows, **Figure 2.13. G**, **J**) at 94h ALH (**Figure 2.13. I**). Therefore, mob4 overexpression in the pNBs leads to early reactivation but does not cause overproliferation.


Figure 2.13. Mob4 overexpression in pNBs does not cause overproliferation: (A, D, G, J) *Insc*-Gal4, *UCD8-GFP* (control; CNTRL) and (B, E, H, K) Mob4 overexpression specifically in pNBs (*Insc*-Gal4, *UCD8-GFP*; UAS-*mob4*, NB>*mob4*) larval brain lobes (BLs) and ventral nerve cords (VNCs) immunostained for Deadpan (Dpn), green fluorescent protein (GFP) and Phospho-Histone-H3 (Ph3) at 48 and 94h ALH. (C) At 48h ALH, pNB divisions in VNCs are higher in pNBs with mob4 over-expression (NB> *mob4*) compared to controls however there was not a difference in divisions in the BLs (Control BL: median 23.5, average 25.5 Dpn/Ph3+ cells n⁼¹⁰ 9 brains; NB> *mob4* BL: median 27.5, average 27.4 Dpn/Ph3+ cells n^{= 10} 10 brains; Control VNC: median 31.5, average 33.4 Dpn/Ph3+ cells n⁼⁸ 8 brains; NB> *mob4* VNC: median 49.5, average 52.1 Dpn/Ph3+ cells n^{= 10} 10 brains). (I) At 94h ALH pNB divisions in the BLs and VNCs are not statistically different from controls (Control BL: median 23.5, average 25.5 Dpn/Ph3+ cells n⁼¹⁰ 10 brains; NB> *mob4* BL: median 27.5, average 25.5 Dpn/Ph3+ cells n⁼¹⁰ 10 brains; NB> mob4 BL: median 27.5, average 25.5 Dpn/Ph3+ cells n⁼¹⁰ 10 brains; NB> *mob4* VNC: median 49.5, average 52.1 Dpn/Ph3+ cells n⁼¹⁰ 10 brains). (I) At 94h ALH pNB divisions in the BLs and VNCs are not statistically different from controls (Control BL: median 23.5, average 25.5 Dpn/Ph3+ cells n⁼¹⁰ 10 brains; NB> *mob4* BL: median 27.5, average 27.4 Dpn/Ph3+ cells n⁼¹⁰ 10 brains; NB> mob4 VNC: median 47, average 48.3 Dpn/Ph3+ cells n⁼¹⁰ 10 brains). *p*=Wilcoxon rank-sum test. *p*>0.05 n.s (non-significant), * *p*<0.05, ****p*<0.001. White arrowheads indicate pNBs. All images are single focal planes, anterior up. Scale bars; 25 µm.

2.5. Activation of the Insulin-like receptor (InR)/PI3K/Akt signalling cascade in pNBs of *mob4* mutants partially rescues reactivation defects

In the absence of nutrition, specifically circulating amino acids, CNS glia do not receive a yet to be defined systemic signal from the larval fat body, and consequently, do not release insulin-like peptides that activate the InR/TOR signalling cascade in pNBs promoting their reactivation. pNB reactivation can be restored by activation of the InR/TOR cascade within pNBs (Chell and Brand, 2010; Sousa-Nunes et al., 2011). As mob4 overexpression in pNBs was able to cause early reactivation, it was first determined if it could also induce pNB reactivation in the absence of the nutritional cue. mob4 was overexpressed in pNBs under nutrition-restrictive conditions (sucroseonly diet) (Figure 2.14). Nutritionally deprived larval brains with Mob4 overexpressing pNBs (insc-Gal4, UCD8-GFP; UAS-mob4, NB>mob4 N.R.) (Figure 2.14. B, E) or controls (*insc*-Gal4, UCD8-GFP, CNTRL N.R.) (Figure 2.14. A, D) were dissected and immunostained for Dpn (red), GFP (green) and Ph3 (blue) at 18h ALH. In the control larvae, pNBs fail to reactivate (Figure 2.14. A, B) and pNBs in NB>mob4 N.R larval brains (white arrows, **Figure 2.14. B**, **E**) showed a minor significant increase in pNB size in the VNC only (Figure 2.14. C) at 18h ALH. Nevertheless, there were no significant differences in pNB divisions in either brain lobes or VNCs (Figure 2.14. F). Overall, *mob4* overexpression in the pNBs is not sufficient to induce pNB mitotic reactivation upon loss of the extrinsic nutrition cue.



Figure 2.14. Overexpression of mob4 is not sufficient to induce pNB reactivation under nutritionrestriction conditions: (A, D) Insc-Gal4, UCD8-GFP (control; CNTRL) and (B, E) mob4 overexpression specifically in pNBs (Insc-Gal4, UCD8-GFP; UAS-mob4) upon nutrition restrictive conditions (sucrose-only diet). Larval brain lobes (BLs) and ventral nerve cords (VNCs) are immunostained for Deadpan (Dpn), green fluorescent protein (GFP) and Phospho-Histone-H3 (Ph3) at 18 hours After Larval Hatching (ALH). Quantification of pNB diameters in BLs shows that mob4 overexpression (pNB>mob4) is not sufficient to overcome reactivation defects observed under nutrition restriction, with exception of a minor increase in pNB sizes in VNCs. (C) pNB diameters quantification (Control BL: median 4.9 µm, average 5.3 µm n³⁹⁸/6 BLs, 5 brains; pNB>mob4 BL: median 4.7 µm, average 5.2 µm n^{= 376}/5 BLs, 5 brains; Control VNC: median 4.2 µm, average 4.2 µm n⁼³²⁵/5 VNCs; pNB>*mob4* VNC: median 4.2 µm, average 4.3 µm n⁼⁴⁴⁰/5 VNCs). Under nutrition restriction, pNBs with mob4 overexpression also did not significantly alter the number of dividing pNBS. (F) pNB divisions (Control BL: median 2, average 1.7 Dpn/Ph3+ cells n⁼⁵⁸ 29 brains; pNB>*mob4* BL: median 1, average 1.5 Dpn/Ph3+ cells n^{= 66} 33 brains). No Dpn/Ph3+ cells were detected in the VNCs at this age. p=Wilcoxon rank-sum test. *p<0.05, p>0.05 ns (non significant). White arrowheads indicate pNBs. Mushroom body pNBs (white asterisks) remain mitotically active. All images are single focal planes, anterior up. Scale bar: 10 µm.

Since active InR/PI3K/Akt signalling is essential for pNB reactivation, and pNBs in $mob4^{\Delta L3}$ mutants are unable to reactivate, it was investigated if activation of this cascade was impaired in *mob4* mutant brains. Using 24h ALH brain lysates from

control wild-type and mutants, I have analysed the levels of activated (phosphorylated) Akt, an essential component of the InR/PI3K/Akt cascade via western blot. Reduced levels of pAkt compared to total AKT were observed in *mob4*^{ΔL3} mutant brains compared to controls (Figure 2.15. A, B). This initial observation was then replicated in our laboratory by Dr Barros, confirming that *mob4*^{ΔL3} mutant brains have impaired insulin signalling as shown by reduced levels of p-Akt (Gil-Ranedo et al., 2019). Next, I tested whether activating the InR/PI3K/Akt cascade in pNBs of mob4 mutants could rescue reactivation defects. Stimulation of TOR signalling by RHEB overexpression activates the InR/PI3K/Akt cascade promoting NSC exit from quiescence (Sousa-Nunes *et al.*, 2011). *mob4*^{ΔL3} mutant larval brains with *rheb* overexpression in the pNBs (*mob4*^{ΔL3}, *insc*-Gal4; UAS-*rheb*, *mob4* $^{\Delta L3}$, *NB*>*rheb*), *mob4* $^{\Delta L3}$ mutants (*mob4* $^{\Delta L3}$) and the *insc- Gal4* controls (CNTRL) were dissected and immunostained for Dpn (red), Dlg (green) and Ph3 (blue) at 18h ALH (Figure 2.15. C-H). As observed before, the *mob4*^{ΔL3} pNBs (white arrow) are unable to reactivate (**Figure 2.15. D, G**). *mob4* $^{\Delta L3}$, NB>*rheb* pNBs show significantly increased pNB enlargement (Figure 2.15. E, H, I) and pNB divisions (Figure 2.15. J) compared to the mutants. However, the pNB>rheb rescue did not reach levels observed in controls (white arrows, Figure 2.15. C, F), and although VNC pNB size increased, the number of divisions detected was not significantly different (Figure **2.15.** I-J). Nevertheless, together these experiments reveal that activation of the InR/PI3K/Akt cascade can partially restore reactivation upon loss of Mob4 (see Discussion), and suggested that Mob4 may act in pNBs upstream or at the same level of Rheb.



Figure 2.15. Expression of the InR/PI3K/Akt signalling cascade activator Rheb in pNBs of mob4 mutants can rescue reactivation defects: (A, B) Western-blot test on mob4^{ΔL3} mutant brain lysate shows a decrease in phosphorylated-AKT (p-AKT) indicating reduced insulin signalling, compared to WT control larval brain lysate. Levels of total AKT remain very similar. β-Actin served as a loading control. Lysates are from 24 ALH brains (B) Quantification of protein signals (n=1 independent assay using 67 WT brains and 90 mob4 mutants). (C-H) pNBs in mob4 mutants fail to enlarge and divide; however, this defect can be rescued by overexpressing ras homolog enriched in brain (rheb) in pNBs (mob4^{ΔL3}, NB>rheb). (C, F) Insc-gal4 (control; CNTRL), (D, G) mob4^{ΔL3}, insc-Gal4 mutants, (E, H,) $mob4^{\Delta L3}$, insc-gal4; UAS-*rheb* ($mob4^{\Delta L3}$, NB>*rheb*) immunostained for **Deadpan** (**Dpn**), **Discs large** (DIg) and Phospho-Histone-H3 (Ph3) at 18 hours After Larval Hatching (ALH). (I) Quantification of the diameters in the BLs and VNCs shows that pNBs in mob4^{ΔL3} mutants overexpressing rheb were significantly larger compared to the mob4^{ΔL3} mutants, however still smaller compared to controls (Control BL: median 7.4 μm, average 7.5 μm n³³⁴/5 BLs, 4 brains; *mob4*^{ΔL3} mutants BL: median 5.2 μm, average 5.4 μ m n³⁵⁷/9 BLs, 9 brains; *mob4*^{ΔL3}, NB> *rheb* BL: median 6.4 μ m, average 6.5 μ m n^{= 466}/5 BLs, 5 brains; Control VNC: median 6.4 µm, average 6.5 µm n¹⁸¹/3 BLs, 3 brains; *mob4*^{ΔL3} mutants VNC: median 3.9 µm, average 4.1 µm n²⁵¹/4 BLs, 4 brains; *mob4*^{ΔL3}, NB> *rheb* VNC: median 5.2 µm, average 5.2 μ m n^{= 454}/5 BLs, 5 brains). (J) The mitotic index of the pNBs in *mob4*^{ΔL3} mutants expressing *rheb* was significantly higher in the BLs compared to that in *mob4*^{ΔL3} mutants, but the results were not statistically different in the VNCs. Dividing pNBs in $mob4^{\Delta L3}$ mutants expressing *rheb* were less compared to the controls (Control BL: median 11, average 10.9 Dpn/Ph3+ cells n⁼⁷ 7 brains; mob4^{ΔL3} mutants BL: median 1, average 0.8 Dpn/Ph3+ cells n⁼³² 16 brains; mob4^{ΔL3}, NB> rheb BL: median 6, average 5.9 Dpn/Ph3+ cells n⁼²⁸ 14 brains; Control VNC: median 11, average 10 Dpn/Ph3+ cells n⁼⁵ 5 brains; *mob4*^{Δ L3} mutants VNC: median 0, average 0 Dpn/Ph3+ cells n⁼¹⁶ 16 brains; *mob4*^{Δ L3}, NB> rheb VNC: median 0, average 1.1 Dpn/Ph3+ cells n=14 14 brains). p=Wilcoxon rank-sum test. p>0.05 ns (non- significant), ***p<0.001. White arrowheads indicate pNBs. All images are single focal planes, anterior up. Scale bar: 10 µm.

2.6. Deactivating the Hippo pathway in pNBs of *mob4* mutants can rescue reactivation defects

The Hippo signalling is another highly conserved pathway and has recently been shown to maintain pNB quiescence (Ding et al., 2016; Poon et al., 2016). As detailed in the Introduction, the canonical Hippo pathway prevents pNB cell growth via a phosphorylation cascade of the core components Hippo (Hpo), Warts (Wts), and the transcriptional co-activator Yorkie (Yki). In guiescent pNBs, the Hippo pathway is active, and Yki is phosphorylated and inactivated in the cytoplasm. However, when the Hippo pathway is deactivated, Yki translocates to the nucleus to activate cell growth. Interestingly, Mob4 has been shown to interact physically and genetically with Hpo: in the *Drosophila* wing, Mob4 knockdown can rescue the overgrowth phenotype caused by Hpo knockdown (Ribeiro et al., 2010). Also, the STRIPAK complex containing Protein Phosphatase 2 A (PP2A), to which Mob4 belongs to, has been shown to inhibit the Hippo pathway in both *Drosophila* and mammals (Liu et al., 2016; Goudreault et al., 2009; Couzens et al., 2013; Shi et al., 2016; Ribeiro et al., 2010; Straßburger et al., 2012; Tumaneng et al., 2012). Since pNBs upon loss of Mob4 are unable to reactivate, it was postulated that Hippo signalling might remain switchedon in pNBs of *mob4* mutant brains. Indeed, this was confirmed in our laboratory by Drs Barros and Gil-Ranedo. I have examined if deactivating Hippo signalling in pNBs of *mob4* mutant brains could rescue mitotic reactivation. Hpo and Wts were knocked down via RNAi in pNBs of *mob4*^{ΔL3} mutants (Figure 2.16.). The RNAi strains used had been previously shown to induce pNB premature reactivation in control pNBs (Ding 2016). Larval brains of Insc-Gal4 (controls; CNTRL, Figure 2.16. A-B), et al.. mob4^{Δ L3}, insc-gal4 mutants (mob Δ L3 mt, Figure 2.16. D-E), mob4^{Δ L3}, insc-gal4; UAShippo-RNAi (mob4^{ΔL3},NB>hpo-RNAi, Figure 2.16. G-H) and mob4^{ΔL3},insc-gal4; UASwarts-RNAi (mob4^{ΔL3}, NB>wts-RNAi, Figure 2.16. J-K) were immunostained for Dpn (red), Dlg (green) and Ph3 (blue) at 18h ALH. Quantifications showed a significant increase of pNB enlargement (white arrows, **Figure 2.16**.) and pNB divisions (yellow arrows, Figure 2.16) in mob4^{Δ L3}, NB>hpo-RNAi and mob4^{Δ L3}, NB>wts-RNAi pNBs compared to those in *mob4* mutants, although not to the control values (Figure 2.16. **C**, **F**). Therefore, deactivating the Hippo pathway within *mob4* mutant pNBs can rescue the pNB enlargement and division defects observed (see Discussion).



Figure 2.16. Deactivating the Hippo pathway in pNBs of mob4 mutants can rescue reactivation defects: (A, B) Insc-Gal4 (controls; CNTRL), (D, E) mob4^{ΔL3}, insc-gal4 mutants (mob4 ^{ΔL3} mt), (G, H) mob4^{ΔL3},insc-gal4; UAS-hippo-RNAi (mob4^{ΔL3}, NB>hpo-RNAi), (J, K) mob4^{ΔL3},insc-gal4; UAS-warts-RNAi (mob4^{ΔL3}, NB>wts-RNAi) larval brain lobes (BLs) and ventral nerve cords (VNCs) immunostained for Deadpan (Dpn), Discs large (Dlg) and Phospho-Histone-H3 (Ph3), at 18h After Larval Hatching (ALH). (C, F) Quantification of pNB diameters in BLs and VNCs shows that deactivating the Hippo pathway within mob4 mutant pNBs can rescue pNB enlargement defects observed, yet not to control levels. (C) BL pNB diameters quantifications (Control BL: median 7.6 µm, average 7.6 µm n²⁸⁹/3 BLs, 3 brains; *mob4*^{ΔL3} BL: median 5 μm, average 5.1 μm n^{= 380}/9 BLs, 9 brains; *mob4*^{ΔL3}, NB>*hpo*-RNAi BL: median 6.2 μ m, average 6.5 μ m n⁵³¹/ 8 BLs, 5 brains; *mob4^{ΔL3}*, NB>*wts*-RNAi BL: median 6 μ m, average 6.2 µm n⁴⁶⁶/8 BLs, 5 brains). (F) VNC pNB diameters quantifications (Control VNC: median 6.2 µm, average 6.3 µm n⁼⁴/4 VNCs; *mob4*^{Δ L3} VNC: median 3.8 µm, average 4 µm n⁼⁵/5 VNCs; *mob4*^{ΔL3}, NB>*hpo*-RNAi VNC: median 4.9 μm, average 5 μm n⁼⁵/5 VNCs; *mob4*^{ΔL3}, NB>*wts*-RNAi VNC: median 5.4 µm, average 5.4 µm n⁼⁵/5 VNCs). (I, L) Quantification of pNB mitosis in BLs and VNCs shows that deactivating the Hippo pathway in mob4 mutant pNBs leads to an increased in pNB divisions, except for VNC pNBs upon knock-down of hippo. However, in all cases, the rescue is partial, not reaching levels seen in controls (I) BL pNB division quantifications (Control BL: median 11, average 10.6 Dpn/Ph3+ cells n⁼¹¹ 8 brains; *mob4^{ΔL3}* BL: median 1, average 1.4 Dpn/Ph3+ cells n⁼³⁴ 17 brains; mob4^{ΔL3},NB>hpo-RNAi BL: median 2, average 2.5 Dpn/Ph3+ cells n⁼⁶⁰ 30 brains; mob4^{ΔL3}, NB>wts-RNAi BL: median 4.5, average 4.8 Dpn/Ph3+ cells n⁼²⁰ 10 brains). (L) VNC pNB division quantifications (Control VNC: median 10, average 10 Dpn/Ph3+ cells n⁼⁵ 5 VNCs; *mob4*^{ΔL3} VNC: median 0, average 0 Dpn/Ph3+ cells n⁼¹⁶ 16 VNCs; mob4^{ΔL3}, NB>hpo-RNAi VNC: median 0, average 0 Dpn/Ph3+ cells n⁼³⁰ 30 VNCs; mob4^{ΔL3},NB>wts-RNAi VNC: median 0, average 1 Dpn/Ph3+ cell n⁼¹⁰ 10 VNCs). p=Wilcoxon rank-sum test. n.s. (non-significant) p>0.05, * p<0.05, ***p<0.001. White arrowheads indicate pNBs; yellow arrowheads indicate Ph3+ pNBs. All images are single focal planes, anterior up. Scale bar: 10 μm.

2.7. Mob4 and another STRIPAK component, Cka, cooperate to promote pNB reactivation

Connector of Kinase to AP-1 (Cka) is another candidate gene which was found to be upregulated in reactivating versus quiescent pNBs on the transcriptome analysis performed in the laboratory. Data acquired previously by the group revealed that, like Mob4, Cka promotes pNB reactivation (Gil-Ranedo et al., 2019). As mentioned in the Introduction, Mob4 and Cka physically interact with each other and are core components of a STRIPAK protein complex that inhibits Hippo signalling (Ribeiro et al., 2010). Thus, the next aim addressed whether Mob and Cka may cooperate to promote pNB reactivation. Larval brains simultaneously overexpressing Mob4 and Cka within pNBs (Insc-Gal4, UCD8-GFP; UAS-cka, UAS-mob4, NB>cka+mob4) and controls (Insc-Gal4, UCD8-GFP; CNTRL) were immunostained for Dpn (red), GFP (green) and Ph3 (blue) at 6h ALH and 18h ALH (Figure 2.17). At 6h ALH, control pNBs are not mitotically active except the mbNBs/vNB (white asterisk, Figure 2.17. **A-B**). NB>*cka*+*mob4* pNBs (white arrows, **Figure 2.17. B, E**) were significantly larger (Figure 2.17. C) compared to the controls (white arrows, Figure 2.17. A, D). However, there were no significant differences in pNB divisions (Figure 2.17. F) at this stage. At 18h ALH, NB>cka+mob4 pNBs (Figure 2.17. H, K) were also significantly larger (Figure 2.17. I) and had more divisions (yellow arrows, Figure 2.17. L) compared to the controls (yellow arrows, Figure 2.17. G, J). The increase in pNB size and divisions is stronger when both Cka and Mob4 are overexpressed compared to the results observed upon overexpression of each alone (Figure 2.11 Gil-Ranedo 2019). These experiments suggest that Mob4 and Cka may cooperate to et al., promote pNB reactivation.



Figure 2.17. Simultaneously overexpressing Cka and Mob4 in pNBs causes stronger premature pNB reactivation than that previously seen with single Mob4 overexpression: (A, D, G, J) Insc-Gal4. UCD8-GFP (controls; CNTRL) and Insc-Gal4, UCD8-GFP; UAS-cka, UAS-mob4 (NB>cka+mob4) larval brain lobes (BLs) and ventral nerve cords (VNCs) immunostained for Deadpan (Dpn), green fluorescent protein (GFP) and Phospho-Histone-H3 (Ph3) at 6h and 18h ALH. (C, F, I, L) Quantification of pNB diameters in BLs and VNCs shows that simultaneous cka and mob4 overexpression specifically in pNBs causes earlier enlargement 6h and 18h ALH, and increased entry into mitosis at 18h ALH. (C) 6h ALH pNB diameters quantifications (Control BL: median 3.6 µm, average 3.9 μ m n³⁰⁶/3 BLs, 3 brains; NB>*cka+mob4* BL: median 4.3 μ m, average 4.8 μ m n^{= 374}/3 BLs, 3 brains; Control VNC: median 3.4 µm, average 3.4 µm n⁼³⁹³/3 VNCs; NB>cka+mob4 VNC: median 4.0 µm, average 4.1 µm n⁼³⁸⁷/3 VNCs). (F) 6h ALH pNB divisions (Control BL: median 1, average 1.1 Dpn/Ph3+ cells n⁼²⁴ 12 brains; NB>*cka+mob4* BL: median 1, average 1.3 Dpn/Ph3+ cells n^{= 32} 16 brains; Control VNC: median 0, average 0 Dpn/Ph3+ cells n=12 12 VNCs; NB>cka+mob4 VNC: median 2, average 2.5 Dpn/Ph3+ cells n^{= 16} 16 VNCs). (I) 18h ALH pNB diameters quantifications (Control BL: median 6.9 μm, average 7.0 µm n⁼³³⁹/4 BLs, 3 brains; NB>*cka*+*mob4* BL: median 8.1 µm, average 8.1 µm n⁼⁴⁰⁰/4 BLs, 4 brains; Control VNC: median 6.1 µm, average 6.2 µm n⁼³²³/3 VNCs; NB>*cka+mob4* VNC: median 6.8 μm, average 6.9 μm n⁼³⁹¹/4 VNCs). **(L)** 18h ALH pNB divisions (Control BL: median 11, average 10.2 Dpn/Ph3+ cells n⁼²⁸ 14 brains; NB>*cka*+*mob4* BL: median 17.5, average 16.9 Dpn/Ph3+ cells n^{= 28} 14 brains; Control VNC: median 8.5, average 8.6 Dpn/Ph3+ cells n⁼¹⁴ 14 VNCs; NB>*cka+mob4* VNC: median 13.0, average 14.4 Dpn/Ph3+ cells n^{= 14} 14 VNCs). p=Wilcoxon rank-sum test. n.s (nonsignificant) p>0.05, ***p<0.001. Overall, effects with double cka/mob4 overexpression are stronger than those obtained by overexpressing just mob4 (compare to Fig. 2.11). White arrowheads indicate pNBs; yellow arrowheads indicate Ph3+ pNBs. All images are single focal planes, anterior up. Scale bar: 10 μm.

2.8. PP2A inhibition promotes pNB reactivation

Together with Mob4 and Cka, the serine/threonine phosphatase PP2A forms part of a STRIPAK complex known to inhibit Hippo signalling (Couzens *et al.*, 2013; Ribeiro *et al.*, 2010). In *Drosophila*, the unique PP2A catalytic subunit is called Microtubule Star (Mts). STRIPAK components act as regulatory subunits of Mts, directing it to Hippo kinase, which leads to its de-phosphorylation and Hippo signalling inhibition. Since Mts was also detected in the transcriptome analysis comparing reactivating with quiescence pNBs performed in our group (Gil-Ranedo *et al.*, 2019), I next analysed pNB reactivation in *mts* trans-heterozygote mutants (*mts*²⁹⁹/*mts*^{XE-2258}; Wang *et al.*, 2009). The reason for using trans-heterozygotes was that *mts* null mutants (*mts*^{XE-2258}) die embryonically but survive until pupae stages if combined with a copy of a

hypomorphic allele (*mts*²⁹⁹; Wang *et al.*, 2009). WT control (CNTRL) and *mts*²⁹⁹/*mts*^{XE-2258} mutant larval brains were dissected and immunostained for Dpn (red), Dlg (green) and Ph3 (blue) at 18h ALH. *mts*²⁹⁹/*mts*^{XE-2258} pNBs (white arrows, **Figure 2.18. B, E**) were significantly larger (**Figure 2.18. C**) than the controls (white arrows, **Figure 2.18. A, D**) at 18h ALH. However, there were no significant differences in the number of pNB divisions in the BLs (**Figure 2.18. A-B**) or the VNCs (yellow arrows, **Figure 2.18. D-E**).



Figure 2.18. *mts* mutants display premature pNB enlargement: (A, D) Wild-type (control; CNTRL) and (B, E) *mts* mutant (*mts*^{299/XE2258}) larval brain lobes (BLs) and ventral nerve cords (VNCs) immunostained for **Deadpan (Dpn)**, **Discs large (Dlg)** and **Phospho-Histone-H3 (Ph3)** at 18h After Larval Hatching (ALH). Quantification of pNB diameters and divisions (Dpn/Ph3+) in BLs and VNCs shows that pNBs in mutants enlarge earlier, but no difference is seen in division numbers compared to controls. (C) pNB diameters quantification (Control BL: median 7.1 µm, average 7.2 µm n^{362/3} BLs, 3 brains *mts*^{299/XE2258} *mt* BL : median 7.9 µm, average 8.1 µm n^{= 235/4} BLs, 4 brains; Control VNC: median 6 µm, average 6.1 µm n⁼²⁰³/2 VNCs; *mts*^{299/XE2258} *mt* VNC: median 7.3 µm, average 7.3 µm n^{=189/2} VNCs). (F) pNB division quantification (Control BL: median 12, average 12.4 Dpn/Ph3+ cells n⁼²⁶ 13 brains; *mts*^{299/XE2258} *mt* BL: median 14, average 14.2 Dpn/Ph3+ cells n^{= 36} 18 brains; Control VNC: median 5, average 6.4 Dpn/Ph3+ cells n⁼¹³ 13 VNCs; *mts*^{299/XE2258} *mt* VNC: median 2.5, average 4.8 Dpn/Ph3+ cells n^{= 18} 18 VNCs). Wilcoxon rank-sum test. n.s (non-significant) *p*>0.05, ****p*<0.001. White arrowheads indicate pNBs. All images are single focal planes, anterior up. Scale bar: 10 µm.

Next, to explore PP2A/Mts function specifically in pNBs, a Dominant-Negative Mts Drosophila UAS line (UAS-mts-DN) was used (Hannus et al., 2002) together with the Inscuteable-Gal4 driver. The expression of Mts-DN results in a truncated Mts form lacking its phosphatase domain, which binds to the endogenous Mts protein and impairs its function (Hannus et al., 2002). Larval brains expressing mts dominantnegative (mts-DN) form specifically in pNBs (Insc-Gal4, UCD8-GFP; UAS-mts-DN; NB>mts-DN) and controls (Insc-Gal4, UCD8-GFP; CNTRL) were dissected and immunostained for Dpn (red), GFP (green) and Ph3 (blue) at 18h ALH (Figure 2.19. **A**, **C**, **D**, **F**). A significant increase in pNB size and divisions was observed compared to the controls (Figure 2.19. G, H). The results strengthen those obtained using the trans-heterozygote *mts* mutants and together indicate that knocking-down PP2A/Mts leads to premature exit from quiescence. The findings were surprising as they contrast with the effect of inhibiting Mob4 (Figures 2.2-2.6, 2.10) or Cka in pNBs (Gill-Renedo *et al.*, 2019), and suggested that PP2A/Mts contributes to maintaining pNBs in quiescence independently from a separate potential role in pNBs as part of STRIPAK (see Discussion).

PP2A is a pleiotropic phosphatase; in addition to a role within STRIPAK, multiple functions have been assigned to it. This variety is provided mainly by the association of the catalytic subunit Mts to possible variable regulatory subunits, which direct PP2A/Mts to different targets (Andreazza et al., 2015; Vereshchagina et al., 2008; Rodgers et al., 2011; Padmanabhan et al., 2009). To date, the Drosophila PP2A regulatory subunits better studied are Well rounded (Wrd), Twins and Widerborst (Wdb; Shi, 2009). Of significance, PP2A-Wdb has been shown to negatively regulate insulin signalling by dephosphorylation of Akt in both Drosophila and mammals (Fischer et al., 2015; Fischer et al., 2016). However, it had not been implicated in pNB quiescence/ reactivation processes. Thus, I next functionally tested Wdb using a Drosophila UAS strain expressing a truncated form of Wdb acting as Dominant Negative (*wdb-DN*; Hannus *et al.*, 2002). Knocking down Wdb (Figure 2.19. B, E) in pNBs leads to a significant increase in the pNB size (Figure 2.19. G) compared to the controls (Figure 2.18. A, D), at 18h ALH. In the BLs, Wdb knockdown in pNBs also lead to significantly more pNB divisions compared to the controls (Figure 2.19. A, B, H). Overall, knocking down PP2A subunits Mts or Wdb causes earlier pNB reactivation, suggesting that PP2A/Mts with its regulatory subunit Wbd acts to maintain

pNB quiescence (see Discussion).



Figure 2.19. Inhibition of the PP2A subunits Microtubule Star (Mts) or Widerborst (Wdb) cause premature pNB reactivation: Larval brain lobes (BLs) and ventral nerve cords (VNCs) of (A, D) Insc-Gal4, UCD8-GFP (control, CNTRL), (B, E) expressing a Wdb Dominant Negative form specifically in pNBs (Insc-Gal4, UCD8-GFP; UAS-wdb-DN; NB>wdb-DN) or (C, F) expressing an Mts Dominant Negative form specifically in pNBs (Insc-Gal4, UCD8-GFP; UAS-mts-DN; NB>mts-DN), immunostained for Deadpan (Dpn), green fluorescent protein (GFP) and Phospho-Histone-H3 (Ph3) at 18h After Larval Hatching (ALH). (G) Quantification of pNB diameters in BLs and VNCs shows that knocking down Wdb or Wdb in pNBs causes earlier pNB enlargement compared to controls. pNB enlargement quantifications (Control BL: median 6.8 µm, average 7.1 µm n⁵⁴¹/8 BLs, 5 brains; NB>wdb-DN BL: median 8.3 µm, average 8.3 µm n^{= 313}/5 BLs, 3 brains; NB>*mts*-DN BL: median 7.7 µm, average 7.9 μm n^{= 263}/5 BLs, 3 brains; Control VNC: median 6.0 μm, average 6.1 μm n⁼³⁶²/4 VNCs; NB>wdb-DN VNC: median 6.5 µm, average 6.6 µm n⁼³¹⁸/3 VNCs; NB>*mts*-DN VNC: median 6.8 µm, average 7 µm $n^{=258}/3$ VNCs). (H) Quantification of pNB divisions shows that knocking down Wdb in pNBs leads to increased pNB divisions in BLs but not a statistically significant difference in VNCs, while knockdown of Mts causes an increase in pNBs of both BLs and VNCs compared to controls. pNB divisions (Control BL: median 11, average 10.8 Dpn/Ph3+ cells n⁼⁴² 21 brains; NB>wdb-DN BL: median 13.5, average 13.4 Dpn/Ph3+ cells n^{= 22} 11 brains; NB>*mts*-DN BL: median 15, average 15.3 Dpn/Ph3+ cells n⁼²⁰ 10 brains; Control VNC: median 8, average 8 Dpn/Ph3+ cells n⁼²¹ 21 VNCs; NB>wdb-DN VNC: median 9, average 8.1 Dpn/Ph3+ cells n^{= 11} 11 VNC; NB>mts-DN VNC: median 13.5, average 14.7 Dpn/Ph3+ cells n^{= 10} 10 VNCs). p=Wilcoxon rank-sum test. p>0.05 ns (non-significant), *p<0.05, ***p<0.001. Yellow arrowheads indicate dividing pNBs. All images are single focal planes, anterior up. Scale bar: 10 µm.

CHAPTER 3

DISCUSSION

3.1 Understanding NSC quiescence and activated states: the significance of the research area and the foundation of this study

NSCs must actively maintain a balance between quiescent and reactivated states; it is crucial for NSC longevity, proper brain homeostasis and functioning (Cheung and Rando, 2013; Chaker *et al.*, 2016; Cavallucci *et al.*, 2016; Tian *et al.*, 2018). As described in chapter 1, the transition between NSC quiescence and NSC reactivation requires a myriad of extrinsic and intrinsic signals, including systemic nutritional and metabolic signals, growth factors, neurotransmitters, epigenetic modifications and transcription factors. How the NSCs orchestrate a barrage of external and internal signals and appropriately decide which fate to choose is a continuing area of research, essential for NSC-based therapies. Studying NSCs has been difficult due to their relatively low number within an intricate environmental niche. However, our understand- ing of NSCs and their complex heterogeneity was enabled with the development of flow cytometry, cell sorting and transcriptomic techniques (Beckervordersandforth *et al.*, 2010; Beckervordersandforth *et al.*, 2017; Codega *et al.*, 2014; Llorens-Bobadilla *et al.*, 2015; Dulken *et al.*, 2017).

Transcriptomic profiling of mammalian quiescent NSCs versus activated NSCs is starting to decipher the functional properties and regulatory mechanisms governing these different states (Martynoga *et al.*, 2013; Llorens-Bobadilla *et al.*, 2015). As described in the Introduction, research exposed various individual molecules and signalling pathways, for example, BMP and InR/PI3K/Akt signalling to be crucial for NSC maintenance (Mathieu *et al.*, 2008; Ziegler *et al.*, 2015). Contrary to the previous idea that NSC quiescence is a passive process, current research indicates that both quiescence and proliferation states, as well as the transition, requires active and specific genomic and proteomic programs (**Figure 3.1**). Understanding how NSCs integrate, interpret and appropriately respond to the variety of information received is critical to the development of safe, and effective NSC-based regenerative therapy for brain disorders (Llorens-Bobadilla *et al.*, 2015; Chaker *et al.*, 2016; Lee *et al.*, 2018; Xiao *et al.*, 2018; Fouad, 2019; Chou *et al.*, 2015; Tang, Y *et al.*, 2017;

Portnow et al., 2017; Baker et al., 2019; Willis et al., 2020).



Figure 3.1. Integration of intrinsic and extrinsic signals influence NSC transit between quiescence and proliferation stages. A simplified overview of influential factors governing NSC behaviour (see Introduction for more details). NSCs orchestrate a barrage of signals to transition between quiescent and proliferative states. Quiescent and activated behaviours require individual intrinsic genomic and proteomic programs and are regulated by a variety of extrinsic factors including systemic and local niche-derived signals (Cheung and Rando, 2013; Chaker *et al.*, 2016; Cavallucci *et al.*, 2016; Tian *et al.*, 2018).

To contribute to this area of research, our laboratory performed a small-scale transcriptome analysis comparing individual quiescent pNBs and reactivating pNBs harvested from live *Drosophila* larval brains (Gil-Ranedo *et al.*, 2019). Analysis of the data revealed 145 genes upregulated and 51 genes downregulated in reactivating versus quiescent pNBs, and the differential expression of a large set was validated. Most identified targets are enriched in the larval CNS (Chintapalli *et al.*, 2007), 63% and 66% have highly conserved mouse and human orthologs, respectively, and some of these had been found in previous NSC quiescence and activated microarray or RNA sequencing data profiles (Martynoga *et al.*, 2013; Llorens-Bobadilla *et al.*, 2015). Thus, the transcriptome data promised the identification of novel and conserved molecular signals potentially involved in NSC quiescence and/or reactivation.

Most relevant to my studies was the identification of transcripts encoding for members of the highly conserved STRIPAK complex - mob4, cka and the catalytic subunit of PP2A, *mts. mts* transcript was found downregulated, and *mob4* and *cka* transcripts were found upregulated in reactivating pNBs versus quiescent pNBs; together forming the foundation of my studies. Using the *in-vivo* model Drosophila, I characterised Mob4 expression in pNBs, performed *mob4* functional analysis during pNB reactivation, and identified a potential mechanism of *mob4* function on pNB reactivation. I also began and made a significant contribution to studies in our group with *cka* and *mts*, also found in the screen, as well as with the regulatory subunit of PP2A, wdb. My results, together with additional data acquired in the laboratory (discussed below) led to a proposed model whereby Mob4, Cka and PP2A/Mts orchestrate the InR/PI3K/Akt and Hippo pathways promoting pNB reactivation. In summary, PP2A/Mts with its regulatory subunit Wdb inhibits the InR/PI3K/Akt signalling pathway via Akt inactivation during NSC guiescence, yet rising levels of Mob4 and Cka recruit PP2A/Mts into a STRIPAK complex to inhibit Hippo, inactivating the Hippo pathway and promoting NSC reactivation (Gil-Ranedo et al., 2019).

3.2. Mob4 acts primarily cell-autonomously in pNBs to promote reactivation mob4 transcript was found upregulated in reactivating versus quiescent pNBs, the result validated by quantitative real-time PCR (Gil-Ranedo et al., 2019). I confirmed these results at the protein level. Mob4 protein expression is present throughout the CNS, including the glia, which are known regulators of pNB reactivation. Mob4 protein in the glia appeared weaker in comparison to surrounding pNBs, although expression in glia was not quantified (see Figure 2.1). Upon analysis of the mob4 mutants, I observed that the pNBs in the BLs and the VNC are of equivalent size compared to the WT controls at 0-1h ALH (Figure 2.2). However, by 4h ALH, I observed that the WT pNBs are beginning to enlarge, whereas the *mob4* mutant pNBs remain small and quiescent. By 24h ALH many of the WT pNBs in the BLs and the VNC are reactivated; fully enlarged and mitotically active. Interestingly, at 24h ALH, I observed that the *mob4* mutant pNBs in the BLs and the VNC increase slightly in size by 1um, however overall they remain small and mitotically quiescent. It is possible that mob4 mutant pNBs can still initiate some response to start pNB cell growth, but the loss of Mob4 prevents progression of this process. In conclusion, pNB reactivation was abolished in *mob4* mutants, indicating a requirement of Mob4 for the transition from quiescence to reactivated states. Upon pNB-specific knockdown of *mob4* using RNAi (Schulte *et al.*, 2010) at 18h ALH, I observed a small but significant delay in pNB cell size growth and a striking reduction in mitosis; however, the overall effect was weaker compared to the *mob4* mutants. The weaker effect is likely due to the efficacy of the RNAi, which causes only partial inhibition, whereas *mob4* mutants are devoid of all *mob4* (Schulte *et al.*, 2010). My results indicated that Mob4 might act cell-autonomously on pNB reactivation.

Upon Mob4 reintroduction specifically into *mob4* mutant pNBs, I observed that the pNB size and divisions in the BLs and the VNC were rescued to control levels at 18h ALH, whereas the *mob4* mutant pNBs retained their characteristic quiescent appearance (**Figure 2.8**). This result is supportive of the mentioned above findings, suggest- ing a cell-autonomous role of Mob4 in pNBs.

Due to the prominent role of glia relaying Insulin-like signalling to pNBs towards pNB reactivation, Mob4 was also reintroduced into glia of mob4 mutants. I observed that the pNB size in the BLs and the VNC showed a minor (0.4um) increase compared to the mob4 mutant pNBs but far from reaching control values. I also observed a significant increase in pNB divisions in the BLs, but not in VNCs; however, the increase was again minor and far from levels in controls (Figure 2.9). Furthermore, upon gliaspecific mob4 overexpression in a WT background, I observed no pNB size differences in the BLs, or differences in the number of dividing pNBs within the BLs or the VNCs, compared to the controls, at 18h ALH. Yet, I observed a very small yet significant increase (0.4um), in VNC pNB diameters compared to the controls (Figure 2.9). A relatively low number of mob4 mt, glia>mob4 and glia>mob4 VNCs was sampled (n=2 and n=2, respectively) and therefore with increased VNCs scored the minor effects may no longer be significant, albeit many pNBs were scored per sample. It is also possible that the small effects observed may be due to inhibition on the Hippo pathway within niche glia cells, which in turn reflects on pNB reactivation, although not as strong as analogous direct effects in pNBs (Ding et al., 2016; discussed later in section **3.5**). Overall, reintroducing Mob4 into glia of *mob4* mutants cannot rescue pNB reactivation (enlargement or division) to control levels, in contrast to the results upon Mob4 reintroduction in pNBs. Furthermore, glia-specific overexpression of mob4 in a WT background produced no effect except for a minor

VNC pNB size increase, in contrast to overexpression of Mob4 in pNBs that leads to accelerated reactivation (see next section). The findings suggest that *mob4* functions primarily in pNBs promoting reactivation, in agreement with the *mob4* mutant and *mob4*-RNAi data.

3.3. Mob4 functions in pNBs during the stage of reactivation

I observed a loss of pNB reactivation in *mob4* mutants. I also observed that overexpression of Mob4 in pNBs leads to premature pNB enlargement in the BLs and VNCs, but no significant differences in the number of dividing pNBs, at 6h ALH (**Figure 2.11**). At 18h ALH, no further pNB enlargement was seen, but an increased number of pNBs dividing in the BL and VNC was observed. These results suggest that Mob4 accelerates pNB reactivation, and I next investigated if its overexpression would cause pNB overproliferation at later ages.

Upon *mob4* overexpression in pNBs at 48h ALH, I observed a small yet significant increase in the number of dividing pNBs in the VNC only (**Figure 2.13**). Despite many pNBs being reactivated by 24h ALH, pNB reactivation occurs up to approximately 48h ALH (Ito and Hotta, 1992; Truman and Bate, 1988; Chell and Brand, 2010; Sousa-Nunes, *et al.*, 2011). The effect observed is likely to be a consequence of accelerated pNB reactivation still at 48h ALH. As mentioned in the Introduction, anterior pNB reactivation begins before the posterior pNBs, and therefore the effect of Mob4 overexpression may still occur in the VNCs at this point. At late larval stages (94h ALH), no differences in dividing pNBs in the BLs or VNCs were observed (**Figure 2.13**).

My Mob4 overexpression data indicates that Mob4 promotes pNB reactivation from early stages throughout the entire reactivation period up to 48h ALH, but seems not involved in controlling the total number divisions pNBs go through during larval stages. Therefore, Mob4 appears to function only within the time-period of pNBs reactivation.

As described in the introduction, during nutrition restriction, the Insulin signalling pathway is inhibited, and pNB reactivation does not occur. However, these defects can be bypassed by reintroducing components or targets of the InR/PI3K/Akt pathway in pNBs and/or in glia (Chell and Brand, 2010; Sousa-Nunes *et al.*, 2011). Owing to

the reliable data showing that Mob4 promotes the pNB reactivation, I assessed if overexpression of Mob4 in pNBs would be strong enough to bypass the lack of external nutritional stimuli and Insulin-signalling. Upon Mob4 overexpression in pNBs in larvae reared on a sucrose only diet, I observed a small but significant increase in pNB enlargement but no re-entry into mitosis in VNCs, and no increase in either size or divisions of pNB divisions in BLs (**Figure 2.14**). I concluded that Mob4 function in pNBs is not sufficient to bypass the lack of extrinsic nutrition signals required for reactivation.

3.4. Mob4 function in pNB reactivation may be evolutionary conserved *Drosophila* Mob4 scores highly in terms of evolutionary orthology conservation to its human match (MOB4, also known as Phocein), using the *Drosophila* RNAiScreening Center Integrative Ortholog Prediction Tool (DIOPT; http://www.flyrnai.org/diopt). The DIOPT score is the number of individual ortholog prediction algorithms used by this resource that predict a specified ortholog pair. A DIOPT score of 3 is considered adequate, and Mob4 scores 13 out of 15. Mob4 and human MOB4/Phocein proteins are 78% identical, demonstrating high conservation (**Table 1.**). Interestingly, in addition to the Mob1/phocein family domain characterising the MOB family of proteins, a Pleckstrin Homology-like (PH-like) domain exists in both *Drosophila* and human proteins with 73% conservation. While the function of the PH-like domain in Mob4/Phocein is unknown, other proteins containing similar domains have been shown to bind phosphatidylinositol lipids and implicated in a variety of processed ranging from cell signalling transduction to cytoskeleton rearrangements (Lemmon *et al.*, 2002; Scheffzek and Welti, 2012; Lemmon *et al.*, 2000).

Excitingly, upon overexpression of human-MOB4/Phocein in *Drosophila* pNBs (**Figure 2.12**), I observed significant increases in pNB enlargement and mitotic activity, similar to the *Drosophila* Mob4 overexpression effect (**Figure 2.11**). In addition to highly conserved amino acid sequences, these results suggest that Mob4/Phocein may also have a conserved function in NSC reactivation, although expression and functional experiments in human cells, tissue and mammalian models would be needed to validate this.

Species	ID	aa	aa	aa	aa	DIOPT	Known	
		length	%	%	%	score	domains	
			similarity	iden-	gap		(% con-	
				tity			served)	
Drosophila	Mob4	227					PH-like	
Melano-							(73%)	
gaster							Mob1	
Ното	MOB4/	225	86%	78%	3%	13	Phocein	
sapiens	Pho-						(84%)	
	cein							
Fly 1 MKMADGSTILRR RPGTKSKDFCRWPDEPLEEMDSTLAVQQYIQQLIKRDPSNVELILTMPEAQD 65								
Human 1 MVMAEGTAVLRRNRPGTKAQDFYNWPDESFDEMDSTLAVQQYIQQNIRADCENIDKILEPPEGQD 65								
Fly 66 EGVWKYEHLRQFCMELNGLAVRLQKECSPSTCTQMTATDQWIFLCAAHKTPKECPAIDYTRHTLD 130								
Human 66 EGVWKYEHLRQFCLELNGLAVKLQSECHPDTCTQMTATEQWIFLCAAHKTPKECPAIDYTRHTLD 130								
Fly 131 GAACLLNSNKYFPSSVSPRVSIKESSVTKLGSVCRRVYRIFSHAYFHHRRIFDEFEAETYLCHRF 195								
Human 131 GAACLLNSNKYFPSRVSIKESSVAKLGSVCRRIYRIFSHAYFHHRQIFDEYENETFLCHRF 191								
Fly 196 T	THFVTKYNLMSKENLIVPINVGENAAPGESEA 227							
Human 192 T	192 TKFVMKYNLMSKDNLI <mark>V</mark> PILEEEVQNSVSGESEA 225							

Table 1. Mob4 and its human orthologue MOB4/Phocein are highly conserved. A table summarising *Drosophila* Mob4 and human MOB4/Phocein amino acid (a.a) length, % alignment, % similarity, % identity, % gaps, DIOPT score, and known protein domains (http://www.flyrnai.org/diopt). The green highlights represent conserved domain regions extracted from RefSeq protein records (Hu *et al.,* 2011). The red box highlights the PH-like domain. The red brackets encompass the sequence that corresponds to the MOB family of proteins.

3.5. Mechanism of Mob4 action in pNBs

3.5.1 Mob4 and the Hippo and InR/PI3K/Akt signalling pathways

The data discussed thus far indicates that Mob4 is required primarily cellautonomously for pNB reactivation; however, the mechanistic action of Mob4 remained elusive and therefore I, and other members of our laboratory, performed a variety of assays to dissect it (see Results and Appendix). Mob4 is part of the highly conserved MOB family of globular scaffold proteins comprising four subgroups (MOB1-4). As mentioned in the Introduction section, MOB1 and 2 have been implicated in the regulation of kinases that contribute to cell cycle control (Gundogdu and Hergovich, 2019).

MOB4 is the most divergent member of the family (BLAST; Trammell *et al.*, 2008; He *et al.*, 2005) but in *Drosophila* S2 cells, it has been shown to contribute to mitotic spindle fibre focusing, independent of centrosome function (Trammell *et al.*, 2008). My data suggest that Mob4 action in pNB reactivation is independent of that reported in spindle fibre focusing as analysis of the *mob4* mutants shows impairment of pNB cell enlargement before mitotic spindle assembly, and in addition, proliferation of mbNBs, which never enter quiescence, was readily observed (**Figure 2.2-2.5**). The next step was to investigate the relation of Mob4 with InR/PI3K/Akt and Hippo pathways in pNBs, which are critical to activate them or maintain quiescence in pNBs, respectively (Sousa-Nunes *et al.*, 2011; Chell and Brand, 2010; Ding *et al.*, 2016; Poon *et al.*, 2016).

As detailed in the Introduction, Insulin signalling is also known to regulate NSC activation in mammals (Chaker *et al.*, 2016; Bracko *et al.*, 2012; Arsenijevic, *et al.*, 2001; Renault *et al.*, 2009; Kippin *et al.*, 2005; Brandhorst *et al.*, 2015). In *Drosophila*, Insulin signalling is relayed to the pNBs via niche glia. Glia secrete Insulin-like peptides (dILPs) that bind the Insulin-like receptor (InR) in pNBs activating the InR/PI3K/Akt cascade leading to pNB cell size growth and re-entry into division. Secretion of dILPs by glia depends on co-ordinated Ca²⁺ oscillations. Glia also enwrap the developing pNBs and their lineages to ensure their survival (see Introduction; Spéder and Brand, 2014; Spéder and Brand, 2018). It is not known if a similar mechanism occurs between mammalian glia and NSCs. However, astrocytes are known to support neurogenesis and display a protective role to NSCs (Gengatharan *et al.*, 2016; Sloan and Barres, 2014). It is possible that mammalian Insulin signalling is also relayed to adult NSCs via niche glia as in *Drosophila*, yet further studies are required to demonstrate it.

On the other hand, although Hippo signalling has not yet been shown to act in mammalian NSC quiescence, it has been associated with maintaining quiescence in liver progenitors (Wang *et al.,* 2017; Zhou *et al.,* 2009). Inactivating Hippo signalling in the mouse liver quiescent progenitor cells by knocking down the mammalian Hippo

kinase (Mst1/2), caused loss of Yap (mammalian Yki) phosphorylation leading to cell cycle activation and hepatocellular carcinoma. This overgrowth could be rescinded by activating Mst1/2 (Zhou *et al.*, 2009). In mammalian epidermal stem cells, the Hippo signalling effector Yap1 was shown to regulate proliferation and tissue expansion and negatively regulated by upstream α -Catenin (Schlegelmilch *et al.*, 2011). α -Catenin modulates Yap1 activity and interaction with 14-3-3 and PP2A (Schlegelmilch *et al.*, 2011). Furthermore, YAP was shown to maintain basal epidermal progenitors and morphogenesis (Zhang *et al.*, 2011). Finally, in the intestine, YAP is required for homeostasis, and YAP inhibition or activation can cause a myriad of phenotypes (Zhou *et al.*, 2011; Barry *et al.*, 2013). For example, transgenic Yap expression causes loss of intestinal crypts and inhibits regeneration by repressing Wnt signals, whereas Yap inhibition caused microadenomas (Barry *et al.*, 2013).

I have shown that the pNB reactivation defects in *mob4* mutants could be rescued by activating InR/PI3K/Akt signalling in pNBs via overexpressing Rheb (**Figure 2.15**), or by inhibiting the Hippo pathway via targeted RNAi expression against the core kinases Hippo and Warts (**Figure 2.16**). The data suggest that InR/PI3K/Akt signalling is inhibited and Hippo signalling is active in *mob4* mutant pNBs, consistent with my data showing that *mob4* mutant pNBs fail to exit quiescence (**Figure 2.2-2.6**). Although deactivating the Hippo pathway (**Figure 2.16**) or activating InR/PI3K/Akt signalling (**Figure 2.15**) improved the enlargement and mitotic activity of the *mob4* mutant pNBs, the rescues were only partial, as the levels did not reach the control levels. Note that the same manipulations in a WT background produce premature pNB reactivation (Chell and Brand, 2010; Sousa-Nunes *et al.*, 2011; Ding *et al.*, 2016). My results could reflect inadequate activation or inactivation of InR/PI3K/Akt signalling or the Hippo pathway signals, and that regulation of both pathways is necessary for effective pNB reactivation.

The idea that the InR/PI3K/Akt signalling is inhibited in *mob4* mutants was supported by a western blot assay I performed on *mob4* mutant CNS lysates, which showed lower p-Akt expression levels in the *mob4* mutant brains compared to the WT controls, yet total Akt levels remain unchanged (**Figure 2.15**). These results were repeated and confirmed in our laboratory (Dr C. Barros) with additional brain lysate samples I prepared (Gil-Ranedo *et al.*, 2019). The downstream target of InR/PI3K/Akt signalling, Akt, is recruited by PIP3 through its pleckstrin homology (PH) PH-domain, and AKT becomes activated by phosphorylation. Akt phosphorylation/activation can be examined using a (PH) domain-GFP fusion protein (PH-GFP) which binds to PIP3; shown as strong membrane-bound PH-GFP in reactivated pNBs (Britton, *et al.,* 2002; Chell and Brand, 2010). Using this tool, Drs Barros and Gil-Ranedo in our team showed that *mob4* mutant pNBs show weak and diffused PH-GFP signal (Gil-Ranedo *et al.,* 2019), confirming InR/PI3K/Akt signalling inhibition upon Mob4 loss.

bantam (*ban*) microRNA expression is known to be regulated by Hippo signalling (Nolo *et al.,* 2006; Thompson and Cohen, 2006). Active *ban* microRNA promotes pNB enlargement and divisions (Ding *et al.,* 2016). During pNB quiescence, activated Hippo pathway signalling inhibits *ban* transcription (Ding *et al.,* 2016). *ban* activity/Hippo pathway signalling can be monitored using a GFP-sensor system, wherein loss of GFP signal equals *ban* activity (Brennecke *et al.,* 2003). No GFP is detected in control re- activated pNBs indicating *ban* activity (Ding *et al.,* 2016). However, using this tool in a *mob4* mutant background, Drs Barros and Gil-Ranedo observed that pNBs express GFP signal, which indicates a lack of *ban* activity and confirms activated Hippo signalling (Gil-Ranedo *et al.,* 2019).

In conclusion, together with the results obtained by other team members, the data I generated suggested that Mob4 affects both Hippo and InR/PI3K/Akt pathways, with the former remaining active and the latter inhibited in pNBs upon *mob4* loss. How can Mob4 directly or indirectly regulate these pathways is what was next investigated. One clue available was that Mob4 genetically and physically interacts with Hippo and is a known member of the STRIPAK complex, known to inhibit Hippo signalling (Shi *et al.,* 2016; Zheng *et al.,* 2017; Couzens *et al.,* 2013; Ribeiro *et al.,* 2010), which I discuss below.

3.5.2. Mob4 and another STRIPAK/PP2A complex member, Cka, combine actions to promote pNBs reactivation

Mob4 is a core member of the STRIPAK protein complex. A large amount of data arose from a seminal combined proteomic and genomic study investigating *Drosophila* STRIPAK/PP2A and its inhibitory action on the Hippo pathway (Ribeiro *et al.*, 2010).

STRIPAK is made up of multiple components, including various kinases and the pleiotropic PP2A phosphatase. It is striking that STRIPAK members can form mutually exclusive complexes, and are therefore able to direct the kinases or PP2A to different targets (Virshup and Shenolikar, 2009; Ribeiro *et al.*, 2010; Shi *et al.*, 2016). STRIPAK complexes are highly conserved from fungi, to fly, to mammals, including humans (Castets *et al.*, 2000) and their pleiotropic potential allows them to regulate a variety of pathways and processes.

The main components of the STRIPAK complexes identified in Drosophila and humans can be found in Table 2. The main components of human STRIPAK are the striatin (STRN) proteins, comprising STRN1, STRN3 (also known as S/G2 nuclear autoantigen; SG2NA) or STRN4 (also known as Zinedin); the pleiotropic PP2A enzyme with α/β variants; a member of Ste20-related germinal center kinases (GCK) family (Stk24/25, Mst4), and adaptor molecules including cerebral cavernous malformations 3 (CCM3), Sarcolemmal membrane-associated protein (SLMAP), STRN-interacting protein 1/2 (Strip1/2, also named FAM40A/B), suppressor of IKBKE 1 (SIKE1) also known as fibroblast growth factor oncogene partner 2 (FGFR1OP2), and MOB4 (Virshup and Shenolikar, 2009; Ribeiro et al., 2010; Glatter et al., 2009; Goudreault et al., 2009; Castets et al., 2000; Chen et al., 2002; Couzens et al., 2013; Hauri et al., 2013; Sakuma and Chihara, 2017; Shi et al., 2016). The STRIPAK complex is implicated in human diseases such as cancer (Shi et al., 2016; Hwang and Pallas, 2014) and is known to associate with mammalian Hippo (Mst1/2) in the Hippo pathway, which is also implicated in human diseases (Polesello et al., 2006; Ribeiro et al., 2010; Zheng et al., 2017; Huang et al., 2013).

STRIPAK subunit	Drosophila	Human
Striatin	Cka	STRN1/3/4
ΡΡ2ΑΑ	PP2A-29B	ΡΡ2ΑΑα/β
PP2AC	Mts	ΡΡ2Αcα/β
Mob	Mob4	MOB4/Phocein
SLMAP	CG17494	SLMAP
GCK	GckIII, Hpo, Msn	Mst1/2, Stk24/25, Mst4
SIKE/FGFR10P2	FGOP2	SIKE/FGFR10P2
CCM3	CG5073	CCM3 or PDCD10
STRIP	CG11526/Strip	STRIP1/2

Table 2. The core subunits of STRIPAK complex in Drosophila and Humans.Table modified from(Ribeiro et al., 2010; Pracheil et al., 2012; Goudreault et al., 2009; Frost et al., 2012; Bloemendal et al.,2012, and Hwang and Pallas, 2014).

In *Drosophila* STRIPAK (**Table 2**), the components comprise PP2A (PP2A-29B structural subunit and Microtubule star, Mts, the *Drosophila* ortholog of the catalytic subunit of PP2A); Cka (the sole *Drosophila* STRN); Mob4 (*Drosophila* ortholog of MOB4/Phocein); CG10158 (the *Drosophila* ortholog of FGFR1OP2); CG11526 (*Drosophila* ortholog of FAM40A); CG5073/Strip (*Drosophila* ortholog of CCM3); CG17494 (*Drosophila* ortholog of SLMAP); and Hpo, Misshapen (Msn) and Germinal centre kinases (GCKIII in *Drosophila*, MST4 and STK24/25 in mammals). Using the previously described DIOPT tool, evolutionary conservation between *Drosophila* and human STRIPAK subunits is evident (**Figure 3.2. B, and E**). The conserved PP2A phosphatase is a trimeric enzyme composed of a catalytic, a structural and regulatory subunit, the latter responsible for directing it to specific targets (**Figure 3.2A, C**). STRIPAK members act as regulatory subunits of PP2A. Yet, PP2A has also a variety of other functions independent of STRIPAK, relying on multiple different proteins functioning as regulatory subunits to directing it to specific targets (Janssens and Goris, 2001; Wang *et al.*, 2009; Zheng *et al.*, 2015).



Figure 3.2. *Drosophila* and Human core STRIPAK/PP2A complex components inhibit Hippo signalling. The *Drosophila* PP2A subunits (A) have human counterparts (C). *Drosophila* STRIPAK subunits (B) also have human counterparts (D). The DIOPT tool was used to obtain conservation scores to human orthologues (E) (http://www.flyrnai.org/diopt; Hu *et al.*, 2011).

In *Drosophila* and in mammals, including in human cells, a STRIPAK complex has been shown to deactivate the Hippo pathway, via recruitment of PP2A, which dephosphorylates Hpo kinase (human MST1/2) (Zheng *et al.*, 2017; Huang *et al.*, 2013; Polesello *et al.*, 2006; Ribeiro *et al.*, 2010; Bae *et al.*, 2017). In *Drosophila* cells, Cka binds to Hpo and Cka overexpression could block Hpo activity, and Cka overexpression was also shown to increase Yki activity (Ribeiro *et al.*, 2010). The Cka-Hpo interaction is reduced when Mts is knocked down by RNAi; therefore, Cka-Hpo interactions depend upon a stable PP2A complex (Ribeiro *et al.*, 2010).

Ribeiro *et al* demonstrated a genetic relationship between Hpo and Cka (Ribeiro *et al.*, 2010). Hpo overexpression in the eye leads to a reduction in size and a rough-like phenotype (Wu *et al.*, 2003; Pantalacci *et al.*, 2003; Harvey *et al.*, 2003; Udan *et al.*, 2003); and this phenotype is enhanced after removing one copy of Cka (Ribeiro, *et al.*, 2010). Hpo knockdown by RNAi causes an overgrowth phenotype in the developing wing, however simultaneously knocking down Cka or FGOP2 by RNAi suppresses the overgrowth phenotype (Ribeiro *et al.*, 2010). Co-depletion of Hpo and Mob4 by RNAi could also suppress this phenotype (Ribeiro *et al.*, 2010). Interestingly, Toll receptor signalling activates Hippo signalling by causing Cka degradation (Liu *et al.*, 2016).

Drosophila PP2A/Mts encodes the catalytic subunit for PP2A and functions as a tumour suppressor by inhibiting self-renewal capacity in pNBs (Wang et al., 2009). In cells, Mts negatively regulates Hpo; Mts inhibition caused increased Hpo and Yki phosphorylation (Ribeiro et al., 2010). Our lab also recently showed that Mts inhibits the Hippo pathway in pNBs (Gil-Ranedo et al., 2019). An upstream component of the Hippo pathway, Salvador (sav) was shown to bind to PP2A and inhibit its activity, which leads to increased Hippo signalling (Bae et al., 2017). In cells, PP2A/STRIPAK inhibits Hippo signalling (Tang *et al.*, 2019). PP2A and STRNS (mammalian Cka) form a complex which recruits other STRIPAK members to secure MST1/2 (mammalian Hpo) with the STRIPAK complex, destabilising the complex diminished the ability of STRIPAK to regulate Hippo signalling (Tang et al, 2019). This complex is mediated by SLMAP (Tang et al, 2019) which has previously been reported to link STRIPAK to MST1/2 in cells and to Hpo in flies (Couzens et al., 2013; Hauri et al., 2013; Bae et al., 2017; Zheng et al., 2017; Goudreault et al., 2009). Moreover, SIKE1 (mammalian FGOP2) and Strip are required to recruit SLMAP to STRN (Tang et al., 2019). Strip can bind to PP2A, and Strip and SLMAP also directly bind to MST1/2 which collectively aids Hippo pathway regulations (Tang et al., 2019).

Within STRIPAK, CCM3 can direct GCK localisation and also function as bridges, connecting STRNS to GCKs (Shi *et al.*, 2016). Human GCK kinases, together with STRIPAK, can phosphorylate MOB1/Lats (*Drosophila* Wts), together with Salvador (*Drosophila* Sav; Yu and Guan, 2013; Pantalacci *et al.*, 2003; Wu *et al.*, 2003; Tapon *et al.*, 2002; Udan *et al.*, 2003). This effect is in contrast to a recent study showing Sav1 inhibiting the STRIPAK/PP2A complex to promote Hippo signalling (Bae *et al.*, 2017). Therefore, a great deal of crosstalk exists between STRIPAK components and Hippo pathway activity to control growth.

Despite a large amount of data on the Hippo pathway and its inhibition by STRIPAK/PP2A complex, STRIPAK had not yet been implicated in NSC quiescence or reactivation. However, three of the core members were identified on the pNB quiescent vs reactivating transcriptome analysis performed in our laboratory: Cka, Mob4 and Mts, the catalytic subunit of PP2A (Gil-Ranedo *et al.,* 2019). Both *mob4*

and *cka* were found upregulated in reactivating versus quiescent pNBs (**Figure 3.2**). I confirmed the Mob4 differential protein expression (**Figure 2.1**) and Cka expression was also confirmed by our group. I found that overexpressing Mob4 and Cka together in pNBs leads to increased pNB reactivation (**Figure 2.17**), generating a stronger increase in pNB cell enlargement as well as the number of pNB divisions than the Mob4 overexpression (**Figure 2.12**), or Cka overexpression, alone. The latter, performed by Dr Gil-Ranedo, resulted in similar results as Mob4 single overexpression. Also, Dr Gil- Ranedo found that depletion of *cka* by RNAi impairs pNB reactivation similarly to Mob4 inhibition (Gil-Ranedo *et al.,* 2019). Interestingly, in mammalian cells, STRN4 knockdown suppressed cancer cell division, migration and invasion (Wong *et al.,* 2014). Together, the data indicated that Mob4 and Cka might work together in pNB reactivation, possibly as part of STRIPAK and suggesting a new function for the complex.

Dr Gil-Ranedo continued investigating the STRIPAK members using *Drosophila* S2 cells in culture to perform co-immunoprecipitation assays and found that depletion of Mob4 and/or of Cka by RNAi impairs the binding of Mts, the catalytic subunit of PP2A, to Hippo, and a consequent increase in Hippo kinase phosphorylation as previously reported (Ribeiro *et al.,* 2010; Couzens *et al.,* 2013). Mts/Hippo binding was almost abolished by simultaneous *mob4* and *cka* RNAi knockdown, suggestive of cooperative action between Mob4 and Cka in bringing Mts and Hippo together to inactivate Hippo (Gil-Ranedo *et al.,* 2019). Finally, to demonstrate an inhibitory function of PP2A/Mts on the Hippo pathway in pNBs, Drs Barros and Gil-Ranedo combined *mts*-DN and *ban*-GFP sensor *Drosophila* lines to visualise *ban* activity as a read-out of Hippo signalling in pNBs. Early pNBs (6h ALH) expressing *mts*-DN were observed to have significantly less *ban*-activity as seen by more GFP signal than controls, indicating that PP2A/Mts can inhibit Hippo pathway signalling in pNBs.

Together, the findings indicate that Mob4 functions together with Cka, forming with PP2A a STRIPAK complex to target Hippo, leading to Hippo pathway inhibition. These results are also consistent with the observations that the Hippo pathway remains active in *mob4* mutant pNBs (Gil-Ranedo *et al.*, 2019), rendering them quiescent.

3.5.3 PP2A with its regulatory subunit Wdb promotes pNB quiescence

If PP2A would only work in pNBs within STRIPAK to inhibit Hippo signalling, prolonged pNB quiescence could be assumed by knocking it down. Interestingly, the expression of the PP2A catalytic subunit, Mts, was found down-regulated at the mRNA level in reactivating versus quiescent pNBs in the transcriptome analysis performed in our team, the opposite to Cka and Mob4 (Gil-Ranedo et al., 2019). I obtained various mutants strains for *mts*; however, it was not possible to examine pNB reactivation in mts^{XE-2258} null mutants, due to embryonic lethality (Snaith et al., 1996). Yet, by combining *mts*^{XE-2258} null mutants with *mts*²⁹⁹ hypomorphic mutants, I was able to examine *mts*²⁹⁹/*mts*^{XE-2258} transheterozygotes which survive to pupal stages (Wang *et* 2009). In contrast to prolonged pNB quiescence as anticipated, I observed al., increased pNB enlargement in *mts*²⁹⁹/*mts*^{XE-2258} transheterozygotes compared to controls, at 18h ALH (Figure 2.18). Next, I used a dominant-negative mts mutant (*mts*-DN, **Figure 2.19**) which lacks the N-terminal region of the phosphatase domain (Hannus et al., 2002) to analyse the effect of Mts inhibition specifically in pNBs. I observed increased pNB enlargement and mitosis compared to controls at 18h ALH (Figure 2.19). In our group, similar assays were performed to analyse pNBs at 6h ALH, and premature pNB reactivation observed (Gil-Ranedo et al., 2019). These results support the $mts^{299}/mts^{XE-2258}$ transheterozygotes mutant data, and interestingly produced a more substantial effect, demonstrating a significant role of PP2A/Mts in pNBs to maintain quiescence.

The *mts* mutant results were initially quite surprising. However, PP2A is notoriously pleiotropic and, in addition to its function within STRIPAK, it is known to use mutually exclusive regulatory B subunits (**Figure 3.2**) to target a variety of different signalling pathways (Ribeiro *et al.*, 2010; Shi *et al.*, 2016; Janssens and Goris, 2001; Wang *et al.*, 2009; Zheng, Y. *et al.*, 2015). In *Drosophila*, known regulatory B subunits of PP2A (**Figure 3.2**) are Widerborst (wdb), B56/Well rounded (Wrd) and Twins (Tws). In dividing *Drosophila* NBs, PP2A/Tws regulates proper aPKC localisation and NB homeostasis (Chabu and Doe, 2009; Ogawa et al., 2009). Also, PP2A was shown to bind and phosphorylate/deactivate Bazooka (Baz) to regulate Apical-Basal polarity (Krahn *et al.*, 2009). Here, immunoprecipitates containing Baz, PP2A, and the catalytic subunit of PP2A/Mts were identified (Krahn *et al.*, 2009). The study found that

the Mts/PP2A/Baz complexes contained the regulatory B subunits of PP2A, Tws and B56, whereas no interaction with Wdb was found (Krahn et al., 2009). Also, Tws was the sole regulatory B subunit shown to assist the PP2A/Mts dephosphorylation of Baz (Krahn *et al.*, 2009). Wdb is known to inhibit Notch signalling (Bose *et al.*, 2014) and can physically and genetically interact with the cell-cycle regulator, Cyclin G (CycG), in-vitro and in-vivo (Johnson and Walker, 1999; McCright et al., 1996; Stanyon et al., 2004; Fischer et al., 2015; Nagel et al., 2016); even rescuing the growth defects observed in cycG mutants (Nagel et al., 2016). Very interesting and in relation to my findings, is the fact that both Wdb and its *C.elegans* orthologue, PPTR-1, were shown to inhibit InR/Pi3K/AKT signalling pathway by directing PP2A to Akt (Padmanabhan et al., 2009; Vereshchagina et al., 2008). Furthermore, wdb overexpression caused decreased organ size (eye and wing), reduced AKT phosphorylation/activation and increased longevity (Funakoshi et al., 2011). In vertebrates including in human cells, inhibition of Akt by PP2A using Wdb orthologues has also been observed (Rodgers, et al., 2011; Funakoshi et al., 2011; Nagel et al., 2016). I therefore performed a analysis of Wdb in pNBs to analyse its possible role on functional quiescence/reactivation. Using a truncated wdb mutant (Hannus et al., 2002) that acts as a dominant-negative (wdb-DN) and expressing it in pNBs, I observed increased pNB enlargement in the BLs and VNCs compared to controls, at 18h ALH (Figure 2.19), consistent with the effects of *mts*-DN expression (Figure 2.19). Further work completed by other members of the laboratory showed that *wdb*-DN expressing pNBs enlarged prematurely at 6h ALH, again similar to the results upon mts-DN expression (Gil-Ranedo et al., 2019). Moreover, using a p-AKT specific antibody, mts-DN pNBs displayed early p-AKT signal compared to the controls at 6h ALH, demonstrating premature InR/PI3K/Akt signalling activity upon PP2A/Mts depletion (Gil-Ranedo et al., 2019). Finally, Dr Gil-Ranedo also demonstrated that depletion of Cka and/or Mob4 in S2 cells did not affect Mts-Akt association nor the levels of phosphorylated/activated AKT (Gil-Ranedo et al., 2019).

Overall my data and the work completed by other laboratory members suggest that PP2A/Mts functions with Wdb to inhibit the InR/PI3K/Akt signalling in pNBs, and this role seems independent of that of PP2A/Mts within STRIPAK and its components Cka and Mob4.

3.5.4. Proposed mechanistic action of Mob4, Cka and PP2A in pNB quiescence to reactivation transition

As discussed above, my results (**Figure 2.19**) together with data obtained from other team members in our laboratory show that PP2A using the Wdb regulatory subunit maintains pNB quiescence by inhibiting the InR/PI3K/Akt signalling cascade via inactivation/ dephosphorylation of Akt (**Figure 3.3**). These findings are consistent with previous reports of PP2A/Wdb inhibiting Akt by in other cells and tissues (Johnson and Walker, 1999; McCright *et al.*, 1996; Stanyon *et al.*, 2004; Fischer *et al.*, 2015; Nagel *et al.*, 2016; Padmanabhan *et al.*, 2009; Vereshchagina *et al.*, 2008; Brownlee, *et al.*, 2011; Funakoshi *et al.*, 2011). Interestingly, PP2A has also been implicated in cellular quiescence in different contexts (Naetar *et al.*, 2014; Kolupaeva *et al.*, 2008; Kolupaeva *et al.*, 2013; Sun and Buttitta, 2015; Kurimchak *et al.*, 2013; Kurimchak and Grana, 2013).

The *Drosophila* eye becomes quiescent ~24h after pupal formation (Buttitta *et al.*, 2007). By testing different PP2A B subunits, Sun and Buttitta demonstrated that PP2A/*wdb* restricts CyclinE/Cdk2 to promote quiescence (Sun and Buttitta, 2015). Knocking down PP2A/*mts* or of PP2A/*wdb* by RNAi in the eye caused an increased S-phase (EdU) positive cells and cell cycle gene expression (Sun and Buttitta, 2015). They determined that loss of PP2A/*wdb* in the *Drosophila* eye and wing, allowed cells to bypass entry into quiescence, which leads to extra divisions (Sun and Buttitta, 2015). In human glioblastoma immortalised cells in culture, PP2A and its regulatory B56γ subunit is required for inducing stable quiescence via Ras signalling (Naetar *et al.,* 2015). Inhibition of PP2A using okadaic acid during G2 interferes with quiescence was thought to occur in G1; however in this study, they showed that Ras inhibition by PP2A/ B56γ occurred during G2 only, and this pathway helps determine the cell-cycle length and quality of the next cell-cycle (Naetar *et al.,* 2014).

The results from my expression and functional analysis of the evolutionarily conserved STRIPAK core component Mob4, demonstrate that it promotes pNB reactivation, primarily in a cell-autonomous fashion. The data also suggest that Mob4 cooperates with Cka, another core STRIPAK complex member. It is known that the Hippo

pathway maintains pNB quiescence (Ding *et al.*, 2016; Poon *et al.*, 2016), whereas InR/PI3K/Akt signalling activation is necessary for pNB reactivation (Chell and Brand, 2010; Sousa-Nunes *et al.*, 2011). Upon loss of Mob4, pNBs are unable to reactivate, with Hippo pathway remaining activated and InR/PI3K/Akt signalling inhibited (Gil-Ranedo *et al.*, 2019). I have further shown that overexpressing the InR/PI3K/Akt signalling activator Rheb, or inhibiting the Hippo pathway core kinases Hpo or Wts specifically in *mob4* mutant pNBs, can rescue reactivation, suggesting that Mob4 is required to regulate Hippo and InR/PI3K/Akt signalling pathways in pNBs. Cka and Mob4 were known to assist PP2A-inhibition of Hippo (Ribeiro *et al.*, 2010), and our team demonstrated that inhibition of Cka and/or Mob4 indeed prevents PP2A/Mts physical association to Hippo kinase, leading to Hippo signalling activation. In accordance, PP2A/Mts inhibition in pNBs also leads to increased Hippo signalling, but the effect is not sufficient to overcome the premature activation of InR/Pi3K/AKT signalling that also occurs, resulting in early reactivation despite Hippo activity (Gil-Ranedo *et al.*, 2019).

Collectively, the data supports the idea that as Mob4 and Cka levels increase in pNBs, a STRIPAK/PP2A complex assembles and acts as a 'molecular switch' to redirect PP2A/Mts to inhibit Hippo signalling and allow activation of InR/PI3K/Akt signalling in pNBs, promoting reactivation (**Figure 3.3**).


Figure 3.3. The STRIPAK members Mob4, Cka and PP2A contribute to pNB transition from **quiescence to reactivation.** In quiescent pNBs, the levels of Mob4 and Cka are low, and PP2A/Mts phosphatase is directed to Akt by the regulatory subunit, Wdb. PP2A/Mts-Wdb dephosphorylates (deactivates) Akt, inhibiting InR/Pi3K/Akt signalling, which is required for pNB reactivation (Chell and Brand, 2010; Sousa-Nunes *et al.,* 2011). The Hippo pathway, which maintains quiescence (Ding *et al.,* 2016; Poon *et al.,* 2016) remains active. During pNB reactivation, the levels of Mob4 and Cka surge, assembling a Mob4/Cka/PP2A STRIPAK complex that redirects Mts to dephosphorylate Hpo, inactivating the Hippo pathway. InR/PI3K/Akt signalling activates, promoting pNB reactivation.

Interestingly, a similar STRIPAK/PP2A-dependent mechanism regulates the *Drosophila* circadian oscillator (Andreazza *et al.,* 2015). The CLOCK (CLK) and CYCLE (CYC) bHL transcription factors temporally activate circadian rhythm genes such as *period, timeless* and *pdp1,* which in-turn repress or activate CLK and CYC

(Hardin, 2011; Ozkaya and Rosato, 2012; Sathyanarayanan *et al.*, 2004). CLK is highly phosphorylated and inactive in the morning, but hypophosphorylated and active in the evening. These temporally controlled oscillating loops of repression and activation, creating *hyper*-or-*hypo*-phosphorylated CLK, are vital for *Drosophila* circadian regulation. Andreazza *et al.*, show that PP2A comprising the regulatory (Wdb) and catalytic (Mts) subunits (PP2A-Mts/Wdb) act to stabilise CLK, without affecting its phosphorylation. However, a complex made of PP2A/Mts and the STRIPAK components Cka and Strip (Strip/Cka/PP2A-Mts) cause CLK dephosphorylation during daytime (Andreazza *et al.*, 2015).

3.6. Conclusion

We are slowly demystifying the brain, NSCs, and how adult NSCs maintain adult brain homeostasis (Morales, 2019; Obernier, 2019; Tian *et al.*, 2018; Chaker *et al.*, 2016; Cheung and Rando, 2013). Improved techniques, such as brain cell sorting, identification, and transcriptome and proteome characterisation, have significantly advanced research efforts (Codega *et al.*, 2014; Beckervordersandforth *et al.*, 2010; Daynac *et al.*, 2015; Mich *et al.*, 2014; Llorens-Bobadilla *et al.*, 2015; Shin *et al.*, 2015; Dulken *et al.*, 2017; Morizur *et al.*, 2018; Berg *et al.*, 2019).

My studies towards a PhD degree aimed to examine candidate genes that arose from a small-scale single-cell transcriptome analysis comparing quiescent and reactivating *Drosophila* NSCs (Gil-Ranedo *et al.*, 2019). Mob4 was a gene identified together with other STRIPAKS members Cka and PP2A. I have investigated Mob4 expression in NSCs, performed functional analysis of Mob4, as well as of Cka and PP2A, and contributed to the identification of a mechanism of Mob4 action together with Cka and PP2A in NSC quiescence to reactivation transition; accomplishing all proposed objectives of this work.

My work described in this dissertation, together with work from other members of our team, indicated in the above Discussion sections, led us to the propose a model of action: In pNBs, PP2A/Wdb functions to inhibit InR/PI3K/Akt signalling promoting quiescence. As Mob4 and Cka levels increase, a STRIPAK/PP2A complex forms and acts as a 'molecular switch' to redirect PP2A to inhibit Hippo signalling, allowing InR/PI3K/Akt signalling activation and promoting pNB reactivation. Thus, Mob4, Cka

and PP2A orchestrate in pNBs Hippo and InR/PI3K/Akt signals, two major pathways controlling quiescence and reactivation, respectively. The studies were recently included in a paper published in Cell Reports in which I am a shared first author (see appendix for full publication).

It is fascinating how PP2A subunits and the STRIPAK complex members collaborate so efficiently to achieve a plethora of functions, across vast cell types, regulating many signalling systems and cellular processes. It is remarkable that PP2A is known to target over 300 substrates and modulate most major signalling cascades such as the Hippo pathway, InR/PI3K/Akt, mTOR, c-Jun N-terminal kinases (JNK), mitogen-activated protein kinase (MAPK), extracellular signal-regulated kinases (ERK), Notch, Wnt and implicated in a variety of human diseases (Remmerie and Janssens, 2019; Eichhorn *et al.*, 2009; Wlodarchak and Xing, 2016; Grech *et al.*, 2016; Ruvolo *et al.*, 2016; Kiely *et al.*, 2015; Westermarck *et al.*, 2008; Sangodkar *et al.*, 2016; Schuhmacher *et al.*, 2019; Clark and Ohlmeyer, 2019).

I have contributed to identifying one mechanism whereby STRIPAK members and PP2A integrate different signalling pathways that are highly conserved from fly to human. This mechanism adds to current knowledge of NSC behaviour and regulation, and may also contribute to the development of future NSC-based therapies against brain-related diseases such as neurodegeneration, autism and even brain tumours. My findings likely have relevance in other stem cells, which also rely on smooth transitions between quiescent and activation states. It will be interesting to examine the expression and function of the highly conserved Mob4, and other STRIPAK members, in mammalian NSCs.

CHAPTER 4. MATERIALS AND METHODS

4.1. MATERIALS

4.1.1. Antibodies and cell proliferation assay kit

The following tables **3** and **4** list the primary and secondary antibodies used for immunostainings and western blotting of *Drosophila* larval brains described in the Results chapter. Antibody name and species made in, source and dilutions used are indicated. Table 5 describes the commercial kit used for EdU incorporation assays shown in section **2.2.** of Results.

Primary Antibodies	Source	Dilution
Rabbit anti-GFP	Gift from U. Mayor	1:1000
	(U. Mayor and A.H. Brand, unpublished)	
Chicken anti-GFP	Millipore AB_310288	1:500
Guinea pig anti-Dpn	Gift from J. Knoblich (Levy and Larsen,	1:2000
	2013) AB_2314299	
Rat anti-Dpn	Abcam AB_195173	1:100
Guinea pig anti-Mob4	Gift from T. Littleton (Schulte, et al., 2010)	1:1000
Mouse anti-Dlg	DSHB AB_528203	1:50
Rabbit anti-pH3	Abcam AB_304763	1:1000
Rabbit anti-pAKT ^{S505}	CST AB_331414	1:400
Rabbit anti-Akt	CST AB_329827	1:400
Rabbit anti-b-Actin	CST AB_330288	1:50
Rat anti-Grainyhead	Gift from Prof. Steven Thor laboratory	1:1000
	(Sweden)	
Mouse anti-Prospero	DSHB AB_528440	1:50
Mouse anti-Repo	DSHB AB_528448	1:50

 Table 3. Primary antibodies used for Immunohistochemistry or western blotting.

Secondary Antibodies	Source	Dilution
Goat Anti-Rat 633	Thermo Fisher A-21094	1:200
Goat Anti-Chicken 488	Thermo Fisher A-11039	1:200
Goat Anti-Guinea pig 568	Thermo Fisher A-11075	1:200
Goat Anti-Rabbit 633	Thermo Fisher A-35562	1:200
Goat Anti-Mouse 488	Thermo Fisher A-10667	1:200
Goat Anti-Rat 568	Thermo Fisher A-11077	1:200
Goat Anti Rabbit IgG–Horse radish	Cell signalling Technologies	1:1000
peroxidase (HRP)	#7074	

Table 4. Secondary antibodies used for Immunohistochemistry or westernblotting.

Resource	Source	Identifier
Click-iT EdU Alexa Fluor 594	Invitrogen	Cat# C10339
Imaging Kit		

Table 5. Cell proliferation kit.

4.1.2. *Drosophila* strains

The table below lists the *Drosophila* strains used in this study, including those enabling the generation of lines via genetic recombination or combination mentioned in the Results chapter. For each line, the genotype, source and identification number (identifier) are indicated.

Strains	Source	Identifier
D. melanogaster: mob4 mu-	Bloomington Drosophila Stock Cen-	BDSC: 36331;
tant:	ter (Schulte, <i>et al.,</i> 2010)	FlyBase:
y[1] w[*]; Mob4[EYDel-		FBst0036331
taL3]/CyO		
D. melanogaster: UAS line	Bloomington Drosophila Stock Centre	BDSC:36488;
expressing mob4 RNAi:	(Schulte, <i>et al.,</i> 2010)	FlyBase:
P{UAS-		FBst0036488
Mob4.RNAi.JS1}attP2		
D. melanogaster: UAS line	Bloomington Drosophila Stock Cen-	BDSC: 36329;
expressing mob4:	ter (Schulte, <i>et al.,</i> 2010)	FlyBase:
y[1] v[1]; P{y[+t7.7]		FBst0036329
v[+t1.8]=UAS-Mob4.S}attP2		
D. melanogaster: UAS line	Bloomington Drosophila Stock Cen-	BDSC: 36330;
expressing hMOB4:	ter	FlyBase:
y[1] v[1]; P{y[+t7.7]	(Schulte, <i>et al.,</i> 2010)	FBst0036330
v[+t1.8]=UAS-pho-		
cein.1}attP2		
D. melanogaster: UAS line	Bloomington Drosophila Stock Centre	BDSC: 55053;
expressing wdb-DN:	(Hannus, <i>et al.,</i> 2002)	FlyBase:
w[*]; P{w[+mC]=UAS-		FBst0055053
wdb.95-524.HA}6		
D. melanogaster: UAS line	Bloomington Drosophila Stock Cen-	BDSC: 41899;
expressing wts RNAi:	ter (Ding, <i>et al.,</i> 2016)	FlyBase:
y[1] sc[*] v[1]; P{y[+t7.7]		FBst0041899
v[+t1.8]=TRiP.GL01331}attP 2		
D. melanogaster: UAS line	Bloomington Drosophila Stock Cen-	BDSC: 33614;
expressing hpo RNAi:	ter	FlyBase:
y[1] v[1]; P{y[+t7.7]	(Ding, <i>et al.,</i> 2016)	FBst0033614
v[+t1.8]=TRiP.HMS00006}att		
P2		

D. melanogaster: Wild type:	Gift from M. Akam (Bossing and	FlyBase:
Oregon-R	Technau, 1994)	FBsn0000276
D. melanogaster: UAS line	Gift from R. Sousa-Nunes (Sousa-	BDSC: 9689;
expressing rheb:	Nunes, <i>et al.,</i> 2011)	FlyBase:
w[*]; P{w[+mC]=UAS-		FBst0009689
Rheb.Pa}3		
D. melanogaster: Gal4 line	Gift from A.H. Brand (Chell and	N/A
under the control of grh: Grh-	Brand, 2010)	
Gal4		
D. melanogaster: UAS line	Gift from S. Eaton (Hannus, et al.,	N/A
expressing mts-DN-HA: UAS-	2002)	
mts.dn181-HA		
D. <i>melanogaster</i> : mts ^{XE225839}	Gifts from H. Wang (Wang, et al.,	BDSC: 5684;
mutant: mts ^{XE2258} /CyO,	2009)	FlyBase:
P{sevRas1.V12}F1		FBst0005684
and mts ²⁹⁹ mutant.		N/A for mts ²⁹⁹
D. melanogaster: Gal4 line	Bloomington Drosophila Stock Cen-	BDSC: 8751;
under the control of insc: w[*];	ter	FlyBase:
P{w[+mW.hs]=GawB}insc[Mz		FBst0008751
1407]		
D. melanogaster: Gal4 line	Gift from A. Hidalgo	N/A
under the control of repo:		
w[1118];		
P{w[+m*]=GAL4}repo/TM6,		
tb		
D. melanogaster: UAS line	(Lee and Luo, 1999)	BDSC: 5137;
expressing CD8-GFP:		FlyBase:
y[1] w[*]; P{w[+mC]=UAS-		FBst0005137
mCD8::GFP.L}LL5, P{UAS-		
mCD8::GFP.L}2		
D. melanogaster: UAS line	(Ding, <i>et al.,</i> 2016)	N/A
expressing dicer2:		
UAS-dicer2		

D. melanogaster: UAS line	Bloomington Drosophila Stock Cen-	BDSC: 53756;
expressing cka-eGFP:	ter	FlyBase:
w[*];P{w[+mC]=UASp-		FBst0053756
Cka.EGFP.C}2		
D. melanogaster: Double Bal-	Bloomington Drosophila Stock Cen-	BDSC: 7197; Fly-
ancer line	ter	Base:
w[*]; I(2)*[*]/CyO; D[1]/TM6B,		FBst0007197
Tb[+]		
D. melanogaster: Double Bal-	Bloomington Drosophila Stock Cen-	BDSC: 24900;
ancer line	ter	FBst0024900
y[1] w[*]; Eps-15[e75]/CyO,		
P{w[+mC]=GAL4-twi.G}2.2,		
P{w[+mC]=UAS-2xE-		
GFP}AH2.2		

Table 6. Drosophila strains.

4.2.METHODS

4.2.1. *Drosophila* husbandry

Fly stocks were maintained in vials for long-term storage at 18°C and at 25°C for ongoing experiments, with the exception of RNAi assays that were performed at 29°C. The fly stocks were kept in standard *Drosophila* fly food. Egg collections and larvae rearing were performed on agar juice plates supplemented with yeast paste. Egg lays were collected in either 30 minutes or 1h time-windows.

Protocol to prepare *Drosophila* food: 750 ml boiling water, 100 g sugar, 50 g yeast, 80 g organic flour, 2 g Ammonium sulphate, (NH4)S04, 5 g tartaric acid, 1.5 g Potassium phosphate monobasic, KH2P04, 22 g agar soaked into 300 ml grape juice (in two batches of 150 ml), 10 ml propionic acid, 1 g Nipagin, Vials, Cotton plugs.

Dissolve the sugar completely in the boiling water. Add the 50 g of yeast and dissolve completely. Mix the 80 g of organic flour into 200-300 ml of water until a smooth paste is formed and then add it to the boiling water to dissolve. Add 2 g of ammonium sulphate, 5 g tartaric acid and 1.5 g potassium phosphate monobasic to the water and boil for 10 minutes. Slowly add one of the agar/grape juice batches (11 g:150 ml) to

the boiling water. Take the other agar/grape juice batch and microwave it until boiling and slowly add it to the mixture and boil for a further 10 minutes. Let the mixture cool down until it reaches ~70°C. Add 1 g of Nipagin using a scalpel to the mixture. On top of the Nipagin pour 10 ml of propionic acid and allow to mix well. Add approximately 1 inch high of food per vial or bottle.

Protocol to prepare Drosophila egg-lay collection plates:

1 I water, 21 g agar, 200 ml grape juice, plates, 2 I beaker.

Pour 1I of water into a 2 I beaker and microwave until boiling (14 minutes). Set the heater to 270°C and the stirrer to 420 rpm. Place the 2 I beaker onto the heater and wait until the water is boiling. Mix half the 21 g of agar and 100 ml of grape juice in a flask and stir. Add the agar solution to the boiling water. Add the rest of the agar in another 100 ml of grape juice and pour into the mixture. Stir and boil the juice mixture in the 2 I beaker for no longer than 2 minutes. Pour juice mixture into small petri dish plates used for egg-lay collections.

For nutritional deprivation experiments, freshly hatched larvae were transferred to agar plates prepared with amino-acid free media (5% sucrose, 1% agar in phosphate buffered saline, PBS).

Prepare 10x PBS (phosphate buffered saline): dissolve 80 g of NaCl, 2 g of KCl, 26.8 g pf Na₂HPO₄-7H2O, 2.4 g KH₂PO₄ in 800 ml of ddH2O, adjust the pH to 7.4 with HCl and bring the total volume to 1 litre.

Prepare 1 x PBS: 1:10 dilution of 10xPBS in ddH2O

4.2.2. Drosophila strains and genetic crosses

Fly stocks were maintained in vials long-term at 18°C and at 25°C for ongoing experiments, with the exception of RNAi assays performed at 29°C. The fly stocks were kept in standard *Drosophila* fly food in vials or bottles. Fly strains needed for experiments were grown in multiple bottles at 25°C to yield maximum female virgin collections. Female virgins were collected every 3 hours (maximum) and stored with the appropriate males at 18°C. The resulting fly cross would be transferred to an egg-lay chamber at 25°C. Egg collections and larvae rearing were performed on agar juice plates supplemented with yeast paste. Embryos were transferred to fresh plates and stored at the appropriate temperature. Embryos with GFP balancers were

selected for, or against, the next day using a fluorescent microscope; the desired embryos transferred to a fresh plate supplemented with yeast.

Drosophila strains used in this study are listed in **Table 6.** Drosophila stocks obtained from the Bloomington Drosophila Stock Center were: mob4^{EYAL3} (36331) (Schulte, et al., 2010) rebalanced over CvO, P(GAL4-twi.G)2.2; UAS-mob4^{RNAi} (36488) (Schulte, et al., 2010); UAS-mob4 (36329) (Schulte, et al., 2010); UAS-hMOB4 (Phocein) (36330) (Schulte, et al., 2010); UAS-cka-eGFP (53756); UAS-wdb-DN (UAS-wdb.95-524.HA; 55053) (Hannus, et al., 2002); UAS-wts^{RNAi} (41899) (Ding, et al., 2016) and UAS-hpo^{RNAi} (33614) (Ding, et al., 2016). Other stocks used were: Wild-type Oregon-R (kind gift from M. Akain); UAS-rheb (Blomington 9689) (Sousa-Nunes, et al., 2011) (kind gift from R. Sousa-Nunes); grh-Gal4 (Chell and Brand, 2010) (kind gift from A.H. Brand); UAS-mts-DN (UAS-mts.dn181-HA) (Hannus, et al., 2002) (kind gift from S. Eaton); mts²⁹⁹ and mts^{XE2258} (Wang, et al., 2009) (kind gifts from H. Wang). NSCspecific RNAi and overexpression assays were performed using *insc-Gal4* (w¹¹¹⁸; p{GAWB}inscMZ1407). Glial-specific expression assays used repo-Gal4 (w¹¹¹⁸; p{GAWB}repo/TM6b, iab-lacZ). The grh-Gal4 driver had been recombined with UAS-CD8-GFP (chromosome II) in the laboratory (Gil-Renado et al., 2019). For rescue experiments, *insc-Gal4* and *repo-Gal4* drivers were recombined or combined with the *mob4*^{EYAL3} mutant strain. For other assays, the *insc-Gal4* driver was recombined with UAS-CD8-GFP or a strain already combining it with UAS-dicer2 and UAS-CD8-GFP was used (Ding, et al., 2016). Genetic combinations or recombinations, balancing and/or re-balancing of *Drosophila* stocks on the II and or III chromosome, followed standard genetic cross procedures (Greenspan, 2004).

4.2.3. Immunohistochemistry and EdU incorporation

Prepare 1xPBST: 1x PBS with 1% Triton X

Prepare fixative solution: 4% formaldehyde in PBS with added EGTA (0.5 M) and MgCL2 (1 M). For 1 ml: 894 μ l PBS 1 x + 100 μ l formaldehyde + 1 μ l EGTA (0.5 M) + 5 μ l MGCl2 (1 M). Vortex.

Prepare blocking solution: 10% goat serum in PBS-1% Triton. For 1 ml: 900 μ l of 1x PBS-1% Triton + 100 μ l 100% Goat serum. Vortex and store at 4C. All chemicals were purchased from SIGMA.

Prior to dissections, larvae were transferred to a silicon dissecting dish in 1xPBS and (if necessary) larvae were checked under the fluorescent microscope. Using a light microscope, larval CNSs were extracted using fine forceps and scissors and fixed for 20 minutes in fixative solution, followed by washes in PBS (2 x 10 min, 3 rinses between washes) and 1 hour block in 1 x PBST (PBS, 1% Triton X-100) with 10% fetal bovine serum, at room temperature on a shaker. Primary antibodies (**Table 3**) were incubated in 1 x PBST overnight or for 2 nights at 4°C. CNSs were washed in 1xPBST (2 x 5 min, 3 rinses between washes) before incubating with secondary antibodies (**Table 4**) for 2 hours at room temperature. Secondary antibodies were washed off with 1 x PBST (2 x 10 min, 3 rinses between washes) and successively embedded in 50% (>30 mins) and 70% glycerol (>3 hours) before carefully mounting on slides in a 1:1 mix of 70% glycerol and Vectashield (Vector Laboratories) and covered by a coverslip.

EdU incorporation assays were performed as previously described (Sousa-Nunes, *et al.*, 2011). Briefly, CNSs were dissected in PBS and incubated in 10 μm EdU/PBS for 1h at room temperature. CNSs were fixed for 15 minutes in 4% formaldehyde/PBS and incorporated EdU detected using Click-iT EdU Imaging kit following manufacturer instructions (Invitrogen).

4.2.4. Image acquisition and processing

Images were obtained on a Leica SP8 confocal laser-scanning microscope using LAS X software. Quantifications were made using z-stacks of 1.5 µm step size, comprising whole-brain lobes, VNCs or CNSs. Representative images shown are single optical sections, with the exception of EdU incorporations (**Fig 2.4**), which are z-projection stacks encompassing whole CNSs. Images were processed in Fiji v2.0 or Adobe Photoshop CS6 and assembled in Adobe Illustrator CS6. NSC sizes (maximum diameters) (Chell and Brand, 2010), pH3 scorings, and Mob4 signal intensities (pixel intensity/ NSC maximum area) and EdU voxel quantification were performed using Fiji v2.0.

4.2.5. Western blotting

Whole larval brain were harvested and lysed in lysis buffer (1% Triton) supplemented with protease inhibitor (Complete, EDTA-free; Sigma) and phosphatase inhibitors (cocktails B+C; Santa Cruz Biotechnology). CNS extracts were spun at 14000 rpm for

30 minutes at 4°C and proteins quantified (BCA protein assay, Pierce). Detection of proteins was performed using standard sodium dodecyl sulfate (SDS) polyacrylamide gel electrophoresis (SDS-PAGE) and western blotting using ECL or ECL Plus chemiluminescent substrate (Pierce) with the following specifications. CNS lysates were run on a 10% polyacrylamide gel at 120 V and transferred onto a nitrocellulose membrane, before blocking with 5% BSA (p-AKT), 3% BSA (β -Actin and AKT), in 0.1% TBS- Tween and incubated with the primary antibody overnight at 4°C under rotation.

4.2.6. Data quantification and statistical analysis

Statistics were performed using SigmaPlot Version 12.5 (Systat software): Shapiro-Wilk and equal variance tests used to evaluate normality; Student's *t*-test applied when data fitted a normal distribution; Wilcoxon rank-sum test used for non-parametric data; p<0.05 considered significant. Data from *Drosophila in vivo* assays were obtained from a minimum of two biological replica sets; sample numbers are indicated in figure legends. Histograms show mean ± standard error of the mean. Box plots represent 25th and 75th percentiles, and black line indicates median, dashed line specifies mean, whiskers indicate 10th and 90th percentiles.

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APPENDIX.

Cell Reports

STRIPAK Members Orchestrate Hippo and Insulin Receptor Signaling to Promote Neural Stem Cell Reactivation

Graphical Abstract



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In Brief

The integration of signals allowing stem cell reactivation from quiescence is unclear. Gil-Ranedo et al. identify STRIPAK members Mob4, Cka, and PP2A/Mts through reactivating versus quiescent neural stem cell (NSC) transcriptional profiling. Their findings suggest that Mob4/Cka/Mts functions as an intrinsic molecular switch coordinating Hippo and InR/PI3K/Akt pathways, enabling NSC reactivation.

Highlights

- Transcriptional profiling of reactivating versus quiescent NSCs identifies STRIPAK members
- PP2A/Mts phosphatase inhibits Akt activation, maintaining NSC quiescence
- Mob4 and Cka target Mts to Hippo to inhibit its activity and promote NSC reactivation
- Mob4/Cka/Mts coordinate Hippo and InR/PI3K/Akt signaling in NSCs





STRIPAK Members Orchestrate Hippo and Insulin Receptor Signaling to Promote Neural Stem Cell Reactivation

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SUMMARY

Adult stem cells reactivate from guiescence to maintain tissue homeostasis and in response to injury. How the underlying regulatory signals are integrated is largely unknown. Drosophila neural stem cells (NSCs) also leave quiescence to generate adult neurons and glia, a process that is dependent on Hippo signaling inhibition and activation of the insulin-like receptor (InR)/PI3K/Akt cascade. We performed a transcriptome analysis of individual quiescent and reactivating NSCs harvested directly from Drosophila brains and identified the conserved STRIPAK complex members mob4, cka, and PP2A (microtubule star, mts). We show that PP2A/Mts phosphatase, with its regulatory subunit Widerborst, maintains NSC quiescence, preventing premature activation of InR/PI3K/Akt signaling. Conversely, an increase in Mob4 and Cka levels promotes NSC reactivation. Mob4 and Cka are essential to recruit PP2A/ Mts into a complex with Hippo kinase, resulting in Hippo pathway inhibition. We propose that Mob4/ Cka/Mts functions as an intrinsic molecular switch coordinating Hippo and InR/PI3K/Akt pathways and enabling NSC reactivation.

INTRODUCTION

Brain homeostasis and damage repair depend on the generation of new neurons and glia by neural stem cells (NSCs). In adult brains, most NSCs are found to be quiescent but can enter proliferation if prompted by extrinsic and intrinsic stimuli. The balance between quiescence and reactivation is critical for the maintenance of an NSC reservoir (Cavallucci et al., 2016; Chaker et al., 2016). Mechanistic insight underlying NSC quiescence and reactivation remains limited—in particular, how regulatory signals are integrated.

In the model organism *Drosophila*, embryonic NSCs give rise to the larval functional CNS. Similar to mammals, NSCs become quiescent at the end of embryogenesis and reactivate postembryonically to generate neurons and glia of the adult brain (Truman and Bate, 1988). Quiescence entry is regulated by Hox proteins, temporal transcription factors, and levels of the homeodomain transcription factor Prospero (Otsuki and Brand, 2019; Lai and Doe, 2014; Tsuji et al., 2008). NSCs are kept guiescent by the canonical Hippo pathway, whereby its core kinases Hippo and Warts prevent the transcriptional co-activator Yorkie from entering the nucleus and triggering growth (Ding et al., 2016; Poon et al., 2016). This signaling can be modulated by niche glia cells via the upstream regulators Crumbs and Echinoid, expressed in both glia and NSCs (Ding et al., 2016). NSC reactivation involves cell size increase, from 4 to 5 μm during quiescence, followed by entry into division (Ding et al., 2016; Chell and Brand, 2010; Prokop and Technau, 1991; Truman and Bate, 1988). NSCs continue to enlarge, reaching up to 10–15 µm when proliferating (Prokop and Technau, 1991; Truman and Bate, 1988). Nutrition stimulates reactivation (Britton and Edgar, 1998): dietary amino acids in the young larvae induce a systemic signal that triggers blood-brain barrier glia to secrete Drosophila insulin-like peptides (dILPs), a process that depends on gap junction proteins and synchronized calcium pulses (Spéder and Brand, 2014). dILPs activate the insulin-like receptor (InR)/phosphoinositide 3-kinase (PI3K)/Akt cascade in neighboring NSCs, promoting quiescence exit (Sousa-Nunes et al., 2011; Chell and Brand, 2010). The conserved heat shock protein 38/90 chaperone associates with InR to promote reactivation, and Spindle matrix proteins, including Chromator, function downstream of InR/PI3K/Akt signaling in this process (Huang and Wang, 2018; Li et al., 2017).

We performed a small-scale transcriptome analysis using single quiescent and reactivating NSC samples obtained directly from live *Drosophila* brains. Members of the evolutionary conserved striating-interacting phosphatase and kinase (STRIPAK) complex (Shi et al., 2016; Ribeiro et al., 2010) were identified and validated: monopolar spindle-one-binder family member 4 (Mob4); connector of kinase to AP-1 (Cka), which is

the sole Drosophila Striatin protein; and the catalytic subunit of protein phosphatase 2A (PP2A; Drosophila Microtubule Star [Mts]). STRIPAK contains multiple components, some of which are mutually exclusive, and STRIPAK members are part of a variety of regulatory proteins that can direct the pleiotropic PP2A to specific targets (Shi et al., 2016; Ribeiro et al., 2010; Virshup, 2000). In Drosophila and mammals, a STRIPAK-PP2A complex containing Mob4 and Cka was reported to inhibit Hippo signaling (Zheng et al., 2017; Couzens et al., 2013; Ribeiro et al., 2010). We show that PP2A/Mts, with its regulatory subunit Widerborst (Wdb), contributes to NSC quiescence via the inactivation of Akt, an essential component of the InR/PI3K/Akt signaling cascade. Conversely, NSC reactivation requires Mob4 and Cka, which are necessary within STRIPAK for Mts association to Hippo and subsequent Hippo pathway inhibition. These findings suggest a mechanism coordinating Hippo and InR/PI3K/Akt signaling in NSCs, enabling the transition from quiescence to proliferation.

RESULTS

Transcriptome Analysis of Reactivating NSCs: Identification of Mob4, Cka, and PP2A/Mts

To identify the mechanisms regulating NSC reactivation, we performed a small-scale analysis comparing single-cell transcriptomes of guiescent and reactivating NSCs from Drosophila larval brains. By combining grh-Gal4 with UAS-CD8-GFP transgenic lines, cell membranes of approximately one-third of all NSCs (Chell and Brand, 2010) were specifically labeled in vivo. NSCs were individually harvested from 17 h after larval hatching (ALH) brains, when both quiescent (small: diameter 4-5 um) (Ding et al., 2016; Chell and Brand, 2010) and reactivating (enlarged) cells can be easily distinguished. Of the enlarged NSCs, only non-dividing cells without any progeny were harvested. Cells were removed from the second and third thoracic segments of the ventral nerve cords (VNCs), minimizing potential differences from spatial positioning and avoiding retrieving a mix of type I and II NSCs, as the latter are absent from VNCs. Using our single-cell transcriptome protocol (Liu and Bossing, 2016; Bossing et al., 2012), cDNA from each NSC was readily obtained. Quantitative real-time PCRs confirmed that guiescent and reactivating cells expressed the NSC markers deadpan (dpn) and asense (ase), with higher levels in the latter. Single NSC transcriptomes were compared in pairs (three reactivating versus quiescent NSC pairs) on whole-genome Drosophila microarrays (Figure 1A). We used a limma moderated paired t test (Ritchie et al., 2015) to shortlist potential candidates, since the limited sample size did not support false discovery rate (FDR) correction. We identified 196 genes with consistent fold expression changes across all 3 replicates (p < 0.05), of which 145 are upregulated and 51 are downregulated (Figure 1B; Table S1; see Method Details). For quality control, we performed quantitative real-time PCR using independent single NSC samples on a subset of candidates classed mainly into nervous system development and neurogenesis Gene Ontology categories. Up- or downregulated expression for all 18 candidates tested in reactivating versus guiescent NSCs was confirmed, including echinoid (ed) and ras homolog enriched in brain (rheb), which are known to maintain NSC quiescence and promote reactivation, respectively (Ding et al., 2016; Sousa-Nunes et al., 2011) (Figures 1B and 1C).

Using FlyAtlas data (Chintapalli et al., 2007), we noted that our dataset (p < 0.05) is mostly enriched in genes expressed in the larval CNS, whereas among adult tissues, the highest enrichment is seen for genes expressed in ovaries, supporting reported gene sets associated with both NSC and germline stem cell maintenance and growth (Yan et al., 2014) (Figure S1A; Table S2). Most genes have highly conserved mouse (63%) and human orthologs (66%), and only 10% have no mammalian counterpart (Figure S1B; Table S1). When comparing the 175 mouse orthologs identified (single best matches) with transcripts found by previous studies as differentially expressed in guiescent versus activated mouse embryonic (Martynoga et al., 2013) or adult NSCs (Llorens-Bobadilla et al., 2015; Codega et al., 2014) and other stem cell types (Fukada et al., 2007; Venezia et al., 2004), we observed that the overlap is always highest (17-21 targets, 10%-12%) with any of the studies examining NSC transcriptomes (Figure S1C; Table S3). These results suggest that our small-scale singlecell transcriptome analysis generated high-quality data exposing conserved genes that are potentially involved in NSC reactivation. The analysis reveals transcripts encoding for some of the core STRIPAK complex members: mob4 and cka upregulated in reactivating versus guiescent NSCs, whereas mts, encoding the catalytic subunit of PP2A, downregulated (Figures 1C and 1D; Table S1). STRIPAK is involved in a variety of cellular functions (Shi et al., 2016), but it has no known role in NSC reactivation. To functionally test the components identified, we focused initially on Mob4.

Loss of Mob4 Prevents NSC Reactivation

Mob4 is highly expressed in the mammalian and Drosophila CNS (Schulte et al., 2010; Baillat et al., 2001). After validating the differential expression of mob4 detected in NSCs (Figure 1C), we examined its protein levels. Immunostaining of NSCs highlighted with membrane-tagged GFP and Deadpan (Dpn) together with Mob4 antibodies in 17 h ALH brains, revealed higher Mob4 levels in reactivating (enlarged) versus quiescent (small) NSCs (Figures 2A-2C). To investigate the potential function of Mob4 in NSC reactivation, we first examined mob4 null mutants (mob4^{EYΔL3}, hereafter $mob4^{\Delta L3}$), of which 10% survive to third-instar stages (Schulte et al., 2010). NSC (Dpn⁺) membranes labeled with anti-Discs large (Dlg) and mitosis with anti-phospho-histone H3 (pH3) antibodies enabled the scoring of size (maximum diameters) and proliferation. In newly hatched larvae (1 h ALH), no differences are detected between NSCs of mob4 mutants and controls in either brain lobes or VNCs (Figures 2D, 2G, 2J, S2A, S2D, and S2G). All NSCs are guiescent, with the exception of four mushroom body NSCs (MbNSCs) per brain lobe that continuously proliferate from embryonic stages (Ito and Hotta, 1992). However, as early as 4 h ALH, while NSCs in controls start to enlarge, those in mob4 mutants remain small. No NSC mitosis re-entry is detected in either group (Figures 2E, 2H, 2J, S2B, S2E, and S2G). At the end of the first-instar larval stage (24 h ALH), when many NSCs in controls are enlarged and dividing, the reduction in both NSC size and proliferation in mutants is striking, with the only mitotic NSCs corresponding to MbNSCs



Figure 1. Single-Cell Transcriptome Analysis of Reactivating NSCs

(A) Workflow: individual quiescent (Q) and reactivating (R) NSCs expressing *CD8-GFP* driven by *grh-Gal4* were harvested from 17 h ALH CNSs, their mRNA reverse transcribed, and resulting cDNA amplified. Quantitative real-time PCRs confirmed higher ase and *dpn* expression in reactivating versus quiescent NSCs (normalized fold change [log₂FC]; n = 3 NSC reactivating/quiescent pairs; error bars: SEMs; Student's t test, **p < 0.01). NSC transcriptomes were compared on whole-genome microarrays (reactivating versus quiescent; three pairs) and a subset of identified targets validated by quantitative real-time PCRs. ALH, after larval hatching; BL, brain lobe; VNC, ventral nerve cord. Scale bars: 10 μm.

(B) Distribution of identified transcripts according to average fold change expression (x axis; log_2FC) and p value (y axis; limma moderated t test; $-log_{10}$ p value; p < 0.05). See also Table S1 and Figure S1.

(C) Normalized expression levels in reactivating versus quiescent NSCs obtained by quantitative real-time PCR for a subset of targets (log₂FC; n = 3 NSC reactivating/quiescent pairs; error bars: SEMs; Student's t test; ***p < 0.001). The results validate the data from the microarray analysis. Most of the targets selected are classified under "nervous system development" and "neurogenesis" GO terms.

(D) STRING-based interaction network of a *Drosophila* PP2A-STRIPAK complex reported to inhibit Hippo signaling (Zheng et al., 2017; Liu et al., 2016; Ribeiro et al., 2010), highlighting (pink) the components identified in our transcriptome analysis and functionally characterized in this study.

(Figures 2F, 2I, 2J, S2C, S2F, and S2G). 5-Ethynyl-2'-deoxyuridine (EdU) incorporation assays monitoring entry into S phase confirmed that NSCs in *mob4* CNSs are not able to re-enter the cell cycle (Figures S2H–S2J). NSC reactivation defects in *mob4* mutants were similarly observed in brain lobes and VNCs. We focused subsequent studies on brain lobes.

Since niche glial cells are involved in NSC reactivation (Sousa-Nunes et al., 2011; Chell and Brand, 2010) and Mob4 is ubiquitous in the larval CNS (Schulte et al., 2010), we next tested whether Mob4 action is cell autonomous. We ectopically expressed *mob4* specifically in NSCs or in glia of *mob4* mutants using *insc-Gal4* and *repo-Gal4* drivers, respectively. NSCs were analyzed at 18 h ALH, when mitotic reactivation is ongoing. Re-introduction of Mob4 in NSCs of *mob4* mutants rescued both NSC size growth and division to the levels observed in controls (Figures 2K–2M and 2P). We observed a small increase in NSC size and division when Mob4 was expressed from glia, but levels are markedly lower than in controls (Figures 2M–2P). Finally, we inhibited Mob4 specifically in NSCs by expressing *mob4-RNAi* (Schulte et al., 2010) using *insc-Gal4*, resulting in a significant, albeit small, reduction in NSC size and a decrease in NSC division at 18 h ALH (Figures S2K–S2M). We conclude that Mob4 functions primarily cell autonomously to promote NSC reactivation.

Overexpression of Mob4 or Its Human Ortholog Accelerates NSC Reactivation

Mob4 is highly conserved (78% identical at the amino acid level) to its human ortholog MOB4 (hMOB4, also known as Phocein), and ubiquitous expression of *hMOB4* fully rescues the lethality of *mob4* null larvae (Schulte et al., 2010). To determine whether increasing Mob4 or hMOB4 levels can promote NSC reactivation, we overexpressed these specifically in NSCs using *insc-Gal4*. Simultaneous expression of membrane-tagged GFP allowed for NSC size examination, and divisions were labeled with pH3 antibodies. At 6 h ALH, NSCs in controls have yet to re-enter mitosis. At this stage, Mob4 or hMOB4 overexpression results in a premature NSC size increase. No re-entry into division was seen upon Mob4 overexpression, but hMOB4 induced a minor but significant increase (Figures 3A–3C and 3G). At 18 h ALH, no difference in NSC size was detected upon Mob4 over-expression, but more NSCs were found in mitosis compared to



Figure 2. Loss of Mob4 Prevents NSC Mitotic Reactivation

(A–C) Mob4 is upregulated in reactivating versus quiescent NSCs. Examples of quiescent (small, A) and reactivating (enlarged, B) NSCs in 17 h ALH CNSs (VNC thoracic region) labeled with *grh-Gal4* driving *CD8-GFP* (GFP, green), Mob4 (red), and Dpn (blue). Mob4 channel also shown in monochrome. Dashed lines: cell bodies. (C) Mob4 protein quantification in reactivating normalized to quiescent NSCs (reactivating NSCs: n = 50, 8 BLs, 8 brains; quiescent NSCs: n = 50, 8 BLs, 8 brains; error bars: SEMs).

(D–J) NSC enlargement and division are impaired in *mob4*^{ΔL3} mutants. Wild-type (WT; D, 1 h ALH; E, 4 h ALH; F, 24 h ALH) and *mob4*^{ΔL3} brain lobes (G, 1 h ALH; H, 4 h ALH; I, 24 h ALH). NSCs (Dpn, red), cell membranes (Dlg, green), and divisions (pH3, blue). Yellow arrowheads: quiescent NSCs; white arrowheads: reactivated NSCs. Mushroom body NSCs (MbNSCs; dashed circles) are large and do not enter quiescence. At 1 and 4 h ALH, there are no NSC divisions, except in MbNSCs.

(J) Quantification of NSC diameters (1 h ALH: WT n = 359 NSCs, 10 BLs, 5 brains; $mob4^{\Delta L3}$ n = 415 NSCs, 10 BLs, 7 brains; 4 h ALH: WT n = 1,121 NSCs, 14 BLs, 7 brains; $mob4^{\Delta L3}$ n = 552 NSCs, 10 BLs, 5 brains; 24 h ALH: WT n = 799 NSCs, 10 BLs, 7 brains; $mob4^{\Delta L3}$ n = 395 NSCs, 18 BLs, 9 brains) and proliferation (1 h ALH: WT n = 24 BLs, 12 brains; $mob4^{\Delta L3}$ n = 24 BLs, 12 brains; 24 h ALH: WT n = 24 BLs, 12 brains; 24 h ALH: WT n = 24 BLs, 12 brains; $mob4^{\Delta L3}$ n = 24 BLs, 12 brains; 24 h ALH: WT n = 24 BLs, 12 brains; 24 h ALH: WT n = 24 BLs, 12 brains; $mob4^{\Delta L3}$ n = 24 BLs, 12 brains; 24 h ALH: WT n = 24 BLs, 12 brains; $mob4^{\Delta L3}$ n = 24 BLs, 12 brains; 24 h ALH: WT n = 24 BLs, 12 brains; $mob4^{\Delta L3}$ n = 24 BLs, 12 brains; 24 h ALH: WT n = 24 BLs, 12 brains; $mob4^{\Delta L3}$ n = 24 BLs, 12 brains; 24 h ALH: WT n = 24 BLs, 12 brains; $mob4^{\Delta L3}$ n = 24 BLs, 12 brains; 24 h ALH: WT n = 24 BLs, 12 brains; $mob4^{\Delta L3}$ n = 24 BLs, 12 brains; 24 h ALH: WT n = 24 BLs, 12 brains; $mob4^{\Delta L3}$ n = 24 BLs, 12 brains; $mob4^{\Delta L3}$ n = 24 BLs, 12 brains; $mb4^{\Delta L3}$ n = 24 BLs, 12 bra



Figure 3. Overexpression of Mob4 or hMOB4 Increases NSC Growth and Division

(A–G) NSC-specific *mob4* or human *MOB4* (*hMOB4*) overexpression leads to premature NSC enlargement and mitosis entry. Brain lobes of control (A and D, *insc-gal4 > CD8-GFP*) and *mob4* (B and E, *insc-gal4 > CD8-GFP*, *mob4*) or *hMOB4* (C and F, *insc-gal4 > CD8-GFP*, *hMOB4*) overexpressing brains at 6 and 18 h ALH. NSCs in green (GFP) and red (Dpn), and divisions in blue (pH3). Dashed circles: MbNSCs; yellow arrowheads: quiescent NSC examples; white arrowheads: prematurely enlarging (B and C) and dividing NSC examples (D–F). Anterior up. Scale bars: 10 µm.

(G) Quantification of NSC diameters (6 h ALH: *insc-gal4* > *CD8-GFP* n = 1,467 NSCs, 20 BLs, 15 brains; *insc-gal4* > *CD8-GFP*, *mob4* n = 639 NSCs, 10 BLs, 5 brains; *insc-gal4* > *CD8-GFP*, *hMOB4* n = 508 NSCs, 7 BLs, 5 brains; 18 h ALH: *insc-gal4* > *CD8-GFP* n = 1,399 NSCs, 18 BLs, 15 brains; *insc-gal4* > *CD8-GFP*, *mob4* n = 508 NSCs, 7 BLs, 5 brains; 18 h ALH: *insc-gal4* > *CD8-GFP* n = 1,399 NSCs, 18 BLs, 15 brains; *insc-gal4* > *CD8-GFP*, *mob4* n = 508 NSCs, 7 BLs, 5 brains; *insc-gal4* > *CD8-GFP* n = 1,399 NSCs, 18 BLs, 15 brains; *insc-gal4* > *CD8-GFP*, *mob4* n = 1,004 NSCs, 13 BLs, 9 brains; *insc-gal4* > *CD8-GFP*, *hMOB4* n = 533 NSCs, 7 BLs, 5 brains) and proliferation (6 h ALH: *insc-gal4* > *CD8-GFP* n = 56 BLs, 28 brains; *insc-gal4* > *CD8-GFP*, *mob4* n = 22 BLs, 11 brains; *insc-gal4* > *CD8-GFP*, *hMOB4* n = 28 BLs, 14 brains; 18 h ALH: *insc-gal4* > *CD8-GFP* n = 32 BLs, 16 brains; *insc-gal4* > *CD8-GFP*, *mob4* n = 30 BLs, 15 brains; *insc-gal4* > *CD8-GFP*, *hMOB4* n = 20 BLs, 10 brains). Wilcoxon rank-sum tests; ***p < 0.001; p > 0.05: ns.

See also Figure S3.

controls. hMOB4 ectopic expression increased NSC size and induced a similar increase in the number of dividing NSCs as seen upon Mob4 overexpression (Figures 3D-3G). We next tested whether Mob4 overexpression leads to NSC overproliferation. Scoring of NSC divisions in late larval brain lobes (94 h ALH) revealed no differences from controls, indicating that Mob4 overexpression effects are restricted to the NSC reactivation process (Figures S3A-S3C). Finally, since NSC reactivation depends on nutritional stimulus (Britton and Edgar, 1998), we inquired as to whether Mob4 overexpression in NSCs could induce reactivation under diet-restriction conditions. In larvae reared in the absence of dietary amino acids, NSCs overexpressing Mob4 remained quiescent (Figures S3D-S3F). We conclude that increased Mob4 levels accelerate NSC reactivation, and this function may be evolutionary conserved. Yet, Mob4 is not sufficient to bypass the extrinsic nutrition stimulus required for NSC reactivation.

Mob4 Regulates InR/PI3K/Akt and Hippo Signaling Activity in NSCs

The Hippo pathway maintains NSCs in quiescence (Ding et al., 2016; Poon et al., 2016), whereas activation of InR/PI3K/Akt signaling cascade triggers reactivation (Sousa-Nunes et al., 2011; Chell and Brand, 2010). To assess how Mob4 function relates to both pathways, we first examined their activity in the absence of Mob4. Upon activation, insulin receptors recruit PI3K to the cell membrane to convert phosphoinositol(4,5)P2 (PIP2) into phosphoinositol(3,4,5)P3 (PIP3), which in turn recruits the Akt protein kinase through its pleckstrin homology (PH) domain, becoming activated by phosphorylation. This process can be monitored using a PH domain-GFP fusion protein (PH-GFP) binding PIP3 (Britton et al., 2002). We confirmed strong membrane-bound accumulation of PH-GFP in reactivated NSCs (Figures 4A and 4A') (Chell and Brand, 2010). In contrast, NSCs in *mob4* mutants show weak and diffused PH-GFP signals

⁽K–P) Mob4 expression in NSCs, but not in glia, rescues NSC reactivation in $mob4^{\Delta L3}$ mutants to control levels. Brain lobes of control (K, *insc-gal4*), $mob4^{\Delta L3}$ expressing Mob4 in NSCs (L, $mob4^{\Delta L3}$, *insc-gal4* > mob4), $mob4^{\Delta L3}$ (M), $mob4^{\Delta L3}$ expressing Mob4 in glia (N, $mob4^{\Delta L3}$, *repo-gal4* > mob4), and control (O, *repo-gal4*) at 18 h ALH.

⁽P) Quantification of NSC diameters (*insc-gal4* n = 219 NSCs, 3 BLs, 3 brains; $mob4^{\Delta L3}$, insc-gal4 > mob4 n = 669, 10 BLs, 5 brains; $mob4^{\Delta L3}$ n = 520 NSCs, 7 BLs, 7 brains; $mob4^{\Delta L3}$, repo-gal4 > mob4 n = 310 NSCs, 6 BLs, 6 brains; repo-gal4 n = 339, 5 BLs, 5 brains) and proliferation (*insc-gal4* n = 12 BLs, 12 brains; $mob4^{\Delta L3}$, insc-gal4 > mob4 n = 20 BLs, 10 brains; $mob4^{\Delta L3}$ n = 10 BLs, 10 brains; $mob4^{\Delta L3}$, repo-gal4 > mob4 n = 57 BLs, 29 brains; repo-gal4 n = 28 BLs, 14 brains) at 18 h ALH.

Wilcoxon rank-sum tests; **p < 0.01, ***p < 0.001; p > 0.05: non-significant (ns). BLs, brain lobes. Anterior up. Scale bars: 10 μ m. See also Figure S2.



Figure 4. InR/PI3K/Akt Pathway Activation or Hippo Signaling Inhibition Rescues NSC Reactivation in mob4 Mutants

(A–C) InR/PI3K/Akt signaling is strongly reduced in *mob4* NSCs. Expression of pleckstrin homology (PH) domain-GFP fusion (GFP, green) does not accumulate at NSC membranes of *mob4* mutants as in controls. Brain lobes of control (A) and *mob4*^{AL3} mutants (B) at 24 h ALH. NSCs in red (Dpn) and divisions in blue (pH3). GFP channel also shown in monochrome (A' and B'). Inset displays higher magnification (B'). Yellow arrowheads: quiescent NSC examples; white arrowheads: reactivated NSC examples.

(C) Phospho-Akt (pAkt^{S505}) is reduced in *mob4* mutant brains, while total Akt levels are comparable to those in controls (24 h ALH brain extracts; β -actin: loading control). Quantification of protein signals (bottom; error bars: SEMs; n = 3 independent assays; Student's t tests; *p < 0.05; p > 0.05: ns.

(D–E') Hippo signaling remains active in *mob4* NSCs. In contrast to controls, NSCs in *mob4* mutants show no *ban* activity, except in MbNSCs (dashed circles). *ban*-activity sensor, in which decreased GFP signal (green) reflects increased *ban* activity, in brain lobes of control (D and D') and *mob4*^{ΔL3} mutants (E and E') at 24 h ALH. NSCs in red (Dpn) and divisions in blue (pH3). GFP channel also shown in monochrome (D' and E'). Inset showing higher magnification (E').

(F–K) NSC-specific expression of *rheb* activating InR/PI3K/Akt signaling and of *warts (wts)*-RNAi or *hippo (hpo)*-RNAi inactivating Hippo signaling can rescue NSC reactivation in *mob4* mutants. Brain lobes of control (F, *insc-gal4*), *mob4*^{$\Delta L3$} (G), *mob4*^{$\Delta L3$} expressing Rheb in NSCs (H, *mob4*^{$\Delta L3$}, *insc-gal4* > *rheb*), *mob4*^{$\Delta L3$} expressing *wts*-RNAi in NSCs (I, *mob4*^{$\Delta L3$}, *insc-gal4* > *wts*^{*RNAi*}), and *mob4*^{$\Delta L3$} expressing *hpo*-RNAi in NSCs (J, *mob4*^{$\Delta L3$}, *insc-gal4* > *hpo*^{*RNAi*}) at 18 h ALH. NSCs in red (Dpn), cell membranes in green (Dlg), and divisions in blue (pH3). Anterior up. Scale bars: 10 µm and 17 µm in insets.

(K) Quantification of NSC diameters (*insc-gal4* n = 286 NSCs, 4 BLs, 3 brains; *mob4*^{Δ L3} n = 524 NSCs, 7 BLs, 7 brains; *mob4*^{Δ L3}, *insc-gal4* > *rheb* n = 466 NSCs, 6 BLs, 5 brains; *mob4*^{Δ L3}, *insc-gal4* > *wts*^{*RNAi*} n = 543 NSCs, 8 BLs, 5 brains; *mob4*^{Δ L3}, *insc-gal4* > *hpo*^{*RNAi*} n = 531 NSCs, 8 BLs, 5 brains) and divisions (*insc-gal4* n = 12 BLs, 12 brains; *mob4*^{Δ L3} n = 10 BLs, 10 brains; *mob4*^{Δ L3}, *insc-gal4* > *rheb* n = 28 BLs, 14 brains; *mob4*^{Δ L3}; *insc-gal4* > *wts*^{*RNAi*} n = 20 BLs, 10 brains; *mob4*^{Δ L3}, *insc-gal4* > *hpo*^{*RNAi*} n = 60 BLs, 30 brains).

Wilcoxon rank-sum tests; *p < 0.05, **p < 0.01, ***p < 0.001.

(Figures 4B and 4B'). We also observed reduced phosphorylated Akt levels in *mob4* whole CNS lysates compared to controls, whereas total Akt levels were equivalent (Figure 4C). To examine Hippo signaling, we tested the activity of *bantam* (*ban*) microRNA that promotes NSC size growth and division (Ding et al., 2016). In quiescent NSCs, active Hippo signaling prevents *ban* transcription (Ding et al., 2016). We used a *ban* GFP-sensor system, in which GFP signal reduction reflects an increase in *ban* activity (Brennecke et al., 2003). No GFP is observed in control reactivated NSCs demonstrating *ban* activity (Ding et al., 2016) (Figures 4D and 4D'). However, the GFP signal is detected in the NSCs of *mob4* mutants, indicating the absence of *ban* activity

and Hippo pathway activation (Figures 4E and 4E'). We next tested whether activating InR/PI3K/Akt cascade or inhibiting Hippo pathways in the NSCs of mob4 mutants could rescue reactivation defects. Stimulation of target of rapamycin (TOR) signaling by Rheb overexpression activates the InR/PI3K/Akt cascade, promoting premature NSC exit from quiescence (Li et al., 2017; Sousa-Nunes et al., 2011). Overexpressing Rheb in the NSCs of mob4 mutants led to NSC size increase and division re-entry (Figures 4F-4H, and 4K). To inhibit Hippo signaling, we used RNAi against warts (wts) or hippo, which induce earlier NSC reactivation (Ding et al., 2016). In mob4 brains, expression of wts-RNAi or hippo-RNAi in NSCs induced cell size growth and mitosis re-entry (Figures 4F, 4G, and 4I-4K). We conclude that the InR/PI3K/Akt signaling cascade is inhibited, while the Hippo pathway stays active in NSCs upon the loss of Mob4, consistent with NSCs in mob4 mutants being unable to exit quiescence. Activation of InR/PI3K/Akt or inhibition of Hippo pathways can restore reactivation. However, the rescues are partial, with the NSC size and proliferation increase observed in mob4 mutant brains not reaching control levels (Figures 4F-4K). The results contrast with the effect of expressing rheb, hippo-RNAi, or wts-RNAi in a control background, where NSC growth and division surpass control levels (Li et al., 2017; Ding et al., 2016; Sousa-Nunes et al., 2011) and may also reflect insufficient activation or inactivation of the respective signals and/or regulation of both pathways that are essential for effective NSC reactivation.

Mob4 and Cka Cooperate to Reactivate NSCs and Assemble a PP2A-Hippo Complex

Our analysis of single NSC transcriptomes also identified the STRIPAK scaffold protein Cka, which is expressed throughout the CNS (Shi et al., 2016; Chen et al., 2002). Similar to Mob4, we found *cka* transcript and protein upregulated in reactivating versus quiescent NSCs (Figures 1C and S4A–S4C). To examine its function, we overexpressed Cka specifically in NSCs and observed premature NSC enlargement at 6 h ALH (Figures 5A, 5B, and 5G), as well as increased NSC size and divisions at 18 h ALH (Figures 5D, 5E, and 5G). Conversely, the expression of *cka*-RNAi resulted in a small but significant reduction in NSC size and decreased NSC mitosis at 18 h ALH (Figures S4D–S4F). Next, we simultaneously overexpressed Mob4 and Cka in NSCs and observed stronger effects compared to those upon single Mob4 or Cka overexpression (Figures 5A–5G; see also Figures 3B, 3E, and 3G).

STRIPAK negatively regulates Hippo signaling via the dephosphorylation of Hippo kinase by PP2A phosphatase (Couzens et al., 2013; Ribeiro et al., 2010). We examined whether the STRIPAK components Mob4 and Cka are essential for mediating the association of PP2A to Hippo. Co-immunoprecipitations (coIPs) were conducted on S2R⁺ cell lysates expressing FLAGtagged Hippo or control FLAG-NTAN, plus Myc-tagged Mts. In addition, we performed RNAi targeting *mob4* and/or *cka*, which effectively depletes the respective proteins (Figure S5A), using RNAi as a control against *DsRed* targeting red fluorescent protein. FLAG-Hippo co-immunoprecipitates Myc-Mts, as reported (Ribeiro et al., 2010), and no association is found with FLAG-NTAN. However, depletion of Mob4, Cka, or both impairs Hippo/Mts binding, with the latter nearly abolishing association (Figures 5H and 5I). We verified that the inhibition of Mob4 and Cka results in increased Hippo activation, as reported (Zheng et al., 2017; Ribeiro et al., 2010), and that chemical inhibition of PP2A with okadaic acid targeting PP2A, and to a lesser extent PP1 (Takai et al., 1992), leads to Hippo hyperphosphorylation as a positive control (Figure S5B). We conclude that Mob4 and Cka cooperate to promote NSC reactivation and are both required for the association of PP2A to Hippo, leading to its inactivation, which is consistent with the Hippo pathway remaining active in NSCs upon *mob4* loss (Figures 4D–4E').

PP2A Inactivates Akt Independently of STRIPAK Cka and Mob4 Members and Maintains Quiescent NSCs

If PP2A/Mts would only function in NSCs to inactivate Hippo signaling via STRIPAK, then a prolonged NSC quiescence could be anticipated upon Mts inhibition. However, PP2A is also a wellestablished negative regulator of the insulin receptor signaling cascade, including by the dephosphorylation of Akt (Padmanabhan et al., 2009; Vereshchagina et al., 2008; Janssens and Goris, 2001). Using S2R⁺ cells, we observed that Mts inhibition with okadaic acid increases Akt phosphorylation, regardless of RNAi-mediated depletion of cka and mob4 (Figure S6A). In addition, in S2R⁺ cells expressing hemagglutinin (HA)-tagged Mts and Myc-tagged Akt, Mts co-immunoprecipitates with Akt. However, unlike for Mts/Hippo interaction, the depletion of cka and mob4 does not disturb Mts/Akt association, nor does it disturb the levels of phosphorylated Akt with or without insulin stimulation (Figures S6A-S6C). Next, we examined whether impaired Mts function affects NSC reactivation. Since mts null mutants (mts^{XE-2258}) are embryonic lethal (Snaith et al., 1996), we analyzed transheterozygotes harboring an mts hypomorphic allele surviving to pupal stages (mts²⁹⁹) (Wang et al., 2009) and mts^{XE-2258}. Reactivating NSCs in mts²⁹⁹/mts^{XE-2258} mutants shows a mild increased cell size as compared to controls (Figures 6A-6C). To knock down mts specifically in NSCs, we expressed a dominant-negative mts mutant (mts-DN) lacking the N-terminal region of the phosphatase domain (Hannus et al., 2002). Premature NSC size increase and entry into division were observed (Figures 6D, 6E, 6G, 6H, and 6J), strengthening the results using mts transheterozygotes. We then examined whether the regulatory PP2A subunit Wdb, shown to modulate Akt downstream of InR/PI3K/Akt signaling in both vertebrates and invertebrates (Rodgers et al., 2011; Padmanabhan et al., 2009; Vereshchagina et al., 2008), may also function in NSCs. Similar to *mts-DN*, expression of a truncated *wdb* mutant form acting as a dominant negative (wdb-DN) (Hannus et al., 2002) leads to increased NSC size growth at 6 h ALH and a higher number of mitotic NSCs at 18 h ALH (Figures 6D, 6F, 6G, 6I, and 6J). Next, we ascertained whether PP2A/Mts inhibition affects pAkt levels. A premature increase in pAkt is seen in NSCs expressing mts-DN, which indicates abnormal InR/PI3K/Akt activation (Figures 6K-6L'). In this condition, a moderate but significant reduction of ban activity (indicated by ban-GFP sensor signal increase) is also observed (Figures 6M-6O). The results suggest that the inhibition of Mts can promote Hippo signaling activity in NSCs, but the effect is insufficient, possibly due to the availability of endogenous Mob4/Cka levels at this stage, or otherwise



Figure 5. Cka and Mob4 Cooperate to Promote NSC Reactivation and Are Required for PP2A/Hippo Interaction

(A–G) NSC-specific *cka* or *cka* and *mob4* double overexpression leads to premature enlargement and increased mitotic NSCs. Double overexpression results in stronger effects (see also Figure 3). Brain lobes of control (A and D, *insc-gal4 > CD8-GFP*), *cka* (B and E, *insc-gal4 > CD8-GFP*), *cka* (B and E, *insc-gal4 > CD8-GFP*, *cka*, *mob4*) overexpressing brains at 6 and 18 h ALH. NSCs in green (GFP) and red (Dpn), and divisions in blue (pH3). Dashed circles: MbNSCs; yellow arrowheads: quiescent NSCs; white arrowheads: prematurely enlarging (B and C) and dividing NSCs (D–F). Anterior up. Scale bars: 10 µm.

(G) Quantification of NSC diameters (6 h ALH: inscgal4 > CD8-GFP n = 1.106 NSCs. 15 BLs. 10brains; insc-gal4 > CD8-GFP, mob4 n = 639 NSCs, 10 BLs, 5 brains; insc-gal4 > CD8-GFP, cka n = 332 NSCs, 5 BLs, 5 brains; insc-gal4 > CD8-GFP, cka, mob4 n = 284 NSCs, 8 BLs, 8 brains; 18 h ALH: insc-gal4 > CD8-GFP n = 1,717 NSCs. 19 BLs, 14 brains; insc-gal4 > CD8-GFP, mob4 n = 1,004 NSCs, 13 BLs, 9 brains; insc-gal4 > CD8-GFP, cka n = 554 NSCs, 8 BLs, 4 brains; inscgal4 > CD8-GFP, cka, mob4 n = 400 NSCs, 6 BLs, 4 brains) and divisions (6 h ALH: insc-gal4 > CD8-GFP n = 56 BLs, 28 brains; insc-gal4 > CD8-GFP, mob4 n = 22 BLs, 11 brains; insc-gal4 > CD8-GFP, cka n = 18 BLs, 9 brains; insc-gal4 > CD8-GFP, cka, mob4 n = 20 BLs, 10 brains; 18 h ALH: insc-gal4>CD8-GFP n = 70 BLs, 38 brains; insc-gal4 > CD8-GFP, mob4 n = 30 BLs, 15 brains; insc-gal4 > CD8-GFP, cka n = 18 BLs, 9 brains; insc-gal4 > CD8-GFP, cka, mob4 n = 20 BLs, 10 brains). Wilcoxon rank-sum tests, ***p < 0.001; p > 0.05: ns. See also Figure S4. (H and I) Depletion of Mob4 and/or Cka inhibits

(H and I) Depletion of Mob4 and/or Cka inhibits PP2A/Mts association to Hippo.

(H) CoIP assays using S2R⁺ cells expressing Myc-Mts and FLAG-Hippo or control FLAG-NTAN, in addition to RNAi against *mob4* and/or *cka* or control *DsRed* (see also Figure S5). Lysates and FLAG-purified immunoprecipitates analyzed by western blot with indicated antibodies.
(I) Quantification of relative binding of Myc-Mts to FLAG-Hippo shown as a mean of the ratio between Myc-Mts and FLAG-Hippo signal intensities relative to

control (*DsRed* RNAi) levels (n = 3 independent assays; error bars: SEMs; Student's t tests; *p < 0.05 and **p < 0.01).

dominated by InR/PI3K/AKT activation, with the final outcome being premature NSC reactivation. Our data indicate that PP2A/Mts may play a dual role in early postembryonic NSCs: first with Wdb to target Akt contributing to quiescence maintenance and second with STRIPAK components Mob4 and Cka targeting Hippo signaling to promote reactivation (Figure 6P).

DISCUSSION

Neural replenishment depends on the ability of NSCs to tightly control the balance between quiescence and proliferation (Tian et al., 2018; Chaker et al., 2016; Cheung and Rando, 2013). Recent advances in profiling quiescent and activated NSCs are increasing our understanding of these cell states. Most approaches have relied on brain tissue dissociation, cell sorting, and culturing procedures (Llorens-Bobadilla et al., 2015; Codega et al., 2014; Martynoga et al., 2013). Here, we reveal a transcript profile of single quiescent versus reactivating NSC samples

obtained directly from live brains. The analysis of identified individual cells taken directly from living tissues at desired time points allows us to precisely examine the transcriptional control at the crossroads of crucial cell fates. Due likely to the reduced sample number and single-cell cDNA amplification variability (Tung et al., 2017; Macaulay and Voet, 2014) (Pearson's r correlations obtained: guiescent NSCs 0.75 < r < 0.81, mean: 0.77; reactivating NSCs 0.58 < r < 0.67, mean: 0.61), our analysis did not support FDR correction. However, the identified genes meeting significance (limma moderated t test, p < 0.05) show consistent expression changes across replicates, and the regulation of all of the targets tested was independently validated. The high conservation with mammalian genes and partial overlap with orthologs reported to be differentially expressed in mouse quiescent versus activated NSCs suggest that our dataset is also a valuable resource for mammalian NSC research.

Adult NSCs must orchestrate extrinsic signals according to the organism's status with intrinsic factors to transit between



Figure 6. Inactivation of PP2A Phosphatase Results in Premature NSC Reactivation

(A-C) *PP2A/mts* hypomorphic mutants show premature NSC size growth. Brain lobes of WT (A) and *mts²⁹⁹/mts^{XE2258}* mutants (B) at 18 h ALH. NSCs in red (Dpn), cell membranes in green (DIg), and divisions in blue (pH3). Arrowheads: NSC examples.

(C) Quantification of NSC diameters (WT n = 362 NSCs, 5 BLs, 3 brains; mts^{299}/mts^{XE2258} n = 324 NSCs, 6 BLs, 5 brains) and divisions (WT n = 25 BLs, 13 brains; mts^{299}/mts^{XE2258} n = 36 BLs, 27 brains). Wilcoxon rank-sum tests; ***p < 0.001; p > 0.05: ns.

(D–J) NSC-specific expression of dominant-negative (DN) forms of PP2A catalytic subunit Mts or regulatory subunit Wdb results in premature NSC size growth and an increased number of mitotically reactivated NSCs. Brain lobes of control (D and G, *insc-gal4* > *CD8-GFP*), *mts-DN* (E and H, *insc-gal4* > *CD8-GFP*, *mts-DN*), and *wdb-DN* (F and I, *insc-gal4* > *CD8-GFP*, *wdb-DN*) expressing brains at 6 and 18 h ALH. NSC in green (GFP) and red (Dpn), and divisions in blue (pH3). Dashed circles: MbNSCs; yellow arrowheads: quiescent NSC examples; white arrowheads: prematurely enlarging (E and F) and mitotically reactivated NSCs (G–I).

(J) Quantification of NSC diameters (6 h ALH: inscgal4 > CD8-GFP n = 219 NSCs, 5 BLs, 5 brains; insc-gal4 > wdb-DN n = 208 NSCs, 5 BLs, 5 brains; insc-gal4 > mts-DN n = 203 NSCs, 5 BLs, 5 brains; 18 h ALH: insc-gal4 > CD8-GFP n = 541 NSCs, 8 BLs, 5 brains; insc-gal4 > CD8-GFP, wdb-DN n = 373 NSCs, 5 BLs, 5 brains; insc-gal4 > CD8-GFP, mts-DN n = 323 NSCs, 5 BLs, 5 brains) and divisions (6 h ALH: insc-gal4 > CD8-GFP n = 16 BLs, 8 brains; insc-gal4 > CD8-GFP, wdb-DN n = 20 BLs, 10 brains; insc-gal4 > CD8-GFP, mts-DN n = 14 BLs, 7 brains; 18 h ALH: insc-gal4 > CD8-GFP n = 42 BLs, 21 brains; insc-gal4 > CD8-GFP, wdb-DN n = 22 BLs, 12 brains; insc-gal4 > CD8-GFP, mts-DN n = 20 BLs, 10 brains). Wilcoxon rank-sum tests; **p < 0.01, ***p < 0.001; p > 0.05: ns.

(K–O) NSC-specific expression of *mts-DN* results in the increased expression of phosphorylated Akt, as well as a decrease in *ban* activity. (K–L¹) Brain lobes of control (K and K', *insc-gal4* > *CD8-GFP*) and *mts-DN* (L and L', *insc-gal4* > *CD8-GFP*, *mts-DN*) expressing brains at 6 h ALH. NSCs in green (GFP) and red (Dpn), and pAkt^{S505} (blue). pAkt^{S505} also shown in monochrome; insets showing higher magnifications (K' and L').

(M–O) *ban*-activity sensor, in which a decrease in GFP signal (green) reflects an increase in *ban* activity, in brain lobes of control (M and M') and *mts-DN* expressing brains (N and N') at 6 h ALH. NSCs in red (Dpn) and Mts-DN (HA-tag, blue). GFP channel also shown in monochrome; insets showing higher magnification (M' and N').

(O) GFP signal (*ban*-activity sensor) quantification in NSCs expressing *mts-DN* normalized to control NSCs (n = 84 *mts-DN* NSCs, 19 BLs, 14 brains; n = 46 control NSCs, 8 BLs, 5 brains; error bars: SEMs; Wilcoxon rank-sum test; **p < 0.01). Anterior up. Scale bars: 10 μ m and 17 μ m in insets.

(legend continued on next page)

quiescence and proliferation. As in mammals, Drosophila NSCs are dependent on niche signals relaying external stimuli for both quiescence and reactivation (Tian et al., 2018; Chaker et al., 2016). In response to a nutritional cue, niche glia cells activate InR/PI3K/Akt signaling in NSCs to promote reactivation (Sousa-Nunes et al., 2011; Chell and Brand, 2010). Niche glia cells also contribute to quiescence by maintaining Hippo signaling activation in NSCs (Ding et al., 2016). In mammals, the insulin and insulin-like growth factor pathway also plays a major role in adult NSC reactivation (Renault et al., 2009; Kippin et al., 2005; Arsenijevic et al., 2001), and while regulation of the Hippo pathway has not yet been implicated in this process, Hippo signaling maintains liver progenitors in quiescence and is indispensable for skin and intestinal regeneration (Wang et al., 2017; Zhou et al., 2009). Here, we show that Mob4, Cka, and PP2A phosphatase, identified in our transcriptome analysis, regulate NSC quiescence to reactivation states, and we propose that they function as an intrinsic integration mechanism of InR/PI3K/Akt and Hippo signals.

We detected the catalytic subunit of PP2A, Mts, downregulated at the transcript level in reactivating versus quiescent NSCs. Mts maintains NSCs in quiescence, preventing premature phosphorylation of Akt, a key component of the InR/PI3K/Akt signaling cascade. PP2A substrate specificity depends on the choice from a variety of regulatory subunits (Shi et al., 2016). In Drosophila, the regulatory subunit Wdb was shown to physically interact and negatively regulate Akt in ovaries (Vereshchagina et al., 2008), and has also been implicated in the inhibition of insulin signaling, controlling organism growth and metabolic regulation (Fischer et al., 2015). Wdb orthologs in Caenorhabditis elegans (PPTR1) and mammals (B56ß) also dephosphorylate Akt to modulate InR/PI3K/Akt, indicating a conserved role (Rodgers et al., 2011; Padmanabhan et al., 2009). We demonstrate that the inhibition of Mts or of Wdb leads to similar premature NSC reactivation effects, suggesting that Wdb/Mts function together to maintain quiescence. PP2A has been linked to cellular quiescence in different contexts. In the developing Drosophila eye and wing, PP2A/Wdb contributes to a quiescent state upon terminal cell differentiation (Sun and Buttitta, 2015); in cycling human cells, PP2A is also required for stable quiescence, a function that is dependent on the B56 γ subunit (Naetar et al., 2014). Thus, PP2A may also have an evolutionary conserved function in maintaining quiescence in NSCs, modulating the InR/PI3K/Akt signaling cascade. PP2A is a pleiotropic phosphatase. In proliferating Drosophila NSCs, it contributes to apicalbasal polarity and prevents excess self-renewal at later larval stages. Here, Wdb was shown to play no role and instead Twins, a B55 subunit ortholog, regulated PP2A/Mts action (Chabu and Doe, 2009; Krahn et al., 2009; Ogawa et al., 2009; Wang et al., 2009).

In contrast to Mts, we found that Mob4 and Cka upregulated in reactivating versus quiescent NSCs. Both are scaffold proteins of STRIPAK, a large molecular complex that is highly conserved from fungi to humans containing PP2A (Shi et al., 2016). We demonstrate that the loss of Mob4 or Cka impairs NSC reactivation, while their overexpression can accelerate it. Furthermore, ectopic expression of human Mob4 also induced premature NSC size growth and mitosis entry, suggesting a conserved function. In cultured S2 cells, Mob4 was shown to focus spindle fibers during mitosis (Trammell et al., 2008). Our Mob4 analysis in NSCs exposed a function in cell size growth before mitosis and additionally showed that MbNSCs, which do not enter quiescence, continue dividing in the absence of Mob4, indicating that the role of Mob4 in NSC reactivation is independent of that reported in spindle fibers.

STRIPAK/PP2A associates with Hippo in Drosophila and mammalian cells, and restricts Drosophila Hippo kinase activity via dephosphorylation (Liu et al., 2016; Couzens et al., 2013; Ribeiro et al., 2010). Previous reports revealed cross-talk inhibition between Hippo and InR/PI3K/Akt pathways in both mammalian and Drosophila tissues (Straßburger et al., 2012; Tumaneng et al., 2012). We demonstrate that Mob4 and Cka are both required for the physical association of Mts to Hippo and its subsequent inhibition, as reported (Ribeiro et al., 2010). We also show that upon loss of Mob4, the Hippo pathway consistently remains switched on in NSCs, and InR/PI3K/Akt signaling is inhibited. Finally, we determined that the inhibition of Mts can enhance Hippo signaling in NSCs but that the effect is overcome by premature activation of InR/PI3K/Akt, resulting in earlier NSC reactivation, despite Hippo activity. Our data suggest that as the levels of STRIPAK members Mob4 and Cka increase in NSCs, a complex with Hippo kinase assembles recruiting PP2A/Mts protein to inactivate Hippo signaling. This may function as an intrinsic molecular switch to turn off Hippo signaling and allow the InR/PI3K/Akt cascade to turn on (Figure 6P). Given their large and versatile composition, it is not surprising that STRIPAK complexes are assigned to an increasing number of functions and linked to clinical conditions, including autism and cancer (Shi et al., 2016). It will be important to determine whether and how STRIPAK proteins contribute to regulating the reactivation of other stem cells.

STAR***METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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- CONTACT FOR REAGENT AND RESOURCES SHARING
- EXPERIMENTAL MODEL AND SUBJECT DETAILS O Drosophila strains and husbandry
- S2R+ cell culture, transfection and drug treatment METHOD DETAILS
- NSC transcriptome analysis
- Immunohistochemistry and EdU incorporation

⁽P) A model of action of STRIPAK Mob4/Cka/PP2A members: in quiescent NSCs, Mob4 and Cka levels are low, while PP2A (Mts/Wdb) phosphatase targets Akt, ensuring that InR/PI3K/Akt signaling is maintained switched off. Hippo signaling is active (Ding et al., 2016; Poon et al., 2016). Mob4 and Cka levels increase, promoting NSC size growth and entry into division; both are required to direct Mts to switch off Hippo signaling, while the InR/PI3K/Akt cascade becomes active (Sousa-Nunes et al., 2011; Chell and Brand, 2010).

- Image acquisition and processing
- dsRNA synthesis
- Co-immunoprecipitations and western blotting
- QUANTIFICATION AND STATISTICAL ANALYSIS
- DATA AND SOFTWARE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/ j.celrep.2019.05.023.

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AUTHOR CONTRIBUTIONS

All of the authors designed and performed the experiments. J.G.-R. and C.S.B. wrote the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR*METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit anti-GFP	Gift from U. Mayor (U. Mayor, personal communication)	N/A
Chicken anti-GFP	Millipore	Cat# 06-896, RRID: AB_310288
Guinea pig anti-Dpn	Gift from J. Knoblich (Levy and Larsen, 2013)	RRID: AB_2314299
Guinea pig anti-Mob4	Gift from T. Littleton (Schulte et al., 2010)	N/A
Mouse anti-Dlg	DSHB	Cat# 4F3 anti-discs large, RRID: AB_528203
Rabbit anti-pH3	Abcam	Cat# ab5176, RRID: AB_304763
Rabbit anti-Cka	Gift from W. Du (Weng and Wei, 2002)	N/A
Rabbit anti-pAKT ^{S505}	CST	Cat# 4054, RRID: AB_331414
Mouse anti-Flag clone M2	Sigma	Cat# F3165, RRID: AB_259529
Rat anti-HA clone 3F10	Roche	Cat# 11867431001, RRID: AB_390919
Rat IgG	Sigma	Cat# I4131, RRID: AB_1163627
Rabbit anti-Akt	CST	Cat# 9272, RRID: AB_329827
Rabbit anti-b-Actin	CST	Cat# 4967, RRID: AB_330288
Mouse anti-Myc clone 9E10	Santa Cruz Biotechnology	Cat# sc-40, RRID: AB_627268
Rabbit anti-pMST1 ^{T183} /pMST2 ^{T180}	CST	Cat# 3681S, RRID: AB_330269
Guinea pig anti-Hippo	Gift from G. Halder (Hamaratoglu et al., 2006)	N/A
Chemicals, Peptides, and Recombinant Proteins		
Okadaic Acid	CST	Cat# 5934
Critical Commercial Assays	'	
Click-iT EdU Alexa Fluor 594 Imaging Kit	Invitrogen	Cat# C10339
Co-Immunoprecipitation Kit	Pierce	Cat# 26149
Deposited Data		
Transcriptome data	This paper	GEO: GSE128646
Experimental Models: Cell Lines		
D. melanogaster: Cell line S2R+	Gift from B. Houdsen	FlyBase: FBtc0000150
Experimental Models: Organisms/Strains		
<i>D. melanogaster: mob4</i> mutant: y[1] w[*]; Mob4[EYDeltaL3]/CyO	Bloomington Drosophila Stock Center (Schulte et al., 2010)	BDSC: 36331; FlyBase: FBst0036331
D. melanogaster: UAS line expressing mob4 RNAi: P{UAS-Mob4.RNAi.JS1}attP2	Bloomington Drosophila Stock Center (Schulte et al., 2010)	BDSC:36488; FlyBase: FBst0036488
<i>D. melanogaster</i> : UAS line expressing <i>mob4</i> : y[1] v[1]; P{y[+t7.7] v[+t1.8] = UAS-Mob4.S}attP2	Bloomington Drosophila Stock Center (Schulte et al., 2010)	BDSC: 36329; FlyBase: FBst0036329
<i>D. melanogaster</i> : UAS line expressing <i>hMOB4</i> : y[1] v[1]; P{y[+t7.7] v[+t1.8] = UAS-phocein.1}attP2	Bloomington Drosophila Stock Center (Schulte et al., 2010)	BDSC: 36330; FlyBase: FBst0036330
D. melanogaster: UAS line expressing cka-eGFP: w[*]; P{w[+mC] = UASp-Cka.EGFP.C}2	Bloomington Drosophila Stock Center	BDSC: 53756; FlyBase: FBst0053756
<i>D. melanogaster</i> : UAS line expressing <i>wdb-DN</i> : w[*]; P{w[+mC] = UAS-wdb.95-524.HA}6	Bloomington Drosophila Stock Center (Hannus et al., 2002)	BDSC: 55053; FlyBase: FBst0055053
<i>D. melanogaster</i> : UAS line expressing <i>wts</i> RNAi: y[1] sc[*] v[1]; P{y[+t7.7] v[+t1.8] = TRiP. GI 01331)attP2	Bloomington Drosophila Stock Center (Ding et al., 2016)	BDSC: 41899; FlyBase: FBst0041899
Continued

REAGENT or RESOURCE	SOURCE	IDENTIFIER	
<i>D. melanogaster</i> : UAS line expressing <i>hpo</i> RNAi: y[1] v[1]; P{y[+t7.7] v[+t1.8] = TRiP.HMS00006}attP2	Bloomington Drosophila Stock Center (Ding et al., 2016)	BDSC: 33614; FlyBase: FBst0033614	
<i>D. melanogaster</i> : UAS line expressing <i>cka</i> RNAi: y[1] v[1]; P{y[+t7.7] v[+t1.8] = TRiP.HM05138}attP2	Bloomington Drosophila Stock Center	BDSC: 28927; FlyBase: FBst0028927	
D. melanogaster: Wild type: Oregon-R	Gift from M. Akam (Bossing and Technau, 1994)	FlyBase: FBsn0000276	
<i>D. melanogaster</i> : UAS line expressing <i>rheb</i> : w[*]; P{w[+mC] = UAS-Rheb.Pa}3	Gift from R. Sousa-Nunes (Sousa-Nunes et al., 2011)	BDSC: 9689; FlyBase: FBst0009689	
D. melanogaster: Gal4 line under the control of grh: Grh-Gal4	Gift from A.H. Brand (Chell and Brand, 2010)	N/A	
D. melanogaster: UAS line expressing mts-DN-HA: UAS-mts.dn181-HA	Gift from S. Eaton (Hannus et al., 2002)	N/A	
<i>D. melanogaster</i> : line expressing pleckstrin homology domain-GFP fusion protein: <i>PH-GFP</i> (tGPH)	Gift from B. Edgar (Britton and Edgar, 1998)	N/A	
D. melanogaster: bantam GFP-sensor line, ban-sensor (db20)	Gift from S.M. Cohen (Brennecke et al., 2003)	N/A	
<i>D. melanogaster: mts^{XE225839}</i> mutant: mts ^{XE2258} /CyO, P{sevRas1.V12}F1 and <i>mts²⁹⁹</i> mutant	Gifts from H. Wang (Wang et al., 2009)	BDSC: 5684; FlyBase:FBst0005684 N/A for <i>mts²⁹⁹</i>	
<i>D. melanogaster</i> : Gal4 line under the control of <i>insc</i> : w[*]; P{w[+mW.hs] = GawB}insc[Mz1407]	Bloomington Drosophila Stock Center	BDSC: 8751; FlyBase:FBst0008751	
<i>D. melanogaster</i> : Gal4 line under the control of <i>repo</i> : w[1118]; P{w[+m*] = GAL4}repo/TM6, tb	Gift from A. Hidalgo	N/A	
D. melanogaster: UAS line expressing CD8-GFP: y[1] w[*]; P{w[+mC] = UAS-mCD8::GFP.L}LL5, P{UAS-mCD8::GFP.L}2	Lee and Luo, 1999	BDSC: 5137; FlyBase:FBst0005137	
D. melanogaster: UAS line expressing dicer2: UAS-dicer2	Ding et al., 2016	N/A	
Oligonucleotides			
Primer: Anchored polyT AAGCAGTGGTATCAAC GCAGAGTACT ₍₂₆₎ VN	Bossing et al., 2012	N/A	
Primer: SM AAGCAGTGGTATCAACGCAGAG TACGCrGrGrG	Bossing et al., 2012	N/A	
Primer: Nested AAGCAGTGGTATCAACGCAGAGT	Bossing et al., 2012	N/A	
Primers used for RT-qPCR	See Table S4	N/A	
dsRNA targeting sequence <i>mob4</i> : Forward: TAATAC GACTCACTATAGGGagatgtggaagtacgagcacctg	Schulte et al., 2010	N/A	
dsRNA targeting sequence <i>mob4</i> : Reverse: TAATACG ACTCACTATAGGGagatgcgagaagatgcgatacac	Schulte et al., 2010	N/A	
dsRNA targeting sequence <i>cka</i> : Forward: TAATACG ACTCACTATAGGGatacgggtccagttctgtgc	This paper	N/A	
dsRNA targeting sequence <i>cka</i> : Reverse: TAATACG ACTCACTATAGGGtgttgtaggccaccacgata	This paper	N/A	
dsRNA targeting sequence <i>DsRed</i> : Forward: TAATAC GACTCACTATAGGGgccgatgaacttcaccttgt	This paper	N/A	
dsRNA targeting sequence <i>DsRed</i> : Reverse: TAATAC GACTCACTATAGGGcgaggacgtcatcaaggagt	This paper	N/A	
Recombinant DNA			
Plasmid: 12XCSL DsRedExpressDL	Hansson et al., 2006	Addgene plasmid #47683	
Plasmid: Flag-NTAN	Gift from P. Ribeiro (Ribeiro et al., 2010)	N/A	

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Continued					
REAGENT or RESOURCE	SOURCE	IDENTIFIER			
Plasmid: Flag-Hippo	Gift from P. Ribeiro (Ribeiro et al., 2010)	N/A			
Plasmid: Myc-Mts	Gift from P. Ribeiro (Ribeiro et al., 2010)	N/A			
Plasmid: HA-Mts	Gift from P. Ribeiro (Ribeiro et al., 2010)	N/A			
Plasmid: Myc-Akt	Gift from W. Hongyan (Li et al., 2014)	N/A			
Software and Algorithms					
R/Bioconductor Limma	Ritchie et al., 2015	https://bioconductor.org/packages/release/ bioc/html/limma.html			
STRING v10.5	Szklarczyk et al., 2015	https://string-db.org/cgi/input.pl?sessionId= QMsQ2cmXKFYZ&input_page_show_ search=on			
FlyAtlas	Chintapalli et al., 2007	http://flyatlas.org/atlas.cgi			
DIOPT – DRSC Integrative Ortholog Prediction Tool	Hu et al., 2011	https://www.flyrnai.org/cgi-bin/DRSC_ orthologs.pl			

CONTACT FOR REAGENT AND RESOURCES SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Claudia Barros (claudia.barros@plymouth.ac.uk).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Drosophila strains and husbandry

Drosophila stocks obtained from the Bloomington Drosophila Stock Center are: mob4^{EYAL3} (36331) (Schulte et al., 2010) rebalanced over CyO, P(GAL4-twi.G)2.2; UAS-mob4^{RNAi} (36488) (Schulte et al., 2010); UAS-mob4 (36329) (Schulte et al., 2010); UAS-hMOB4 (36330) (Schulte et al., 2010); UAS-cka-eGFP (53756); UAS-wdb-DN (UAS-wdb.95-524.HA; 55053) (Hannus et al., 2002); UASwts^{RNAi} (41899) (Ding et al., 2016); UAS-hpo^{RNAi} (33614) (Ding et al., 2016) and UAS-cka^{RNAi} (28927). Other stocks used are: Wildtype Oregon-R (kind gift from M. Akain); UAS-rheb (Blomington 9689) (Sousa-Nunes et al., 2011) (kind gift from R. Sousa-Nunes); grh-Gal4 (Chell and Brand, 2010) (kind gift from A.H. Brand); UAS-mts-DN (UAS-mts.dn181-HA) (Hannus et al., 2002) (kind gift from S. Eaton); PH-GFP (tGPH) (Britton and Edgar, 1998) (kind gift from B. Edgar); ban-sensor (db20) (Brennecke et al., 2003) (kind gift from S.M. Cohen), mts²⁹⁹ and mts^{XE2258} (Wang et al., 2009) (kind gifts from H. Wang). NSC-specific RNAi and overexpression assays were performed using insc-Gal4 (w¹¹¹⁸; p{GAWB}inscMZ1407) and glial-specific expression assays used repo-Gal4 (w¹¹¹⁸; p{GAWB}repo/TM6b, iab-lacZ). grh-Gal4 driver was recombined with UAS-CD8-GFP. For rescue experiments, insc-Gal4 and repo-Gal4 drivers were recombined or combined with the mob4^{EYAL3} mutant strain. For other assays, the insc-Gal4 driver was recombined with UAS-CD8-GFP and/or combined with UAS-dicer2 (Ding et al., 2016). Fly lines were kept in standard Drosophila fly food. Egg collections and larvae rearing were performed on agar juice plates (21 g agar, 200ml of grape juice per I of water) supplemented with yeast paste. Egg lays were collected in either 30min or 1h time-windows. For nutritional deprivation experiments, freshly hatched larvae were transferred to agar plates prepared with amino-acid free media (5% sucrose, 1% agar in phosphate buffered saline, PBS).

S2R+ cell culture, transfection and drug treatment

S2R+ cells (kind gift from B. Houdsen) were maintained in 25- or 75-cm² T-flasks at 25°C in Schneider's Medium (GIBCO) with 10% heat-inactivated FBS (One Shot, GIBCO) and antibiotics. For transient transfections, 1.6x10⁶ cells/well were seeded in 6-well plates. Effectene transfection reagent (Quiagen) was used to transfect 1 µg and/or 2 µg of each appropriate plasmid and/or dsRNA, respectively, following manufacturer guidelines. Cells were incubated 72 hours before harvest. Plasmids used are Flag-NTAN, Flag-Hippo, Myc-Mts, HA-Mts (kind gifts from P. S. Ribeiro) and Myc-AKT (kind gift from W. Hongyan). For okadaic acid experiments, cells were transfected as above, incubated 70 hours and treated with 50 nM okadaic acid (CST) or 0.005% DMSO (vehicle; Corning) for 2 hours prior harvest.

METHOD DETAILS

NSC transcriptome analysis

Single NSC harvest, mRNA isolation, cDNA generation and microarray hybridization were performed essentially as previously described (Bossing et al., 2012). Single quiescent (small; 4-5µm) and reactivating (enlarged) NSCs were individually removed from freshly dissected 17 ALH CNS expressing membrane-tagged GFP specifically in NSCs (grh-Gal4, UAS-CD8-GFP). Samples with any trace of non-fluorescent material were rejected. Each single cell was expelled in its own Eppendorf tube containing annealing mix: 0.3 µL anchored polyT primer (5'-AAGCAGTGGTATCAACGCAGAGTACT(26)VN-3', 10pM), 0.3 µL SM primer (5'-AAGCAGTGG TATCAACGCAGAGTACGCrGrGrG-3', 10pM), 0.4 µL RNase inhibitor (Superase, Ambion) and 2 µL Lysis Mix (10% Nonidet P-40, 0.1M DTT in DEPC-treated ultrapure water), and processed in less than 20 min. Each sample was spun (14000rpm, 1min, 4°C), primers annealed (3min, 70°C) and snap-frozen in dry ice/ isopropanol. 1.5 µL of mix 1 (1 µL Invitrogen first strand buffer, 0.5 µL 10mM dNTPs) and 0.5 µL of mix 2 (3 µL Invitrogen Superscript II reverse transcriptase, 0.5 µL Ambion Superase RNase inhibitor) were added per sample. Samples were thawed during centrifugation (14000rpm, 1 min, 4°C) and reverse transcribed (37°C, 90min), followed by enzyme thermal inactivation (65°C, 10 min). RNA was digested in 2 µL digestion mix (0.7 µL Roche RNase H buffer, 0.5 µL Roche RNase H, 0.8 µL ultrapure water) for 20min at 37°C, followed by enzyme thermal inactivation (65°C, 15 min). For cDNA PCR amplification, 2 μL of nested primer (5'-AAGCAGTGGTATCAACGCAGAGT-3'), 2 μL dNTPs (10mM), 5 μL Roche buffer, 0.5 µL Roche Long Expand polymerase and 34.5 µL of ultrapure water were added. PCR program: one cycle (95°C 3min, 50°C 5min, 68°C 15min) followed by 25 cycles (95°C 20 s, 60°C 1min, 68°C 7min). 3 pairs of NSC quiescent/ reactivating samples showing clear banding patterns on agarose gels were sent for microarray analysis (FlyChip, University of Cambridge). 1 µg of each sample were Klenow-labeled using BioPrime DNA Labeling System (Invitrogen) in the presence of Cy3- or Cy5-dCTP (GE Healthcare) for 2 hours 37°C. Unincorporated dye and nucleotides were removed using AutoSeg G-50 columns (GE Healthcare), following manufacturer instructions. Cy3- and Cy5-labeled pairs of samples were combined with salmon sperm DNA as blocking agent and co-hybridized (16 hours, 51°C) in a HybStation hybridization station (Digilab Genomic Solutions) on long oligonucleotides FL003 microarrays (International Drosophila Array Consortium; Gene Expression Omnibus accession number GPL14121). Post-hybridization washes were performed according to Full Moon Biosystems protocols. Detailed protocols for labeling, hybridization and washing can be requested from the Cambridge Systems Biology Centre UK (https://www.sysbiol.cam/ac.uk/CSBC). Arrays were scanned at 5 µm resolution (GenePix scanner, Axon Instruments) using optimized PMT gain settings for each channel.

RT-qPCR validation of selected genes was done using SYBRGreen on a StepOnePlus thermal cycler (Applied Biosystems) and primers indicated in Table S4. *ribosomal protein 49 (rp49)* was used as internal calibrator for all reactions. Single NSC cDNA samples used were obtained as described above. Candidates validated were also selected based on their Gene Ontology (GO) *Nervous system development* and *Neurogenesis* classification (STRING v10.5) (Szklarczyk et al., 2015).

Tissue-specific expression of identified targets was performed using FlyAtlas (Chintapalli et al., 2007). Gene orthology analysis used DIOPT (DRSC Integrative Ortholog Prediction Tool) (Hu et al., 2011). Protein-protein interaction network of *Drosophila* PP2A-STRIPAK components was performed using STRING (v10.5) (Szklarczyk et al., 2015) with experimental-based data only as source, and as previously described (Zheng et al., 2017; Liu et al., 2016; Ribeiro et al., 2010).

Immunohistochemistry and EdU incorporation

Immunohistochemistry assays were performed as previously described (Chell and Brand, 2010), with minor modifications. Briefly, larval CNSs were dissected in PBS and fixed for 20 min in 4% formaldehyde/PBS with 5 μ M MgCl₂ and 0.5 μ M EGTA or 10 μ M MgCl₂ and 1 μ M EGTA (3rd instar larvae), followed by washes in PBS (2 × 10 min, 3 rinses between washes) and block for 1h in PBST (PBS, 1% Triton X-100) with 10% fetal bovine serum (FBS). Primary antibodies were incubated in PBST overnight or for 2 nights at 4°C. CNSs were washed in PBST and secondary antibodies incubated 2h at room temperature, followed by PBST washes and sequentially embedding in 50% and 70% glycerol before mounting in a 1:1 mix of 70% glycerol and Vectashield (Vector Laboratories). Antibodies used are: rabbit anti-GFP (1:1000, kind gift from U. Mayor), chicken anti-GFP (1:500, Millipore), guinea pig anti-Dpn (1:2000, kind gift from J. Knoblich), guinea pig anti-Mob4 (1:1000, kind gift from T. Littleton), mouse anti-Dlg (1:50, DSHB), rabbit anti-pH3 (1:1000, Abcam), rabbit anti-Cka (1:1000, kind gift from W. Du), rabbit anti-pAKT^{S505} (1:50, CST) and rat anti-HA clone 3F10 (1:1000, Roche). EdU incorporation assays were performed as previously described (Sousa-Nunes et al., 2011). Briefly, CNSs were dissected in PBS and incubated in 10 μ M EdU/PBS for 1h at room temperature. CNSs were fixed for 15min in 4% formaldehyde/PBS and incorporated EdU detected using Click-iT EdU Imaging kit following manufacturer instructions (Invitrogen).

Image acquisition and processing

Images were obtained on a Leica SP8 confocal laser-scanning microscope using LAS X software. Quantifications were made using z stacks of 1.5 µm step size, comprising whole brain lobes, VNCs or CNSs. Representative images shown are single optical sections, with the exception of Figure 1A, which is a z-projection stack (3 steps, 0.5 µm each), and EdU incorporations, which are z-projection stacks encompassing whole CNSs. Images were processed in Fiji v2.0 or Adobe Photoshop CS6 and assembled in Adobe Illustrator CS6. NSC sizes (maximum diameters) (Chell and Brand, 2010), pH3 scorings, Mob4 and Cka signal intensities (pixel intensity/ NSC maximum area), *ban*-GFP signal intensity (pixel intensity/ NSC maximum area outlined by Dpn staining) and EdU voxel quantification were performed using Fiji v2.0 or Adobe Photoshop CS6.

dsRNA synthesis

For *cka*, *mob4* and *DsRed* dsRNA, DNA templates of target genes were PCR amplified from larval genomic DNA or 12XCSL-DsRedExpressDL plasmid (Addgene) to include the T7 promoter sequence on both ends. Primers used are: dsRNAmob4_Fwd: 5'-TAATACGACTCACTATAGGGagatgtggaagtacgagcacctg-3'(Schulte et al., 2010), dsRNAmob4_Rev: 5'-TAATACGACTCACTA TAGGGagatgcgagaagatgcgatacac-3'(Schulte et al., 2010), dsRNAcka_Fwd: 5'-TAATACGACTCACTATAGGGatacgggtccagttct gtgc-3', dsRNAcka_Rev: 5'-TAATACGACTCACTATAGGGtgttgtaggccaccacgata-3', dsRNADsRed_Fwd: 5'-TAATACGACTCACTA TAGGGgccgatgaacttcaccttgt-3', dsRNADsRed_Rev:

5'-TAATACGACTCACTATAGGGcgaggacgtcatcaaggagt-3'. The size of DNA bands was confirmed, purified using QIAquick Gel Extraction Kit (QIAGEN) and used as template for dsRNA synthesis. *In vitro* transcriptions were performed using MEGAscript T7 kit (Invitrogen), incubated for 6 hours at 37°C and treated with TURBO DNase (Invitrogen) for 15 min at 37°C. RNA was precipitated using LiCl precipitation solution (Invitrogen) and re-hydrated in ultrapure water. dsRNA was annealed by incubation at 65°C 30 min and cooled down to room temperature.

Co-immunoprecipitations and western blotting

S2R+ cells were harvested and lysed in lysis buffer (25mM Tris, 0.15M NaCl, 1mM EDTA, 1% NP-40, 5% glycerol; pH 7.4) supplemented with protease inhibitor (Complete, EDTA-free; Sigma) and phosphatase inhibitors (cocktails B+C; Santa Cruz Biotechnology). Cell extracts were spun at 14000 rpm for 30 min at 4°C and proteins quantified (BCA protein assay, Pierce). Using the Pierce Co-immunoprecipitation kit (Pierce), 20 µg of anti-Flag M2 (Sigma), anti-HA (3F10; Roche), or rat IgG (Sigma) were immobilized in 50 µL of AminoLink Plus Coupling resin slurry following manufacturer instructions. Protein lysates were incubated in the resin on a rotator at 4°C overnight, washed 4 times with PBS and eluted following manufacturer instructions. Detection of proteins was performed using standard SDS-PAGE and western blotting using ECL or ECL Plus chemiluminescent substrate (Pierce). Antibodies used are: rabbit anti-Akt (1:500, CST), rabbit anti-pAkt^{S505} (1:500, CST), rabbit anti-β-Actin (1:1000, CST), mouse anti-Flag Clone M2 (1:3000, Sigma), mouse anti-Myc clone 9E10 (1:500, Santa Cruz Biotechnology), rabbit anti-Cka (1:5000, kind gift from W. Du), guinea pig anti-Mob4 (1:5000, kind gift from T. Littleton), rabbit anti-pMST1^{T183}/pMST2^{T180} (1:500, CST), guinea pig anti-Hippo (1:5000, kind gift from G. Halder) and rat anti-HA clone 3F10 (1:3000, Roche).

QUANTIFICATION AND STATISTICAL ANALYSIS

Transcriptome data: all genes with raw signal intensity values below 150 were removed from the analysis, generating a matrix containing 2455 genes. Quantile normalization (Bolstad et al., 2003) across all samples was performed using R/Bioconductor limma package (Ritchie et al., 2015). Remaining genes were analyzed with limma, by fitting a linear model. Adjusting *p*-values with False Discovery Rate (FDR) did not reach statistical significance. Instead, a limma moderated paired t test was employed. Targets with expression fold changes with associated p < 0.05 values were used for subsequent analysis, including expression validation. Expression of selected candidate genes assayed by RT-qPCR was quantified using the Livak method (Livak and Schmittgen, 2001).

Other statistics were performed using SigmaPlot Version 12.5 (Systat software): Shapiro-Wilk and equal variance tests used to evaluate normality; Student's t test applied when data fitted a normal distribution; Wilcoxon rank-sum test used for non-parametric data; p < 0.05 considered significant. Data from *Drosophila in vivo* assays were obtained from a minimum of two biological replica sets; sample numbers are indicated in figure legends. Cell culture/ biochemistry results derive from a minimum of three independent assays. Histograms show mean \pm standard error of the mean. Boxplots represent 25th and 75th percentiles, black line indicates median, red line specifies mean, whiskers indicate 10th and 90th percentiles.

DATA AND SOFTWARE AVAILABILITY

Processed transcriptome data is shown in Table S1. Raw transcriptome data has been deposited in the Gene Expression Onmibus (GEO) public database under ID code GSE128646.

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Supplemental Information

STRIPAK Members Orchestrate Hippo and Insulin

Receptor Signaling to Promote

Neural Stem Cell Reactivation

Jon Gil-Ranedo, Eleanor Gonzaga, Karolina J. Jaworek, Christian Berger, Torsten Bossing, and Claudia S. Barros

Supplemental Figures and Figure Text



Figure S1, related to Figure 1. Tissue expression enrichment and orthology conservation of identified targets in transcriptome analysis. (A) Heatmap depicting identified larval and adult tissue-specific gene sets enriched by a minimum of 2-fold to expression in whole fly. gl: gland. Sperm: spermatheca; acc: accessory; TAG: thoracicoabdominal ganglion. See also Table S2. (B) Number of human and mouse orthologues (single best matches) of identified genes grouped by orthology score (DIOPT) (Hu, et al., 2011). See also Table S1. (C) Number of identified genes with mouse orthologues (single best matches) reported upregulated in quiescent or proliferating mouse embryonic or adult NSCs, skeletal muscle satellite stem cells (MuSCs) and hematopoietic stem cells (HSC). DIOPT score groups indicated. See also Table S3. The data used for the above assays comprise targets identified in our transcriptome analysis as up- or downregulated in reactivating (R) versus quiescent (Q) NSCs (limma moderated *t*-test, p<0.05).



Figure S2, related to Figure 2. NSC reactivation defects upon Mob4 loss or inhibition. (A-G) NSC enlargement and division is impaired in *mob4* mutant ventral nerve cords (VNCs). VNCs of WT (A, 1h ALH; B, 4h ALH; C, 24h ALH) and *mob4*^{Δ L3} (D, 1h ALH; E, 4h ALH; F, 24h ALH). NSCs in red (Dpn), cell membranes in green (Dlg), divisions in blue (pH3). Yellow and white arrowheads indicate quiescent and reactivated NSC examples, respectively. Scale bar: 10µm. Anterior up. (G) Quantification of NSC diameters (1h ALH: WT n=316 NSCs, 7 VNCs; *mob4*^{Δ L3} n=171 NSCs, 5 VNCs. 4h ALH: WT n=700 NSCs, 6 VNCs; *mob4*^{Δ L3} n=331 NSCs, 5 VNCs. 24h ALH: WT n=584 NSCs, 5 VNCs; *mob4*^{Δ L3} n=88 NSCs, 8 VNCs) and proliferation (1h ALH: WT n=12 VNCs; *mob4*^{Δ L3} n=12 VNCs). (H-J) NSCs in *mob4* mutant larval brains do not enter S-phase, except the MbNSCs. WT (H) and *mob4*^{Δ L3} (I) CNSs at 24hph Edu-labelled (red). (J) Quantification of Edu⁺

voxels from CNSs, normalized to controls (WT n=7 CNSs, $mob4^{\Delta L3}$ n=8 CNSs; error bars: s.e.m). Scale bar: 50µm. (K-M) NSC-specific expression of mob4-RNAi results in a small reduction in cell size and decreased number of NSCs in division. Brain lobes (BLs) of control (K, *insc-gal4>CD8-GFP*) and *mob4*-RNAi expressing brains (L, *insc-gal4>CD8-GFP, mob4*^{RNAi}) at 18h ALH. NSCs in green (*CD8-GFP*, GFP) and red (Dpn), divisions in blue (pH3). Arrowheads: NSC examples. Anterior up. Scale bar: 10µm. (M) Quantification of NSC diameters (*insc-gal4>CD8-GFP*, n=684 NSCs, 9 BLs, 9 brains; *insc-gal4>CD8-GFP, mob4*^{RNAi} n=614 NSCs, 8 BLs, 8 brains) and divisions (*insc-gal4>CD8-GFP*, n=46 BLs, 23 brains; *insc-gal4>CD8-GFP, mob4*^{RNAi} n=28 BLs, 14 brains). Wilcoxon rank sum tests, ***p<0.001, *p*>0.05: non-significant (ns).



Figure S3, related to Figure 3. Mob4 overexpression does not lead to NSC overproliferation nor induces NSC reactivation under nutrition restriction. NSC-specific *mob4* overexpression does not affect NSC proliferation in late larval brains (A-C) nor promotes reactivation of NSCs in larvae deprived of amino acids (sucrose-only diet; D-F). Brain lobes (BLs) of control (A, D, *insc-gal4>CD8-GFP*) and *mob4* overexpressing brains (B, E, *insc-gal4>CD8-GFP, mob4*) at 94h (A, B) and 18h ALH (D, E). NSCs in green (GFP) and red (Dpn), divisions in blue (pH3). Dashed line: central brain region. White arrowheads: dividing NSC examples. Yellow arrowheads: quiescent NSC examples. Dashed circles: MbNSCs. Scale bar: 10µm. Anterior up. (C) Quantification of NSC divisions (94h ALH: *insc-gal4>CD8-GFP* n=10 BLs, 10 brains; *insc-gal4>CD8-GFP*, *mob4* n=10 BLs, 10 brains). (F) Quantification of NSCs, 5 BLs, 5 brains) and divisions (18h ALH: *insc-gal4>CD8-GFP* n=200 NSCs, 6 BLs, 5 brains; *insc-gal4>CD8-GFP* n=22 BLs, 16 brains; *insc-gal4>CD8-GFP*, *mob4* n=30 BLs, 16 brains). Wilcoxon rank sum tests, *p*>0.05: non-significant (ns).



Figure S4, related to Figure 5. Cka inhibition delays NSC growth and division. (A-C) Cka is upregulated in reactivating (R) compared with guiescent (Q) NSCs. Examples of guiescent (small; A) and reactivating (enlarged, B) NSCs in 17h ALH brains (VNC thoracic region) labelled with grh-Gal4 driving CD8-GFP (GFP, green), Cka (red) and Dpn (blue). Cka channel also shown in monochrome. Dashed lines: cell bodies. (C) Cka protein quantification in reactivating normalised to quiescent NSCs (n= 20 reactivating NSCs and n=20 quiescent NSCs, 20 BLs, 10 brains; error bars: s.e.m.; Wilcoxon rank sum test, *p<0.05). (D-F) NSC-specific expression of cka-RNAi results in reduced cell size and decreased number of NSCs in division. Brain lobes (BLs) of control (D, insc-gal4>CD8-GFP) and cka-RNAi expressing brains (E, insc-gal4> CD8-GFP, cka^{RNAi}) at 18h ALH. NSCs in green (CD8-GFP, GFP) and red (Dpn), divisions in blue (pH3). Arrowheads: NSC examples. Anterior up. Scale bar: 10µm. (F) Quantification of NSC diameters (insc-gal4>CD8-GFP, n=584 NSCs, 8 BLs, 4 brains; insc-gal4> CD8-GFP, cka^{RNAi} n=658 NSCs, 8 BLs, 4 brains) and divisions (insc-gal4>CD8-GFP, n=17 BLs, 9 brains; insc-gal4> CD8-GFP, cka^{RNAi} n=20 BLs, 10 brains). Wilcoxon rank sum tests, ***p<0.001.



Figure S5, related to Figure 5. Inhibition of Mob4 and Cka increase Hippo phosphorylation. (A) Efficiency of mob4 and cka RNAi-mediated depletions used in Drosophila S2R+ assays. dsRNAs targeting mob4 or cka, but not DsRed control, lead to depletion of Mob4 and Cka proteins. Lysates analysed with indicated antibodies. β-Actin: loading control. Quantification of Mob4 and Cka signal intensities normalised to control (*DsRed* RNAi) levels (lower panel; n=3 independent assays; error bars: s.e.m; Student's t-tests, ***p<0.001). (B) Mob4/ Cka depletion leads to increased levels of activated (phosphorylated) Hippo (pHippo) in S2R+ cells, consistent with published studies (Zheng, et al., 2017; Liu, et al., 2016; Ribeiro, et al., 2010). Drosophila S2R+ cells treated with dsRNAs targeting mob4 and cka or control DsRed, as well as in the presence of vehicle (0.0005% DMSO) or Okadaic acid (OA) as a positive control for hippo phosphorylation. Lysates analysed by western-blot with indicated antibodies. β-Actin: loading control. Note the total Hippo band mobility shift due to hyperphosphorylation in OA-treated samples. Quantification of pHippo levels shown as mean of the ratio between pHippo and total Hippo signal intensities relative to control (*DsRed* RNAi) levels (lower panel; n=3) independent assays; error bars: s.e.m; Student's t-tests, p*<0.05, p>0.05: nonsignificant, ns).



Figure S6, related to Figure 5. Inhibition of Mob4 and Cka does not affect Akt phosphorylation nor the association of PP2A/Mts to Akt. (A, B) Depletion of *mob4* and *cka* has no effect in the levels of activated (phosphorylated) Akt (pAkt^{S505}) in S2R+ cells (A) with or without stimulation with Insulin (B). (A) *Drosophila* S2R+ cells treated with dsRNAs targeting *mob4* and *cka* or control *DsRed*, as well as in the presence of Okadaic acid (OA) as a positive control for Akt phosphorylation or vehicle (V). Lysates analysed with indicated antibodies. β-Actin: loading control. Quantification of pAkt levels shown as mean of the ratio between pAkt and total Akt signal intensities relative to control (*DsRed* RNAi) levels (lower panel; n=3 independent assays). (B) RNAi-mediated depletion of *mob4* and *cka*, or control *DsRed*, in S2R+ cells treated with Insulin or vehicle. Lysates analysed by westernblot with indicated antibodies. β-Actin: loading of pAkt levels shown as mean of pAkt/ Akt signal intensity ratios relative to control (*DsRed* RNAi) levels (right panel; n=3 independent assays). (C) Co-IP assays using S2R+ cells expressing Myc-Akt and HA-Mts, in addition to RNAi against *mob4* and *cka* or control *DsRed*. Lysates and HA-purified immunoprecipitates analysed by westernblot with indicated antibodies. Negative control co-IP performed using rat IgG instead of rat anti-HA antibody. Quantification of relative binding of Myc-Akt to HA-Mts shown as mean of the ratio between Myc-Akt and HA-Mts signal intensities relative to control (*DsRed* RNAi) levels (lower panel; n=3 independent assays). Error bars: s.e.m. Wilcoxon rank-sum tests, p>0.05: non-significant (ns).

Supplemental Table

Gene (Symbol)	Forward Primer (5'→3')	Reverse Primer (5'→3')	Source
ase	CACCTACCAACTGCTGACG	GCTGCTGCTGCTAATGTTG	This paper
dpn	CGCTATGTAAGCCAAATGGATGG	CTATTGGCACACTGGTTAAGATGG	(Berger, et al., 2012)
rheb	TGAGGTGGTGAAGATCATATACGAA	GCCAGCTTCTTGCCTTCCT	(Zitserman, et al., 2012)
cka	GGAGACGGAAGGCGTCAT	TCTCGTCGTCGGACATC	This paper
CG10903	GAGTCTCGGTTGATTTTGGACA	TCTCCCAGAATGACATCCCCA	This paper
asf1	GGGCGACACATCTTTGTCTTC	GCAGGTAAGCAGAACAATGGTAA	This paper
Gbeta13F	TGGTGGCTATCTATCGTGCTG	GCCCAAAAACGAGGTTACCTG	This paper
phax	ATGATGGAACTGCACGCAAAT	CAGGTGGTAAGGGGACTGG	This paper
NiPp1	ATGGCTAACAGCTACGACATACC	TGTTGCGACCAAATAGATAGCAT	This paper
mob4	TGGGCACGATCAGATTCTCC	CATCTTCTCGCACGCCTACT	This paper
CrC	GAAAACTGGGAGGATACGTGG	GAGAGGTCTGAATGCCTTTGTC	This paper
bet3	ATGTCACGACAAGCCTCTCG	GAGTGCTCCGTAGGTGAGT	This paper
ed	GATGAGCTCCTGTTCTCCGG	GTTGGAATCGCAATGGTCGG	This paper
pdp1	AATCCCCATTACCAGCGCAA	GGCATTCCCATTCGATCCCT	This paper
how	AACTTTGTCGGTCGCATTTT	CGTCCTCCTTCTTCTTGTCG	This paper
p120ctn	AACATGGACCTTTCATTGACGC	ATATCCTGCTGCCGAAAATTGA	This paper
Rip11	TGGAGTCCGACGCACTGTA	CAATGGTGACGAAGCAGTTGT	This paper
l(2)35Df	CATCGAAAGAAGCTACATCCTCC	GTGGGTTCGTCATCTGCATTAT	This paper
nito	ACAAGAAGTTTGGCGATTTTAGC	CTTCAGGCGTTCGGAAGCAA	This paper
mts	TCCAGTTCCATAAGAGCCGC	CACGATCGCAATGTGGTCAC	This paper
rp49 (calibrator)	GCTAAGCTGTCGCACAAATG	GTTCGATCCGTAACCGATGT	(Kohyama- Koganeya, et al., 2008)

Table S4, related to STAR Methods. Primers used for real-time quantitativePCR assays.