Investigating the Genetic Interaction of dLRRK/LRRK2 with the Visual System and the Stability of LRRK2 Protein in a Drosophila model of Parkinson’s Disease

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Abstract

The G2019S mutation within the kinase domain of Leucine-rich repat kinase 2 (LRRK2) is the most common reported and studied cause of familial Parkinson’s Disease (PD). LRRK2 has been associated with many cellular pathways, but its exact role is still to be elucidated. Using Drosophila as a model for PD, the role of the protein is examined.

In this project, we found that dLRRK loss-of-function (dLRRK$^{LOF}$) flies had an age-dependent visual deficit that was rescued with the expression of the native gene (dLRRK) in non-neuronal cells (fat body and pigment cells) bolstering the suggestion of protein’s ability to transport cell-cell. Furthermore, placing mutations of different eye colour genes encoding proteins normally involved in vesicle trafficking in the dLRRK$^{LOF}$ background caused a loss of viability in combination which suggests a role for dLRRK in regulating pigment granules specifically and in the lysosomal pathway in general.

In a gain-of-function approach, young flies expressing LRRK2-G2019S in dopaminergic (DA) neurons and pigment cells demonstrated an aberrant electroretinogram (ERG) indicating a failure in their ability to adapt to light. This was partially complemented by anatomical assessment where expressing LRRK2-G2019S in pigment cell caused flies to have larger deep pseudopupil (DPP) than the controls indicating defective eye function.

Finally, mutations in LRRK2 are thought to potentially affect the stability of the protein in fly model as well as in cells. We used two kinase inhibitors (vitamin B$_{12}$ and MLi-2) with different mode of actions to test their effect on LRRK2 stability. We showed that these two inhibitors might variably affect LRRK2 stability dependent on the mutation on the protein.
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Declaration

I hereby declare that, unless explicitly stated, all of the work presented within this thesis is a presentation of original work and I am the sole author. This work has not previously been presented for an award at this, or any other, University. All sources are acknowledged as References.
## List of Abbreviations

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<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>♀</td>
<td>Female</td>
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<tr>
<td>♂</td>
<td>Male</td>
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<tr>
<td>3-HK</td>
<td>3-hydroxykynurenine</td>
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<tr>
<td>4E-BP</td>
<td>4E-binding protein</td>
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<td>6-OHDA</td>
<td>6-hydroxydopamine</td>
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<td>6-PTP</td>
<td>6- pyruvoyltetrahydropterin</td>
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<td>a.u.</td>
<td>arbitrary unit</td>
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<td>Alternating Current</td>
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<td>ALS</td>
<td>Amyotrophic Lateral Sclerosis</td>
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<td>ANK</td>
<td>ankyrin repeat</td>
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<td>ANOVA</td>
<td>Analysis Of Variance</td>
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<td>Adaptor Protein 3</td>
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<td>Armadillo-like Repeat</td>
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<td>Adenosine Triphosphate</td>
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<td>Blood Brain Barrier</td>
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<td>BCA</td>
<td>Bicinchoninic Acid</td>
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<td>BDSC</td>
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<td>BH4</td>
<td>Tetrahydrobiopterin</td>
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<td>Biogenesis of Lysosome-related Organelles Complex</td>
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<td>BMP</td>
<td>Bone Morphogenetic protein</td>
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<tr>
<td>MPP+</td>
<td>1-methyl-4-phenylpyridinium</td>
</tr>
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</table>
MPTP 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine
ms milliseconds
mV millivolt
MW Molecular Weight
NHP Non-Human Primate
NM Neuromelanin
NSAIDs Non-Steroidal Anti-Inflammatory Drugs
NSF N-ethylmaleimide Sensitive Fusion
OMIM Online Mendelian Inheritance in Man
ONL Outer Nuclear Layer
OPL Outer Plexiform Layer
or orange
PBS Phosphate Buffered Saline
PCR Polymerase Chain Reaction
PD Parkinson's Disease
PDA Pyrimidodiazepine
PINK1 PTEN Induced Kinase 1
PKA Protein Kinase A
PPA Pseudopupil Analysis
pr purple
PRDX-3 Peroxiredoxin-3
pu punch
PVDF Polyvinylidene Fluoride
R Photoreceptors
rb ruby
RGCs Retina Ganglion Cells
Rh Rhodopsin
RIPA Radioimmunoprecipitaion Assay
RNA Ribonucleic Acid
RNAi RNA interference
ROC Ras of Complex Proteins
ROS Reactive Oxygen Species
RPE Retinal Pigment Epithelium
RT Room Temperature
RU486 Mifepristone (Roussel-Uclaf 486)
s scarlet
SDS-PAGE Sodium Dodecyl Sulfate-Polyacrylamide Gel Electrophoresis
se sepia
SEM Standard Error of the Mean
SNCA α-synuclein
SNpc Substantia Nigra pars compacta
sptr sepiapterin reductase
STN Subthalamic Nucleus
t1/2 half-life
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Full Form</th>
</tr>
</thead>
<tbody>
<tr>
<td>TAE</td>
<td>Tris-Acetate-EDTA</td>
</tr>
<tr>
<td>TBS-T</td>
<td>Tris Buffered Saline-Tween</td>
</tr>
<tr>
<td>TC</td>
<td>Tetracycline</td>
</tr>
<tr>
<td>TGFbeta</td>
<td>Transforming Growth Factor beta</td>
</tr>
<tr>
<td>TGN</td>
<td>Trans-Golgi Network</td>
</tr>
<tr>
<td>TH</td>
<td>Tyrosine Hydroxylase</td>
</tr>
<tr>
<td>$T_m$</td>
<td>Melting Temperature</td>
</tr>
<tr>
<td>TRIM1</td>
<td>Tripartite Motif Family 1</td>
</tr>
<tr>
<td>UAS</td>
<td>Upstream Activated Sequence</td>
</tr>
<tr>
<td>UCA</td>
<td>Urscholanic Acid</td>
</tr>
<tr>
<td>UDCA</td>
<td>Ursodeoxycholic Acid</td>
</tr>
<tr>
<td>$v$</td>
<td>vermilion</td>
</tr>
<tr>
<td>$w$</td>
<td>white</td>
</tr>
<tr>
<td>WB</td>
<td>Western Blot</td>
</tr>
<tr>
<td>WT</td>
<td>Wild Type</td>
</tr>
</tbody>
</table>
1 Introduction

1.1 Parkinson’s Disease (PD)

PD was first described by James Parkinson, an English physician, in his 66-page monograph “An Essay on the Shaking Palsy” published in 1817. The disease was later named after him in respect of his pioneering work by Jean-Martin Charcot who added more details to Parkinson’s observations and established it as a neurological disorder distinctive from palsy or paralysis (Goetz, 1986).

PD is the second most common neurodegenerative disorder after Alzheimer’s disease (AD). It affects around 145,000 people in the UK (Parkinson’s UK, 2018) and around 6 million globally (Dorsey et al., 2018). Age is the main risk factor and disease prevalence increases by up to 4% in people aged 60 and older. Males are more prone to develop PD than females with no clear explanation; however, some reports suggest a neuroprotective effect for oestrogen (de Lau and Breteler, 2006).

1.1.1 Clinical manifestations

PD is still being diagnosed by the subject manifesting what is known as classical motor features that were defined over 200 years ago. These are bradykinesia (slowing of movement), resting tremor and rigidity. The first symptom is present in all patients while the latter two are present in the majority. Other features might include postural reflex disturbances and postural instability (Obeso et al., 2017). Although PD is considered a motor disorder, surprisingly, almost all patients report at least one non-motor symptoms (e.g., visual and/or olfactory dysfunction) and currently these symptoms contribute toward the certainty of the diagnosis (Kim et al., 2013b) (Figure 1.1). In theory, these symptoms can help in predicting the probability of an individual developing PD because some of them (e.g., alterations in olfactory acuity) appear years before the manifestation of the motor symptoms and such symptoms can help as well in the possibility of discovering biomarkers for PD. However, the difficulty is in distinguishing PD-related symptoms from normal individuals that have them as a normal part of aging (Krishnan et al., 2011). The range of reported non-motor symptoms vary from as high as 90% such as those observed with sleep disorders, constipations and olfactory deficits to as low as 20% - 40% in depression, anxiety and visual disturbances (Pfeiffer, 2016).
Figure 1.1 PD non-motor symptoms

Some of the non-motor symptoms reported in PD patients with the red symptoms indicating higher incidence than others. Data obtained form (Davidsdottir et al., 2005, Fasano et al., 2015, Pfeiffer, 2016).

1.1.2 Neuropathology

Underlying the motor symptoms of PD is a progressive selective moderate to severe loss of dopaminergic (DA) pigmented neurons in the substantia nigra pars compacta (SNpc) in the midbrain region. The substantia nigra connects to the striatum, which is part of the basal ganglia and includes the caudate and putamen. In doing so, they establish the nigrostriatal dopamine pathway, which is hypothesized to be important in movement facilitation. Moreover, the substantia nigra performs activities other than motor control. It is also thought to be involved in a variety of other activities and behaviours, such as learning, drug addiction, and emotion (Kandel et al., 2014). The cause of this loss is still under investigation but it is the most consistent neuropathological observation found in all individuals diagnosed clinically with PD and can be examined clearly macroscopically or microscopically post mortem. Interestingly, the ventral tegmental region (VTA), where the cell bodies of the mesolimbic DA neurons are located next to the SNpc, is significantly less impacted in PD (Uhl et al.,
Another diagnostic feature reported in PD, though with less frequency than neuronal degeneration, is the presence of neuronal inclusions called Lewy bodies (LBs). These consists of misfolded protein aggregates with \(\alpha\)-synuclein making up the majority of the aggregate. The case for LBs as a major cellular pathology in PD is a little controversial. On one hand, (Braak et al., 2003) has developed a staging method for PD progression based on the spread of LB lesions in different parts of the brain and this is very helpful in diagnosis. On the other hand, LB is not specific for PD because it can be seen in other neurodegenerative disorders and normal individuals. Interestingly, the majority of individuals with Lewy pathologies in their brains are Alzheimer’s patients, where up to 60% have these inclusions (Dickson et al., 2009, Halliday et al., 2011). (Figure 1.2) illustrates the difference in DA neurons loss in SNpc in healthy and PD patients. Nonetheless, mis-sense mutations in \(\alpha\)-synuclein that pre-dispose the protein to aggregation, or mutations that increase the concentration of \(\alpha\)-synuclein via gene duplication or triplication cause PD (reviewed in (Waxman and Giasson, 2009)).
Figure 1.2 PD neuropathology in the substantia nigra

Schematic representation of a normal nigrostriatal pathway (A) and a PD one (B). The substantia nigra pars compacta (SNpc) contain DA neurons that extend to the basal ganglia and synapse in the striatum i.e., putamen and caudate nucleus. Transverse brain sections indicate that the SNpc is normally pigmented in control brains while the loss of the dark-brown pigment neuromelanin is seen in PD brains. (C) Immunohistochemical labelling of midbrain dopaminergic neurons for α-synuclein and ubiquitin, identifying Lewy bodies. Taken from (Dauer and Przedborski, 2003).

An important question that the PD community is still looking for an answer to is why DA neurons, particularly in the SNpc, are more prone to insult than other neurons in PD? One hypothesis suggests that their anatomy could be the reason. A lengthy and heavily branched unmyelinated axon with an unusual number of transmitter release sites distinguishes DA neurons in the SNpc. In rodents, SNpc DA neurons contain axons that branch abundantly in the striatum and up to 200 000 vesicular release sites (Matsuda et
This unique morphology put them at a disadvantage in that they simply have “too many mouths to feed” and thus making them especially vulnerable to stressors of any sort, including environmental and genetic factors (Bolam and Pissadaki, 2012). For example, mitochondrial oxidant stress, one of the potential drivers of neurodegeneration, is increased in the axon of SNpc DA neurons and this stress is alleviated by reducing the size of the axonal arborization (Pacelli et al., 2015). In addition, α-synuclein is primarily a synaptic protein, the extraordinarily extensive axonal arborization of SNpc DA neurons is extremely likely to increase its expression, increasing the risk of α-synuclein aggregation and incidence of disease such as PD (Zharikov et al., 2015). However, some additional factors may be at play as not all neurons with long, branching axons are sensitive in PD, such as striatal cholinergic interneurons (Zhou et al., 2002). Another potential cause of the selective degeneration of SNpc DA neurons might be the surrounding environment around their cell bodies. The substantia nigra neuropil, which is made up of axon projections from the striatum and globus pallidus, stains highly for calbindin D28K, and most dopaminergic cell bodies are found inside this calbindin-rich neuropil. However, in PD, vulnerable neurons tend to be in calbindin-poor portions of the substantia nigra (Dauer and Przedborski, 2003). Other suggested cause of this selectivity might include dopamine toxicity, iron content, and autonomous pacemaking (Giguère et al., 2018).

1.1.3 Aetiology

PD is a complex disorder with less than 10% of cases attributed to hereditary origin while the majority of cases are sporadic (Thomas and Beal, 2007). Although aging could be the greatest risk to eventually developing PD, at present, the evidence supports an interplay of environmental and genetic factors in disease development. Many environmental and genetic factors have been linked to PD since the accidental discovery of a PD-like illness caused by 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP) in 1983 (Langston et al., 1983) and the discovery of the first gene causing a familial form of PD which is α-synuclein (SNCA) in 1997 (Polymeropoulos et al., 1997). MPTP itself is not toxic but when it enters the brain it is converted into 1-methyl-4-phenylpyridinium (MPP+) which is taken up by the DA neurons. In the DA neurons, MPP+ is toxically concentrated in the mitochondria which leads to cell death, eventually (Langston, 2017). Exposure to pesticides such as paraquat and rotenone have been reported to increase PD risk. This is not surprising since good animal models were produced using paraquat and rotenone but this increased risk has not been reported in all studies. Additionally, head
injury (especially in those with repeated injury) and those involved in farming and living in rural areas which might increase the likelihood of pesticide exposure have been found to have an increased PD risk. Interestingly, drinking coffee and alcohol and cigarette smoking reduce the risk of PD. This could be due to the presence of caffeine in coffee and nicotine in cigarette which were found to be neuroprotective in animal models (Goldman, 2014, Hernán et al., 2002, Noyce et al., 2012).

Since the discovery of the first gene associated with PD i.e., SNCA, more genes and loci have been discovered and associated with PD. They have been designated with the names PARK genes and were numbered in chronological order of discovery. For example, SNCA is called PARK1; however, this naming scheme is a bit misleading because PARK4 refers to SNCA as well but with different type of mutation. Furthermore, some of the PARK genes have an unconfirmed link to PD like PARK2, 5, 11, 13, 18, 21. Hence, the Movement disorder Society Task Force for the Nomenclature Genetic Movement Disorders have recommended a new naming scheme based on clinical grounds into i) classical PD, ii) early-onset PD with similarity to classical PD and iii) atypical parkinsonism (Marras et al., 2016) to alleviate some of the confusion that might rise due the old system (Table 1.1).
Table 1.1 List of monogenic PD causes

<table>
<thead>
<tr>
<th>Old Name</th>
<th>New Name</th>
<th>Inheritance</th>
<th>OMIM*</th>
<th>Clinical clues</th>
</tr>
</thead>
<tbody>
<tr>
<td>PARK1 &amp; 4</td>
<td>PARK-SNCA</td>
<td>Autosomal Dominant</td>
<td>168601</td>
<td>Missense mutations cause classical parkinsonism. Duplication or triplication of this gene cause early onset parkinsonism with prominent dementia.</td>
</tr>
<tr>
<td>PARK8</td>
<td>PARK-LRRK2</td>
<td></td>
<td>607060</td>
<td>Clinically typical PD</td>
</tr>
<tr>
<td>PARK17</td>
<td>PARK-VPS35</td>
<td></td>
<td>614203</td>
<td>Clinically typical PD</td>
</tr>
<tr>
<td>PARK2</td>
<td>PARK-PARKIN</td>
<td>Autosomal Recessive</td>
<td>600116</td>
<td>Often presents with dystonia, often in a leg.</td>
</tr>
<tr>
<td>PARK6</td>
<td>PARK1-PINK1</td>
<td></td>
<td>605909</td>
<td>Psychiatric features common.</td>
</tr>
<tr>
<td>PARK7</td>
<td>PARK-DJ1</td>
<td></td>
<td>606324</td>
<td></td>
</tr>
<tr>
<td>PARK9</td>
<td>PARK-ATP13A2</td>
<td>Autosomal Recessive</td>
<td>606693</td>
<td>Kufor-Rakeb syndrome with parkinsonism and dystonia.</td>
</tr>
<tr>
<td>PARK14</td>
<td>PARK-PLA2G6</td>
<td>Autosomal Recessive</td>
<td>612953</td>
<td>PLA2G6-associated neurodegeneration (PLAN): dystonia, parkinsonism, cognitive decline, pyramidal signs, psychiatric symptoms (adult phenotype), ataxia (childhood phenotype)</td>
</tr>
<tr>
<td>PARK15</td>
<td>PARK-FBXO7</td>
<td>Autosomal Recessive</td>
<td>260300</td>
<td>Early onset parkinsonism with pyramidal signs.</td>
</tr>
<tr>
<td>PARK19</td>
<td>PARK-DNAJC6</td>
<td></td>
<td>615528</td>
<td>Occasional mental retardation and seizures.</td>
</tr>
<tr>
<td>PARK20</td>
<td>PARK-SYNJ1</td>
<td></td>
<td>615530</td>
<td>May have seizures, cognitive decline, abnormal eye movements, and dystonia.</td>
</tr>
</tbody>
</table>

Adapted from (Marras et al., 2016). *Online Mendelian Inheritance in Man

Genome-wide association studies (GWAS) have detected risk genes for PD, demonstrating that monogenic and sporadic PD are not distinct entities, and that many genes may cooperate to influence downstream shared targets. Variation in the glucocerebrosidase gene (GBA1) is the prototypic genetic risk factor for PD. The gene was originally linked to autosomal recessive Gaucher’s disease, a lysosomal storage disorder. This was discovered because family members of Gaucher’s disease patients had an elevated risk of PD, and many were found to be heterozygous GBA1 variant carriers (Halperin et al., 2006, Tayebi et al., 2001). Adaptive immune system genes, mitochondrial genes, and genes involved in dopamine metabolism are among the other routes for PD that have been discovered by genetic study. Five genes make up the
PARK16 locus, but RAB7L1 has received the most attention due to its expression in the brain and PD susceptibility. It has been demonstrated that RAB7L1 interacts with LRRK2 and VPS35, and it appears to be involved in endosomal-lysosomal trafficking (MacLeod et al., 2013). The genetic variation of the microtubule-associated protein tau (MAPT) and idiopathic PD have been linked, according to many genome-wide association studies of PD (Pankratz et al., 2012, Skipper et al., 2004). Figure 1.3 shows the molecular function of PD-associated genes and their neuronal subcellular locations.

**Figure 1.3 Molecular mechanisms implicated in PD according to the genetic findings**

This figure shows a DA neuron (green), an axon of a presynaptic glutamatergic cortical neuron (blue), and a dendritic spine of a medium spiny neuron (yellow). The subcellular locations of genes reported to be associated in sporadic PD and associated pathogenic models. Genes linked to sporadic PD are most frequently found in the mitochondria, vesicular trafficking organelles, the Golgi network, and lysosomes and endosomes. Taken from (Trinh and Farrer, 2013).
1.1.4 Pathogenesis

There has been a number of mechanisms implicated in PD pathogenesis such as protein misfolding, mitochondrial dysfunction, and neuroinflammation. α-synuclein misfolding and aggregation which lead to the formation of LBs is a pathological feature found in most PD. The mechanism of this misfolding and aggregation is elusive but perhaps the native structure of α-synuclein in brains, that is largely an unstructured monomer, make it more liable to aggregate than a protein with defined stable structure (Burré et al., 2013). Moreover, α-synuclein has been shown to act in a prion-like manner capable of spreading from cell to cell and cause toxicity (Ma et al., 2019). As the illness advances, the PD-brain experiences progressive modifications, with LBs impacting susceptible neural circuits, according to research done in 2003 by Braak and colleagues (Braak et al., 2003). Based on the progression of neuropathological alterations they noticed, they developed a staging system, arguing that neuronal damage doesn’t occur randomly but rather develops after a predefined chain of pathological incidents. They classified the pathological evolution associated with PD into six phases (Figure 1.4), stage 1 includes the regions afflicted at the onset of the illness. Stage 1 lesions begin in the anterior olfactory nucleus and dorsal motor nucleus of the glossopharyngeal and vagal nerves (lower brainstem). The illness then advances stepwise in an upward direction toward the midbrain and forebrain nuclei, eventually impacting the neocortex in its latter stages (Braak et al., 2003).
Figure 1.4 Braak staging system of PD pathology

The staging suggests that α-synuclein pathology starts in the olfactory bulb and brainstem regions (Braak stages 1 and 2) and, subsequently, the Lewy pathology spreads into the midbrain and the basal forebrain (Braak stages 3 and 4) before finally reaching cortical regions (Braak stages 5 and 6). Taken from (Doty, 2012).

Mitochondrial dysfunction is a recurrent theme in sporadic and familial PD pathology and has been linked to genetic and environmental risk factors (Figure 1.5). A defect in the mitochondrial complex-I has been shown in studies of MPTP, paraquat, and rotenone exposure and has given a clue to the association of mitochondrial dysfunction with PD. Also, mitochondrial complex-I abundance has been reported to be reduced in post-mortem PD patients’ brain. Death in DA neurons due to energy depletion caused by this dysfunction could explain this link (Moon and Paek, 2015). Additionally, the discovery of PTEN induced kinase 1 (PINK1) and the PRKN involvement in mitophagy gave another clue of mitochondrial dysfunction theme in PD. A loss of function mutation in either genes have led to impaired mitochondria production and degradation which might lead to developing PD (Pickrell and Youle, 2015).
Figure 1.5 Multiple roles of mitochondria in Parkinson’s disease

Mitochondrial integrity may change as a result of familial PD genes (autosomal recessive or dominant). Through various mitochondrial functions like bioenergetics, dynamics, transport, and quality control, genetic and environmental factors may be involved in the pathogenesis as well. Taken from (Exner et al., 2012).

An interesting mechanism that could be involved in PD is neuroinflammation. Studies of PD risk factors and in patients pre and post-mortem have suggested an involvement of neuroinflammatory process in PD pathology either directly in neurons damage or indirectly in respond to the damage (Hirsch and Hunot, 2009). For example, an outbreak can occur within the brain, then stressed neurons activate glial cells (neurons immune cells) which would cause an upregulation of neuroinflammatory processes with subsequent neurodegeneration as a result or systemic inflammation could propagate into the brain and activate on site inflammatory events, which in the end will lead to neurodegenerative processes (Tansey and Goldberg, 2010). Interestingly, the density of microglial cells is very high in the region of the substantia nigra which makes the neuroinflammation mechanism even more plausible (Kim et al., 2000). Having said that, neuroinflammation is commonly reported in other neurodegenerative disorders such AD (Hirsch and Hunot, 2009) so it is not a mechanism exclusive to PD. Another mechanism that might be implicated in PD includes dysfunction in protein degradation.
systems either via the ubiquitin-proteasome system (UPS) or lysosome system where failure to clear misfolded α-synuclein or defective mitochondria leads to the possibility of LBs formation (Kouli et al., 2018).

1.1.5 Treatment

To date, there is no cure for PD; however, management and treatment have improved significantly. Pharmacologically, Levodopa (L-DOPA) is still the standard to treat PD for more than 50 years since it was first used in 1967 (Cotzias et al., 1967). L-DOPA, a dopamine precursor, and unlike dopamine, is able to cross the blood-brain barrier (BBB) can then convert into dopamine by DOPA decarboxylase to replenish the depleted neurotransmitter lost due to the disease. Despite its clear effectiveness, it comes with significant side effects. Some of the early treatment side effects include nausea and drowsiness while chronic treatment includes motor fluctuations, dyskinesia, hallucinations, and other psychiatric side effects. These side effects could be managed with dose control and a change in route of administration such as injection instead of oral (Zahoor et al., 2018). Beside the side effects, another major issue with L-DOPA that has been reported is the possibility of being neurotoxic itself (Zahoor et al., 2018). Other drugs used to treat PD, mostly in the early stages, include dopamine agonists which work by simulating the activity of the dopamine system without the need to be converted into dopamine. These include monoamine oxidase B (MAO-B) and Catechol-O-methyl transferase inhibitors which work by stabilising the level of endogenous dopamine, and anticholinergics which work by restoring the normal balance between dopamine and acetylcholine (Zahoor et al., 2018). Apart from using drugs, neurosurgery and cell therapy could offer good options to treat PD. Deep brain stimulation (DBS) is the most used surgical tool to treat PD and it had been used to treat a number of disorders since the 1960s (Bishop et al., 1963) and used in PD treatment in the 1990s by stimulating the subthalamic nucleus (STN) (Pollak et al., 1993). Despite its effectiveness, the mechanism of action is still not clear (Lozano and Lipsman, 2013).

In summary, PD is a very complex neurodegenerative disorder that has an effect on a selective part of the brain which manifest mainly as an impaired locomotor activity in patients. The disease is heterogeneous with multiple non-motor symptoms and different risk factors reported and other than symptoms modifying treatments, no cure has been found.
1.2 Leucine-rich repeat kinase 2 (LRRK2)

The first clue of LRRK2 association to PD was discovered in 2002 via a genome-wide linkage analysis of a Japanese family which displayed a phenotype similar to sporadic PD (Funayama et al., 2002). Later, more studies have confirmed this pathogenic link and found it to cause a late onset autosomal dominant PD (Paisán-Ruíz et al., 2004, Zimprich et al., 2004a). At that time, dardarin (from the Basque word dardara which means tremor) was the name given by (Paisán-Ruíz et al., 2004) to the encoded protein but today it is mostly referred to as LRRK2 for Leucine-Rich Repeat Kinase 2. Mutations in LRRK2 are reported to cause 5-13% of familial and 1-5% of sporadic PD cases (Kumari and Tan, 2009).

1.2.1 LRRK2 structure and localisation

LRRK2 gene is located on chromosome 12 and encodes a large multifunctional protein of 2527 amino acids with a molecular weight of approximately 280 kDa. LRRK2 protein harbours presumably four domains for protein-protein interactions, two GTPase domains, and one kinase domain (Figure 1.6). The protein-protein interaction domains are located at the ends of protein terminals while the enzymatic domains are in the core. On the N-terminal of the protein there are armadillo repeats (ARM), ankyrin repeats (ANK), and leucine-rich repeats (LRR) domains while the tryptophan-aspartic acid 40 repeat (WD40) domain is present on the C-terminal. These domains are mostly involved in protein structure and signalling. For example, the WD40 domain has been shown to be involved in membrane binding and protein localization while LRR domain was involved in inter/intramolecular interactions (Gilsbach and Kortholt, 2014). The core domains of LRRK2 are the functional enzymatic domains with multiple pathogenic mutations found linked to PD. The GTPase function is mediated by Ras of complex (ROC) and C-terminal of ROC (COR) domains. These domains throughout evolution have been always linked together and some consider them as one functional unit even though they are structurally different. A diagram of LRRK2 structure can be found in Figure 1.6. The kinase function of LRRK2 is mediated by a serine/threonine kinase domain which has the ability to phosphorylate many substrates and to autophosphorylate as well (Berwick et al., 2019). The presence of these GTPase bi-domain and kinase domain classifies LRRK2 as a member of ROCO family proteins, review by (Marín et al., 2008), which include among others a mammalian homologue LRRK1. Despite the great homology between the two proteins, no links to PD have yet been found in LRRK1 (Taylor et al., 2007).
Figure 1.6. LRRK2 structure, mutations, and interactors

The structure of LRRK2 protein with different mutations sites and several upstream and downstream interactors. Domains implicated in protein-protein interactions are in brown, domains involved in GTPase function are in purple, and the kinase domain is in blue. LRRK2 mutations are colour coded according to pathogenic (red), risk factor (yellow), and possibly neuroprotective (green) reports. The LRRK2 interactors are just a sample of many reported, for more see (Jeong and Lee, 2020, Marchand et al., 2020).

LRRK2 is a cytosolic protein found mostly as a monomer with low kinase activity. While the activation mechanism is not completely understood, it is believed that membrane-recruitment and confirmational activations reactions lead to LRRK2 dimerization and activation (Soliman et al., 2020). The expression levels of LRRK2 are variable throughout the body (Rideout et al., 2020). It is highly expressed in the kidney, lung, and cells of the peripheral immune system. It is also found expressed in the intestine, multiple regions of the brain and extracellularly through exosomes (Rideout et al., 2020). In the brain, the expression was detected in the neurons of cerebral cortex, caudate–putamen and SNpc regions (Higashi et al., 2007). Inside cells, LRRK2 has been shown to colocalize with various cellular structures including endosomes, lysosomes, multivesicular bodies, the mitochondrial outer membrane, lipid rafts, microtubule-
associated vesicles, the Golgi apparatus and the endoplasmic reticulum (ER) (Hatano et al., 2007). The association to a variety of membrane structure indicates multifunctional role in multiple cellular pathways for LRRK2.

1.2.2 LRRK2 Mutations

Since its discovery in the early 21st century, many mutations have been reported throughout the LRRK2 gene. The most well-studied PD-associated pathogenic mutations are missense mutations located within the enzymatic domains i.e., GTPase and kinase domains. For example, $LRRK2$-$R1441C/G/H$ and $LRRK2$-$Y1699C$ mutations were reported on domains ROC and COR respectively by (Mata et al., 2005, Paisán-Ruiz et al., 2004, Zimprich et al., 2004b) in families with probable English and German origins. On the kinase domain, $LRRK2$-$G2019S$ and $LRRK2$-$I2020T$ were reported by (Di Fonzo et al., 2005, Funayama et al., 2005, Gilks et al., 2005, Nichols et al., 2005) and found in families of European, American, and Japanese origins. Moreover, many other mutations have been identified with various links to PD such as $LRRK2$-$G2385R$ in the WD40 domain which is reported to be a risk factor (Jaleel et al., 2007) and interestingly $LRRK2$-$N551K$ and $LRRK2$-$R1398H$ are reported to be neuroprotective in some cohorts (Ross et al., 2011, Tan et al., 2010).

The most common mutation reported and studied in LRRK2 is $G2019S$. It accounts for 4% of familial and 1% of sporadic PD cases and it has been reported worldwide (Healy et al., 2008) in many populations with some incidence of higher frequencies in populations such as in Ashkenazi Jewish and North African Berbers which accounts for 29% and 37% of familial cases of PD in these populations respectively (Lesage et al., 2006, Ozelius et al., 2006). The second most frequent mutation is $LRRK2$-$R1441C/G/H$ and it is mostly reported in European populations. In fact, $LRRK2$-$R1441G$ particularly has been found to be a founder mutation responsible for 46% of familial cases in the Basque population (Gorostidi et al., 2008, Simón-Sánchez et al., 2006).

The penetrance of LRRK2 mutations vary depending on populations and age. For instance, $LRRK2$-$G2019S$ has been found to cause 60% of PD cases in Tunisian people by age 60 whereas only 20% of Norwegians who carried the mutation developed PD (Hentati et al., 2014). Furthermore, the penetrance in Basque population ranges from 13% at age 65 to 83% at age 80 for $LRRK2$-$R1441G$ mutation(Ruiz-Martínez et al., 2010).
1.2.3 LRRK2 Interactors

As mentioned earlier, LRRK2 is a large multi-domain protein expressed in many tissues with varying degree which makes elucidating its function very difficult. Despite this complexity, much research has been done and is still going to uncover the mysteries of this protein and which effectors it acts through. Not surprisingly, the majority of research was focused on the kinase activity because of the high prevalence of G2019S mutation which affects kinase activity and the reports of altered kinase level found in different pathogenic mutations.

In terms of phosphorylation of LRRK2, the protein is a highly phosphorylated one with at least 74 phosphorylation sites determined by phosphosite mapping studies and 60% of these sites have been identified as autophosphorylation sites (Marchand et al., 2020). The autophosphorylation process is still a matter of investigation where the GTPase, kinase, and even WD40 domains were reported to play interactive roles in the process in addition to dimerization and phosphorylation of other substrates like Rab29 (Berwick et al., 2019). Beside autophosphorylation, there are proteins that have been found to phosphorylate and dephosphorylate LRRK2. Protein kinase A (PKA) was the first protein reported to be able to phosphorylate LRRK2 on sites S910 and S935 with the latter one showing altered levels of phosphorylation reported in familial mutations LRRK2-G2019S, LRRK2-R1441G and LRRK2-Y1699C (Ito et al., 2007, Li et al., 2011, Muda et al., 2014). Other kinases shown to be acting on the same phosphosites include Ikappa B kinases (IKKa & β) and Casein Kinase 1-alpha (CK1α) (Chia et al., 2014, Dzamko et al., 2012). On S910 and S935, only the alpha catalytic subunit of Protein Phosphatase 1 (PPP1CA) has been confirmed to be able to dephosphorylate LRRK2 in several types of cells (Lobbestael et al., 2013). Moreover, 14-3-3 proteins have attracted attention due to their abundant presence in the brain, their binding ability to LRRK2 and the possible promotion of cell survival associated with them found in PD cellular models (Shimada et al., 2013). The full interaction cascade is still not established but it is reported to be at sites S910 and S935. Dephosphorylation of these sites have shown disrupted binding of 14-3-3 with subsequent influence on LRRK2 cellular localization. This disruption was markedly demonstrated in five of the most common pathogenic mutations (LRRK2-R1441C/G/H, LRRK2-Y1699C and LRRK2-I2020T). Binding of 14-3-3 has also been shown to prevent the self-association of LRRK2 into dimers which might affect the kinase activity by reducing the amount of the dimerised activated kinase form of LRRK2 (Mamais et al., 2014, Nichols et al., 2010).
Downstream substrates of LRRK2, many substrates have been identified to be phosphorylated. The list includes proteins in different tissues and pathways such as moesin, β-tubulin, tau, Futsch, endophilin A1, a subset of Rab GTPases and many others (Jeong and Lee, 2020). Among these different LRRK2 substrates, Rab GTPases have been the focus of investigation recently perhaps because they have been validated to be phosphorylated in vivo by some groups compared to most of the others (Jeong et al., 2018) and they have been found in abundance in CNS (D’Adamo et al., 2014). Rab GTPases are key organizers of intracellular membrane trafficking and were initially discovered in brain tissue (Touchot et al., 1987). Among more than 60 Rab GTPases that are encoded by the human genome, 24 are specific or enriched in CNS (Brighouse et al., 2010, Wilson et al., 2014). Many of these have been implicated into neurodegenerative disorders either directly or indirectly (Steinert et al., 2012, Wilson et al., 2014); however, the exact mechanism by which LRRK2 regulates Rab GTPase function is still not clear.

1.2.4 LRRK2 and cell biological functions

LRRK2 has been reported to be involved in many cellular processes (Figure 1.7). In the next sections, we’ll be discussing the role of LRRK2 in biological functions.

1.2.4.1 Autophagy

Autophagy is a crucial process to maintain protein homeostasis. It is a physiological process by which cells transport long-lived proteins or organelles such as mitochondria, that are unnecessary or even detrimental for cell survival to the lysosome for degradation. Autophagy is classified into three types: macroautophagy, microautophagy, and Chaperone-Mediated Autophagy (CMA). Macroautophagy is a nonselective process that targets macromolecules or subcellular organelles in bulk. Cytoplasmic material is sequestered into an autophagosome and delivered to the lysosome (or endolysosome) for degradation. Microautophagy involves the capture of cytoplasmic components through direct invagination of endolysosome membranes and can be nonspecific (in bulk) or highly specific. The CMA is a selective process of targeting proteins that contain a KFERQ pentapeptide-related motif by HSC70 and other co-chaperones such as HSP40 (Aman et al., 2021).

Alterations in autophagy have been observed in brain tissue of PD patients as well as in different models. An analysis of macroautophagy protein markers, namely p62 and
LAMP1, in LRRK2-G2019S-PD patients’ brains with LB pathology have shown significant low levels compared to sporadic PD brains suggestive of a divergence of effect in terms of autophagy between sporadic and LRRK2 carriers (Mamais et al., 2018). Furthermore, LRRK2-G2019S has been found to be poorly degraded by CMA than the wild-type LRRK2 in cell culture, mouse and DA neurons of PD patients suggesting a disturbance of the typical CMA process caused by pathogenic mutations in LRRK2 (Orenstein et al., 2013). A knock-in mice with LRRK2-R1441G have shown marks of impaired CMA accompanied with accumulation of SNCA oligomers and increase in CMA markers like LAMP2a and HSPA8 in striatal tissue (Ho et al., 2020). The previous observations point out to a physiological role to LRRK2 in autophagy with a mechanism that is still needs to be investigated further.

1.2.4.2 Mitochondria

Impairment in mitochondrial functions and morphology is a recurrent theme in PD. This impairment is well-established prior to any discovery of a genetic link to PD where mitochondrial complex I was found to be deficient in SNps of PD patients (Schapira et al., 1990). Observations have been reported of direct and indirect effect of LRRK2 on mitochondrial function. Directly, an impairment of mitochondrial complexes III and IV was reported in primary human fibroblasts from manifesting and non-manifesting LRRK2-G2019S carriers (Mortiboys et al., 2015). In addition, other studies have reported abnormalities in mitochondrial morphology caused by LRRK2 mutations (Smith et al., 2016, Yue et al., 2015). Indirectly, LRRK2-G2019S has been reported to increase phosphorylation of peroxiredoxin-3 (PRDX-3), a scavenger of hydrogen peroxide produced by mitochondria, compared to the wild-type which leads to the increase of inhibition of endogenous peroxidase and promoting mitochondrial dysregulation and oxidative damage (Angeles et al., 2011, Angeles et al., 2014). Interestingly, LRRK2 has been reported to be localised to the mitochondrial membrane (Biskup et al., 2006) and this localisation might have an effect on mitochondrial biogenesis. For example, LRRK2 has been reported to interact with an outer mitochondrial protein called Miro that is important in mitochondrial transportation. This interaction triggers the transport of damaged mitochondria along axonal microtubules which lead to mitophagy of damaged mitochondria and this response is lost in cells with the LRRK2-G2019S mutation (Hsieh et al., 2016). Similar results have been reported for R1441C and Y1699C mutations as well (Godena et al., 2014, Thomas et al., 2016).
1.2.4.3 Endocytosis

Endocytosis is the process of engulfing and trafficking of plasma membrane associated proteins through a series of intracellular membrane-limited compartments that leads to degradation by lysosomes or recycling into other intracellular locations. LRRK2 has been reported to interact with different protein associated with endocytosis. The mechanism of this interaction is still to be explained; however, Rab GTPases have been associated with many of these interactions (Steger et al., 2017, Steger et al., 2016). Rab29, also known as Rab7L1, has been found to recruit LRRK2 to enlarged lysosomes in chloroquine stressed cells. This recruitment led to the phosphorylation of Rab8a and Rab10 by LRRK2 and eventually aided lysosomal function by promoting lysosomal secretion (Eguchi et al., 2018). Similar results have been reported by (Herbst et al., 2020, Lee et al., 2020) on the effect of LRRK2 phosphorylation of Rab8a and Rab10 in stressed endosomes and lysosomes. Expressing LRRK2-G2019S and LRRK2-R1441C/G mutations in iPSC-derived DA neurons from PD patients have impaired lysosomal function (Ysselstein et al., 2019) suggesting a crucial role for LRRK2 in regulating lysosomal functions.

Moreover, LRRK2 has been implicated in synaptic vesicle endocytosis. Overexpression of LRRK2-G2019S and LRRK2-R1441G in primary midbrain, primary hippocampal neurons, and iPSC-derived DA neurons from PD patients impaired endocytosis of synaptic vesicles (Matta et al., 2012, Nguyen and Krainc, 2018, Pan et al., 2017). Many proteins have been reported to interact with LRRK2 in regulating synaptic vesicle endocytosis that include auxilin, Dynamin GTPases, and synaptojanin-1 (Gad et al., 2000, Stafa et al., 2014, Yim et al., 2010). In addition, Rab5 and Rab7 have been reported to be involved with LRRK2 in early and late endosome respectively (Dodson et al., 2012, Shin et al., 2008).

1.2.4.4 LRRK2 function and Trans-Golgi network (TGN) and endoplasmic reticulum (ER)

LRRK2 interacts with several proteins that reside at the Golgi body to regulate its integrity and function. For example, it has been found that Rab29, LRRK2, BAG5 and GAK proteins form a complex localised to TGN in cultured cells and mouse brain. Overexpression of pathogenic variants of LRRK2 or any of these individual proteins causes Golgi fragmentation. This induced fragmentation can be blocked by the knockdown of any one of the other three proteins which indicates the importance of
these proteins in maintaining Golgi integrity (Beilina et al., 2014). Additionally, LRRK2 has been reported to phosphorylate and form a complex with N-ethylmaleimide sensitive fusion (NSF) protein, which is an essential factor for vesicle trafficking in the Golgi body. NSF knockdown has shown similar results to LRRK2 knockdown in terms of causing Golgi fragmentations and the formation of enlarged late endosomal structures. Disruption of plasma membrane receptor recycling pathways and trafficking of lysosomal hydrolysates from TGN to lysosomes were observed as well resulting in an impaired lysosomal degradation capacity (Lanning et al., 2018).

Additionally, LRRK2 has been found to form a complex with Sec16a protein, a key protein in the formation of endoplasmic reticulum exit sites (ERES). The knockdown of LRRK2 in primary hippocampal neurons led to the disruption of LRRK2-Sec16a complex and impaired the transport of vesicles to the Golgi. Similar results were found when a knock-in of LRRK2-R1441C was expressed (Cho et al., 2014).

1.2.4.5 Microtubules and ribosomes

LRRK2 has been associated with the cytoskeleton, particularly microtubules. This interaction was identified first in 2006 and was shown to be an association of LRRK2 with β-tubulin (Biskup et al., 2006). A post translational modification known to induce microtubule stability i.e. acetylation was reported to be increased in LRRK2 knockout MEF cells and mouse kidney cells and associated with this is a strong localization of LRRK2 to microtubules that are dynamic i.e. not acetylated and less stable suggesting a negative effect of LRRK2 on microtubule stability (Law et al., 2014, Pellegrini et al., 2018). Interestingly, the activity of the association of LRRK2 with dynamic microtubules seems specific to the GTPase domain since it is only observed in LRRK2-R1441C and LRRK2-Y1699C but not in LRRK2-G2019S mutations (Godena et al., 2014).

Furthermore, LRRK2 has been implicated in protein synthesis. This implication is a bit controversial because it was only replicated in fruit flies and weakly in vitro but not mammalian models (Imai et al., 2008, Kumar et al., 2010, Trancikova et al., 2012). The eukaryotic initiation factor 4E binding protein (4E-BP), a repressor of translations, was found to be phosphorylated by the LRRK2 homologue in the Drosophila LRRK (dLRRK) and was able to inactivate 4E-BP (Imai et al., 2008).
Figure 1.7. The involvement of LRRK2 in different organelles and cellular mechanisms.

LRRK2 is involved in the regulation of several biological processes, some of which are intertwined. Rab proteins are involved in several of these processes and potentially mediate some effects of LRRK2. Taken from (Esteves et al., 2014).

In conclusion, LRRK2 is a large protein with several mutations discovered throughout the protein that are linked to PD. It has been found in many tissues and has been associated with many cellular processes. This introduces complexity to our understanding of the protein which makes it very difficult to elucidate its function and pathology in PD. Thus, further investigations are encouraged to uncover the mysteries of this protein.

1.3 Modelling PD

Potentially the best way to study human biology is to study it in humans themselves; however, the ethical and technical limitations are too significant to disregard. Thus, finding the best model that would recapitulate human biology is very crucial. Throughout the years, many models have been developed for PD, but none have been
able to fully recapitulate all disease characteristics. Nevertheless, the available models are still beneficial with different advantages and disadvantages.

### 1.3.1 Cellular models

Cellular models offer a controlled environment to simplify experimental variables compared to what is present in animal models. This type of modelling is very useful when investigating single cellular events or pathways; however, it fails to recapitulate non-cell autonomous effects that are essential in disease pathology. Compared to animal models, cellular models are fast and reproducible with the opportunity for specific genetic and environmental manipulations that are devoid of many ethical issues (Lazaro et al., 2017).

Many cellular models have been developed to study PD. Human embryonic kidney 293 (HEK 293) and human neuroglioma (H4) cells are established cell lines that are commonly used as heterologous cellular models. These cell lines are quite useful because they are easy to maintain, provide unlimited supply of homogenous cells and grow rapidly which enable large scale studies, but being derived from cancer cells, many proteins are produced at abnormal (higher or lower) levels. HEK 293 and H4 cells have been used to study the effect of *SNCA* mutations (Lázaro et al., 2016, Lazaro et al., 2014) and *LRRK2* mutations (Nichols et al., 2010) on different cellular events found in PD. The contributions of these cells are great; however, they lack the neuronal phenotype which limit their usefulness. This can be overcome by the use of cell lines that are capable of differentiating into neuronal-like cells, specifically, DA cells. For example, Lund Human Mesencephalic (LUHMES) cells are capable of differentiating into strong DA phenotype neuron-like cells (Scholz et al., 2011). These cells have been used to study the different cytotoxic effects of α-synuclein (Tong et al., 2017, Xiang et al., 2013).

Recently, human induced pluripotent stem cells (iPSCs) have been instrumental in biomedical research since Shinya Yamanaka pioneered the technology of reprogramming somatic cells in 2006 (Takahashi and Yamanaka, 2006). Compared to the established cell lines, these cells are capable of being differentiated into DA neurons when taken from PD patients, which provide the best genetic background to study the disease. Over the past decade, many iPSCs models have been developed for neurodegenerative diseases including PD. Familial and sporadic PD have been modelled in these cells demonstrating the different pathological events of the disease such as
protein aggregation, mitochondrial dysfunction, and susceptibility to autophagy and oxidative stress. For more information see review by (Avazzadeh et al., 2021). In addition to modelling PD to understand the pathway of the disease, the use of iPSCs has a great potential in drug discovery and cell therapy (Ke et al., 2019). For example, an autologous transplant of DA progenitor cells in induced PD rhesus monkeys have shown alleviation of motor and depressive behaviours (Tao et al., 2021). Moreover, the use of a DA progenitor cells (MSK-DA01) in PD patients is being assessed for its efficacy and safety after proving safe and effective in rats (Piao et al., 2021).

A great advancement in cellular modelling of PD is the use of human organoids. These are small, self-organized, 3D tissue cultures that are derived from human stem cells. They have great potential in disease modelling, developmental biology and drug discovery. Compared to the traditional monolayered 2D cell cultures, human physiology is recapitulated better due to cellular heterogeneity. Additionally, they offer a better modelling for biological processes that are specific to the human body than animal models (Corro et al., 2020, Kim et al., 2020). In PD modelling, human midbrain organoids (hMOs), have shown a great potential to recapitulate many aspects of the disease and they have the capability to bridge the gap between cellular and animal models. Since their first introduction in 2014 (Tieng et al., 2014), organoids have been successfully used to model PD genetically and phenotypically (Galet et al., 2020).

### 1.3.2 Animal models

Although cellular models give great insights into the cellular pathology of PD, they lack the non-autonomous aspects of the disease progression. In addition, the physical and behavioural manifestations that occur in patients are impossible to demonstrate in cell culture. Thus, the next best option is animal models. They provide an excellent opportunity to understand the full picture of disease progression in a form mimicking human biology.

When looking for the ideal model for PD, one should look for what recapitulate or mimics the pathology of the disease such as a selective DA neurodegeneration, the presence of neuronal inclusions like LBs and a display of motor deficits. There have been two main types of animal models used in PD research, toxin treated and genetic models. The focus in the next sections will be on rodents, while the fruit fly will be discussed in later sections (section 1.4).
1.3.2.1 Toxin-treated models

This type of model utilises neurotoxins to induce the symptoms of PD. In 1968, Urban Ungerstedt injected 6-hydroxydopamine (6-OHDA) into rats’ brains which led to the degeneration of neurons with high selectivity of DA neurodegeneration and marked motor deficits in these rats (Ungerstedt, 1968). Currently, 6-OHDA is a widely used neurotoxin to induce the symptoms of PD in animals. It demonstrates most of the known PD symptoms such as neurodegeneration and mitophagy; however, LBs seem to be absent, which may be a function of time (the injection of 6-OHDA may not allow time for aggregates to form). A drawback for this drug is its inability to cross the BBB which means it requires an injection into the brain to be administered (Hernandez-Baltazar et al., 2017).

Another highly selectivity neurotoxin to DA neurons is MPTP. The parkinsonism effect of this drug was discovered by accident in 1983 in drug abusers (Langston et al., 1983). Unlike 6-OHDA, MPTP can cross the BBB. MPTP itself is not toxic but when it enters the brain it is converted into 1-methyl-4-phenylpyridinium (MPP+) which is taken up by the DA neurons. In the DA neurons, MPP+ is toxic to the mitochondria which leads to cell death, eventually (Langston, 2017). Similar to 6-OHDA, MPTP treated brain fail to show LBs which is a cellular hallmark in PD.

The reports of increased risk of PD from environmental agents such as pesticides have resulted in the use of rotenone and paraquat as neurotoxin-based models (Tanner et al., 2011). Rotenone and paraquat are similar to MPTP in that they are able to cross the BBB which enable them to be administered systemically. Additionally, they are able to mimic PD in terms of forming LBs unlike 6-OHDA and MPTP. Rotenone is a natural compound found in many plants and has been used as pesticide. Similar to MPTP, it causes DA neurons degeneration via mitochondrial dysfunction. The models produced are generally selective to the DA neurons but some reports showed multisystem degeneration (Hoglinger et al., 2003) and the high mortality observed in some rat models (Fleming et al., 2004) is a drawback. Paraquat was identified as a possible candidate to induce parkinsonism due to the structural similarity to MPP+. Unlike rotenone and MPTP, paraquat causes DA neuron degeneration via the production of mitochondrial derived reactive oxygen species (ROS) which causes oxidative stress that leads to cell death. Paraquat selectivity to DA neurons is questionable since some have
found it to be selective (McCormack et al., 2002) while others have not or have but only when paraquat is conjugated with other chemicals (Thiruchelvam et al., 2000).

There have been a range of neurotoxin-based animal models generated throughout the years; however, rodents and non-human primates (NHP) are the most used and the use of MPTP is considered to be the gold standard due to its ability to demonstrate almost all of the PD hallmarks (Jackson-Lewis et al., 2012, Pingale and Gupta, 2020).

1.3.2.2 Genetic models

Although the majority of PD cases are idiopathic, they share many characteristics with the familial ones. Thus, genetic models offer great opportunity to understanding the pathogenesis of the disease and enable therapeutic screening. Just like in neurotoxin-based models, there isn’t a perfect genetic model that would demonstrate all PD hallmarks. Among many genes that have been linked to PD, five have been mostly studied in different animal models and these are SNCA, LRRK2, PINK1, PRKN, and DJ-1. These models have been developed by either generating knockouts, transgenic animals or injecting viral vectors carrying the gene of interest or preformed fibrils into the brain (Konnova and Swanberg, 2018).

Overexpression of SNCA in rodents and NHPs causes α-synuclein aggregates but the motor deficits and DA neurodegeneration can be variable depending on the route of administration. In transgenic rodents when SNCA is overexpressed, some motor deficits were observed with no DA neurodegeneration (Cannon et al., 2013, Fleming et al., 2005, Recasens et al., 2014, Van der Perren et al., 2015, Volpicelli-Daley et al., 2016, Wakamatsu et al., 2008).

The large size of LRRK2 gene has limited its use in viral vector models (Van der Perren et al., 2015); however, some attempts to express LRRK in mice and rats were able to induce some motor deficit with mild or no DA neurodegeneration (Dusonchet et al., 2011, Lee et al., 2010, Li et al., 2009). Interestingly, injecting helper-dependent canine adenovirus type 2 (HD CAV-2) with LRRK2-G2019S in the striatum of Microcebus murinus, a NHP model, has induced clinical and histological symptoms similar to PD (Mestre-Francés et al., 2018). The majority of transgenic mouse and rat models of LRRK2 have little effect on DA neurodegeneration and most lack motor dysfunction (Blesa and Przedborski, 2014, Dawson et al., 2010).
The nature of mutations in PINK1, PRKN, and DJ-1 genes (autosomal recessive loss of function) have resulted in developing knockout models. Unfortunately, most did not to display motor deficits or DA neurodegeneration (Dawson et al., 2010). Inactivation of PINK1 and PRKN did not produce better models in mice but the DA neurons were more sensitive to subsequent insult by MPTP (Haque et al., 2012, Van der Perren et al., 2015).

Interestingly, one key observation that remains to be explained due to its unknown physiological significance and potential contribution to the pathogenesis of the disease is the loss of neuromelanin (NM) pigmentation normally produced by DA neurons in the brain, specifically in the SNpc. The intensity of this pigmentation in the substantia nigra is the greatest in humans where it can be examined macroscopically (Marsden, 1961). This key observation couldn’t be replicated in any laboratory animals commonly used in PD research because it seems unique only to humans. However, (Carballo-Carbajal et al., 2019) were able to generate a rat model that is able to produce human-like intensity of NM by overexpressing human tyrosinase in rat SNpc. This model was able to produce age dependent neurodegeneration associated with neuronal dysfunction and formed PD-like neuronal inclusions similar to that found in PD. The pathology presented in this model sounds promising, however, no report of motor deficit was revealed.

Rodents and NHPs comprise the majority of animal models in PD research. This is undeniably due to their close similarity to human biology in terms of genes and symptomatic and pre-symptomatic phenotypes that can be observed and measured (Konnova and Swanberg, 2018). Having said that, the variability in cellular and physical phenotypes opens the door for other animal models to be developed to better understand the full picture of the disease. Non-mammalian models such as C. elegans, zebrafish and D. melanogaster have well-defined neuropathology and behaviour, low maintenance costs, rapid life cycle and a genome easy to manipulate. Many models have been generated in C. elegans and zebrafish that displayed variable phenotypes similar to rodents. The number of animals produced and the rapid life cycle of C. elegans and zebrafish is an advantage in terms of drug screening and discovery (Cooper and Van Raamsdonk, 2018, Vaz et al., 2018). As for D. melanogaster, a more detailed dive into its contribution in PD research can be found in the next section.

Finally, the attempts, so far, to generate a comprehensive model for PD that would recapitulate all hallmarks of the disease have not been successful; however, these
variance in models might be a reflection of the heterogeneity of the disease. Therefore, studying a combination of models or choosing one that would best recapitulate an aspect of the disease, I think, would be an optimal approach to understanding and studying PD pathology.

1.4 *Drosophila as a powerful model organism*

Among many models available to use, the common fruit fly (*Drosophila melanogaster*) has emerged as a powerful candidate for PD modelling. The use of fly as a model started at the beginning of the 20th century with the work of William E. Castle and his students at Harvard University (Castle *et al.*, 1906). Influenced by his work, Thomas Hunt Morgan worked further on *Drosophila* to establish it as a robust model, especially in understanding the basis of genetics. His pioneering works led to the discovery of the *white* (*w*) mutation, which resulted from a natural mutation occurring on the X chromosome that change the colour of a wild type fly eye from red to white (Kenney and Borisy, 2009). Using *Drosophila* in research labs is easy and cheap. Its size allows it to be maintained in high numbers with small space used and the relatively less complex structure enables the ease of experimental manipulations. It has a short lifespan (~90 days) with a rapid life cycle (it takes ~10 days to generate adult flies at 25°C). The low redundancy of its genome and the wide range of genetic tools available allow forward and reverse genetics to be made with efficiency (Roote and Prokop, 2013). In addition, the sequencing and publication of the whole genome of the *Drosophila* was a breakthrough in demonstrating similarities to the human genome. *Drosophila* has around 13,600 genes spread across four chromosomes (one sex chromosome, the first one, and three autosomes). Surprisingly, more than 75% of human-associated disease genes have orthologues in *Drosophila* (Adams *et al.*, 2000, Reiter *et al.*, 2001).

Using *Drosophila* as a model is still proving strong in a sense that six Noble prizes have been awarded to ground-breaking works based on fruit fly and the most recent one was awarded in 2017.

1.4.1 *The genetic toolbox of Drosophila*

The genetic toolbox in *Drosophila* sets it apart from other model organisms. Prior to its genome sequencing, unbiased randomly generated forward genetics were used to achieve genetic manipulations. This type of genetics works by introducing random mutations into the genome and then identifying or observing the effect on the
phenotypes. Such mutations can occur naturally or be generated artificially using chemicals, radiation, or insertional mutagens e.g., transposable elements. Chemically, flies can be fed ethyl methane sulphonate (EMS) that can introduce point mutations via alkylation (Sega, 1984). Ionising radiation e.g., X-ray has been known to be mutagenic and can introduce chromosomal rearrangements and deletions (Pastink et al., 1987). Insertional mutagens utilise the ability of mobile genetic elements such as P-elements to jump randomly around the genome to cause mutations. These elements are not as strong as the previous two in term of introducing a wide range of mutations; however, the mutated genes can be rapidly and easily identified using the element as a genomic tag (St Johnston, 2002). In reverse genetics, on the other hand, there is no screening for mutations because the target is a known gene and changes to a phenotype can be observed when that particular gene is manipulated. Many methods have been developed to alter the gene of interest. P-elements can be modified and inserted within or near the gene of interest to cause mutations. Transgenic RNAi and CRISPR/Cas9 can be used for a more targeted way of altering the function of a gene (Adams and Sekelsky, 2002, Fire et al., 1998, Gratz et al., 2013). These tools, in combination with Gal4/UAS system and balancer chromosomes (see next sections), help Drosophila recapitulate human biology.

1.4.1.1 The Gal4/UAS system

One significant advantage of using Drosophila as a model organism is the Gal4/UAS system (Brand and Perrimon, 1993). It is originally based on an enhancer-trap method where the ability to express a gene of interest in specific cells or tissues is achieved. The system consists of two elements, a transcriptional activator derived from Saccharomyces cerevisiae (Gal4) placed under the control of a cell or tissue-specific promotor and a transgene placed under the control of the upstream activator sequence (UAS), which Gal4 binds to. These individual elements are made separately in two different flies and can only be expressed in the desired way when flies are mated to produce a progeny that carries both (Figure 1.8). Throughout the years, many flies have been produced to drive expression in different parts of the fly, e.g., GMR-Gal4 (eye), elav-Gal4 (neurons), and TH-Gal4 (DA neurons), as well as many flies have been produced to express different genes of interest.
Figure 1.8. The Gal4/UAS system in Drosophila

The Gal4/UAS system allows ectopic expression of the gene of interest in specific tissues or cells when its two components are present in a fly. This figure illustrates the ability of the progeny to express gene X in a tissue-specific fashion. This occurred as a result of a crossing scheme between a tissue-specific Gal4 activator fly (Driver) and a UAS fly that has Gal4 binding sites (Responder).

The system has undertaken many modifications since its inception. The generation of Gal4-hormone receptor (GeneSwitch) elements allows temporal gene expression control (Osterwalder et al., 2001). This temporal expression is achieved due to the ability of these lines to be activated only in the presence of the appropriate ligand such as mifepristone (RU486) that can be administered when required (Figure 1.9). Other modifications include, the addition of extra elements, such as FLP recombinase and Gal80, that add an extra level of regulation and the addition of RNA-mediated interference (RNAi), making the system capable of targeted gene knockdowns (Duffy, 2002).
Figure 1.9. The Gal4/UAS system (GeneSwitch) in Drosophila

This modification to the conventional Gal4/UAS system (described previously) allows temporal expression of the gene of interest. This figure illustrates gene expression is dependent upon the presence of RU486.

1.4.1.2 Balancer chromosomes

The existence of balancer chromosomes makes fly genetics extremely versatile. Throughout the years, many balancer chromosomes have been developed for the main three chromosomes (the 4th chromosome is effectively ignored for these purposes being too small to undergo regular recombination events). These chromosomes are structurally rearranged with multiple aneuploid chromosomal inversions that inhibit homologous recombination during meiosis. If recombination does occur, the deletion of genomic regions or duplication of genomic regions will result in lethality. In addition, balancer chromosomes carry dominant markers that produce a distinct observable phenotype (Figure 1.10). All these properties aid in maintaining fly stocks, particularly those with deleterious mutations, and tracking mutations when designing crossing schemes (Kaufman, 2017).
Balancer chromosomes aid in fly selection of the desired genotype during crossing schemes. In this example, the desired genotype is the one carrying mutations X and Y. In order to acquire this, parent flies each carrying one of the desired mutations on one copy of the chromosome and the balancer one (the dominant marker for CyO is curly wings) on the other copy of the chromosome are crossed together. The results are two flies similar to the parents and one that would be the desired fly (red square). Of note, the fourth possible option won’t appear in the offspring due to homozygous lethality caused by the balancer chromosomes.

1.4.2 *Drosophila* as a model for neurodegenerative disease

Neurogenerative diseases are the result of neuronal death in the nervous system which leads to disruption in cognition, motor functions and eventually death. The nervous system is one of the most, if not the most, complex system in humans. Therefore, studying these disorders in human poses technical and ethical challenges. Human genetics and genomic studies have helped linked many of these disorders to candidate genes e.g. *apolipoprotein E (APOE)* to Alzheimer disease (AD) (Harold et al., 2009), *LRRK2* to PD (Funayama et al., 2002), and *Huntingtin (HD)* gene to Huntington’s disease (Gusella et al., 1983). However, identifying the pathological mechanism and progression of the disease is still difficult because the diagnosis usually occurs post-mortem or late when most of the initially affected neurons are dead. These challenges are best tackled with animal models and *Drosophila* has been shown to be very useful for modelling neurodegenerative diseases (McGurk et al., 2015).
Technically, the short lifespan and the large isogeneic number of flies it can produce are valuable in studying human ailments that are age-related (Hirth, 2010). Additionally, basic aspects of cell biology are quite similar in humans and flies, including regulation of gene expression, membrane trafficking, neural connectivity, cell signalling, and cell death. Many genes and pathways have been later discovered in mammals that were originally studied in flies. For example, Wnt protein that is crucial in many aspects of mammalian cell biology was studied first identified and studied in mutant wingless flies. This phenotype is caused by loss of the Drosophila wingless gene, an orthologue of the Wnt gene in mammals (Sang and Jackson, 2005).

There are very few ethical and safety issues in using Drosophila in scientific research compared to other animals such as mice. For example, using Drosophila doesn’t require certain licensing from governments.

Drosophila have already proven highly successful in the study of neurodegenerative disorders including AD, PD, HD, Amyotrophic Lateral Sclerosis (ALS), and many others (Afsari et al., 2014, Cao et al., 2008, Watson et al., 2008, Weiss et al., 2012).

1.4.3 Drosophila models of PD

As mentioned earlier, a comprehensive one model that would recapitulate all hallmarks of PD is not yet available. Therefore, choosing the best model that fit the researcher needs is crucial. Most Drosophila models of PD have shown selective age dependent DA neurodegeneration that mimic what is found in humans which make them a great candidate to uncover the mechanism of this degeneration (Xiong and Yu, 2018). This is interesting because transgenic mice models, apart from viral mediated ones, have failed to replicate this aspect of the disease. A different dopamine metabolism and the expression of intrinsic protective factors or other genetic modifications that prevent the toxic effects of these PD-related mutations could explain this failure in mice (Burbulla et al., 2017, Dawson et al., 2010). Additionally, a large-scale drug screening and discovery is much more rapid and cost-effective in Drosophila compared to mice.

Despite lacking the homologous gene, expressing human α-synuclein in Drosophila was revealed to be successful in generating a PD disease model. In 2000, (Feany and Bender, 2000) managed to generate a fly model that expresses α-synuclein in neurons which then displayed age dependent DA neurodegeneration, neuronal inclusions and locomotor dysfunction. These phenotypes were replicated by others with varying
degrees of similarity (Chen and Feany, 2005, Periquet et al., 2007). It is worth re-stating that the Drosophila genome contains no apparent ortholog of SCNA which would contribute to the generation of LBs.

PRKN and PINK1 have been shown to be involved in mitochondrial abnormalities, which is one of the key observations found in PD. Mutations in both genes in flies exhibited similar phenotypes of mitochondrial abnormalities, reduced lifespan and locomotive impairment. Moreover, the overexpression of human PINK1 in mutant PINK1 flies have managed to restore normal mitochondrial morphology which suggest a conserved functionality. Also, both genes seem to be essential in regulating mitochondrial function (Clark et al., 2006, Greene et al., 2003).

Two orthologs of the DJ-1 gene, that when mutant can cause PD in humans, exist in flies, DJ-1α and DJ-1β. The expression of DJ-1β is more wide-spread than DJ-1α which is mostly expressed in male testis. Flies with double knockout genes looked fairly normal in terms of viability, fertility and lifespan; however, when exposed to oxidative stress such as H₂O₂ and paraquat they displayed sensitivity that was rescued by re-introducing the genes into the flies (Meulener et al., 2005). In another model, (Menzies et al., 2005) were able to demonstrate that overexpression of DJ-α protected DA neurons from the insult of paraquat. These results suggested a possible role of fly DJ-1 in protection against environmental and oxidative stress that can be seen in PD.

1.4.3.1 LRRK2 Drosophila models of PD

By producing and analysing LRRK transgenic alleles and loss of function mutants in Drosophila, (Lee et al., 2007) developed a fly model for the fly homolog of LRRK2 (dLRRK) and convincingly established an endogenous role for LRRK2 in preventing the degeneration of DA neurons. No DA neurons loss or impairments in locomotive function were seen in this investigation. Pathogenic mutants and wild-type dLRRK transgenic expression did not demonstrate any noticeable abnormalities, whereas dLRRK loss-of-function mutants have significantly reduced locomotor activity. Additionally, the morphology and significant loss in tyrosine hydroxylase (TH) immunostaining of DA neurons in dLRRK mutants suggested a neurodegeneration in the mutants (Lee et al., 2007).

The human LRRK2-WT and the mutant form LRRK2-G2019S were overexpressed to construct a gain-of-function LRRK2 Drosophila model by (Liu et al., 2008). Expression
of both forms of \textit{LRRK2} resulted in premature mortality, locomotor impairment, retinal degeneration, and selective loss of DA neurons in the brain. Furthermore, compared to \textit{LRRK2-WT}, \textit{LRRK2-G2019S} induced a more severe parkinsonism-like phenotype. L-DOPA treatment alleviated the \textit{LRRK2-G2019S} induced locomotor impairment but did not prevent the loss of TH-positive neurons (Liu et al., 2008).

(Imai et al., 2008) employed \textit{Drosophila} to study how LRRK2 functions normally in the body and how its dysfunction results in DA neurodegeneration. They showed genetic and biochemical data that \textit{dLRRK} controls protein synthesis in order to maintain DA neurons. In addition, they observed that LRRK2 primes 4E-BP phosphorylation, and that action plays a significant role in mediating the pathogenic consequences of mutant \textit{dLRRK}. Lastly, they found that behavioural abnormalities are accompanied with dopamine and dopaminergic cell death (Imai et al., 2008).

The expression of the transgenics \textit{G2019S}, \textit{Y1699C}, and \textit{G2385R} variant in flies showed late-onset loss of DA neurons in specific clusters and exhibited locomotor abnormalities compared to the \textit{LRRK2-WT} (Ng et al., 2009). Additionally, no retinal degeneration was observed in flies aged 20 or 60 day old. This discovery is in line with the findings of (Imai et al., 2008, Lee et al., 2007) where transgenic flies expressing \textit{LRRK2-WT} are protected from this age-associated phenotype, whereas mutant \textit{LRRK2}-mediated degeneration appeared to be late-onset and is limited to specific clusters of DA neurons. However, the findings in (Liu et al., 2008) were in contrast to (Ng et al., 2009) where degeneration in \textit{Drosophila} expressing either \textit{LRRK2-WT} or \textit{LRRK2-G2019S} was observed to occur non-selectively across all the DA neuronal clusters and a significant retinal degeneration at as early as 3 week old flies.

Using \textit{Drosophila} electroretinograms (ERGs), (Hindle et al., 2013) examined the impact of \textit{LRRK2} mutations. They reported that in flies expressing \textit{LRRK2-G2019S} mutation in the DA neurons, the photoreceptor function was gradually deteriorating. After 28 days, it was found that there had been a loss of vision and that the photoreceptors had high levels of autophagy, apoptosis, and mitochondrial disorganisation. Furthermore, even in areas not directly innervated by DA neurons, fly head dissections showed widespread neurodegeneration across the visual system. Other mutations tested didn't show any photoreceptors degeneration or loss of vision. Lastly, they manipulated the levels of \textit{dLRRK} expression simultaneously with expressing
LRRK2-G2019S in DA neurons and came to the conclusion that the LRRK2-G2019S mutation is a gain of function rather than dominant negative (Hindle et al., 2013).

In summary, Drosophila is one of the most powerful organisms to use as a model in PD and neurodegenerative diseases in general due to its relatively cheap cost, short and rapid lifespan, having less ethical issues, ease of handling and keeping, similarity to humans in many biological aspects and wide experimentation for over 100 years. However, as in all animal models, there are some drawbacks in using Drosophila. It has to be transferred into fresh media on a regular basis to keep stocks viable i.e., it can’t be frozen for long term storage. Moreover, the small size of Drosophila may introduce some difficulty in terms of handling and experiment design.

1.5 Investigating PD through Drosophila eyes

When modelling PD in animals, one might wonder how eye and vision relates to PD. In the next sections, the case for using Drosophila’s visual system to investigate PD will be discussed.

1.5.1 The retina

The retina is the neural part of the eye responsible for receiving the outside light and processing it into signals that are transmitted via the optical nerves to the brain where these signals are converted into perceived images. This processing is mediated by different types of neural cells, namely photoreceptors, horizontal cells, bipolar cells, amacrine cells and ganglion cells with metabolic and homeostatic support provided by different glial cells and retinal pigment epithelial cells. These retinal cells are organised in distinctive histological layers and starting from where light comes in i.e., outside to inside, there are the ganglion cell layer (GCL), inner plexiform layer (IPL), inner nuclear layer (INL), outer plexiform layer (OPL), and outer nuclear layer (ONL). These layers are sandwiched between a nerve fibre layer and a retinal pigment epithelium (RPE) layer (Sung and Chuang, 2010) (Figure 1.11) demonstrates the structure of the retina. ONL contains photoreceptors which are responsible for sensing light and transmitting it as an electrical signal to the next order of neurons. Photoreceptors are dominated by rods comprising 95% of all photoreceptors compared to 5% of cones that are concentrated in the foveola. Rods are responsible for low-light (scotopic) vision while cones are specialized for bright-light (photopic) colour vision (Lamb, 2016). The location of these photoreceptors (at the back of the retina) might seem counterintuitive at first but RPE
that set in proximity to photoreceptors help to absorb any scattered light that would cause image distortion (Archibald et al., 2009). OPL contains horizontal cells which feedback and feedforward signals from photoreceptors and bipolar cells (Masland, 2012a). Bipolar cells somas are found in INL and are responsible for receiving information from the photoreceptors and projecting it to the retina ganglion cells (RGCs). Depending on the signal received from photoreceptors, bipolar cells can be divided into ON-bipolar cells (depolarize in response to light) or OFF-bipolar cells (hyperpolarize in response to light). Cones can be connected to either ON or OFF types of bipolar cells while rods are only connected to ON ones (Euler et al., 2014). The next layer (IPL) contains amacrine cells and the synapses of bipolar cells and RGCs. Like horizontal cells, amacrine cells modulate the information transfer between bipolar cells and RGC and they are extremely diverse in function and level of axon stratifications (Masland, 2012b). The last layer (GCL) as the name suggests contains RGCs. These cells are the last stop of information modulation before it travels to the brain through the optic nerve. The GCL transmit image and non-image forming signals that help in the function of circadian rhythm, melatonin release modulation, and pupil size regulation (Mahabadi and Al Khalili, 2020).
1.5.2 Visual dysfunctions in PD

Visual dysfunction is one of the non-motor symptoms reported by PD patients. The reported symptoms include visual hallucination, double vision, difficulty in reading, and deficit in colour discrimination and contrast sensitivity (Price et al., 1992, Urwyler et al., 2014). Using assays to evaluate the visual function such as electroretinogram (ERG), visual dysfunctions have been reported in patients (Bodis-Wollner et al., 1987) and animal models (Hindle et al., 2013, Onofrj and Bodis-Wollner, 1982). The cause for this dysfunction is not clear but likely involves the degeneration of DA neurons, just like in the SNpc, could be one. Among the different types of retinal cells, A18 amacrine cells have been identified as DA neurons and are located in the INL (Frederick et al., 1982). Studies have shown that the INL was significantly thinner in PD patients (Hajee et al., 2009) and the DA innervation was reduced (Nguyen-Legros, 1988). Additionally, post-mortem analysis has revealed a significant reduction of retinal DA neurons in PD patients. Interestingly, this reduction was comparable to non-PD patients for patients who received L-DOPA treatment (Harnois and Di Paolo, 1990). Similar results have been observed in animals as well. For example, a decrease in DA amacrine cells in mice (Marrocco et al., 2020), a reduction in dopamine level in rats (Meng et al., 2012), and
neurodegeneration in the retina of flies (Hindle et al., 2013). These results suggest a crucial role for dopamine in maintaining the normal function of the retina. In addition to the DA degeneration, α-synuclein aggregates have been identified in the retina of PD patients and could explain the visual abnormalities (Ortuño-Lizarán et al., 2018). Visual dysfunction can result from abnormalities beyond the retina such as abnormalities in the visual cortex (Weil et al., 2016).

1.5.3 The structure of the Drosophila visual system

The visual system in Drosophila consists of the retina and the optic lobe. The retina, or the compound eye, is made up of about 800 individual units known as ommatidia that are organised in hexagonal crystalline array. Each ommatidium is insulated by pigment cells that prevent light passing between ommatidia. The core of the ommatidium contains 8 photoreceptor (R) cells with 6 in the outer region (R1-R6) while the other two (R7-R8) in the inner one. These cells are arranged in an asymmetrical trapezoid and only 7 can be seen (R1-R7) because R8 is located below R7 (Kumar, 2012) (Figure 1.12). Each R cell contains a microvillar structure known as rhabdomeres which express a light absorbing molecule called rhodopsin (Rh). On the one hand, R1-R6 express Rh1 and are similar to the mammalian rods in that they are involved in dim light vision and motion perception. On the other hand, R7 and R8 are similar to the mammalian cones in that they involved in colour vision and express different types of Rh. R7 expresses UV-sensitive Rh3 and Rh4 while R8 expresses blue-sensitive Rh5 and green-sensitive Rh6 (Behnia and Desplan, 2015). Rh2 is an UV-sensitive Rh that is expressed in the ocelli which is a simple eye found in the upper part of the fly head responsible for light detection rather than image formation and will not be discussed any further here (Montell, 2012, Salcedo et al., 1999).
Figure 1.12. *Drosophila* eye with a schematic structure of the ommatidium.

The left image shows a *Drosophila* eye with the white square focusing on a group of ommatidia that are arranged in hexagonal crystalline array. The right image shows a schematic structure of a single ommatidium with the asymmetrical trapezoid arrangement of the photoreceptors above it.

The retina projects its synapses to the optic lobe which consists of the lamina, the medulla, and the lobula complex (lobula and lobula plate) (Figure 1.13). The modular organisation of the retina is maintained in the lamina where around 750 independent units called cartridges receive signals from the outer photoreceptors (R1-R6) that are involved in motion processing (Paulk *et al.*, 2013). The lamina contains five types of monopolar neurons that connect the retina to the medulla and amacrine cells that have connection within the lamina. The inner photoreceptors (R7 and R8) project directly to the medulla passing the lamina. The medulla is arranged in 10 layers (M1-M10) with the regions from M1 to M6 called the distal medulla and the regions from M7-M10 called proximal medulla. The distal medulla receives signals from the retina and the lamina while the proximal one receives information from the distal medulla. The medulla is the largest part of the optic lobe with around 40,000 neurons (Neriec and Desplan, 2016). The lobula complex is the last stop for visual information to go through before going to the central brain for further processing.
Figure 1.13. The anatomy of *Drosophila* visual system

The upper image shows a schematic structure of *Drosophila* visual system as described in (Nerice and Desplan, 2016). The lower image is adapted from (Afsari *et al.*, 2014) and shows photoreceptors (R1–R8), second-order lamina neurons (L1 and L2), amacrine neurons (A), and two kinds of third-order medulla neurons (C and T). The figure also highlights dopaminergic neurons (DA) that are intrinsic to the medulla and others that branch into the lamina.

1.5.4 Similarities between the *Drosophila* and human visual systems

In a quest to find a simpler neural circuitry than the retina of vertebrate, Santiago Ramón y Cajal and his colleagues turned to the fly visual system. To their surprise, they found cellular diversity and complexity similar to that observed in the vertebrate retina (Ramon y Cajal and Sanchez, 1915). Even though it is more than a century ago (Figure...
1.14), their detailed drawing is still relevant in illustrating the similarities between the two systems.

**Figure 1.14. Cajal’s drawing of fly and vertebrate visual systems**

A) A drawing of the fly visual system with the different types of neurons and layers indicated with letters and Roman numerals, respectively. B) A drawing of fly visual system with the neurons cells bodies moved to correspond to their position in vertebrates. C) A drawing of vertebrate visual system with the different types of neurons and layers indicated with letters and Arabic numerals, respectively, that correspond with the fly system. Adapted from (Sanes and Zipursky, 2010)

The similarities between the two systems are interior rather than exterior. These similarities include the arrangement of neurons in distinctive layers, the small number of main neuronal types (5 in vertebrate and 6 in fly), the presence of multiple contact synapses with a single presynaptic terminal in close contact to multiple postsynaptic elements, and the arrangement of neurons in orderly mapping from one level to the next (review by (Sanes and Zipursky, 2010)’s Figure 1.15). Furthermore, like humans, *Drosophila* has DA neurons that innervate the visual system (Hindle *et al.*, 2013).
Figure 1.15. Similarities between vertebrate and fly visual systems

A and B show the similar neurons and layers between the two systems in colour coded manner. C and D show the similar steps in information transfer through the visual system. Adapted from (Sanes and Zipursky, 2010)

The similarities between the two systems can be observed in the phototransduction which is the process of converting light into electrical signals. Both vertebrates and invertebrates carry out phototransduction via a specialised form of a G protein-coupled receptor cascade (Figure 1.16). When a photon is absorbed by a molecule of visual pigments, it causes the isomerisation of the light-sensitive vitamin A derivative 11-cis 3-hydroxyretinal to all-trans 3-hydroxyretinal; in vertebrates, the isomerisation of the chromophore 2-dehydroretinal is needed instead of 3-hydroxyretinal. In both cases, this isomerisation results in the activation of rhodopsin, which forms metarhodopsin. In flies, metarhodopsin binds to the alpha subunit of a heterotrimeric G-protein (Gq), causing GDP to be exchanged for GTP and the G-alpha subunit to be activated. Phospholipase C (PLC) is activated by the active G-alpha subunit, which cleaves
phosphatidyl inositol 4,5 bisphosphate into inositol triphosphate (InsP$_3$) and diacyl glycerol (DAG). There is a calcium influx into the photoreceptors, and two hypotheses have been proposed to explain how this influx happens. According to one of these hypotheses, when InsP3 binds to the InsP$_3$-receptors found in intracellular Ca$^{2+}$ storage, the cation channels TRP and TRPL are activated. As a result, Ca$^{2+}$ is released via a store-operated process. According to the second concept, DAG indirectly gates the TRP and TRPL channels, resulting in an influx of Ca$^{2+}$ and Na$^+$ into the photoreceptors. The influx of Ca$^{2+}$ leads photoreceptors to depolarize, resulting in histamine release at the synapse. This subsequently causes downstream neurons to hyperpolarize. In contrast to Drosophila, vertebrate phototransduction does not make use of the inositol phospholipid signalling pathway. Instead, the G-protein effector is a phosphodiesterase that converts 3'-5' cyclic guanosine monophosphate (cGMP) to 5' GMP. This causes cGMP-gated channels to close and hyperpolarisation to occur (Hardie and Juusola, 2015, Hardie and Raghu, 2001).

![Phototransduction cascades in vertebrates and Drosophila](image)

**Figure 1.16 Phototransduction cascades in vertebrates and Drosophila**

Numbers in circles (1-6) refer to the following steps: (1) Photoisomerisation; rhodopsin (R) is photoisomerised to metarhodopsin (M). Long-wavelength light can convert M back to R in Drosophila; in vertebrate, M releases the bound chromophore all-trans retinal. (2) GTP/GDP exchange; On the heterotrimeric G-protein (transducin in vertebrate rods, Gq in Drosophila), M catalyses the exchange of GDP for GTP, leading to the dissociation of the active GTP-bound α-subunit. (3) Activation; Ga binds to and activates the effector enzyme (PDE in vertebrate rods, PLC in Drosophila). (4) Substrate resynthesis; in vertebrates, cGMP is resynthesised by guanylate cyclase (GC) and GC-activating protein (GCAP), which is inhibited by Ca$^{2+}$. In Drosophila, DAG is converted to phosphatidylic acid (PA) by DAG kinase (DGK). PA is converted to PIP$_2$ via a multienzymatic pathway. (5) Metarhodopsin inactivation; rhodopsin kinase (RK) phosphorylates M, which is then capped by arrestin (A). In the presence of Ca$^{2+}$,
recoverin (Rec) inhibits RK in vertebrates. (6) Inactivation of G protein and effector; The GTPase activity of the G protein inactivates the effector enzyme and Ga, resulting in reassociation with Gβ, γ. This is accelerated by the activity of RGS9, Gβ5 and PDE in vertebrate rods, and by PLC in Drosophila (Hardie and Raghu, 2001).

In conclusion, the retina is highly organised tissue and shares key neurons and features with the brain. It provides an accessible platform to study the CNS and this accessibility led to the development of many experimental non-invasive tools. Using the fly visual system to model PD can help shed some light on visual dysfunction and DA neurons loss found in PD.

1.6 Aims

Since the discovery of LRRK2 association to PD in the early 2000s, many studies have been undertaken to investigate the cellular basis of this link. No precise mechanism has been identified and studies are still on going to elucidate how mutations in LRRK2 lead to PD pathology. This project aims to further our knowledge on the mechanism of action of LRRK2 and how mutations in this gene may lead to neurodegeneration. Drosophila melanogaster will be used mainly as the animal model. Loss-of-function (LOF) and gain-of-function approaches will be used to examine the effect of LRRK2 in the visual response of flies. Examining the visual response of flies provides results that can be quantifiable and help us understand the consequences of producing too little or too much LRRK2 in dopaminergic neurons in a functioning nervous system. Additionally, this project will aim to establish the stability and half-life of LRRK2 protein. This will help us understand the functional window over which the protein functions. The specific contributing aims of this investigation are outlined below:

1. To test the autonomy of LRRK2 activity by assessing the ability of dLRRK, the Drosophila orthologue of human LRRK2, to act in neuronal tissues when expressed in non-neuronal tissues to rescue visual dysfunction in old flies. This study will be linked an examination of the genetic interaction between eye colour genes and dLRRK where previously a synthetic lethality has been observed.
2. To determine the effect of increased LRRK2-G2019S activity on eye anatomy and physiology.
3. To test the stability of LRRK2 protein in vitro and vivo and the effect of mutations and kinase inhibitors on its stability.
2 Materials and Methods

2.1 Drosophila husbandry and genetics

2.1.1 Drosophila stocks

*Drosophila* stocks used in this project were purchased from Bloomington *Drosophila* Stock Centre (BDSC, Indiana University, Bloomington, USA), or were kindly donated from members of the *Drosophila* community. Rebalancing or recombination stocks also generated additional stocks. A summary of primary stocks used in this project are listed in Table 2.1. Stocks were quanrantined before first use for at least 2 generations to confirm that they were mite free.

Stocks were raised at either 18°C or 25°C and were transferred to fresh media every 4 or 2 weeks, respectively. Experimental crosses performed were raised at either 25°C or 29°C, giving a generation time of about 10-12 days.

**Table 2.1 Fly Stocks**

<table>
<thead>
<tr>
<th>Stock (referred to as)</th>
<th>Description</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Wild type</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Canton-S (+)</em></td>
<td>Wild-type, red eyes</td>
<td>Elliott/Sweeney Lab Stock</td>
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<tr>
<td>w*&lt;sup&gt;1118&lt;/sup&gt;</td>
<td>Wild-type, white eyes</td>
<td>Elliott/Sweeney Lab Stock</td>
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<tr>
<td><em>w</em>&lt;sup&gt;apricot&lt;/sup&gt;</td>
<td>Wild-type, orange eyes</td>
<td>BDSC #148</td>
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<td><strong>Balancer Stocks</strong></td>
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<td></td>
</tr>
<tr>
<td><em>w</em>&lt;sup&gt;+&lt;/sup&gt;; +/+; TM3/TM6B</td>
<td>Third Chromosome Balancer</td>
<td>Elliott/Sweeney Lab Stock</td>
</tr>
<tr>
<td><em>W</em>&lt;sup&gt;+&lt;/sup&gt;;CyO/Iff; TM6B/MKRS (DB)</td>
<td>Second and Third Chromosome Balancer</td>
<td>Elliott/Sweeney Lab Stock</td>
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<tr>
<td><strong>Mutant Stocks</strong></td>
<td></td>
<td></td>
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<tr>
<td><em>w</em>&lt;sup&gt;+&lt;/sup&gt;; +/+; dLRRK&lt;sup&gt;e03680&lt;/sup&gt; (dLRRK&lt;sup&gt;LOF&lt;/sup&gt;)</td>
<td>PBac(Burbulla <em>et al.</em>) PiggyBac-element disruption of dLRRK, generation of dLRRK loss of function mutant</td>
<td>(Wang <em>et al.</em>, 2008a)</td>
</tr>
<tr>
<td><em>cn</em>&lt;sup&gt;35k&lt;/sup&gt;</td>
<td>cinnabar hypomorphic allele mutant</td>
<td>BDSC #268</td>
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<tr>
<td><em>pr</em>&lt;sup&gt;l&lt;/sup&gt;</td>
<td>purple mutant</td>
<td>BDSC #370</td>
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<td><strong>Gal4 Stocks</strong></td>
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<td>P(Gal4)56C (Pigment cell)</td>
<td>Eye pigment cells specific driver.</td>
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<td>P(GawB)c564 (Reproductive)</td>
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<td>Lsp2-Gal4 (Fat body)</td>
<td>Fat body specific driver.</td>
<td>BDSC #6357</td>
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<tr>
<td>rh1-Gal4/CyO</td>
<td>Rhodopsin 1 promoter; photoreceptor specific driver</td>
<td>(Gonzalez-Bellido et al., 2009)</td>
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<td>TH-Gal4</td>
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### UAS Stocks

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<td>UAS-hLRRK2</td>
<td>Human LRRK2 transgene</td>
<td>(Liu et al., 2008)</td>
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<tr>
<td>UAS-hLRRK2-G2019S</td>
<td>Human LRRK2-G2019S mutant transgene</td>
<td>(Lin et al., 2010)</td>
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<td>Human LRRK2-R1441C mutant transgene</td>
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<tr>
<td>UAS-hLRRK2-I2020T</td>
<td>Human LRRK2-I2020T mutant transgene</td>
<td>(Venderova et al., 2009)</td>
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<td>UAS-hLRRK2-I1122V</td>
<td>Human LRRK2-I1122V mutant transgene</td>
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<tr>
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<td>(Kim et al., 2017)</td>
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<td>Second chromosome <em>sepi</em> insertion</td>
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<tr>
<td>UAS-pu</td>
<td>Third chromosome <em>punch</em> insertion</td>
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</table>

### 2.1.2 Drosophila media

Stocks were maintained in 25x95 mm plastic vials (Dutscher Scientific, UK) plugged with cotton wool (Fisher Scientific, UK) containing about 7 ml standard yeast-sucrose-agar media: (25 g/l sucrose, 3.75 g/l agar, 0.125 g/l CaCl₂, 0.125 g/l FeSO₄, 0.125 g/l MnCl₂, 0.125 g/l NaCl, 2 g/l KNaC₄H₄O₆.4H₂O); following boiling and cooling for 1 hr to about 45°C, the antifungal agents Bavistin (1.5 mg/l in 100% ethanol [EtOH]; BASF, Auckland, New Zealand) and Nipagin (0.7 mg/l in 100% EtOH; Sigma, UK) were added. Experimental flies kept on this media were transferred to fresh vials every 3-4 days.
For drug experiments, 4-24® instant *Drosophila* medium (Carolina Biological Supply Company, USA) was utilised. The instant media was prepared by mixing 50:50 with dH2O. Stocks solutions of pharmaceuticals (10 mg/ml) were prepared by dissolving the drugs into 100% ethanol or dH2O with subsequent storage at 4°C or -20°C. The appropriate volumes of these stocks were added to dH2O to give the desired concentration according to the experiment before it was mixed with the instant food media.

In some instances, dried yeast is added to the medium to encourage flies to lay eggs. Experimental flies were aged at 29°C in complete darkness. When required in large numbers, stocks were raised in 1/3 pint bottles.

### 2.1.3 *Drosophila* anaesthesia

Adult flies were anaesthetised on a porous gas pad with continuous CO2 administration to identify gender and genotype. Anaesthetised flies were observed using a dissecting microscope (Zeiss Stemi-2000, Carl Zeiss AG, Germany).

### 2.1.4 *Drosophila* crossing techniques

A crucial step in performing *Drosophila* crosses is to isolate virgin females in order to have a controlled genetic cross. *Drosophila* males and females are easily distinguished. Males have blackened tip to their abdomen and a patch of bristles on their foreleg, known as sex combs, compared to the females. Newly eclosed flies exhibit a pale pigmentation, unexpanded wings and the presence of a meconium, visible through the abdominal cuticle (Figure 2.1). At 25°C, female flies will not mate within 8 hours of eclosion. Based on this criterion, virgin females were collected in the morning through completely emptying vials and isolating obvious virgins based on the presence of the meconium and lack of cuticle tanning and then collecting any further females that eclosed within the following 8 hrs. Adult males and female virgins were then crossed in a fresh food vial. The parent flies were then removed after 7 days and transferred into a new vial/bottle or discarded in order to prevent over-crowding and allow specific selection of first generation for further crosses or experiments. All the experimental crosses performed were raised at either 25°C or 29°C, giving a generation time of 10-12 days (egg to adult).
Figure 2.1 Selection criteria for Drosophila crossing

A-C Views of male (top) and female (bottom) Drosophila. A) Lateral whole body view with a magnified view of the front legs showing the presence of sex combs in male. B) Dorsal abdomen view showing a slightly larger female with dark separated stripes at the posterior tip that is merged in male. C) Ventral abdomen view showing a male with darker and more complex anal plates than the pin-like extension in females. D) Lateral whole body view of a virgin female showing pale pigmentation and unexpanded wings. The arrow indicates the presence of a meconium (larval food remains) which is visible in the early hours after eclosion.

2.1.5 Recombination

To allow the use of two or more genetic components on a single chromosome, Homologous chromosomal recombination in female flies was utilised to generate stocks in which two genetic components were present on the same chromosomal arm. For example, this approach was used to recombine the dLRK<sup>e03680</sup> piggyBac mutation with both Lsp2-Gal4 (fat body driver) (Figure 2.2). Recombination is achieved through mating individual stocks with each other and selecting against balancers to identify male offspring that carry the two desired genetic components. Chromosomal recombination does not occur in male flies, so identified males can be then crossed to appropriate balancer stocks to capture the required synthetic chromosome containing the desired elements. Individual flies that carry two transgenic elements often have a darker eye colour due to the presence of two copies of the mW<sup>e</sup> (mini-White) eye colour cassette,
one being present on each element. These flies can be easily recognised and selected. These offspring were then crossed to virgin females from balancer stocks for the relevant chromosome and potential recombinant offspring were selected. Where possible, these selections were made based on eye colour as well as other characteristic phenotypes. The presence of the desired genetic components was confirmed via PCR for \( dLRRK^{e03680} \) (see Figure 2.7) and the expression of GFP for \( Lsp2\)-Gal4 which was carried out by crossing the Gal4 fly to a UAS-GFP fly and then observing the expression in the progeny.

![Figure 2.2. Recombination scheme in a Drosophila to allow the use of 2 desired elements on the same chromosome](image)

**Figure 2.2. Recombination scheme in a Drosophila to allow the use of 2 desired elements on the same chromosome**

The first cross is carried out to produce a fly that carry the 2 elements of interest (\( Lsp2\)-GAL4 and \( dLRRK^{e03680} \)) on the same chromosome. Then, only female flies are crossed with a balancer fly to aid in appropriate offspring selection of flies with recombined chromosomes (darker eye, \( 2x\ mW^+ \), and extra hairs on the thorax, \( Hu \) for \( TM6B \)). The final cross is carried out with only male flies to balancer flies to produce the stock and experimental flies.

2.2 Cell culture

2.2.1 Cell lines

Cell lines used in this investigation are Flp-In T-REx 293 cells that stably express different \( LRRK2 \) mutations (\( G2019S \), \( D2017A\)-Kinase dead) and the WT in inducible
and controlled fashion. These cells were obtained from Dario Alessi’s lab (Dundee) and were made by transfecting HEK293 cells with plasmids and vector containing \textit{LRRK2} gene (Nichols \textit{et al.}, 2010). This fashion of expression is achieved due to the presence of tetracycline repressor that prevent \textit{LRRK2} expression unless tetracycline (TC) or doxycycline (Doxy) is added (Figure 2.3).

\textbf{Figure 2.3. Flp-In T-REx system mechanism}

Tetracycline repressor (TC\textsubscript{R}) protein is expressed and binds to tetracycline operator 2 (TetO\textsubscript{2}) sequence resulting in suppression of gene X expression. When tetracycline (TC) is added, it binds to TC\textsubscript{R} and causes a conformational change that leads to its release from TetO\textsubscript{2} and de-suppression of gene X expression.

\subsection{2.2.2 Growth and maintenance of cells}

Cells were cultured in 1X Dulbecco’s Modified Eagle Media (DMEM) (gibco, Invitrogen; + 4.5g/L D-Glucose, L-Glutamine, + Pyruvate) supplemented with 10\% Foetal Bovine serum (FBS), 1:100 Penicillin/Streptomycin, 15 \textmu g/ml blastocidin, and 100 \textmu g/ml hygromycin.

To maintain cells for experiments, they were passaged every 3-4 days with a ratio of 1:5 in a 25cm culture flask. When cells reached around 90\% in confluence, they were passaged by removing the media and then washing the flask with sterile Phosphate-
Buffered Saline (PBS). Cells were detached from the surface of the flask using either pre-warmed 1X trypsin/EDTA solution (Sigma; 0.5g porcine trypsin, 0.2g EDTA 4Na/L Hanks) (37°C for 3-5 mins) or a cell scraper. When using trypsin, the flask was hit on the sides firmly to dislodge cells after the incubation. After that, medium was added to neutralize trypsin and the cell mix transferred into 15 ml falcon tube. The tube was centrifuged at 1000 rpm for 5 mins and the supernatant is discarded. A pellet was left at the bottom of the tube that was resuspended in 1ml of media and passaged into new flaks according to appropriate ratios.

2.2.3 Counting and plating of cells

From the 1 ml resuspended pellet (see previous section), about 15 μl is pipetted and added under a coverslip in a haemocytometer. Using a 10X objective in inverted microscope, cells were counted in one of the 4 large corner squares (4 X 4 grid, Figure 2.4). If there was fewer than 50 cells counted, another couple of squares were counted and an average is taken. The number of cells per ml was calculated by the number of cells in a 4 X 4 square multiplied by 10,000 e.g., a count of 100 = 1 X 10⁶ cells/ml. Cells will be diluted according to the typical plating densities (5-50,000 cells/well for a 24 well plate, 100,000-250,000 cells/well for a 6 well plate and 1-2 million cells/well for a 10 cm dish).

Figure 2.4. Haemocytometer gridlines

A haemocytometer with a zoomed-in image of the gridlines seen under the microscope to count cells. Cells were counted in any one of the 4 X 4 grid (red squares).
For LRRK2 stability and drug experiment, 6-well plates were used. After plating, cells were left for 24 hours to stick to the bottom of the flasks before doxycyclin was added (1 μg/ml) to induce the expression of LRRK2. It was then removed to stop the expression and the drug of choice was then added. Stocks solutions (10 mg/ml) were prepared by dissolving the drugs into DMSO (for MLi-2) or dH₂O (for B12) and were stored at 4°C or -20°C. The appropriate volumes of these stocks were added to the media to give the desired concentration according to the experiment.

2.3 Physiological and anatomical analyses

2.3.1 Flash Electroretinogram (fERG)

Unanaesthetised flies were pushed to the narrow end of a trimmed 200 μl yellow Gilson pipette tip so that just the head was left protruding. The fly then was fixed in place using nail varnish (La Femme, England, UK). In the ERG device, one glass electrode filled with simple Drosophila saline (130 mM NaCl, 4.7 mM KCl, 1.9 mM CaCl₂; Heisenberg, 1971) was placed on the surface of the eye for recording. Another glass electrode filled with the same saline is placed on the mouthparts was used as a reference. In one setup, flies were dark-adapted for 2 mins and then ERGs were recorded in response to five stimuli (10 seconds apart, 0.5 seconds long) of multiple channels blue LED light (Kingbright, KAF-5060PBESEEVG, maximum emission wavelength 465 nm, Taipei, Taiwan) placed about 6 cm in front of the fly. The five stimuli were averaged for each fly. In another setup (eye adaptation experiment), a single LED channel was used to emit rapid flashes (100ms) of blue light at 5 seconds interval for 4 minutes in the dark. The amplitude of around 25 ERGs were averaged for each fly. An example of an ERG trace is shown in Figure 2.5.
Figure 2.5. Recording the visual response of *Drosophila* using the flash ERG

A. Flies were restrained in shortened yellow pipette tips and exposed to light from the blue component of an LED light. An electrode filled with a simple saline solution was placed on the surface of the eye to record the response of the visual network, whilst a second electrode was placed within the mouthparts to act as a reference. B. A typical recording from a WT fly with the three main ERG components indicated. The on-transient was determined as the potential difference between the starting potential and the maximum value of the ERG trace. The photoreceptor response was determined as the potential difference between the starting level and the potential about halfway along the recording. The off-transient was determined as the potential difference between the end of the photoreceptor response and the minimum value of the ERG trace.

2.3.2 Deep Pseudopupil (DPP) and Pseudopupil Analysis (PPA)

For DPP, flies were prepared as described in the previous section (section 2.3.1) with an external white light positioned at an angle that would produce DPP (antidromic) in the eye of the fly. Footage was taken by an eyepiece camera (Dino-Eye Digital Eye Piece Camera, Dino-Lite, USA) fitted onto a dissecting microscope (Zeiss Stemi-2000 Carl Zeiss AG, Germany). For PPA, fly heads were decapitated on CO₂ pad and mounted onto a microscope slide using clear nail polish to fix the heads in place. Decapitated heads were held carefully from the proboscis using a tweezer to keep the eyes intact and heads should be fixed on the edges of the clear nail polish (Figure 2.6). When nail polish was dry, slides were examined under glycerol or oil 40X objective lens (Nikon) to analyse individual ommatidium and its rhabdomeres.
2.4 Molecular Biology

2.4.1 Genomic DNA isolation

Genomic DNA was isolated from single adult flies by homogenisation in 50 μl of extraction buffer (25 mM NaCl, 10 mM Tris-HCL pH 8.2, 1 mM EDTA) with the addition of Proteinase K (200 μg/ml). Proteinase K was added fresh on the day of the extraction. The homogenate was incubated at 37°C for 30 mins, followed by an incubation at 85°C for 10 mins in order to inactivate the proteinase K. To allow separation of particulates, the homogenate was centrifuged at 13000 rpm for 3 mins. 1-2 μl of supernatant was used as a PCR template. To increase the amount of isolated DNA, more than one fly was used.

2.4.2 Polymerase Chain Reaction (PCR)

The total volume of a PCR reaction was 20 μl and was run using 10 μl PCR mastermix (Promega, UK; 25 U/ml Taq DNA polymerase, Taq Reaction buffer, 200 μM of each dNTP, 1.5 mM MgCl₂), 1 μl of Forward primer and 1 μl of Reverse primer, 1-2 μl of genomic DNA and 6-7 μl of nuclease free water. A list of primers used in this investigation can be found in Table 2.2. Primers were ordered from Sigma-Aldrich, UK.
Table 2.2. Primer sequences

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5’ – 3’)</th>
</tr>
</thead>
</table>
| UAS-pu   | Forward TCTACGGAGCGACAATTCAATTC  
Reverse CAGTGGTGTTTTCGTTGTG | |
| UAS-dLRRK| Forward TCTACGGAGCGACAATTCAA  
Reverse AGAAGGGTGTTTCTGCCTGA | |
| dLRRKe03680| Forward CGATAAAACACATGCAGTC  
Reverse GGCTAACCGATGCAGAGGA | |

Reactions were run in a Techne PRIME PCR thermocycler for 30 cycles. The PCR program was run as 95°C for 5 mins for an initial denaturation followed by 30 cycles of denaturation at 95°C for 30 sec, annealing temperature was calculated as 5°C lower than the lowest primers melting temperature I for 30-40 sec, and extension at 72°C for 1 min per kb. A final step of extension at 72°C for 10 mins was performed before the reaction was cooled down to 4°C to prevent DNA decomposition.

### 2.4.3 DNA agarose gel electrophoresis

To analyse the product of PCR, agarose gel electrophoresis was utilised. 1 g of agarose was heated using a microwave oven in 100 ml of 1X TAE buffer to produce 1% agarose gel. The distance travelled by a PCR fragment in agarose in a gel is inversely related to the size of PCR product. To visualise DNA in the gel using a blue light transilluminator, SYBR® safe (Invitrogen, UK; 10 μl/100 ml) was added to the gel when it was sufficiently cool prior to pouring 6X DNA loading dye (0.25% bromophenol blue (w/v) and 30% glycerol (v/v)) was added to PCR product in order to assist in loading the samples into the wells of the gel. A 1 kb or 100 bp DNA ladder (0.5 ug/lane; NEB, UK) was run alongside the DNA products in order to visualize the DNA band sizes. All gels were run at 100V for 30-45 mins.

PCR followed by gel electrophoresis was used to screen stocks for the presence of the dLRRKe03680 piggyBac P-element following a recombination. An example gel with wild-type (w^{118}1), homozygous dLRRKe03680 and successfully recombined Lsp2-GAL4, dLRRKe03680/dLRRKe03680 stocks is shown in Figure 2.7.
Figure 2.7. PCR of Mutant $dLRRK^{e03680}$ recombinants

Primers were designed to detect the presence of $dLRRK^{e03680}$. The gel shows a successful recombination of $dLRRK^{e03680}$ with $Lsp2$-Gal4. The first lane is $w^{1118}$ (negative control), the second lane is the homozygous $dLRRK^{e03680}$ (positive control), and the third and fourth lanes are the successful recombinants in the homozygous and heterozygous forms. The presence of $Lsp2$-Gal4 was confirmed by GFP expression.

2.5 Western blotting

The method used for western blotting was adapted from Abcam (2016).

2.5.1 Protein extraction

Protein was extracted from adult fly heads. Flies were collected in 1.5 ml Eppendorf tubes and snap frozen on dry ice or liquid nitrogen. To separate the heads from the rest of the body, the 1.5 ml Eppendorf tube was placed inside a 50 ml falcon tube containing dry ice and vortexed 2-3 times for 30-60 secs. Fly heads were then collected into a new 1.5 ml Eppendorf and kept on dry ice or stored in -20°C till ready for extraction. RIPA buffer (150mM NaCl, 1.0% IGEPAL® CA-630, 0.5% sodium deoxycholate, 0.1% SDS, 50 mM Tris, pH 8.0; Sigma, UK) was used for protein extraction with the addition of protease inhibitor cocktail tablets (cOmplete tablet, Mini EDTA-free, EASYpack;
Roche). 1μl of buffer per head was added to the heads and they were homogenized by a sterile plastic pestle and left on ice for 30 mins. After the incubation, tubes were centrifuged at 14000 rpm for 15 mins at 4°C. The extracted protein in the supernatant was removed into a new tube.

For haemolymph extraction of adult flies, an adapted protocol from protocol.io (https://www.protocols.io/view/Haemolymph-extraction-of-adult-Drosophila-dkn4vd) was used. A hole was made at the bottom of a 0.5 ml Eppendorf tube and put into a 1.5 ml Eppendorf tube with the lid removed. Flies were anesthetised on a CO₂ pad and their thorax were speared with a needle. 40 speared flies were collected into the 0.5 ml Eppendorf tube on ice and then centrifuged for 1 min at 5000 rpm at 4°C. Around 1μl of haemolymph was collected.

For cell culture, the same RIPA buffer used in fly protein extraction was used here as well. Cells were washed with ice cold PBS and then RIPA buffer was added. For a 6-well plate, 60 μl of RIPA was used and cells were scraped off of the plate using cell scraper. Then, lysates were transferred into 1.5 ml Eppendorf and incubated for 30 mins on ice. After the incubation, tubes were centrifuged at 14000 rpm for 5 mins at 4°C. The extracted protein in the supernatant was removed into a new tube.

2.5.2 Quantification of protein concentration

Protein concentration was calculated using BCA kit (Pierce, Fisher Thermo Scientific, UK). A standard curve of bovine serum albumin (BSA) was produced by diluting a known BSA concentration (2 μg/ml) serially (2, 1. 0.5, 0.25, 0.125, 0.0625, 0.03125, 0 μg/ml) in RIPA buffer. To increase the quantification accuracy, lysates were diluted 2X and 10X and the average is taken. The assay was performed in a 96-well plate according to the manufacturer instructions. Absorption values were measured at 562 nm using a spectrophotometer (BMG Labtech Clariostar Plate Reader, Ortenberg, Germany). Absorption spectra values were used to determine protein concentration of samples in μg/μl.

2.5.3 Sodium Dodecyl Sulphate Polyacrylamide Gel Electrophoresis (SDS-PAGE)

Samples were mixed with 5X laemml buffer (0.25M Tris-HCL, 10% SDS w/v, 50% glycerol v/v, 25% β-mercaptoethanol v/v, and 0.02% bromphenol blue w/v of 6.8 pH)
to a working concentration of 1X and boiled for 5 min at 85°C to denature protein. Protein samples (20 µl in a loading concentration of 7.5-10 µg/µl) and a protein ladder (6 µl PageRuler™ Plus Protein Ladder, Thermo Fisher Scientific, UK) were loaded into a handcasted gel of 7.5% resolving gel and 4% stacking gel. Gels were run in Criterion™ Vertical Electrophoresis Cell (Bio-Rad, UK) connected to a PowerPac™ Basic Power Supply (Bio-Rad, UK) filled with running buffer (25 mM Tris, 192 mM glycine, 0.1% SDS pH 8.3) for 30 min at 80V followed by ~ 1.5 hrs at 100V in cold conditions.

2.5.4 Protein transfer to polyvinylidene fluoride (PVDF) membrane

After protein separation in SDS-PAGE, a Mini-Trans-Blot Cell (Bio-Rad) was used to transfer proteins from the gel to a PVDF membrane (Amersham Hybond 0.45 µm PVDF; GE Healthcare, UK). Foam pads, Whatman gel blot paper and the SDS-PAGE gel were soaked in transfer buffer (25 mM Tris, 192 mM glycine, 20% methanol v/v). PVDF membrane was cut according to the gel size and then activated in 100% methanol for 30-60 secs followed by a quick dip into the transfer buffer. In the Mini Gel Holder Cassette, one foam pad is placed on the negative (black) side followed by a one Whatman gel blot paper and then the gel. On top of that, the activated PVDF membrane with a one Whatman gel blot paper and one foam pad. Avoiding any bubbles trapped between the gel and the PVDF membrane, the cassette is closed carefully and placed into the transfer tank filled with the transfer buffer. Proteins were transferred by applying 30V overnight increasing this to 60V the next morning for 30 mins or applying 80-100V for 1-2 hrs in cold conditions.

2.5.5 Probing of PVDF membrane

After protein transfer to the PVDF membrane, the membrane was washed in Tris buffered saline-Tween (TBS-T; 10 mM Tris, pH 7.6, 150 mM NaCl supplemented with 0.1% (v/v) Tween® 20) 2 times for 5 min at RT followed by the addition of a blocking solution, either 5% (w/v) BSA or skimmed milk in TBS-T, for 1 hr to prevent any non-specific binding. This was followed by an overnight incubation at 4°C of the primary antibody in 5% (w/v) BSA in TBS-T. Excess antibody was removed via washing the membrane 3 times for 10 min at RT. The suitable species of secondary antibody (conjugated to HRP) was added in either 5% (w/v) BSA or skimmed milk in TBS-T for 1-2 hrs at RT followed by removing excess antibody via washing the membrane in TBS-T 3 times for 10 min at RT. All washes and antibody incubation were performed on a rocking platform. The signal was visualised with chemiluminescent substrate
(Amersham ECL Prime Western Blotting Detection Reagent, GE Healthcare, Germany) according to the manufacturer instructions and was imaged either by exposing the blot in the dark to UltraCruz® X-Ray autoradiography films (Santa Cruz Biotechnology, Germany) or visualised using iBright™ FL1000 imaging system (Invitrogen, Thermo Fisher Scientific, UK). The list of antibodies used in this investigation can be found in Table 2.3.

**Table 2.3. List of antibodies used in western blot**

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Species</th>
<th>Dilution</th>
<th>Source</th>
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<tr>
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<td>NeuroMab</td>
</tr>
<tr>
<td>hLRRK2</td>
<td>Rabbit</td>
<td>1:10,000</td>
<td>Abcam</td>
</tr>
<tr>
<td>dSynaptotagmin1</td>
<td>Rabbit</td>
<td>1:1000</td>
<td>(West et al., 2015b)</td>
</tr>
<tr>
<td>β-Tubulin</td>
<td>Mouse</td>
<td>1:1000</td>
<td>Proteintech</td>
</tr>
<tr>
<td>GAPDH</td>
<td>Mouse</td>
<td>1:10,000</td>
<td>Abcam</td>
</tr>
<tr>
<td>pRab10</td>
<td>Rabbit</td>
<td>1:1000</td>
<td>–</td>
</tr>
<tr>
<td>panRab10</td>
<td>Mouse</td>
<td>1:1000</td>
<td>–</td>
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<table>
<thead>
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<th>Antibody</th>
<th>Species</th>
<th>Dilution</th>
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<td>Jackson ImmunoResearch</td>
</tr>
<tr>
<td>Rabbit IgG HRP linked</td>
<td>Goat</td>
<td>1:10,000 – 20,000</td>
<td>–</td>
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</tbody>
</table>

**2.6 Statistical analysis and bioinformatics**

All statistical analyses were carried out in GraphPad Prism. Student’s t-test was performed to test for statistical significance between two groups; univariate ANOVA followed by a post-hoc Tukey comparison was performed when comparing genotypes to control. Statistical significance was defined as p<0.05 throughout. Error bars are either SD (±SD) or SEM (±SEM). P values are indicated graphically with ****p<0.00001 ***p<0.001, **p<0.01, *p<0.05, ns (no significant difference). The half-life (t_{1/2}) calculation for protein degradation was carried out in GraphPad Prism using non-linear regression analysis (curve fit) one phase decay model. The fitting method used was (Least square regression) with plateau = 0. Confidence Intervals (CI) of parameters are either 95% or 90%.

During the course of this investigation the following software and online resources were used. Unless otherwise stated default settings, assumptions and parameters were used.

For data management, analysis, and graphs plotting:
• Microsoft Excel
• GraphPad Prism v9.

For graphic design and image processing:
• Microsoft PowerPoint
• Photoshop (template from Roote and Prokop, 2013 Genotype Builder)
• BioRender.com
• Fiji
• DinoXcope software

For primer design:
• Primer3 (http://bioinfo.ut.ee/primer3-0.4.0)

For western blots quantification:
• Image Studio Lite

For ERG data processing:
• MATLAB (full code can be found at https://github.com/wadelab/flyCode)
• DASYLab for ERG recording
• DASYView for ERG analysis (customed software, C. J. H. Elliott, University of York).
3 The effects of dLRRK<sup>e03680</sup> on eye physiology and eye pigment

3.1 Introduction

3.1.1 Drosophila LRRK (dLRRK)

The *Drosophila* genome has one orthologue to human LRRK1/LRRK2 which is dLRRK. (CG5483) found on the right arm of chromosome 3. The dLRRK protein is around 2400 amino acids in length and structurally appears more similar to LRRK1 than LRRK2 as dLRRK does not have the ANK and WD40 domains (Figure 3.1). The GTPase domain of dLRRK shows 33% and 30% identity to the GTPase domain of hLRRK2 and hLRRK1, respectively, and the dLRRK kinase domain shows 36% and 38% identity to the respective kinase domains of hLRRK2 and hLRRK1. Like LRRK2, dLRRK is ubiquitously expressed in *Drosophila* tissues and has been found largely in the cytoplasm associated with membranous structures such as lysosomes, endosomes, and synaptic vesicles (Langston *et al.*, 2016).

![Figure 3.1 Comparison of the domain structure of LRRK proteins in human and Drosophila.](image)

The predicted domains are indicated by labelled boxes. Percentage identities based on a Clustal 2.1 multiple sequence alignment are given relative to human LRRK2 (left, bolded) and relative to human LRRK1 (right) for LRR, Roc-COR and kinase domains. Adapted from (Langston *et al.*, 2016).

As the structural homology implies, dLRRK is functionally an active kinase capable of autophosphorylation (Imai *et al.*, 2008) and phosphorylation of many substrates. These diverse substrates may include 4E-BP, Futsch, endophilin A and ribosomal protein S15
In addition, dLRRK has been shown to interact with different membrane traffic-regulating Rab proteins in *Drosophila* follicle cells (Dodson et al., 2012). The similarities between *dLRRK* and *hLRRK2* can also be observed in the similar effects they produce when manipulating their expression. The transgenic expression of mutant *dLRRK* in *Drosophila* DA neurons has shown a severe age dependent cell loss when expressed with *dLRRK*-Y1383C (equivalent to *LRRK2*-Y1699C) or *dLRRK*-I1915T (equivalent to *LRRK2*-I2020T) are expressed compared to the wild type *dLRRK* expression (Gehrke et al., 2010, Imai et al., 2008). The systemic loss of function mutant *dLRRKe03680* or the overexpression of *hLRRK2*-G2019S in neuronal cells of the *Drosophila* have previously been shown to have caused an age dependent loss of visual function (Furmston, 2016, Hindle et al., 2013). Although these results have shown a close resemblance between *dLRRK* and *hLRRK2* function, other reports showed irreproducible or conflicting outcomes. For example, the DA neurodegeneration was not reported in all studies (Wang et al., 2008a) and the synaptic transmission in *dLRRK* mutant flies was found to be significantly depleted in one study (Lee et al., 2010) while in another one no significant change was reported (Matta et al., 2012). These results show the conservation between *dLRRK* and human *LRRK2* which makes investigating *dLRRK* even further for PD modelling a valuable tool.

### 3.1.2 Vision and *dLRRK*

Of non-motor symptoms reported by PD patients, a common one is disturbances in the visual function. This could be due in part to the loss of retinal DA neurons (Harnois and Di Paolo, 1990) such as the DA amacrine and inter-plexiform cells (Zhang et al., 2021) that eventually affects the normal neural circuitry of the eye. Moreover, an age dependant visual dysfunction was observed in flies expressing human *LRRK2*-G2019S in DA neurons (Hindle et al., 2013) and flies carrying the loss of function mutation *dLRRKe03680* (Furmston, 2016). This visual dysfunction was assessed using an electroretinogram (ERG) technique which measures field potentials from the neuronal activity from the photoreceptors and the underlying second-order lamina neurons in response to pulses of light (see Figure 3.2). The visual dysfunction observed in these two *Drosophila* models might look similar, however, it is well to note that the decline of visual function was observed in all elements of the ERG (on-transient, photoreceptors, and off-transient) in *LRRK2*-G2019S model while in the loss of function *dLRRK*, it was mostly observed in the off-transient element.
Figure 3.2 Age dependant loss of visual function in flies expressing human *LRRK2-G2019S* in DA neurons and *dLRRK*\(^{e03680}\) flies.

Representative ERG traces from young (left, 3 days) and old (right, 28 or 21 days) flies expressing human *LRRK2* in DA neurons (*TH>*h*LRRK2*), *LRRK2-G2019S* in DA neurons (*TH>*G2019S*), wild-type (*dLRRK*), and *dLRRK* loss of function (*dLRRK*\(^{e03680}\)) flies. ERGs are recorded in response to 0.5 second blue light pulses. The blue lines drawn below the bottom ERG traces represent the duration of the light pulse. Scale bars for time (seconds) and potential (mV) are shown. Adapted and modified from (Furmston, 2016, Hindle et al., 2013).

Interestingly, the loss of visual function observed in *dLRRK*\(^{e03680}\) flies was rescued with the expression of the native gene (*dLRRK*) autonomously in CNS (Furmston, 2016). The ability of the wildtype *dLRRK* transgene to rescue loss-of-function *dLRRK* mutants was tested in the lamina neurons, DA neurons and photoreceptors (Figure 3.3).
3.3 $dLRRK$ expression in the lamina neurons, DA neurons or photoreceptors in $dLRRK^{LOF}$ background rescues the age-dependant loss of visual function.

Representative ERG traces from 3 days (left) and 21 days (right) of wild-type ($dLRRK^+$; green), $dLRRK^{LOF}$ mutant (red) and flies expressing $dLRRK$ specifically in the lamina neurons, DA neurons or photoreceptors in a $dLRRK^{LOF}$ background (shades of blue). ERGs are recorded in response to 0.5 second blue light pulses. The blue lines drawn below the bottom ERG traces represent the duration of the light pulse. Scale bars for time (seconds) and potential (mV) are shown. Each trace is the average of the fly’s response to at least three flashes of light. Adapted and modified from (Furmston, 2016).

This previous data in addition to subsequent analysis from others like (Afsari et al., 2014) demonstrate the essential role of dLRRK/hLRRK2 for the maintenance of normal visual function.

3.1.3 *Drosophila* eye pigments biosynthesis

Genes regulating eye pigmentation have been linked to neurodegeneration including PD and Huntington’s Disease (Campesan et al., 2011, Cunningham et al., 2018, Maddison and Giorgini, 2015, Tan and Guillemin, 2019). Observing the effects of manipulating the different genes involved in the synthesis of these pigments is a valuable tool to uncover the link between PD and pigmentation regulation. It is of interest to note that the substantia nigra pars compacta, where the PD sensitive population of DA neurons is
found, is a pigmented neuronal structure containing neuromelanin (Nagatsu et al., 2022).

Wild type flies have red-brown eye colour due to a combination of two classes of pigments found in their pigment cells. These pigments are ommochromes (brown) and pteridines (red) and their production goes through distinct pathways with different enzymes for each. Figure 3.4 outlines the different eye colour genes involved in the kynurenine, pteridine and granule pathways.

Ommochromes are produced via a pathway known as kynurenine synthesis. It involves a series of oxidations that starts with tryptophan and end with the production of the brown pigment xanthommatin (a type of ommochromes found in most insects). Formylkynurenine, kynurenine and 3-hydroxykynurenine (3-HK) are intermediates of this synthesis pathway. The Vermilion (v) gene encodes tryptophan oxygenase (Searles and Voelker, 1986) which convert tryptophan into formylkynurenine and the cinnabar (cn) gene encodes kynurenine-3-hydroxylase (Warren et al., 1996) which convert kynurenine into 3-hydroxykynurenine. Scarlet (s) and white (w) genes are localised at the membrane of the pigment granules and are involved in the transportation of 3-HK to the inside of the pigment granules (Mackenzie et al., 1999). Once in the granule, phenoxazinone synthetase converts 3-HK to xanthommatin (Summers et al., 1982).

The pathway of synthesising pteridines (or drosopterins) is more complex and less well understood than the synthesis of ommochromes. There are five types of pteridines produced in the fly, drosopterin and isodrosopterin are the major species. A series of enzymatic and non-enzymatic reactions are involved in the biosynthesis of pteridines derived from guanosine triphosphate (GTP). The Punch (p) gene encodes GTP cyclohydrolase I (GTPCH I) (McLean et al., 1993) which converts GTP to 7,8-dihydroxopterin triphosphate (H2-NTP). H2-NTP is then converted by 6-pyruvoyltetrahydropterin (6-PTP) synthase, which is encoded by purple (pr) gene, into 6-PTP (Park et al., 1990, Yim et al., 1977). The Sepia (se) gene encodes pyrimidodiazepine (PDA) synthase (Kim et al., 2006) that converts 6-PTP into PDA (a type of pteridines) which then would be transported into pigment granules (Kim et al., 2013a). In addition to the white (w) gene, the brown (bw) gene is responsible for transporting pteridine precursors into pigment granules (Borycz et al., 2008, Dreesen et al., 1988). Interestingly, the biosynthesis of tetrahydrobiopterin (BH₄), which is an essential cofactor in the synthesis of the rate-limiting enzyme for dopamine tyrosine
hydroxylase (TH) (Homma et al., 2013), is shared with pteridines synthesis and include three consecutive enzymatic reactions catalyzed by GTPCH I, PTPS, and sepiapterin reductase (Kim et al., 2013a). Heterozygous mutations in the \textit{GTPCH1} gene in humans causes an L-DOPA responsive dystonia, and is a known risk-factor in the inheritance of PD (Ichinose et al., 1994, Nalls et al., 2014).

The colour of a fly eye depends on its ability to synthesis the two pigments. For example, a bright red eye colour in \textit{v} mutants is due to their inability to synthesis ommochromes (Searles and Voelker, 1986) and a dark brown in \textit{se} mutants is due to their inability to synthesis pteridines (Kim et al., 2006). In case of \textit{white} flies, no pigments are being transported into pigment granules resulting in lack of any colour (Mackenzie et al., 1999).

Eye colour genes are also involved in vesicular trafficking and pigment granules synthesis. Pigment granules are a type of lysosome-related organelle. Ommochromes are carried into type I pigment granules, a process which occur in primary and secondary pigment cells of ommatidia while pteridines are carried into type II pigment granules which occur exclusively in secondary pigment cells (Shoup, 1966). Among many genes, \textit{Ruby} (\textit{rb}), \textit{garnet} (\textit{g}), \textit{carmine} (\textit{cm}) and \textit{orange} (\textit{or}) genes are involved in lysosome-related organelle synthesis (Grant et al., 2016). Intriguingly, \textit{dLRRK} has been suggested to have a role in trafficking cargo to the pigment granules. A genetic interaction between \textit{dLRRK} and some pigments colour genes in the kynurenine and pteridine pathways lead to synthetic lethality that might be due to \textit{dLRRK} interference (Furmston, 2016). Many stages of eye pigment synthesis occur in other, non-neuronal tissues such as the Malpighian tubule (the Drosophila equivalent of the kidney) and how different tissues and pigment movement from tissue to tissue contribute to the final eye colour are not completely known.
Figure 3.4 An illustration of the pathways involved in the biosynthesis of pigment granules

The biosynthesis of pigment granules involves the interaction of kynurenine, pteridine and granule pathways. Some pigments colour genes have been found to be synthetically lethal in dLRRK<sup>LOF</sup> flies suggesting that dLRRK might play a crucial role in this interaction. Taken from (Furmston, 2016).

3.2 Aims

1. Test if expressing dLRRK in non-neuronal tissues could rescue the loss of visual function observed in dLRRK<sup>LOF</sup> flies via ERG recording.
2. Test the effect of manipulating eye pigment genes on their interaction with dLRRK through genetic and physiological approaches.

3.3 Results

3.3.1 Investigating the ability of dLRRK to rescue the visual dysfunction in dLRRK<sup>LOF</sup> old flies when expressed in non-neural tissues

It is thought that dLRRK/hLRRK2 can move from cell to cell and act in a non-cell autonomous manner (Wang et al., 2017). We therefore wished to test if the visual dysfunction observed in dLRRK<sup>LOF</sup> flies could be rescued via expression of a dLRRK transgene in non-neuronal tissue. Prior to testing the rescue ability of dLRRK expression in non-neuronal tissues, an ERG test was used to confirm the finding from (Furmston, 2016) of reduced off-transient signal in old dLRRK<sup>LOF</sup> flies. This was carried out because the amplifier used in this investigation is different in terms of having an AC-
coupling, instead of DC, which would enhance transient signals and improve the signal to noise ratio. My data confirm the severe decline in the off-transient element of the ERG in old \textit{dLRRK}^{LOF} flies. However, I also observed a severe decline to the on-transient element of the ERG as well (Figure 3.5).

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{Figure3.5.png}
\caption{A reduction in ERG amplitude in \textit{dLRRK}^{LOF} old flies (21 days) compared to control.}
\end{figure}

Representative ERG traces from 3 days (left) and 21 days (right) of control flies (\(w^1/w^2\)) and \textit{dLRRK} loss of function mutant (\textit{dLRRK}^{e03680}). The red arrows indicate the observed reduction in the on-transient and off-transient elements of the ERG in old mutant flies compared to control. ERGs are recorded in response to 0.5 second blue light pulses. Each trace is the average of the fly’s response to five flashes of blue light. ERG potential is measured in millivolt (mV) per milliseconds (ms).

\subsection{3.3.1.1 A possible rescue of visual function in \textit{dLRRK}^{LOF} flies when \textit{dLRRK} is expressed in non-neuronal tissues}

Next, the ability of restoring the age-dependant visual dysfunction observed in \textit{dLRRK}^{LOF} old flies (21 days) was tested by expressing the native gene (\textit{dLRRK}) in non-neural tissues using the Gal4/UAS system. Three Gal4 constructs that drive expression in different tissues have been used to express the native \textit{dLRRK} in fat body (\textit{Lsp2-Gal4}; subsequently referred to as Fat body), pigment cells (\textit{P(GAL4)54C}; subsequently referred to as Pigment cell), and male reproductive system (\textit{P(GawB)c564}; subsequently referred to as Repro tract). These lines were chosen based on the expression pattern of LRRK detected in fat body by (Lazareva \textit{et al.}, 2007), pigment cells by (Nagaraj and Banerjee, 2007), and male reproductive tract by (Hrdlicka \textit{et al.}, 2002) and they were
chosen based on their distal location from the visual system (pigment cell are the closest while the reproductive tract is the farthest).

In order to express \textit{dLRRK} in these locations, a multi-step fly crosses supported by balancer chromosomes have been performed and the summary of the crosses can be found in Figure 3.6.

![Figure 3.6 Crossing summary of flies expressing \textit{dLRRK} gene in non-neural tissues using Gal4/UAS system for the assessment of visual system rescue.](image)

The left side shows the different Gal4 lines used to cross with flies (in the middle) expressing \textit{dLRRK} to produce flies expressing \textit{dLRRK} in non-neuronal tissues (the right side). * The \textit{Lsp2-Gal4} line was produced using recombination by the method described in section 2.1.5.

Examining the ERG traces in flies expressing the native gene (\textit{dLRRK}) in non-neuronal tissues in \textit{dLRRK}^{LOF} background revealed no reduction in amplitude (fat body and pigment cells rescues) or a modest reduction (Repro tract rescue) in old flies compared to the young flies (Figure 3.7A). The quantification of the components of ERG (on transient, photoreceptors, and off transient) showed a significant difference between young and old flies in Repro tract rescue flies similar to that seen in \textit{dLRRK}^{LOF} flies. This difference is not seen in fat body rescue or pigment cell rescue flies (Figure 3.7 B). The initial visual response (on-transient) observed in the tested young flies (fat body, pigment cells, and repro tract rescues) appeared to be lower than the WT which might be due to the darker eye colours in these transgenes. These data suggest the possibility
that \textit{dLRRK} has the ability to rescue the age-dependent loss of visual function in \textit{dLRRK}^{LOF} old flies when expressed in fat body or pigment cells, but this ability is limited when the rescue is attempted from the reproductive tract. On-transient and off-transient data of the ERG are reflective of the lamina neuron response which contains DA neurons. Focusing on these data revealed a possible rescue for the age-dependant visual loss based on dLRRK function in fat body and pigment cells (Figure 3.7 C).
A

3 days

21 days

On-transient response

Off-transient response

B

C

Mean (mV)

Mean (mV)
Figure 3.7 *dLRRK* expression in the fat body or pigment cells in *dLRRK*\textsuperscript{LOF} background revealed a possible rescues of the age-dependent loss of visual function.

A. Representative ERG traces from 3 days (left) and 21 days (right) of wild-type (w\textsuperscript{1}/w\textsuperscript{a}), *dLRRK*\textsuperscript{LOF} mutant and flies expressing *dLRRK* specifically in fat body, pigment cells, or reproductive tract in a *dLRRK*\textsuperscript{LOF} background. The red arrows indicate the observed reduction in the on-transient and off-transient elements of the ERG in the old mutant flies. ERGs are recorded in response to 0.5 second blue light pulses. Each trace is the mean of the fly's response to five flashes of blue light. ERG potential is measured in millivolt (mV) per milliseconds (ms). B. Quantification of the individual ERG components. The ERG of on-transient (top), photoreceptors (middle), and off-transient (bottom) responses were quantified for WT flies (w\textsuperscript{1}/w\textsuperscript{a}), *dLRRK*\textsuperscript{LOF}, and flies expressing *dLRRK* in fat body, pigment cells, or reproductive tract at 3 days (blue bars) and 21 days (red bars). C. Quantification of 21 day old flies on-transient (top) and off-transient (bottom) components of ERG. The ERG response was quantified for WT flies (w\textsuperscript{1}/w\textsuperscript{a}, orange bar), *dLRRK*\textsuperscript{LOF} (grey bar), and flies expressing *dLRRK* in fat body, pigment cells, and reproductive tract (green bars). A significant difference was found between *dLRRK*\textsuperscript{LOF} flies and flies expressing *dLRRK* in fat body and pigment cells which suggest the ability of *dLRRK* to rescue the visual dysfunction when expressed in these tissues. All statistical values are ANOVA with post-hoc Tukey: ****p<0.00001 ***p<0.0001 **p<0.001, *p<0.05, ns (no significant difference). Data presented are mean ± SD bars; n numbers are displayed inside bars. N.B. The presence of the mW\textsuperscript{+} cassette in the Gal4 and UAS transgenes increase eye colour to a deep red which might lead to lower visual response.

3.3.1.2 Does dLRRK leak to other tissues?

UAS-transgenes can sometimes be expressed in low levels or leak to other tissues in the absence of Gal4 protein. We also wished to see if Gal4 expression alone in the absence of the UAS-*dLRRK* transgene could contribute to the rescue of the *dLRRK* mutant phenotype. Therefore, an experiment was carried out to record ERGs of flies carrying only a single component of the UAS/Gal4 system in *dLRRK*\textsuperscript{LOF} background. An age-dependent loss of visual function in old flies should indicate no leak of dLRRK from UAS-transgene. The data obtained showed a reduction in ERG amplitude in old flies of the *dLRRK*\textsuperscript{LOF};UAS-*dLRRK* genotype, however, the reduction was not as significant as the one found in the *dLRRK*\textsuperscript{LOF} mutant flies alone (Figure 3.8 A). The quantification of the components of ERG (on transient, photoreceptors, and off transient), on the other hand, showed a possible leak of dLRRK when the UAS-*dLRRK* transgene is present, leading to a rescue of the off-transient response in 21 day old flies. Similarly, when the pigment cells-Gal4 construct is present there is not a statistical significance difference found in the on-transient and off-transient responses (Figure 3.8 B-C). In addition to the ERG experiment, a western blot was performed to test for secreted hLRRK2 for extracted proteins from the haemolymph in flies expressing the hLRRK2 transgene.
using fat body Gal4 driver to assess the presence of the protein of interest (anti-hLRRK2 was used due to the availability of this antibody in the lab) which might indicates a leak of protein from the haemolymph to the CNS. Unfortunately, the experiment was not successful due to the inability to find a loading control protein and possibly the ineffective technique of protein extraction (see section 2.4.1). The genetic and physiological data indicate a possible leak of dLRRK from the rescue transgene while the result found in the presence of the pigment cell-Gal4 construct suggests that eye colour may contribute to a protection of the eye function over time.
Figure 3.8 A possible leak of dLRRK is detected in pigment cells Gal4 flies when only one component of the UAS/GAL4 system is expressed in dLRRK^{LOF} flies background.
A. Representative ERG traces from 3 days (left) and 21 days (right) of \textit{dLRRK}^{LOF} mutant, flies with the UAS \textit{dLRRK} component only in a \textit{dLRRK}^{LOF} background, and flies with the GAL4 component only (Fat body and pigment cell). ERGs are recorded in response to 0.5 second blue light pulses. Each trace is the mean of the fly’s response to five flashes of blue light. ERG potential is measured in millivolt (mV) per milliseconds (ms). B. Quantification of the individual ERG components. The ERG of on-transient (top), photoreceptors (middle), and off-transient (bottom) responses were quantified for \textit{dLRRK}^{LOF} mutant, flies with the UAS \textit{dLRRK} component only in a \textit{dLRRK}^{LOF} background, and flies with the Gal4 component only (Fat body and pigment cell). C. Quantification of 21 day old flies on-transient (top) and off-transient (bottom) components of ERG. All statistical values are ANOVA with post-hoc Tukey: ****p<0.00001 ***p<0.0001, **p<0.001, *p<0.05, ns (no significant difference). Data presented are mean ± SD bars; n numbers are displayed inside bars.

3.3.2 \textit{dLRRK}^{LOF} interaction with eye pigments

Many flies bearing eye pigment mutations have been crossed with \textit{dLRRK}^{LOF} to test for the presence of synthetic lethality. This synthetic lethality indicates a crucial role of the tested genes in the viability of the fly and a possible genetic interaction between eye pigment genes and \textit{dLRRK}. To date, it has been found that loss-o-function mutations in \textit{cinnabar}, \textit{brown}, \textit{scarlet}, \textit{vermillion}, and \textit{punch} are synthetically lethal with \textit{dLRRK}^{LOF} (Cording & Furmston, unpublished). In this investigation, additional eye pigment genes have been tested to establish a better understanding of the possible interaction between \textit{dLRRK} and eye pigments gene function in flies. Genetically, additional eye pigment mutations were examined to determine their potential synthetic lethality with \textit{dLRRK}^{LOF}. We initially tested a \textit{cinnabar} hypomorph (\textit{cn}^{35k}) (Warren et al., 1996) which works in the kynurenine pathway and \textit{purple} hypomorph (\textit{pr}^{l}) (Kim et al., 1996) which works in the pteridine pathway. Physiologically, additional eye pigment genes were examined to determine their ability to rescue the age dependent visual loss in \textit{dLRRK}^{LOF} flies. These eye pigment genes include \textit{sepia} (\textit{se}), \textit{sepiapterin reductase} (\textit{sptr}) and \textit{punch} (\textit{pu}).

3.3.2.1 The combination of \textit{dLRRK}^{LOF} with \textit{pr}^{l} showed no synthetic lethality.

To test for any genetic interaction between \textit{dLRRK}^{LOF} and \textit{pr}^{l} we performed the following cross outlined in (Figure 3.9). All possible progenies were viable from the experimental cross of \textit{dLRRK}^{LOF} with \textit{pr}^{l} indicating the absence of synthetic lethality. However, the viability for the desired genotype (homozygous for \textit{pr}^{l} and \textit{dLRRK}^{LOF}) was noticeably affected obstructing further experiment on them such as ERG and
survivability suggesting an evident physiological effect caused by the combined loss of 
*dLRRK* and *pr*.

![Figure 3.9](image)

**Figure 3.9 Homozygous or hetrozygous *pr* mutation is NOT synthetically lethal with homozygous *dLRRK*LOF mutation**

An experimental cross with adult virgin male and female flies that have the genotype 
*pr*/CyO; *dLRRK*LOF/TM6B was established. The possible progenies are shown with the 
associated balancer chromosomes markers (curly wings for CyO and extra hairs on the 
thorax for TM6B). The desired genotypes are the two from the right, in particular, the 
first one from the right (homozygous for *pr* and *dLRRK*LOF). The desired genotypes 
have shown to be viable indicating an absence of synthetic lethality. The viability was 
recorded in the first generation, but the number produced was later recorded from later 
generations that had the same genotype indicated by the asterisk (*). However, overtime 
the flies started to lose CyO marker which made counting flies with curly wings not 
possible, hence, the absence number in these genotypes. NA= not available.

3.3.2.2 The combination of *dLRRK*LOF with *cn*35k showed no synthetic lethality.

To test for any genetic interaction between *dLRRK*LOF and *cn*35k we performed to 
following cross outlined in (Figure 3.10). Like *pr*, *cn*35k produced all possible progenies 
from the experimental cross with *dLRRK*LOF indicating the absence of synthetic lethality. 
However, unlike *pr*, the viability was severely affected for all genotypes including the 
desired one (homozygous for *cn*35k and *dLRRK*LOF) restricting further experiment on
them such as ERG and survivability. This also suggests a physiological effect caused by the combined loss of \( dLRRK \) and \( cn \).

\[ \text{cn}^{35k}; \ dLRRK^{LOF} \quad \text{CyO} \quad \text{TM6B} \]

\[ \text{cn}^{35k}; \ dLRRK^{LOF} \quad \text{CyO} \quad \text{TM6B} \]

\[ \sigma \text{ or } \varphi \]

\[ \sigma \text{ or } \varphi \]

\[ \sigma \text{ or } \varphi \]

\[ \sigma \text{ or } \varphi \]

\[ \text{Viable} \quad n=21 \quad \text{Viable} \quad n=14 \quad \text{Viable} \quad n=8 \quad \text{Viable} \quad n=3 \]

**Figure 3.10 Homozygous or hetrozygous \( cn^{35k} \) mutation is NOT synthetically lethal with homozygous \( dLRRK^{LOF} \) mutation**

An experimental cross with adult virgin male and female flies that have the genotype \( cn^{35k}/\text{CyO}; \ dLRRK^{LOF}/\text{TM6B} \) was established. The possible progenies are shown with the associated balancer chromosomes markers (curly wings for \( \text{CyO} \) and extra hairs on the thoracis for \( \text{TM6B} \)). The desired genotypes are the two from the right, in particular, the first one from the right (homozygous for \( \text{cn}^{35k} \) and \( dLRRK^{LOF} \)). The desired genotypes have shown to be viable indicating an absence of synthetic lethality.

3.3.2.3 The effect of overexpressing of \( se \), \( sptr \), and \( pu \) in the DA neurons on visual function

It has previously been found that overexpression of \( \text{cinnabar (cn)} \) and \( \text{brown (bw)} \) in DA neurons and glial cells can rescued the loss of visual function in \( dLRRK^{LOF} \) flies (Furmston, 2016). These rescues suggested that \( \text{cn} \) and \( \text{bw} \) work within the same pathways as \( dLRRK \), and by increasing \( \text{cn} \) or \( \text{bw} \) in a \( dLRRL^{LOF} \) background a restoration of the age-dependent loss of synaptic signaling is achieved. Here, we examined the expression of three additional pigment genes in DA neurons to look for a rescue of the \( dLRRK^{LOF} \) loss of visual function and a possible interaction with \( dLRRK \). These genes are \( \text{sepi} (se) \), \( \text{sepiapterin reductase (sptr)} \) and \( \text{punch (pu)} \), which encode enzymes
necessary for the pteridine pathway of pigment formation. A summary of experimental crosses can be found in Figure 3.11.

Figure 3.11 Crossing summary of flies over-expressing sptr, se, and pu in DA neurons in dLRRK<sup>LOF</sup> background

The left side shows the different UAS lines used to cross with flies (in the middle) carrying Gal4 specific for DA neurons to produce flies expressing sptr, se, and pu in DA neurons in dLRRK<sup>LOF</sup> background (the right side).

Examining the ERG traces of sptr, se and pu expression in combination with dLRRK<sup>LOF</sup> showed a reduction in amplitude for old flies (Figure 3.12 A) and the quantification of ERG components (on transient, photoreceptors, and off transient) confirm this reduction by showing a significant age dependant decline in all genotypes tested indicating an inability to restore the visual dysfunction to the levels of young flies except in the off-transient element for sptr expression flies which shows no significant difference from the wild type control (Figure 3.12 B). Looking into the on-transient and off-transient data in old flies revealed no statistically significant difference is observed between tested and dLRRK<sup>LOF</sup> mutant flies except in the on-transient element in sptr and se flies (Figure 3.12 C). The majority of the data from this experiment suggest that sptr, se, and pu expression were unable to rescue the age-dependent loss of visual function which indicate that they may not function within the same pathways as dLRRK.
A

3 days

21 days

B

On-transient response

C

On-transient response

Off-transient response
Figure 3.12 Over-expression of sptr, se, and pu had no effect on restoring normal ERG in old flies

A. Representative ERG traces from 3 days (left) and 21 days (right) of wild-type (w¹/w¹), dLRRK<sup>LOF</sup> mutant and flies over-expressing sptr, se, and pu in DA neurons in a dLRRK<sup>LOF</sup> background. ERGs are recorded in response to 0.5 second blue light pulses. Each trace is the mean of the fly’s response to five flashes of blue light. ERG potential is measured in millivolt (mV) per milliseconds (ms). B. Quantification of the individual ERG components. The ERG of on-transient (top), photoreceptors (middle), and off-transient (bottom) responses were quantified for WT flies (w¹/w¹), dLRRK<sup>LOF</sup>, and flies over-expressing sptr, se, and pu in DA neurons in a dLRRK<sup>LOF</sup> background. C. Quantification of 21 day old flies on-transient (top) and off-transient (bottom) components of ERG. All statistical values are ANOVA with post-hoc Tukey: ****p<0.00001 ***p<0.0001, **p<0.001, *p<0.05, ns (no significant difference). Data presented are mean ± SD bars; n numbers are displayed inside bars.

3.4 Discussion

In this investigation, we attempted to determine if the presence of dLRRK in other tissues could rescue the synaptic deficits observed in visual function of dLRR<sup>LOF</sup> mutant flies. We tested expression of dLRRK in fat bodies, reproductive tract and pigment cells for ability to rescue elements of the ERG in the dLRR<sup>LOF</sup> mutant. The expression of dLRRK in non-neuronal tissues (pigment cells or fat body) was potentially able to rescue the age dependent loss of visual function in dLRRK<sup>LOF</sup> old flies.

The observation that the visual deficits in dLRR<sup>LOF</sup> can be rescued by non-neuronal dLRRK raises the question of how the cells signal to each other - is it by cell-cell transfer of dLRRK or a downstream mechanism (e.g., TGF-β signalling)? The precise action of LRRK2 is still uncertain and has been linked to different organelles and cellular mechanisms (see section 1.2.4). A subset of late endosomal pathway extracellular vesicles known as exosomes are of particular interest because of their role in spreading misfolded proteins associated with neurodegenerative diseases. These exosomes contain materials such as DNA, RNA, or proteins, that are not normally secreted or are not stable extracellularly and can travel throughout the body and pass specialised compartments including the blood brain barrier (BBB) (Coleman and Hill, 2015). The findings in this investigation potentially support the notion of the ability of LRRK2 ability to travel throughout the body. The LRRK2 orthologue in the flies (dLRRK) when expressed outside the fly’s visual system (a neuronal tissue) was able to rescue the age dependant loss of visual function found in dLRRK<sup>LOF</sup> old flies. The rescue was seen in flies expressing the native gene in pigment cells and fat body (non-neuronal tissue). Although
the quantification of ERG elements revealed no rescue was observed when \textit{dLRRK} is expressed in male reproductive accessory organs, I would argue that it was because no gender selection was made and if the experiment were to be repeated just with male flies, I suspect a different result would come out.

Furthermore, the assertion that the severe reduction in the off-transient element of the ERG observed by (Furmston, 2016) in the \textit{dLRRK}^{LOF} old flies was proposed to be due to the dysfunction in lower order neurons of the visual system (lamina neurons) is supported by the observation of a severe reduction in both the on-transient and off-transient found in this investigation. This is because the on-transient and off-transient elements of the ERG reflect the function of the lamina neurons that lie beneath the photoreceptors.

Mutations and imbalances in the levels of metabolites in the kynurenine and pteridine pathways have been linked to neurodegenerative disorders including PD. For example, in the kynurenine pathway, the levels of kynurenine and 3-HK have been shown to be higher in the CNS of PD patients (Iwaoka et al., 2020). In addition, mutation in \textit{cinnabar} (\textit{cn}^{l}) mutant flies have shown to affect the dynamic of mitochondria (Maddison et al., 2020), which is a recurring theme in PD (Moon and Pack, 2015, Mortiboys et al., 2015, Pickrell and Youle, 2015, Schapira et al., 1990). Also, mutations in \textit{scarlet} (\textit{st}^{l}) mutant flies have revealed progressive loss of DA neurons, shortened lifespan, locomotor dysfunction and elevated levels of ROS. Furthermore, over-expression of \textit{st} in the DA neurons of a PD fly model showed a neuroprotective effect (Cunningham et al., 2018).

In the pteridine pathway, BH$_4$ is produced from GTP by the action of \textit{GTPCH I} (fly orthologue of \textit{punch}) and \textit{sepiapterin reductase} (fly orthologue of \textit{sptr}). BH$_4$ is an essential cofactor for TH which regulates the biosynthesis of dopamine (Daubner et al., 2011). A mutation in \textit{punch} or \textit{sepiapterin reductase} could have a cascading effect on BH$_4$ and therefore on dopamine levels. In fact, BH$_4$ levels have been found to be low in the CSF of PD patients and MPTP-treated monkeys (Ichinose et al., 2018). That being said, the interaction mechanism between the different eye pigment genes in these pathways with dLRRK and the cause of synthetic lethality that it produces is still unknown. Although the crosses in this investigation (\textit{cn}^{35k} and \textit{pr}^{l}) didn’t produce any lethality, the severe reduction in number and viability still indicate an interaction affecting physiology. Interestingly, the \textit{cn}^{35k} mutation is a hypomorph mutation and has probably produced some protein that affected viability rather than lethality compared to the null mutation (\textit{cn}^{l}) used in (Furmston, 2016) which led to a lethal interaction. In
addition, the overexpression of three eye pigment genes *punch, sepiapterin reductase,* and *sepia* in *dLRRK*LOF old flies were not able to rescue the loss of visual function. This inability might indicate no interaction with the *dLRRK* mutation in the visual system or these genes function upstream of dLRRK. However, in the case of *punch,* data from the Elliott lab has demonstrated that *puR1* is synthetically lethal in combination with *dLRRK*LOF (Cording and Elliott, unpublished) which might indicate an interaction is occurring but outside the visual system. Outside of the kynurenine and pteridine pathways, pigment genes can be involved in the trafficking and biosynthesis of pigment granules. They are involved with other sorting complexes proteins such as AP-3 and BLOC (Cheli et al., 2010, Lloyd et al., 1998) and RAB GTPase (Fukuda, 2021) in the movement of membrane bound vesicles carrying cargo. It is believed that dLRRK/LRRK2 might function in this pathway and any dysfunction in the protein might lead to a negative effect on the granule pigment function which may lead to cell dysfunction and subsequent death. This idea is supported by the documented interaction of LRRK2 with RAB proteins (Kuwahara and Iwatsubo, 2020) and the role of dLRRK and LRRK2 in regulating the secretory dynamic of the trans Golgi network (Bonet-Ponce and Cookson, 2021, Lin et al., 2015).

The pigment granules in the fly eye are a type of cell specific organelle known as lysosome-related organelles (LROs) generated by retinal cells. These LROs are derived from the endosomal system and their content varies based on the cell type where they originated from. In humans, LROs include melanosomes (produced by melanocytes in the skin), lamellar bodies (produced by alveolar type II cells in the lung), and alpha granules (produced by platelets cells in the blood) just to name a few (Bowman et al., 2019). PD patients have high prevalence of melanoma than the general population (Bertoni et al., 2010). Despite the association between PD and melanoma being well established, the mechanism underlying this association is still to be elucidated. For some time, it has been suggested that levodopa, the most effective treatment for PD, was the culprit (Sandyk, 1992). But recent studies such as (Dalvin et al., 2017) found that the chance of developing melanoma for PD patients is almost the same as developing PD for melanoma patients which suggested no role for levodopa. *LRRK2-G2019S* patients have high cancers risk (Agalliu et al., 2015) including an association with melanoma among others (Inzelberg et al., 2016). Moreover, DA neurons and melanocytes are both pigmented and share common pathways that include L-dopa. If LRRK2 plays a role in
regulating LROs such as melanosomes, then the increased risk of developing melanoma in LRRK2-PD patients could be attributed to LRRK2 dysfunction.

Another interesting type of LRO that might be affected in LRRK2-PD is lamellar bodies. These are produced by alveolar type II epithelial cells in the lungs which are responsible for synthesis and secretion of pulmonary surfactants that are required for normal lung function (Schmitz and Müller, 1991). It has been found that using LRRK2 inhibitors such as PFE-360 and MLi-2 induced abnormal changes to lamellar bodies in vivo or in vitro (Baptista et al., 2020, Harney et al., 2021). However, these changes were reversible and didn’t affect lung function suggesting a safe and on target efficacy for these inhibitors. Interestingly, a recent study by (Lebovitz et al., 2021) could tell a different story. Instead of using LRRK2 inhibitors, they used LRRK2 knockout mice to reduce the expression of the gene. They have found that reduction in LRRK2 promoted tumorigenesis in the lung which was associated with dysfunctional surfactant metabolism. These reports suggest an interaction between LRRK2 and lamellar bodies, however, the mechanism of that interaction is still to be elucidated. Our findings of genetic interactions between pr, cn and dLRRK\textsuperscript{LOF} further indicate and expand a potential role of dLRRK/LRRK2 in the regulation of membrane traffic and pigment synthesis in neurons.
4 The effect of dopaminergic \textit{LRRK2-G2019S} expression on eye adaptation to light

4.1 Introduction

As mentioned in Chapter 1, visual dysfunction is one of the non-motor symptoms reported by PD patients. Although this dysfunction manifests mostly in the retina, PD-related neurological dysfunction can also be observed in the visual circuits beyond the retina e.g., in the visual cortex (Weil \textit{et al.}, 2016). Investigating these visual symptoms is crucial because of their prevalence, as observed in one report by (Davidsdottir \textit{et al.}, 2005) found 78% of PD patients showed at least one visual dysfunction, suggesting the possibility of using this system as a marker for early diagnosis (Diederich \textit{et al.}, 2010).

The ability of the eyes to adjust to different light intensity depends on the feedback coming from the photoreceptors. In dark conditions, photoreceptors are very sensitive to light. This sensitivity decreases when illumination increases to avoid photoreceptors being saturated and extend their ability to detect a range of light intensity (Purves \textit{et al.}, 2001). The process of converting light into neural signal is called phototransduction which involves a series of molecular reactions occurring within photoreceptors. These reactions start with activating the G-protein coupled receptor rhodopsin and working through multiple steps eventually leading to (in mammals) the closing of the ions channels in the photoreceptors membrane which leads to neurotransmitter being released and the initiation of neural signal to the next neurons. For more details on phototransduction see review by (Lamb and Pugh, 2006). This feedback will lead to eye adaptation which include change in pupil sizes.

Flies, just like humans, must adapt to light in order to respond to the variation of light levels that come from their environment. In fact, they adapt to changes in light intensity in a circadian fashion (Nippe \textit{et al.}, 2017) where axons have been found to change in size and shape at the start of the day and the start of the night (Damulewicz \textit{et al.}, 2020, Pyza and Meinertzhagen, 1999). In addition, they adapt to light over a shorter period of time (100 ms) (Juisola and Hardie, 2001), 10-100 times faster than vertebrates (Hardie and Juusola, 2015).

We investigate abnormalities in eye adaptation to light is investigated here as a model of visual symptoms observed in PD. Similar dysfunction has been reported in PD patients where after a period of light adaptation, PD patients demonstrated a
significantly larger pupil diameters with unequal pupil sizes (anisocoria) compared to controls (Micieli et al., 1991). Moreover, irregular pupil reflexes to light (Giza et al., 2011, Stergiou et al., 2009) and supersensitivity to eye-drop test were observed as well (Yamashita et al., 2010). Similarly, in our PD mimic flies, expressing LRRK2-G2019S in dopaminergic neurons produces abnormalities in the neural response of the eye: young flies showed higher neural responses, but old flies had lost their photoreceptors responses (Afsari et al., 2014). This is a sign of failure of the eye to adapt. It was suggested that this was due to the high energy demand at an early age followed by an increased apoptosis and degradation of photoreceptors and their mitochondria (Afsari et al., 2014, West et al., 2015a). This led to the hypothesis that the overresponsive Steady State Visually Evoked Potential (SSVEP), which is an assay to evaluate visual function, in young flies expressing LRRK2-G2019S in the dopaminergic neurons might be due to a failure of adaptation.

In this investigation, eye adaptation will be tested in flies expressing LRRK2-G2019S in multiple neuronal cell types. Adaptation will be evaluated physiologically and anatomically. The physiological evaluation will utilize ERG (see section 2.3.1) to detect abnormalities in eyes neural response. In this approach, the neural response of the eye to brief flashes of dim light before and after a prolonged bright light is investigated.

Anatomically, deep pseudopupil (DPP) formation will be used to assess the integrity of the eye structure. DPP acts in a similar way to the change in pupil size in humans during light adaptation and was first observed in flies by (Franceschini and Kirschfeld, 1971). When the eye of a WT fly is illuminated from above, the DPP manifests as a dark spot where around 6 ommatidia absorb light. If the eye is illuminated from below, the DPP manifests as a bright spot where light travels along the 6 ommatidia towards the observer. The size of the DPP decreases with increased illumination. This is due to the pigment granule migration from the photoreceptor cell body towards the rhabdomeres which facilitates the reduction of the amount of light coming into the eyes to prevent harmful effects. When illumination is reduced, pigment granules migrate back to the cell (Fellgett et al., 2021, Stavenga et al., 2017).

In addition to the DPP which looks at the entire eye, pseudopupil analysis (PPA) is also used to look at the retinal anatomy in more detail. This is This technique has been used by different research groups (Campesan et al., 2011, Song et al., 2013) to examine individual ommatidia and their 7 visible rhabdomeres to assess any damage to them.
4.2 Aim

The aim of this chapter is to test if expressing \textit{LRRK2-G2019S} in different parts of the fly’s eye (photoreceptors, DA neurons, or pigment cells) has an effect on eye adaptation to light and this will be done

1. Physiologically using ERG to detect any abnormalities in eye neural response and
2. Anatomically using DPP and PPA to detect any structural abnormalities in the eye.

4.3 Results

4.3.1 Investigating the effect of expressing \textit{LRRK2-G2019S} in components of the visual circuit on eye adaptation using ERG

The effect of expressing \textit{LRRK2-G2019S} in sub-structures of the visual circuit on eye adaptation to light is assayed physiologically using ERG. Flies will be subjected to flashes of dim light in the dark before and after a period of prolonged bright light. The mean ERG amplitude before (referred to as dark adapted) and after (referred to as light adapted) will be quantified and compared to look for any differences that might suggest dysfunction of eye adaptation to light. As shown in Figure 4.1. The fly was acclimatised to the rapidly flashing blue light for 4 minutes and then exposed to bright white light for 6 minutes, before the rapid blue flashes were resumed. In some cases, the DC offset of the trace had to be adjusted to keep the recording on the screen – as shown by the red arrow.

In addition to \textit{LRRK2-G2019S} (a hyperactive kinase) expression, \textit{LRRK2-WT} and \textit{LRRK2-G2019S-K1906M} (kinase dead, KD) were tested as controls. Ideally, \textit{WT} flies (\textit{CS} or \textit{w}) and flies with only one component of the Gal4/UAS should have been tested as well. The measurement for these flies is at young (4 hours and 3 days) and old ages (21 days) and these time points were selected based on the work of (Afsari \textit{et al.}, 2014, Hindle \textit{et al.}, 2013) which showed abnormal ERG for \textit{TH>G2019S} flies at these ages.
Figure 4.1 An example of a recorded ERG for eye adaptation to light test

Rapid blue flashes of 100ms were given at 5 seconds intervals and the ERG recorded as in section 2.3.1. The peak-peak amplitude of the ERG was measured before and after the bright light (green bars). To keep the trace within range, the DC offset was adjusted as required, as in this example (red arrow).

4.3.1.1 The effect of expressing LRRK2-G2019S in the photoreceptors

The first tissue to be tested for the effect of LRRK-G2019S expression on visual responses was the photoreceptor. To facilitate this expression, the Gal4 driver to be utilised is rh1-Gal4, which drives expression in the first layer of the fly retina, the photoreceptors. This transgene is specific for the R1-6 photoreceptors. Across all the different ages and genotypes of tested flies, no change in ERG amplitude has been observed in the ERG traces (Figure 4.2 A) and their quantifications (Figure 4.2 B).
Figure 4.2 No changes in ERG amplitude is observed in flies expressing *LRRK2-G2019S* in the photoreceptors

A. Representatives ERG traces with their dark (in black) and light (in grey) adapted means next to them of flies expressing *LRRK2-WT, LRRK2-G2019S,* and *LRRK2-G2019S-K1906M* respectively in photoreceptors. The blue, red, and green boxes indicate the representative ages when the ERG was recorded which is 4 hours, 3 days,
and 21 days respectively. B. The quantification of ERG amplitude of 8-10 flies from each genotype in 4 hours, 3 days, and 21 days respectively. All statistical values are ANOVA with post-hoc Tukey: ****p<0.00001 ***p<0.0001, **p<0.001, *p<0.05, ns (no significant difference). Data presented are mean ± SD bars; n numbers are displayed inside bars.

4.3.1.2 The effect of expressing LRRK2-G2019S in DA neurons

Since no changes were observed in visual responses when LRRK2-G2019S was expressed in the photoreceptors, TH-Gal4 was next used to drive the expression of LRRK2-G2019S in DA neurons. These DA neurons are found in the CNS and they can also be found in the lamina cortex which is the second layer of neurons found beneath the photoreceptors. In addition, the expression of LRRK2-G2019S in DA neurons has already been found to have an effect on fly visual function in aged flies (Hindle et al., 2013). In this investigation, the expression of LRRK2-WT, LRRK2-G2019S, or LRRK2-G2019S-K1906M (KD) in DA neurons induced a significant difference in ERG amplitude before and after exposure to bright light in young flies (4 hours). The dark adapted flies had much larger ERGs than the light-adapted flies. By day 3, this difference was only seen in flies expressing LRRK2-G2019S. This is likely because in the LRRK2-G2019S expressing flies the dark adapted response was kept at almost the same level as the 4 hours one. By 21 days, the difference in ERG amplitude has disappeared but a decline in the overall amplitude for all genotypes is observed (Figure 4.3). This is interesting because a failure in light adaption was expected to be at an old age rather than at young one.
Figure 4.3 A failure in adaptation is observed in ERG amplitude in young flies expressing G2019S in DA neurons indicated by the decline in ERG amplitude in light adapted condition

A. Representatives ERG traces with their dark (in black) and light (in grey) adapted means next to them of flies expressing LRRK2-WT, LRRK2-G2019S, and LRRK2-G2019S-K1906M respectively in DA neurons. The blue, red, and green boxes indicate
the representative ages when the ERG was recorded which is 4 hours, 3 days, and 21 days respectively. B. The quantification of ERG amplitude of 8-10 flies from each genotype in 4 hours, 3 days, and 21 days respectively. All statistical values are ANOVA with post-hoc Tukey: ****p<0.00001 ***p<0.001, **p<0.01, *p<0.05, ns (no significant difference). Data presented are mean ± SD bars; n numbers are displayed inside bars.

4.3.1.3 The effect on visual responses of expressing LRRK2-G2019S in pigment cells

The process of eye adaptation to light involves changes in phototransduction and migration of pigment granules toward or away from rhabdomere. Expressing LRRK2-G2019S in photoreceptors and DA neurons tested the changes in phototransduction effect on adaptation while expressing the protein it in the pigment cells would test the pigment migration effect on adaptation. Expressing LRRK2-G2019S in pigment cells (P(Gal4)56C) in the 4 hour old flies demonstrated a decline in ERG amplitude from the dark adapted state to the light adapted state, however; only the LRRK2-G2019S expressing flies show statistical significance (Figure 4.4). In the 3 day old flies, the decline is again only seen in the LRRK2-G2019S expressing flies. Lastly, in the 21 day old, there is no effect, though the data suggests a larger sample might reveal a difference.
Figure 4.4 A failure in adaptation is observed in young flies expressing \textit{LRRK2-G2019S} in pigment cells indicated by the decline in ERG amplitude in light adapted condition.

A. Representatives ERG traces with their dark (in black) and light (in grey) adapted means next to them of flies expressing \textit{LRRK2-WT}, \textit{LRRK2-G2019S}, and \textit{LRRK2-G2019S-K1906M} respectively in pigment cells. The blue, red, and green boxes indicate
the representative ages when the ERG was recorded which is 4 hours, 3 days, and 21 days respectively. B. The quantification of ERG amplitude of 8-10 flies from each genotype in 4 hours, 3 days, and 21 days respectively. All statistical values are ANOVA with post-hoc Tukey: ****p<0.00001 ***p<0.0001, **p<0.001, *p<0.05, ns (no significant difference). Data presented are mean ± SD bars; n numbers are displayed inside bars.

Focusing on the data from the young flies, particularly the 3 days old data, an ANOVA test for the different genotypes showed that the expressing of LRRK2-G2019S in DA neurons and pigment cells seems to affect significantly flies ability to adapt to light (Figure 4.5).

**Figure 4.5** A significant difference in ERG amplitude is observed when LRRK2-G2019S is expressed in DA neurons and pigment cells in 3 days old flies

The quantification of ERG amplitude from 8-10 flies of LRRK2-WT, LRRK2-G2019S, and LRRK2-G2019S-K1906M in photoreceptors, DA neurons, and pigment cells respectively. All statistical values are one way ANOVA: ****p<0.00001 ***p<0.0001, **p<0.001, *p<0.05, ns (no significant difference). Data presented are mean ± SD bars; n numbers are displayed inside bars.
4.3.2 Investigating the effect of expressing \textit{LRRK2-G2019S} on eye adaptation using Deep Pseudo-Pupil (DPP) analysis

In addition to investigating the effect of expressing \textit{LRRK2-G2019S} expression on eye adaptation physiologically using ERG, DPP was utilised to examine any effect on the eye anatomy that may be development or degenerative. Flies are again subjected to a bright light before and after a period of prolonged darkness. The mean sizes of DPP of the before period (referred to as light adapted) and the after period (referred to as dark adapted) will be quantified and compared to look for any differences that might suggest dysfunction of eye adaptation to light. The same ages and genotypes from the ERG experimental analysis will be applied here as will.

4.3.2.1 The effect of expressing \textit{LRRK2-G2019S} in photoreceptors on DPP development

We analysed DPP size and presence in flies expressing \textit{LRRK2-G2019S} in photoreceptors. Overall, the mean size of the DPP in light adapted was $0.33 \text{ mm}^2 \pm 0.08$ SD and $0.32 \text{ mm}^2 \pm 0.07$ SD in dark adapted which corresponds to the area of 8-12 ommatidia. Assessing the formation of DPP in flies expressing \textit{LRRK2-WT}, \textit{LRRK2-G2019S}, and \textit{LRRK2-G2019S-K1906M} in photoreceptors using \textit{rh1-Gal4} revealed no structural differences observed between light adapted and dark adapted states of any age or genotypes. Furthermore, the quantification of DPP sizes confirm this observation where no significant statistical differences were found within the same genotype. Interestingly, however, an analysis of variance performed showed a significant difference among the different genotypes across all ages (Figure 4.6). At 4 hours and 21 days, \textit{LRRK2-G2019S} expressing flies had an increased DPP of approximately 30% - 50% from either \textit{LRRK2-WT} or \textit{LRRK2-G2019S-K1906M} flies. At 3 days, DPP in \textit{LRRK2-G2019S} expressing flies was around 35% more than \textit{LRRK2-G2019S-K1906M} flies. Approaching this result was the observation that DPP in \textit{LRRK2-WT} expressing flies was nearly the same size as the effect in \textit{LRRK2-G2019S} expressing flies.
Figure 4.6 A larger DPP is formed in flies expressing LRRK2-G2019S in photoreceptors

A. Representative images of DPP formation in flies expressing LRRK2-WT, LRRK2-G2019S, and LRRK2-G2019S-K1906M in photoreceptors. B. The quantification of the size of DPP for 8-10 flies from each genotype in 4 hours, 3 days, and 21 days respectively. Abbreviations WT (LRRK2 wild type), GS (G2019S), and KD (kinase dead). The white dotted line in light adapted WT flies indicate a sample of the DPP area of measurement. Scale bar 1 mm. All statistical values are ANOVA with post-hoc Tukey: ****p<0.00001  ***p<0.0001, **p<0.001, *p<0.05, ns (no significant difference). Data presented are mean ± SD bars; n numbers are displayed inside bars.

To add an extra layer in investigating the effect of expressing LRRK2-G2019S in subcomponents of the fly eye on DPP, PPA was utilised to visualise the rhabdomic
structure and arrangement in the ommatidium. Unfortunately, the analysis for these flies (rh1-Gal4) was unsuccessful in acquiring any clear images. This failure, most likely, was not due to abnormalities in the structure of the ommatidia but rather to the unoptimized settings of the microscope used to perform the task and the dark red nature of these flies’ eyes because a WT fly with red eye colour showed similar results.

4.3.2.2 The effect on DPP of expressing \textit{LRRK2-G2019S} in DA neurons

The assessment of DPP formation in flies expressing \textit{LRRK2-G2019S} in DA neurons (\textit{TH}-Gal4) was unsuccessful. The very pale-pigmented nature of their eyes made it very difficult to tell whether they are forming DPP or not, especially in young flies. Figure 4.7 demonstrates the best samples obtained from these flies where older flies seem to have some DPP formed as their eye pigments intensify with age. Moreover, it is possible that the camera used in this experiment was not sensitive enough to capture DPP in these pale-pigmented flies. Therefore, the quantification of DPP was not performed.

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure4.7.png}
\caption{Inconclusive results for DPP formation in flies expressing \textit{LRRK2-G2019S} in DA neurons}
\end{figure}

On the other hand, PPA of the TH-Gal4 was more successful than for the rh1-Gal4 flies, because the eyes were orange rather than dark red. The PPA of all three crosses showed a regular structure and clustering. There were no abnormalities in rhabdomere numbers (in each case 7 cells were counted) and the hexagonal pattern was preserved (Figure 4.8) indicating no anatomical damage brought by expressing \textit{LRRK2-G2019S} in DA neurons.

**Figure 4.8 No degeneration of ommatidia observed in flies expressing \textit{LRRK2-G2019S} in DA neurons**

Representative images of pseudopupil analysis in flies expressing \textit{LRRK2-WT}, \textit{LRRK2-G2019S}, and \textit{LRRK2-G2019S-K1906M} in DA neurons. A 25-30 ommatidia were examined per fly. Scale bar 50 µm.

4.3.2.3 The effect on DPP of expressing \textit{LRRK2-G2019S} in pigment cells

Similar to expressing \textit{LRRK2-G2019S} in photoreceptors, expressing \textit{LRRK2-G2019S} in pigment cell didn’t reveal any structural differences between light adapted and dark adapted states for the DPP formation for any age or genotype. However, the
quantification of the size of the DPP formed revealed a significant difference among the different genotypes in mostly the young flies (4 hours and 3 days) compared to all ages in the photoreceptors of expressing flies. In addition, the size of DPP in the 4 hour old flies expressing LRRK2-G2019S was around 40% and 70% more than the WT and KD respectively. At 3 days, DPP in flies expressing LRRK2-G2019S was around 20% and 60% more than the wild type and kinase dead respectively. This increase was almost lost at 21 days (Figure 4.9).

Figure 4.9 A larger DPP formed in young flies expressing LRRK2-G2019S in pigment cells

A. Representative images of DPP formation in flies expressing LRRK2-WT, LRRK2-G2019S, and LRRK2-G2019S-K1906M in pigment cells. B. The quantification of the size of DPP for 8-10 flies from each genotype in 4 hours, 3 days, and 21 days
respectively. Abbreviations WT (LRRK2 wild type), GS (G2019S), and KD (kinase dead). Scale bar 1 mm. All statistical values are ANOVA with post-hoc Tukey: ****p<0.00001 ***p<0.0001, **p<0.001, *p<0.05, ns (no significant difference). Data presented are mean ± SD bars; n numbers are displayed inside bars.

In regard to PPA of the for flies expressing LRRK2-G2019S in pigment cells, the data showed similar results to flies expressing LRRK2 variants in the the DA neurons flies i.e., no abnormalities in rhabdomeres numbers and structure have been observed (Figure 4.10).

![Figure 4.10 No degeneration of ommatidia observed in flies expressing LRRK2-G2019S in pigment cells](image)

Representative images of pseudopupil analysis in flies expressing LRRK2-WT, LRRK2-G2019S, and LRRK2-G2019S-K1906M in pigment cells. A 25-30 ommatidia were examined per fly. Scale bar 50 μm.

4.4 Discussion

In this investigation, adaptation was assessed physiologically by comparing the amplitude of the ERG before and after a period of intense bright light. The key observation is that young flies (3 day old) expressing LRRK2-G2019S in DA neurons or
pigment cells fail to adapt to the dark, as indicated by their relatively larger ERGs before exposure to bright light. This was partially complemented by anatomical assessment where the DPP in flies with $LRRK2\text{-}G2019S$ expressed in pigment cells was larger than the DPP of flies expressing $WT$ or $KD$ forms of $LRRK2$ (Figure 4.7). This raises the question of the mechanism(s) behind this observation. First, as $LRRK2\text{-}G2019S$ expression in fly DA neurons reduces dopamine release (Cording et al., 2017) and this will affect the shape of the photoreceptor response in the ERG as flies treated with octopamine or dopamine had reduced photoreceptors signals (Chyb et al., 1999), this may lead to aberrant ERGs. Secondly, I have shown earlier in chapter 3 that $dLRRK$ (the *Drosophila* orthologue of $LRRK2$) mutants genetically interacts with mutations in genes regulating the formation of the pigment granules, therefore $LRRK2\text{-}G2019S$ expression might result in disruption to the ERG and DPP through regulation of pigment granule formation and function. Both dopamine function and pigment granule migration affect the detection of light on the photoreceptor, so that, in either case $LRRK2\text{-}G2019S$ expression leads to hyperactivity. The aberrant, oversize ERGs would indicate an excessive ionic flux, and resulting demand for ATP during each light flash. Synthesis of the extra ATP will potentially incur mitochondrial damage and start a degenerative cascade (Hindle et al., 2013). It is reassuring that the $LRRK2\text{-}G2019S$ expression in DA neurons and pigment cells have similar effects - as both of these processes control the adaptation of the eye, though probably by different mechanisms. Here, failure to adapt to light was observed in young flies (3 days) expressing $LRRK2\text{-}G2019S$ in DA neurons. This observed physiological abnormality might be due to the degeneration of DA neurons which was reported in humans (Nguyen-Legros, 1988) and animal models (Marrocco et al., 2020, Meng et al., 2012). The larger neural response seen in young flies is similar to those observed by (Afsari et al., 2014) which support the hypothesis of increased ATP demand in young flies because the neural response in old flies was unaffected but smaller than the young ones. The DPP-related anatomical assessment for these flies was unattainable due to low pigment content in their eyes. Thus, crossing them out with flies that have dark pigment might be a reasonable solution. Furthermore, using a different method for assessing the development of DPP than the one used in this project i.e., measuring the size of the DPP, might help obtaining quantifiable results. Examples of these methods include measuring light temperature from light scattering and absorption through DPP (Lo and Pak, 1981), counting the number of glowing ommatidia in the DPP (Fellgett et al., 2021), or using slow motion camera to measure the time taken to form DPP (Katz and Minke, 2009).
In contrast, expressing \textit{LRRK2-G2019S} in fly photoreceptors has no observed physiological effects on their ability to adapt to light. One possible explanation for this is that photoreceptors in flies are histaminergic neurons (Hardie, 1987) rather than dopaminergic which are the main type of neurons affected in PD. Anatomically, no structural damage was observed in the eye of the \textit{rh1>G2019S} flies indicated by the formation of DPP but the DPP was larger than the controls in which other forms of \textit{LRRK2} were expressed (Figure 4.6). This observation is in contrast with multiple studies that demonstrated signs of eye degeneration in the presence of Parkinson’s related manipulations (Feany and Bender, 2000, Hindle \textit{et al}., 2013, Liu \textit{et al}., 2008, Venderova \textit{et al}., 2009). However, it is crucial to note that these reports were in flies expressing PD-related genes (\textit{α-synuclein}, \textit{parkin} and several \textit{LRRK2} transgenes) in the eye using the \textit{GMR-Gal4}, which expresses in all retinal cells, rather than just in photoreceptors. Moreover, the degree of pigmentation in these \textit{rh1>G2019S} flies might have affected their adaptation to light as (Miller \textit{et al}., 1981) has reported different sensitivity to light between red-eyed and white-eyed flies.

On a longer timescale, disruption of adaptation in the eye may also affect other physiological processes. One of these is sleep dysfunction, which affects up to 60% of PD patients (De Cock \textit{et al}., 2008) and the aetiology of this dysfunction is still under investigation. Sleep-wake cycle is controlled by the circadian system which is regulated by endogenous processes that respond to environmental cues and the most important one is light (Ashbrook \textit{et al}., 2020). Disturbance in light exposure has been associated with alteration in sleep-cycle such as increased evening fatigue and low sleep quality (Figueiro \textit{et al}., 2017, Viola \textit{et al}., 2008). Abnormalities to circadian rhythm in PD have been reported by multiple studies (Ahsan Ejaz \textit{et al}., 2006, Placidi \textit{et al}., 2008, Videnovic and Golombek, 2013). That being said, eye adaptation to light works in a circadian fashion (Damulewicz \textit{et al}., 2020) which raise the question whether an impairment in light adaptation, as seen in this investigation, would have an effect on the circadian system and in turn affect the sleep-cycle and give us further insight into the pathophysiology of sleep disorders observed in PD.
5 Stability of LRRK2 Mutants Proteins \textit{in vitro} and \textit{in vivo}

5.1 Introduction

As set in detail in Chapter 1, Parkinson’s disease (PD) is the second most common neurodegenerative disorder caused by the loss of dopaminergic (DA) neurons. Mutations in the \textit{LRRK2} gene are genetic factors that have been associated with the autosomal dominant form of the disease. Many allelic variants associated with PD have been found throughout the gene; some are pathogenic (e.g., \textit{LRRK2-R1441C/H}, \textit{LRRK2-G2019S} and \textit{LRRK2-I2020T}), others are risk factors (\textit{LRRK2-G2385R}) (Kumari and Tan, 2009) and some may be neuroprotective (\textit{LRRK2-N551K} and \textit{LRRK2-R1398H}) (Gopalai \textit{et al.}, 2019). The pathogenic mutations mostly affect the kinase or GTPase functions of LRRK2. Although there is little consensus about which other proteins are phosphorylated by LRRK2, novel therapies are being developed based on inhibition of LRRK2 kinase. For example, DNL-151 and DNL-201 are kinase inhibitors developed by Denali Therapeutics have shown promising results in clinical trials (GlobeNewswire).

In this context, knowing the stability and half-life of LRRK2 is crucial to help understand the functional window over which the protein functions. This is particularly important as (in cell culture) the breakdown of LRRK2 is affected by both mutations and kinase inhibitors (De Wit \textit{et al.}, 2018). Steady state measurement of protein amount using western blot from cell transfection experiments indicate less \textit{LRRK2-R1441C} and \textit{LRRK2-G2385R} than \textit{LRRK2-WT} protein, interpreted as an increase in the rate of degradation (Greene \textit{et al.}, 2014, Rudenko \textit{et al.}, 2017). In vivo, namely in \textit{Drosophila}, less \textit{LRRK2-R1441C} protein was also detected (Cording \textit{et al.}, 2017). However, this reduction in protein could also be due to the differences in protein synthesis. Effects of LRRK2 expression on overall cellular protein synthesis have been demonstrated by (Martin \textit{et al.}, 2014a). To determine if there are differences in LRRK2 stability, a pulse chase protocol is more appropriate. In a pulse chase experiment, the synthesis of LRRK2 can be controlled by a drug which is then washed away, and the decay of LRRK2 is followed. Only one mutant of LRRK2, \textit{LRRK2-G2385R} has been tested in a pulse-chase experiment. This suggested that the mutant protein decayed faster than the wild type (Rudenko \textit{et al.}, 2017). The availability of cell lines in which the \textit{LRRK2-WT} and mutant proteins can be induced by the addition of a drug facilitates this approach.
Additionally, it is important to test if results from cell culture are reflected in vivo. The availability of inducible Gal4 constructs and Geneswitch technology (Osterwalder et al., 2001) permits the design of pulse-chase experiments in *Drosophila*. Here the inducible GAL4 is used to drive expression of *hLRRK2-WT* and of mutant forms.

In line with the suggestion that LRRK2 autophosphorylation controls the rate of breakdown, kinase inhibitors also affect the level of protein recorded in steady state assays (Lobbestael et al., 2016). However, it is sometimes difficult to mechanistically reconcile the effects of the pathogenic mutation and kinase inhibitors due to the different level of phosphorylation observed at different sites of the protein (De Wit et al., 2018), and it is not known if these cell culture overexpression assay data conducted in heterologous cells are typical for neurons in vivo.

Since observing that toxic effects of increased kinase activity in *LRRK2* mutants causes toxic effects, inhibiting this activity has been a therapeutic target. Many kinase inhibitors have been tested to look for a potent and selective LRRK2 kinase inhibitor (Taymans and Greggio, 2016). Cis-2,6-dimethyl-4-(6-(5-(1-methylcyclopropoxy)-1H-indazol-3-yl) pyrimidin-4-yl) morpholine (MLi-2) is a highly potent and selective kinase inhibitor with CNS activity and well tolerated in PD mice (Fell et al., 2015). These characteristics make it an attractive candidate for further investigation. However, as the case in many kinase inhibitors, MLi-2 works in a non-allosteric manner i.e., it is an ATP competitive compound. Interestingly, in a screening of 2080 FDA-approved compounds for their ability to reduce LRRK2 activity, an enzymatically active form of vitamin B12 (5′-deoxy-adenosylcobalamin or AdoCbl) was found to modulate LRRK2-G2019S kinase activity in an allosteric fashion (Schaffner et al., 2019). This is interesting because vitamin B12 has been associated with PD (Shen, 2015) and can be found naturally in foodstuffs based on animal products.

In this investigation, LRRK2 stability will be tested in a pulse-chase approach. HEK293 Flp-In T-REx cells (for more information see section 2.2.1) will be used to examine in vitro effects of LRRK2 mutations on protein stability. These cells have already been transfected with different *LRRK2* mutations (*LRRK2-G2019S*, *LRRK2-R1441C*, and *LRRK2-D2017A*) or the *LRRK2-WT* gene. These cells have the advantage that the *LRRK2*-construct is always landed at the same site, minimizing discrepancies between the mutant lines. Cells will be induced to express *LRRK2* by the addition of doxycycline (Doxy). In vivo, the Gal4 Geneswitch/UAS technique (for more information see section
1.4.1.1) will be used in *Drosophila melanogaster* to express different human LRRK2 mutations (*LRRK2-G2019S*, *LRRK2-G2019S-K1906M*, *LRRK2-I2020T*, *LRRK2-R1441C*, *LRRK2-I1122V*, and *LRRK2-G2385R*) or the *LRRK2-WT* pan-neuronally (*elav*<sup>GS</sup>-Gal4). These genes will be induced by the addition of Mifepristone (RU486) in the food. Unfortunately for these flies, the LRRK2 transgene landing site is not the same for each mutant which might introduce some observed differences.

Moreover, the effect of kinase inhibition on the stability of LRRK2 mutants will be tested. The active form of vitamin B<sub>12</sub> (5’-deoxy-adenosylcobalamin) will be utilised as an allosteric compound (the term vitamin B<sub>12</sub> will be used subsequently) and MLi-2 will be utilised as a non-allosteric compound. The stability of the protein will be examined using western blot (WB) and Image Studio Lite software will be used to quantify protein density.

### 5.2 Aims

1. To test the effect of multiple LRRK2 mutations on the stability of the protein, using a pulse-chase approach.
2. To test the effect of kinase inhibition on LRRK2 stability using vitamin B<sub>12</sub> (allosteric inhibition) and MLi-2 (non-allosteric inhibition).

### 5.3 Results

We attempted to determine the half-life of LRRK2 and PD associated variants in mammalian cell culture and in *Drosophila* neurons. In our cell culture experiments, we used the Human Embryonic Kidney 293 (HEK293) cell line containing the Flp-In T-REx system to express LRRK2 and PD associated variants in a Doxycyclin inducible manner (see Figure 2.3). In *Drosophila*, we used inducible Gal4 constructs and Geneswitch technology (see Figure 1.9). The timeline of the experiment (Figure 5.1) includes the induction of cells or flies with doxycyclin or RU486 respectively and quantifying the amount of protein using WB for 6 days. However, for half-life determination, the starting point (designated as day 0) of comparison for LRRK2 stability between the different mutations and treatments (untreated, B<sub>12</sub>, or MLi-2) was set to be 1 day post media change and removal of inducing agents. Moreover, cells and flies which had not been exposed to the inducing agents were tested in WB for any possible leakage from the expression construct in the absence of the inducer.
The inducing agents (Doxy for cells or RU486 for flies) were added for 2 days to induce the expression of LRRK2. At the end of day 2, media or food was changed and B12 or MLi-2 were added. The next day and for 4 days, proteins were quantified using WB for each treatment. The numbers within the red rectangle represents the set days of quantification and comparison between the different treatment.

5.3.1 In vitro evaluation of LRRK2 stability

5.3.1.1 No evident leak of expression was detected in the tested cell lines

Immunoblotting for LRRK2 abundance in Flp-In T-REx HEK293 uninduced cells revealed no leak of expression which is demonstrated by the absence of the LRRK2 band (Figure 5.2). The LRRK2 mutations to be evaluated in cells here were LRRK2-WT, LRRK2-G2019S, LRRK2-D2017A (kinase dead), and LRRK2-R1441C.

Figure 5.2 In the absence of inducing agent, no LRRK2 protein is expressed by the FLP-In T-REx system

Representative blots for LRRK2-WT, LRRK2-G2019S, LRRK2-D2017A (kinase dead), and LRRK2-R1441C in Flp-In T-REx HEK293 cells when induced or not with Doxy (1 µg/ml). β-tubulin/GAPDH lane represents the housekeeping loading protein. MW = Molecular Weight
5.3.1.2 Stability of LRRK2 protein in vitro

Examination of pulse-chased LRRK2 protein variants in cells revealed a shorter half-life ($t_{1/2}$) for LRRK2-WT than LRRK2-G2019S and LRRK2-D2017A (kinase dead) ($t_{1/2}$-WT of 2.257 days, 95% CI of 1.714 to 3.181 days vs $t_{1/2}$-G2019S of 3.377 days, 95% CI of 2.089 to 7.884 days and $t_{1/2}$-D2017A of 3.335 days, 95% CI of 2.007 to 8.506 days) while LRRK2-R1441C appeared to have the shortest half-life ($t_{1/2}$-R1441C of 0.9 days, 95% CI of 0.5734 to 1.438 days). Interestingly, it appeared that LRRK2-R1441C is the only mutation to have lost most of the protein by the last day of measurement while the rest are still maintaining some protein (Figure 5.3). The loading control changed from β-tubulin to GAPDH between experiments due to antibody availability. However since protein amounts of GAPDH or tubulin do not change throughout each experiment the results are reasonably internally controlled.
Figure 5.3 LRRK-R1441C appears to be potentially the least stable PD associated LRRK2 protein variant when expressed in cell culture

A. Representative blots and B. the time courses of the decay of LRRK2-WT, LRRK2-G2019S, LRRK2-D2017A (kinase dead), and LRRK2-R1441C in Flp-In T-REx HEK293 cells when induced with Doxy. Error bars indicate SEM with N = 3.
5.3.1.3 Kinase inhibitors effects on protein stability in *vitro*

After determining relative protein for LRRK2 and PD associated variants in untreated conditions, cells were treated with kinase inhibitors (vitamin B_{12} and MLI-2) to investigate their effects on LRRK2 protein stability. Inhibitors were added after the removal of inducing agents. When vitamin B_{12} was added to cells (1.9 \mu g/ml), the stability of LRRK2-WT (t_{1/2-WT+B_{12}} of 2.194 days, 90% CI of 1.299 to 5.168 days vs t_{1/2-WT} of 2.257 days), LRRK2-G2019S (t_{1/2-G2019S+B_{12}} of 3.66 days, 90% CI of 1.923 to 19.86 days vs t_{1/2-G2019S} of 3.377 days), and LRRK2-R1441C (t_{1/2-R1441C+B_{12}} of 0.7179 days, 90% CI of 0.586 to 0.8736 days vs t_{1/2-R1441C} of 0.9 days) were comparable to untreated cells. Interestingly, vitamin B_{12} appear to statistically reduce the stability of LRRK2-D2017A (kinase dead) by almost 2 days (t_{1/2-D2017A+B_{12}} of 1.398 days, 90% CI of 0.7684 to 3.217 vs t_{1/2-D2017} of 3.335 days), however, the protein doesn’t appear to decay totally with time but there is less abundance than the untreated cells (Figure 5.4).
Figure 5.4 Vitamin B$_{12}$ appears to affect LRRK2-D2017A protein stability

A. Representative blots and B. the time courses of the decay of LRRK2-WT, LRRK2-G2019S, LRRK2-D2017A (kinase dead), and LRRK2-R1441C in Flp-In T-REx HEK293 cells when treated with or without vitamin B$_{12}$. Error bars indicate SEM with N = 3.
When the LRRK2 inhibitor MLi-2 is added to cells (1.3ng/ml), the stability of LRRK2-WT ($t_{1/2}$-WT+MLi-2 of 2.149 days, 95% Cl of 1.166 to 6.777 days vs $t_{1/2}$-WT of 2.257 days), LRRK2-D2017A ($t_{1/2}$-D2019A+MLi-2 of 3.676 days, 95% Cl of 1.736 to 325.2 days vs $t_{1/2}$-D2019A of 3.335 days), and LRRK2-R1441C ($t_{1/2}$-R1441C+MLi-2 of 1.102 days, 95% Cl of 0.688 to 1.895 days vs $t_{1/2}$-R1441C of 0.9 days) were comparable to untreated cells. The case for LRRK2-G2019S is similar to that of LRRK2-D2017A and vitamin B12 where MLi-2 appear to statistically reduce the stability of the protein by almost a day ($t_{1/2}$-G2019S+MLi-2 of 2.565 days, 95% Cl of 1.6 to 5.504 vs $t_{1/2}$-G2019S of 3.377 days), however, the amount of protein appeared in blot to be very similar to the untreated cells and the effect is subtle. (Figure 5.5).
Figure 5.5 MLi-2 appears to have no effect on LRRK2 stability

A. Representative blots and B. the time courses of the decay of LRRK2-WT, LRRK2-G2019S, LRRK2-D2017A (kinase dead), and LRRK2-R1441C in Flp-In T-REx HEK293 cells when treated with or without vitamin MLi-2. Error bars indicate SEM with N = 3.
5.3.2 *In vivo* evaluation of LRRK2 stability

5.3.2.1 Testing for leak of expression of LRRK2 in uninduced transgenic flies

To test for LRRK2 and PD associated LRRK2 variant stability in an *in vivo* system we employed the *Drosophila* Geneswitch system where the Gal4 expression of LRRK2 and PD associated variants are induced by the addition of RU486 to the diet of the experimental flies. We initially tested for protein leakage expression in the absence of inducing agent we tested animals for LRRK2 expression in the absence of RU486. WB for transgenic uninduced flies revealed no leak of expression which is demonstrated by the absence of the LRRK2 band except for the *LRRK2-I2020T* mutation (Figure 5.6). Therefore, the available mutations in flies that can be evaluated here were *LRRK2-WT*, *LRRK2-G2019S*, *LRRK2-G2019S-K1906M* (kinase dead), *LRRK2-R1441C*, *LRRK2-I1122V*, and *LRRK2-G2385R*.

![Representative blots for LRRK2 and Synaptotagmin](image)

**Figure 5.6** No *LRRK2* expression from transgenes is observed in uninduced animals indicating absence of expression except for the *LRRK2-I2020T* transgene.

5.3.2.2 Stability of LRRK2 protein in vivo

We carried out a full pulse-chase analysis for the LRRK2 protein and the LRRK2 PD associated variants to test for LRRK2 protein stability in in vivo flies. WB analysis of pulse chase expressed LRRK2 and PD-associated variants revealed comparable stability level for LRRK2-WT, LRRK2-G2019S-K1906M (kinase dead), LRRK2-G2019S, and LRRK2-I1122V (t1/2-WT of 0.905 days, 95% Cl of 0.569 to 1.431 vs t1/2-G2019S-K1906M of 0.746, 95% Cl of 0.523 to 1.044 days, t1/2-G2019S of 1.049 days, 95% Cl of 0.6343 to 1.773 and t1/2-I1122V of 0.851 days, 95% Cl of 0.578 to 1.241). The stability for LRRK2-R1441C appeared to be reduced by half a day than the LRRK2-WT (t1/2-R1441C of 0.563 days, 95% Cl of 0.4811 to 0.6520 vs t1/2-WT of 0.905 days). While LRRK2-G2385R appeared statistically to be more stable than the WT by half a day (t1/2-G2385R of 1.606 days, 95% Cl of 0.7581 to 5.424 vs t1/2-WT of 0.905 days), this stability looks questionable because the calculation started from a low level of protein to a very low level pointing to a limited reliability in this result. It is good to note that it looks like LRRK2-R1441C and LRRK2-G2385R are the only two to have lost all of the protein by the second to last day of measurement (Figure 5.7).
Figure 5.7 R1441C is potentially the least stable of LRRK2 protein variants


5.3.2.3 Kinase inhibitors effects on protein stability in vivo

To test for LRRK2 protein stability in the presence of kinase inhibitors in vivo vitamin B$_{12}$ and MLi-2 were added to the diet of experimental flies. When vitamin B$_{12}$ was added
(4 μg/ml), the stability of LRRK2 for LRRK2-WT (t1/2-WT+B12 of 0.727 days, 90% CI of 0.606 to 0.869 vs t1/2-WT of 0.905 days), LRRK2-G2019S (t1/2-G2019S+B12 of 1.421 days, 90% CI of 1.073 to 1.954 days vs t1/2-G2019S of 1.049 days), LRRK2-G2019S-K1906M (t1/2-G2019S-K1906M+B12 of 0.614 days, 90% CI of 0.433 to 0.835 days vs t1/2-G2019S-K1906M of 0.746 days), and LRRK2-R1441C (t1/2-R1441C+B12 of 0.714 days, 90% CI of 0.531 to 0.943 vs t1/2-R1441C of 0.563 days) are comparable to the untreated flies. The LRRK2-I1122V stability appeared to be improved by a day (t1/2-I1122V+B12 of 2.106 days, 90% CI of 1.118 to 6.821 vs t1/2-I1122V of 0.851 days) when vitamin B12 is added. Although the stability of LRRK2-G2385R appeared to be reduced by vitamin B12 (t1/2-G2385R+B12 of 0.325 days, 90% CI 0.164 to 0.457 vs t1/2-G2385R of 1.606 days), this reduction compared to the untreated flies is questionable because the t1/2 calculation of the latter ones was not accurate (Figure 5.8).
Figure 5.8 Vitamin B_{12} might affect LRRK2 stability of LRRK2-I1122V

A. Representative blots and B. the time courses of the decay of LRRK2-WT, LRRK2-G2019S, LRRK2-G2019S-K1906M (kinase dead), LRRK2-R1441C, LRRK2-I1122V, and LRRK2-G2385R in *Drosophila melanogaster* when treated with or without vitamin B_{12}. Error bars indicate SEM with N = 3.

When MLi-2 is added (1 µg/ml), compared to uninduced flies, the stability of LRRK2-G2019S (t_{1/2-G2019S+MLi-2} of 1.173 days, 95% CI of 0.773 to 1.894 vs t_{1/2-G2019S} of 1.049 days), LRRK2-G2019S-K1906M (t_{1/2-G2019S-K1906M+MLi-2} of 1.03 days, 95% CI 0.56 to
2.043 vs t1/2-G2019S-K1906M of 0.746 days, and LRRK2-R1441C (t1/2-R1441C+MLi-2 of 0.801 days, 95% CI of 0.577 to 1.106 vs t1/2-R1441C of 0.563 days) are fairly comparable to untreated flies. The stability of LRRK2-WT (t1/2-WT+MLi-2 of 1.495 days, 95% CI of 0.871 to 3.165 vs t1/2-WT of 0.905 days) and LRRK2-I1122V (t1/2-I1122V+MLi-2 of 1.259 days vs t1/2-I1122V of 0.851 days) are improved by almost half a day. Lastly, the reduced stability of LRRK2-G2385R (t1/2-G2385R+MLi-2 of 0.397 days, 95% CI 0.174 to 0.619 vs t1/2-G2385R of 1.606 days) is still questionable similar to results observed with vitamin B12 (Figure 5.9).
Figure 5.9 MLi-2 might effect LRRK2 stability for LRRK2-WT and LRRK2-I1122V


5.3.3 Examining LRRK2 stability via Rab10 and phosphorylated Rab10 expression

Rab10 is a protein that belongs to the Ras family of small GTPases that regulate intracellular vesicle trafficking. It has been found to be a direct target for phosphorylation by LRRK2 (Steger *et al.*., 2016) which, therefore, makes it an attractive downstream target for further investigation. Recently, the Elliott lab has published papers demonstrating a functional association between LRRK2-G2019S and Rab10 in DA neurons (Fellgett *et al.*, 2021, Petridi *et al.*, 2020). In line with this suggestion, the stability of Rab10 and phosphorylated Rab10 should be disrupted if LRRK2 stability is disrupted as well. Due to the ongoing pandemic, the work on these experiments were cut short and unfortunately only one replicate of WB assay was performed which prevented any useful interpretations or conclusions.

5.4 Discussion

In this investigation, the effect of LRRK2 PD-associated mutations and kinase inhibitors on LRRK2 protein stability was tested in cell culture (*in vitro*) and *Drosophila melanogaster* (*in vivo*). In both situations, pulse-chase experiments potentially showed that LRRK2 mutations had differences in the rate of degradation, and that this stability might be impacted by both inhibitors tested. In *vitro*, the order of stability (most stable to least stable) was LRRK2-D2017A (kinase dead), LRRK2-G2019S, LRRK2-WT, and LRRK2-R1441C. In *vivo*, the order of stability was similar for LRRK2-WT, LRRK2-G2019S-K1906M (kinase dead), LRRK2-I1122V, and LRRK2-G2019S then LRRK2-R1441C. Vitamin B<sub>12</sub> potentially improved the stability of LRRK2-I1122V in flies while it potentially reduced the stability of LRRK2-D2019A (kinase dead) in cells. MLi-2 had no effect on cells while it potentially improved the stability of LRRK2-WT and LRRK2-I1122V in flies.
The findings in this investigation suggest the notion that mutations and the presence of kinase inhibitors have an impact on LRRK2 stability. The half-life of LRRK2-WT found here was 0.9 days in flies which converts to around 21 hours. These findings are within the range found in other reports (Ohta et al., 2009, Rudenko et al., 2017, Wang et al., 2008b) that have used tagging methods in pulse-chasing the protein degradation. LRRK2 degradation has been linked to lysosomal and proteasomal pathways. In the lysosomal pathway, Chaperone Mediated Autophagy (CMA) has been found to regulate LRRK2 abundance and mutations to the protein \( \text{LRRK2-G2019S, LRRK2-D1994A, and LRRK2-R1441C} \) compromised the ability of this mechanism to degrade the protein (Orenstein et al., 2013). For degradation via the proteasomal pathway, the C-terminus of Hsp70-Interacting Protein (CHIP) has been found to mediate LRRK2 degradation and like in CMA mutations \( \text{LRRK2-G2019S, LRRK2-D1994A, and LRRK2-R1441C} \) affected the ability of this system to degrade the protein. CHIP regulates this degradation via ubiquitination of LRRK2 and it has been found to bind to different LRRK2 domains (ARM, ANK, ROC, LRR, WD40) (Ding and Goldberg, 2009, Ko et al., 2009, Rudenko et al., 2017). The presence of the mutations potentially alters the ability of CHIP to mediate ubiquitination of LRRK2. Some of the results obtained in this investigation agree with these previous findings while other do not. For instance, LRRK2-R1441C degradation curve suggests a less stable protein in cells and flies while the kinase dead was found to be less stable just in flies. These discrepancies might simply be due to the different models used (cells vs flies) and the different type of mutations e.g., the kinase dead used here is LRRK2-D2017A compared to LRRK2-D1994A (Orenstein et al., 2013). It is possible the effect of difference in models used can be seen clearly in the amount of LRRK2 protein measured at last day of measurement where in cells all transgene except R1441C have still some proteins left compared to flies (see Figure 5.3). In addition, in our cell-based experiment, all LRRK2 constructs are inserted at the same loci in cells while this is not the case in flies.

In addition to PD-associated mutations, kinase inhibition has been reported to affect LRRK2 stability. Here, MLi-2, a highly potent and selective kinase inhibitors, was found to reduce LRRK2-WT stability significantly in cells and slightly in flies which is similar to observations reported by (Lobbestael et al., 2016) in cells, (Kelly et al., 2018) in rats, and (Baptista et al., 2020) in primates. In terms of LRRK2 mutations, G2019S was shown to be resilient to inhibition by MLi-2 in rats (Kelly et al., 2018), similar to what has been observed in flies in this investigation.
MLi-2 and many other kinase inhibitors developed for LRRK2 works in a non-allosteric manner i.e., they are ATP competitive compounds. Although very effective, they have to be highly specific to overcome the high degree of similarities in other ATP binding kinase in cells and to be able to compete with the high amount of ATP found in cells without being toxic. Vitamin B_{12} is a very interesting compound that inhibits LRRK2 kinase activity by blocking its dimerization (Schaffner et al., 2019). The different levels of stabilization by vitamin B_{12} effects on different LRRK2 mutations found in this investigation further our understanding on the mechanism of action of this compound. This is interesting because (Schaffner et al., 2019) reported that vitamin B_{12} interacts with the kinase domain of LRRK2 while we see a vitamin B_{12} stabilization effect LRRK2 bearing mutations in other domains e.g., LRRK2-R1441C in ROC domain and LRRK2-G2385R in WD40 domain. Another interesting allosteric approach that has shown promising results in modulating LRRK2 kinase activity and abundance is the use of small and stable single-domain fragments derived from camelid heavy chain–only antibodies called nanobodies. Expression of nanodoies that bind LRRK2 have been shown to inhibit LRRK2 from phosphorylating Rabs while leaving autophosphorylation intact. Moreover, they can target multiple domains of LRRK2 to potentially modulate activity (Singh et al., 2022).

Investigating kinase inhibition on LRRK2 stability has implications on other forms of familial PD. *GBA1*, which encodes the lysosomal enzyme glucocerebrosidase (GCase) activity, has been associated with an increased risk of developing PD (Avenali et al., 2020). PD patients carrying LRRK2-R1441C or LRRK2-R1441G mutations have shown increased accumulation of oxidized dopamine, decreased lysosomal GCase activity, and increased α-synuclein protein levels in their iPSC-derived DA neurons (Nguyen and Krainc, 2018, Ysselstein et al., 2019). Likewise, knockdown of Rab10, reduces lysosomal GCase activity in human fibroblasts and iPSC-derived DA neurons (Steger et al., 2016). Surprisingly, overexpression of Rab10-WT was sufficient to restore lysosomal GCase activity in iPSC-derived DA neurons LRRK2 mutations (LRRK2-G2019S or LRRK2-R1441C) or GBA (GBA-N370S or GBA-E326K) mutations of iPSC-derived DA neurons. Also, pharmacological LRRK2 kinase inhibition can restore lysosomal GCase activity, and reduce oxidized dopamine and phosphorylated α-synuclein levels, in iPSC-derived DA neurons carrying GBA mutations (N370S or E326K) and primary astrocytes derived from D409V GBA knock-in mice (Sanyal et al., 2020, Ysselstein et al., 2019). While the previous observations clearly suggest an impact
of LRRK2 and its inhibitions on lysosomal abnormalities in PD, the precise mechanism has still to be elucidated.
6 General discussion and future works

6.1 Introduction

The overall aim of this research was to further our understanding on the mechanism of action of LRRK2 mutations and how they may lead to neurodegeneration using the fruit fly as an animal model. In order to achieve this, the following questions were investigated:

1. Can we test for non-autonomy of LRRK2 function by expressing dLRRK in non-neuronal tissues to rescue the visual dysfunction observed in dLRRK\textsuperscript{LOF} old flies? Related to this, given a synthetic lethal genetic interaction between eye colour mutations and dLRRK we wished to investigate the viability of the flies mutant for combinations of eye colour genes and dLRRK.
2. Does dopaminergic LRRK2-G2019S expression have an effect on the ability of the eye’s ability to adapt to light?
3. Do LRRK2 mutations and kinase inhibition affects protein stability \textit{in vivo} as well as \textit{in vitro}?

This chapter reviews our answers to these questions, offering a comprehensive, and concise overview of the key data generated from this investigation. It sets the findings in the general context of LRRK2 biology and also raises further questions and topics for further investigation.

6.2 dLRRK/LRRK2 mode of action

The outcomes from this thesis builds on previous findings from (Furmston, 2016) which found a specific early loss of the ERG off-transient component in a dLRRK\textsuperscript{LOF} model. The findings in this thesis strengthen this observation by adding the loss of the ERG on-transient component which strongly supports that the defect was in the adaptation of lower-order visual neurons (lamina neurons) rather than in the photoreceptors. Moreover, the expression of dLRRK in different neuronal tissues led to the rescue of visual response in dLRRK\textsuperscript{LOF} flies (Furmston, 2016). In addition, the extensive neurodegeneration observed throughout the visual system caused by DA expression of LRRK2-G2019S in flies (Hindle \textit{et al.}, 2013) indicated a transcellular signalling of unknown mechanism. The possibilities for this cell-cell transfer could be alterations in the release of dopamine or another neurotransmitter, changes in secretion of growth factors, secretion of endosomal intermediates such as exosomes or diffusion of free
radicals. However, due to dLRRK/LRRK2 close association with the endocytic pathway, it is more likely to presume that dLRRK/LRRK2 is secreted by exosomes perhaps in a mis-folded manner. The findings in this thesis strongly support the exosomes theory because non-neuronal (pigment cells and fat body) expression of dLRRK was able to rescue the visual deficit in dLRRKLOF flies which is not surprising since they can travel throughout the body and pass specialised compartment including the BBB. Indeed, this is made more likely by the position of the pigment cells and fatty tissue near to the retina. The exosomes theory is supported even further by LRRK2 interaction with different membrane proteins critical in the biosynthesis and traffic of pigment granules found in this thesis and previously (Furmston, 2016, Petridi, 2017). The role of dLRRK in the pigment granules (a type of lysosomal related organelle) regulation supports the idea that LRRK2 interacts with and regulates the lysosome rather than the mitochondria. For example, (Sanyal et al., 2020) have found that LRRK2 inhibition can restore some of the impaired lysosome function observed in GBA heterozygous iPSC neurons. This interaction between LRRK2 and lysosomes was also observed by (Bonet-Ponce et al., 2020, Kluss et al., 2021) where LRRK2 sequesters Rab8a to damaged lysosomes and vesicle sorting from lysosomes.

Intriguingly, another approach that might explain the non-autonomous action of LRRK2 is the link found between inflammatory diseases and LRRK2. Inflammatory bowel disease (IBD) including Crohn’s disease (CD) have been linked to increased risk of developing PD (Park et al., 2019, Wan et al., 2020) and LRRK2 has been shown to play a crucial role in the gut immune cells (Santos et al., 2019) which when affected could in turn lead to neuroinflammation. Neuroinflammation causes the build-up of cytokines and ROS in the brain which are particularly harmful to DA neurons in the SNpc that is known for having the highest density of microglia (Cabezudo et al., 2020). This idea is not new to PD since Braak described the staging of PD (see section 1.1.4) by the level of α-synuclein spreading (Braak et al., 2003). Our data points to another mechanism of spreading in addition to the non-autonomy of α-synuclein driving PD progression, we suggest that PD may also spread via LRRK2 function, potentially via exosomes. The findings in this thesis and others may provide novel therapeutic targets for preventing disease progression e.g., in the lysosomal pathway or immune cells.
6.3 LRRK2 and non-motor deficits in PD

The findings in chapter 4 support even further the theory of LRRK2 spreading from cell to cell since the aberrant ERG was found in flies expressing LRRK2-G2019S in pigment cells that do not contribute to the ERG trace, as well when LRRK2-G2019S is expressed in DA neurons. During each light flash, the initial larger ERGs observed when LRRK2-G2019S is expressed generate an excessive ionic flow and resulting in increased demand for ATP (Hindle et al., 2013). The excess ATP production will potentially cause mitochondrial damage, triggering a degenerative cascade of photoreceptors indicated by the smaller ERG amplitude observed in old flies. Mitochondrial damage is a theme in PD however it appears to be an indirect consequence in case of LRRK2 mutations. The involvement of LRRK2 in lysosomal function may point to dysfunction in lysosomes leading to failure to degrade damaged and senescent lysosomes, thus causing excessive ROS generation by degenerate mitochondria. This is not the case with parkin or PINK1 linked PD which are known to be involved more directly in mitochondrial function (Quinn et al., 2020). Therefore, therapies targeting mitochondria function in LRRK2 mutations linked PD, such as using ursocholanic acid (UCA) or ursodeoxycholic acid (UDCA) (Furmston, 2016), would not obviously be the favourable approach, but may have some benefits.

Failure of neuronal adaptation might also contribute to other PD deficits. For example, the reward system which is suggested to be altered in PD in a dopamine-dependant manner (Frank et al., 2004). Reward and motivation in healthy individuals cause pupillary dilation, presumably due to effects of arousal and attention. Dopamine-induced dilation of the pupil may occur as a result of increased sympathetic outflow in the locus coeruleus, energising upcoming actions (Murphy et al., 2011, Varazzani et al., 2015). A reduced pupillary dilation has been observed in PD patients (Manohar and Husain, 2015, Muhammed et al., 2016) similar to the light adaptation abnormalities in the fly eye found in this thesis. Assessment and treatment of the visual deficits in PD is often neglected probably because PD patients and their family members are often oblivious of the relationship between visual deficits and PD, the movement deficits being a primary observation and concern. Thus, development of portable ERG/SSVEP setups for human use would be helpful, both in assessing the progress of the disease, but also in monitoring any therapy.
6.4 LRRK2 stability and kinase inhibitors

A key finding in chapter 5 was that LRRK2 stability appeared to be affected by mutations and kinase inhibitors. The cause of this instability is still not clear but recent experiments with LRRK2 and ubiquitination might give us some clues. Ubiquitination is a common post-translational protein modification that involves the covalent attachment of a single (monoubiquitination or multi-monoubiquitination) or a chain of ubiquitin proteins (polyubiquitination). The link occurs between the carboxyl group of C-terminal residue of ubiquitin (G76) and the side chain amino group of a lysine (K) residue of the protein substrate (Buneeva and Medvedev, 2022). According to the Protein Lysine Modification Database (PLMD), 4 sites on LRRK2 have been identified as ubiquitination sites (K905, K906, K412, and K1132) so far. LRRK2 has been found to be promoted for proteasomal degradation by 2 ubiquitin ligases, CHIP and tripartite motif family 1 (TRIM1). The exact ubiquitination sites for these are still not clear. For TRIM1, it is reported to be between the ARM and LRR domains while CHIP is reported to act on multiple domains (Ding and Goldberg, 2009, Stormo et al., 2022). In addition, dephosphorylation of LRRK2 can induce its ubiquitination by binding to PP2A phosphatase, eventually leading to degradation (Drouyer et al., 2021). The previous reports give multiple targets to consider for more investigation and targeted novel therapies.

Investigating the effect of kinase inhibitors on LRRK2 stability can lead to potential therapeutic interventions for PD. The most common mutation, LRRK2-G2019S is proposed to have an increase in intrinsic kinase activity (Lee et al., 2019). In this investigation, 2 kinase inhibitors of different mechanism of action on kinase activity and different potential effects on LRRK2 protein stability, MLi-2 (non-allosteric) and vitamin B₁₂ (allosteric) were tested for LRRK2 protein stability. The latter one is of interest because it is found naturally in many animal products but not in plant foods, therefore it is reasonable to imply that it is potentially safe as a therapeutic. Deficiency in B₁₂ either due to diet or inability to absorb the vitamin as observed in some cases of IBD can increase the risk of developing PD and other neurological symptoms among many (Hunt et al., 2014, Shen, 2015). In this project, we have found that Vitamin B₁₂ was potentially able to improve the stability of LRRK2-I1122V in flies and LRRK2-D2019A (kinase dead) in cells. Vitamin B₁₂ in this investigation and in another study (Schaffner et al., 2019) has shown promising results on LRRK2 linked PD but it has also been shown to affect other types of familial PD specifically PINK1 linked PD.
Vitamin B₅ (a family member of vitamin B) has been shown to rescue the mitochondrial dysfunction found in PINK1 loss of function flies (Huang et al., 2022). Our results and previous results give a good platform to consider using vitamin B₁₂ in more research with regard to LRRK2 function.

### 6.5 Future works

In chapter 3, dLRRK has been suggested to be involved in the biosynthesis and function of pigment granules in addition to genetic interaction with different small molecule synthesis pathways (kynurenine, pteridine and granule pathways) that are important to neurotransmitter, antioxidant and tissue pigment generation. We have found in this project that expressing eye pigment genes se, sptr, and pu were not able to rescue the age-dependent loss of visual function in dLRRK^{LOF} old flies. In addition, we have found that no synthetic lethality was produced when cn^{5k} and pr′ individually were crossed with dLRRK^{LOF}. It would be useful to test the interaction of dLRRK with more eye colour genes to add the data we have generated here to form a clear picture for this interaction. Furthermore, to expand on the data of testing the genetic interaction in null or loss of function mutations, to test for interaction at the tissue and cellular level by knocking down eye pigment gene function using targeted RNAi in various tissues to locate the critical tissue where this interaction between these genes and LRRK2 occurs.

In chapter 4, a critical issue in the inability to obtain good results for DPP or PPA was due to the variation in eye colour pigmentation of flies. The degree of pigmentation of the eye is also known to affect the size of the ERG due to the absorbance of light by the pigment (Stark, 1973). Thus, testing ERGs in PD models would be optimal where eye colour pigmentation is uniform between the different genotypes. Flies expressing LRRK2 in pigment cells have shades of orange and gave the best results in all three assays in this project. Incorporating w^{Apricot}, which has an orange eye, into flies might be a good strategy to unify eye colour of flies into shades of orange. Moreover, some statistical significance might not appear in graphs even though they look significant due to the small sample size. This could be easily addressed by increasing the number of examined sample.

In chapter 5, we observed that mutations in LRRK2 was potentially affecting the protein stability, specifically LRRK2-R1441C where it appeared to be less stable than the wild type in cells and in flies. LRRK2 degradation in cells appeared to be still ongoing by the end of the measurement period for all genotypes except for LRRK2-R1441C.
compared to the flies which LRRK2 appeared to be mostly degraded by the end of the measurement period which might not give a complete picture of the protein decay in cells. An extension of the measurement period could resolve this issue. However, in case of cell culture, when I began the project, the hypothesis that LRRK2 was associated with microtubules was not well supported and considered unlikely. But several recent papers have emphasised that LRRK2 can interact with microtubules. Evidence has been accumulating from structural studies which supports this interaction (Deniston et al., 2020, Leschziner and Reck-Peterson, 2021). This potential interaction suggests that using GAPDH instead of β-tubulin for normalisation would be more appropriate. A different approach in examining LRRK2 stability is to investigate strong and direct interactors upstream (e.g., protein 14-3-3) or downstream (Rab10) of LRRK2 which for the latter I had some preliminary data that was cut short by Covid. In flies, it would be good to use a set of flies where the transgenes are all inserted in the same genomic landing sites, and an inducible TH-Gal4 so that the precise effects and timing of LRRK2 expression in DA neurons can be established.

6.6 Conclusion

Throughout this project, the function and dynamic of dLRRK/LRRK2 have been tested in a Drosophila model of PD and in tissue culture. Multiple assays have been utilised to investigate the interaction of LRRK2 and dLRRK with other mutations in pigment synthesis and pigment granule biogenesis on viability, effects on vision, and general stability of the protein in flies as well as in cells. The findings provided in this thesis supplement previously published LRRK2 data and raise new research concerns that may help us learn more about how mutations in this gene cause Parkinson's disease.

Summary of key findings:

1. Expressing dLRRK in non-neuronal tissue was able to rescue visual deficit found in dLRRK^LOF old flies.
2. Expressing eye pigment genes se, sptr, and pu were not able to rescue the age dependant loss of visual function in dLRRK^LOF old flies.
3. No synthetic lethality was observed for cn^tsk and pr^ individually when crossed with dLRRK^LOF.
4. Young flies expressing LRRK2-G2019S in DA neurons or pigment cells failed to adapt to the light.
5. The potential order of LRRK2 stability in flies (most stable to least stable) was similar for LRRK2-WT, LRRK2-G2019S-K1906M (kinase dead), LRRK2-I1122V, and LRRK2-G2019S then LRRK2-R1441C while in cells it was LRRK2-D2017A (kinase dead), LRRK2-G2019S, LRRK2-WT, and LRRK2-R1441C.

6. The LRRK2 inhibitor Vitamin B$_{12}$ potentially improved the stability of LRRK2-I1122V in flies while it potentially reduced the stability of LRRK2-D2019A (kinase dead) in cells.

7. The LRRK2 inhibitor had no effect on the stability of LRRK2 while it potentially improved the stability of LRRK2-WT and LRRK2-I1122V in flies.


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