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Early Mechanisms of Retinal Degeneration in the harlequin Mouse

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Supervisor: Dr. Kathleen Hill, *The University of Western Ontario* A thesis submitted in partial fulfillment of the requirements for the Master of Science degree in Biology © Eric Dolinar 2014

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Early Mechanisms of Retinal Degeneration in the harlequin Mouse

(Thesis Format: Monograph)

by

Eric A Dolinar

Graduate Program in Biology

A thesis submitted in partial fulfillment

of the requirements for the degree of

Master of Science

The School of Graduate and Postdoctoral Studies

The University of Western Ontario,

London, Ontario, Canada

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Abstract

Retinal diseases are personally debilitating and expensive, yet many early disease mechanisms leading to their onset and progression remain poorly understood. The *harlequin* mouse is a model of human mitochondrial dysfunction and parainflammation leading to subsequent cerebellar and retinal degeneration. Diagnosis of retinal degeneration can be tracked *in vivo* and is associated with AIF dysfunction. Here, retinal dysfunction in the *harlequin* mouse was first quantified using electroretinography followed by assay of blood-retinal-barrier integrity and transcriptome alterations in young adulthood. Nonmetric multidimensional scaling of oscillatory potentials provided a novel, comprehensive assessment of inner-retinal health and can detect shifts in OP parameters. Barrier integrity ruled out confounding exogenous antigens and confirmed an endogenous source of retinal tissue malfunction. In addition, transcriptome alterations support the necessity of the hq retina to maintain metabolic demands. Alternative metabolism pathways are hypothesized to be important for hq complex I mitochondrial-dysfunction associated retinal degeneration.

Keywords: *harlequin, Apoptosis-inducing factor*, retinal degeneration, mitochondrial dysfunction, electroretinography, oscillatory potentials, nonmetric multidimensional scaling, blood retina barrier, Evans Blue, ATP deficiency, quantitative gene expression, glycerol metabolism.

Co-authorship Statement

Eric Ammon Dolinar performed the following work under the supervision and financial support of Dr. Kathleen Allen Hill. This thesis is presented in monographic format. Eric Ammon Dolinar performed the experimental research presented in this thesis and will be a co-author on resulting publications. Dr. Kathleen Hill will be a senior author on all publications produced from this research due to her role in project design, supervision, literature research, data analysis and assistance with publication writing. Alex Laliberté, Thomas MacPherson, Anita Prtenjaca, Anson Li and Dr. C.M.L Hutnik will be co-authors on specialized papers produced from the research due to assistance in experimental design, gene expression array data or electroretinography and clinical relevance.

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List of Abbreviations

Adcy10	Adenylate cyclase 10
ADP	Adenosine Diphosphate
AGEs	Advanced Glycation End-products
Aif	Apoptosis-inducing factor
Aifm1	Apoptosis-inducing factor 1
Aldh2	Aldehyde dehydrogenase 2
AMD	Age-related Macular Degeneration
ANOVA	Analysis of Variance
ATP	Adenosine Triphosphate
bp	base pair
BRB	Blood Retinal Barrier
Ca ²⁺	Calcium Ion
cAMP	cyclic Adenosine Monophosphate
Cdkn1a	Cyclin-dependent kinase inhibitor 1A
cDNA	complementary Deoxyribonucleic Acid
cGMP	cyclic Guanosine Monophosphate
CO_2	Carbon Dioxide
ddH ₂ O	double-distilled water
DNA	Deoxyribonucleic Acid
DR	Diabetic Retinopathy
EB	Evans Blue Dye
ERG	Electroretinogram/Electroretinography
FADH ₂	Flavin Adenine Dinucleotide

- FU Fluorescence
- G3P Glycerol-3-Phosphate
- G-protein Guanosine nucleotide-binding protein
- GABA Gamma-Aminobutyric Acid
- Gapdh Glyceraldehyde 3-phosphate dehydrogenase
- GCL Ganglion Cell Layer
- gDNA genomic Deoxyribonucleic Acid
- Gk5 Glycerol Kinase 5
- *Glut4 Glucose transporter, member 4*
- *Gpd2 Glycerol-3-phosphate dehydrogenase 2*
- HCl Hydrochloric Acid
- HCO₃ Bicarbonate Ion
- Hk1 Hexokinase 1
- HMGB1 High Mobility Group Box 1
- hq harlequin
- *hqX* Moderate disease *harlequin* female, heterozygote genotype
- *hqY* Severe disease *harlequin* male, hemizygous genotype
- INL Inner Nuclear Layer
- IPL Inner Plexiform Layer
- IS Inner Segment (Photoreceptors)
- ISCEV International Society for Clinical Electrophysiology of Vision
- K⁺ Potassium Ion
- kb kilobase
- *Ldhc Lactate dehydrogenase C*
- Mct Monocarboxylate Transporters

- mRNA messenger Ribonucleic Acid
- Na⁺ Sodium Ion
- NAD+ Nicotinamide Adenine Dinucleotide
- NaOH Sodium Hydroxide
- NC No DNA Control
- NMDS Nonmetric Multidimensional Scaling
- nt nucleotide
- NTC No Template Control
- OAG Open-angle Glaucoma
- ONL Outer Nuclear Layer
- OP Oscillatory Potentials
- OPL Outer Plexiform Layer
- OS Outer Segment (Photoreceptors)
- **OXPHOS** Oxidative Phosphorylation
- PAMPS Pathogen-Associated Molecular Patterns
- PBS Phosphate Buffered Saline
- PCR Polymerase Chain Reaction
- *Pdhx Pyruvate dehydrogenase, component X*
- *Pfkfb3 Phosphofructose-2-kinase/fructose-2,6-bisphosphatase-3*
- PRP Pattern Recognition Receptors
- qPCR quantitative Polymerase Chain Reaction
- RAGEs Receptor for Advanced Glycation End-products
- RIN Ribonucleic Acid Integrity Number
- Rn18s 18S Ribosomal subunit
- RNA Ribonucleic Acid

- ROS Reactive Oxygen Species
- RPE Retinal Pigmented Epithelium
- rRNA ribosomal RNA
- Samd4 Sterile alpha motif domain containing 4
- Scn4b Sodium channel beta 4 subunit
- Sdk2 Sidekick-2
- Slc4a10 Sodium bicarbonate transporter, member 10
- STD Size Standard
- Tbc1d4 TBC1 domain family, member 4
- TCA cycle Tricarboxylic Acid cycle
- TNFα Tumor Necrosis Factor *alpha*
- VEGF Vascular Endothelial Growth Factor
- WT Wild Type, specifically XY
- X^{hq}X harlequin Heterozygous Disease Mouse
- X^{hq}X^{hq} harlequin Homozygous Disease Mouse
- X^{hq}Y harlequin Hemizygous Disease Mouse

Chapter 1. Introduction

1.1 Increased prevalence of neurodegenerative disorders in an increasingly aged population

Neurodegenerative diseases are characterized by progressive functional deficiencies and cell death of neurons resulting in nervous system dysfunction. Medical advances in cardiovascular diseases, cancers and other common illnesses have increased human longevity and thereby increased prevalence of neurodegenerative disorders.¹ Predispositions from familial and environmental risk factors may also play a role in neurodegenerative disease onset and progression, however, the highest propensity of disease risk occurs with age.² Despite high prevalence and extreme negative consequences to quality of life, the causes and early disease mechanisms of such diseases remain unknown.

In Canada, it is expected that within 25 years the number of seniors will double to well over 9.9 million people and surpass the age demographic of 0-14 year olds for the first time in Canadian history (Population Projections for Canada, Provinces and Territories. 2009-2036).³ Common retinal degenerative diseases such as: age-related macular degeneration (AMD), diabetic retinopathy (DR) and open-angle glaucoma (OAG) affect well over 1.75 million Canadians and the prevalence of such diseases is expected to rise proportionally (CNIB Report, 2009).⁴ These three diseases fall under the umbrella term of "vision loss" and have the highest direct healthcare costs of any disease in Canada at \$8.6 billion annually.⁴ Despite this high level of expenditure, little is known about early disease mechanisms and prevention leaving the diseases personally debilitating and expensive. Vision research remains critical for increasing the quality of

life of affected individuals and also for reducing expenditures as the increasingly aged population brings alarming financial demands.

1.2 Mechanisms of common retinal degenerative diseases remain unclear

As the Canadian population shifts towards an older demographic cohort, the prevalence of common retinal degenerative disorders is expected to rise.⁴ These common diseases are progressive and require continual and enhanced treatment as the disease advances. Retinal degenerative disorders can lead to losses of central, peripheral, and even complete vision in the later stages of disease progression. Unfortunately, disease mechanisms and triggers leading to neuronal death are numerous and little is known about initiating mechanisms that cause retinal degeneration.

Age-related macular degeneration (AMD) is a prevalent retinal degenerative disorder affecting over one million Canadians and is the leading cause of irreversible blindness in the developed world.^{4,5} AMD begins with the deposition of acellular protein and lipid debris, known as drusen, in the space behind the retina around the macula.⁶ The macula contains the highest density of cone photoreceptors in the primate eye and is responsible for the central vision acuity that we appreciate as human beings.⁷ Excess drusen can cause damage to the retina and lead to hypoxia. Hypoxia induces the expression of angiogenic cytokines, such as vascular endothelial growth factor (VEGF), causing neovascularization in the macula and leading to subsequent central vision loss.⁵

Diabetic retinopathy (DR) is the second-most prevalent retinal degenerative disorder affecting over 500,000 Canadians.⁴ It is a common complication of diabetes caused by hyperglycemia, hypoxia and neovascularization.⁸ The exact mechanisms by

which DR causes neovascularization and progressive retinal degeneration remain poorly understood.

Open-angle glaucoma (OAG) is the third-most prevalent retinal degenerative disorder affecting over 250,000 Canadians.⁴ Glaucoma is often caused by an increase in intraocular pressure leading to optic neuropathy.⁴ Increases in intraocular pressure caused by reduced aqueous humor drainage lead to progressive retinal neuron axon loss, particularly along the optic nerve.⁹ Mechanisms that drive retinal cell loss remain unknown and lead to peripheral vision loss with no preventative treatment available.¹⁰

The progressive symptoms associated with these common retinal degenerative diseases are clearly characterized and various treatments are available in the later advancements of the disease. However, early mechanisms and triggers leading to retinal degenerative disorders are poorly understood and treatment is only available after cell damage has already begun. Therefore, to increase the quality of life, an understanding of early disease mechanisms is critical to limit disease onset and progression.

1.3 The physiology and anatomy of a healthy retina is complex

The retina is one of the most complex structures of the vertebrate eye and a key component of the visual pathway (Figure 1.1A). The retina is a light-sensitive tissue that lies in the posterior region of the vertebrate eye and is divided into two regions known as the neural retinal and retinal pigmented epithelium (RPE) (Figure 1.1B).⁷ These two anatomical regions develop from the embryonic forebrain and undergo complex differentiation and ultimately function as an outcropping of the central nervous system.¹¹

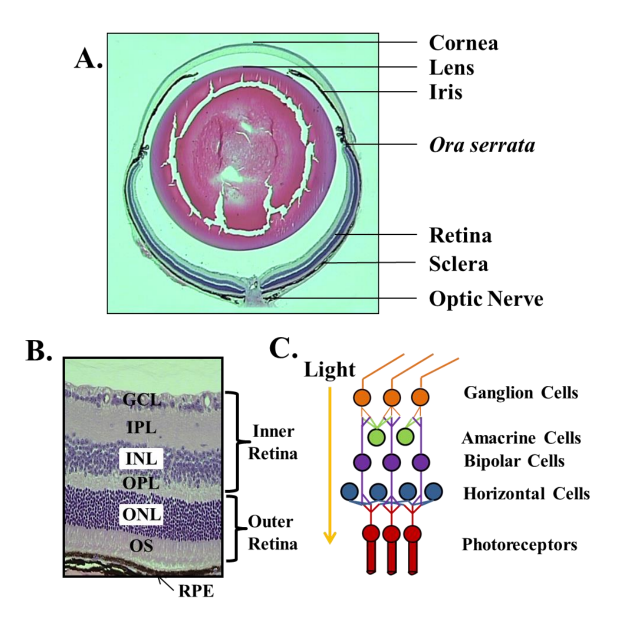


Figure 1.1 A cross-section of the mouse eye and retina stained with Hemotoxylin and Eosin. (A) A C-cut section of the mouse eye demonstrates similar anatomical features of the human eve including the cornea, lens, pupil, iris, retina, scalera and optic nerve. Representative wild-type mouse eve adapted from MacPherson (2009).¹⁹ (B.C) A crosssection of the mouse retina indicates the distinct retinal cells layers. Light must transverse all layers of the retina prior to excitation of the photoreceptors which are positioned in the outer segment (OS). The retinal pigment epithelium (RPE) lies at the base of the retina and absorbs any stray photons of light not absorbed by photoreceptors. In addition, the RPE separates the retina from the posterior choroid and acts as a nutrient shuttle. Phototransduction begins in the OS as the electrochemical signal is conveyed through the outer nuclear layer (ONL) and synapses through the outer plexiform layer (OPL) onto the horizontal cells and bipolar cells of the inner nuclear layer (INL). The electrochemical signal is then relayed to the amacrine and ganglion cells of the ganglion cell layer (GCL) through the inner plexiform layer (IPL). The electrochemical signal leaves the GCL via the optic nerve where it is transmitted to the visual cortex for interpretation.

The design of the mammalian retina is counter-intuitive in nature because light must cross all neural retinal layers prior to the excitation of photoreceptors. Thereby, photoreceptors are positioned at the most posterior region of the neural retina, only posteriorly superseded by the RPE (Figure 1.1B).¹² This is due to the fact that the RPE continuously supplies nutrients to the highly metabolic photoreceptors.¹¹ The RPE is a monolayer of pigmented cells that separates the internal choroid from the neural retina and acts as a nutrient shuttle, providing a steady stream of 11-*cis* retinal, glucose, and fatty acids from the choroid to the outer segment of the photoreceptors.^{13–15} The RPE also plays an essential role in the recycling of all-*trans* retinal back into 11-*cis* retinal once it has been isomerized by light.^{14,13} The RPE is highly pigmented, containing a high proportion of melanin granules that absorb stray photons of light in the retina to enhance vision acuity and reduce exposure to light radiation.⁷ In addition, the RPE creates an outer blood-retinal-barrier (BRB) through epithelium tight junctions which maintains optimal homeostatic conditions for retinal neurons.¹⁵

Anterior to the RPE lies the neural retina composed of six differentiated layers (posterior to anterior) (Figure 1.1B,C). The outer segment (OS) contains the familiar rod and cone photoreceptors responsible for the initial absorption of light and relay of electrochemical signal.⁷ All vertebrate retinas contain both rod and cone photoreceptors which differ by their absorption wavelength and cell morphology. Rods are thin, rod-shaped structures that are highly sensitive to blue-green light (500 nm) and are densely packed with the visual surface photopigment rhodopsin. Rods are generally used for low-light vision (scotopic vision) and make up 95-97% of all photoreceptors in the mammalian eye (human 95%).^{7,16} Cones are large, conical structures that are densely

packed with opsin photopigments and are sensitive to daylight and bright colours (photopic vision). Three types of cones with different associated opsin photopigments exist in the human retina, each responsible for the absorption of a different wavelength of light: red (559 nm), green (431 nm) or blue (419 nm).¹⁶ This provides the trichromatic vision that we appreciate as humans. The inner segment (IS) contains the cell bodies of the photoreceptors and is responsible for maintaining metabolism.¹⁶ In the outer nuclear layer (ONL) lie the nuclei of the photoreceptors. The outer plexiform layer (OPL) is a thin region of synaptic connections between the photoreceptors and the neural cells of the inner nuclear layer (INL). The outer plexiform layer (OPL) is a region densely populated with axons from the rod and cone photoreceptors and neurites of bipolar and horizontal cells.¹¹ Eleven types of human bipolar cells exist in the INL, the most common being bipolar cells connecting photoreceptors with ganglion cells through the release of glutamate.¹¹ Horizontal cells have a large lateral extension across the retina and synapse back onto rod and cone photoreceptors in an inhibitory manner.¹⁷ This provides a level of local gain in the retina and allows image processing to begin through inhibiting lightsaturated photoreceptors to prevent scattering of light in the retina.¹⁸ The INL also contains the nuclei of amacrine cells which mediate the effect of the ON-,OFF- bipolar cells on ganglion cells through the release of neurotransmitters, glycine and gammaaminobutyric acid (GABA).¹⁸ These mechanisms assist with the early building of images in the retina.⁷ Synapsis of bipolar and amacrine with ganglion neurons occurs in the inner plexiform layer (IPL). The innermost layer of the retina is the ganglion cell layer (GCL) which contains both the axons and nuclei of the ganglion cells and is responsible for relaying electrochemical signals to the visual cortex via the optic nerve.

1.4 Phototransduction is compromised in retinal degeneration

Phototransduction is the process in which photons of light excite photopigments in the photoreceptor cells to generate an electrical response.²⁰ First, light must travel through all layers of the retina and will only activate photoreceptors in the field of In rod photoreceptors, rhodopsin is the primary photopigment and is reference. covalently linked to the chromophore, 11-cis retinal.²¹ During the absorption of light, 11*cis* retinal is converted to all*-trans* retinal through photoisomerization.²⁰ Conformational changes of rhodopsin are capable of activating photoreceptor-specific G-protein, transducin.²⁰ Transducin stimulates the activity of cGMP phosphodiesterase, resulting in a reduction of cytoplasmic cGMP concentration and subsequent closure of cGMP gated channels.²² Closure of the cGMP channels results in a decrease of Na^+ and Ca^{2+} influx triggering membrane hyperpolarization and reduced release of synaptic neurotransmitter, glutamate.²⁰ Glutamate is the principal neurotransmitter involved in electrochemical transmission in the retina.⁷ Unlike common nerve cell function, photoreceptors release glutamate in the dark as Na⁺ and Ca²⁺ flow freely across the phospholipid membrane.²⁰ Upon light stimulus, glutamate release is hindered in photoreceptors and leads to subsequent depolarization of bipolar, horizontal and amacrine cells of the INL.²² This signal is further processed by the GCL before travelling to the visual cortex via the optic nerve. Phototransduction occurs in a similar fashion in cone photoreceptors, however, opsin proteins vary slightly in their optimal absorption wavelength. Phototransduction is imperative to normal retinal function and vision. Common retinal degenerative diseases can lead to damage of the photoreceptors and phototransduction pathway by various

disease factors including: age, light toxicity, diabetes, hypertension, genetic mutations, vitamin A deficiency, angiogenesis, parainflammation and mitochondrial dysfunction.^{23,24}

1.5 Neuronal homeostasis is maintained by glial cells of the retina

Three types of resident glial cells reside in the retina and are essential for retinal function and homeostasis, these include Müller cells, astroglia, and microglia cells. Müller cells are the most abundant glia of the retina and provide architectural support across all neural layers.²⁵ Müller cell bodies sit in the INL and perform a variety of functions including: the breakdown of glycogen for anaerobic metabolism,²⁶ removal of excess neurotransmitters from the subretinal space,²⁷ assistance with phagocytosis of apoptotic neurons and neuronal debris,²⁸ assistance with the formation of the BRB²⁹ as well as maintaining ionic homeostasis in the retina by recycling $K^{+,30}$ Astroglia are the supportive glia that originate in the optic nerve head and are subsequently present in the GCL.^{31,32} Similar to Müller cells, they assist with the formation of barrier properties by the breakdown of glycogen for neuron nutritive vascular endothelial cells.^{29,33,34} support,²⁶ and regulation of ionic homeostasis³⁰ as well as GABA metabolism.²⁵ The resident macrophage of the retina is the microglial cell. The retina is a region of immune privilege protected by the BRB to hinder the movement of pathogens and microbes into the highly sensitive neural retina. Microglia cells act as part of the innate immune system and partake in immune survelliance³⁵, neuronal support³⁶ and phagocytosis of cellular debris and pathogens.³⁷ Under necrotic or apoptotic pathways, cellular debris such as ATP, uric acid and other alarmins are released and stimulate the innate immune system.³⁸

Microglial cells become activated and elicit an inflammation response to clear cytotoxic debris from the retina. Together, these three types of glial cells all play a vital and distinct role in the retina to maintain homeostasis and proper retinal function.

1.6 Retinal neurons rely heavily of oxidative phosphorylation for ATP generation

Neurons of the retina require high amounts of ATP to maintain ionic potentials through the active transport of Na⁺ influx and K⁺ efflux by Na⁺K⁺ATPases.³⁹ In addition, the continuous recycling of neurotransmitters places additional metabolic pressure on the cell.⁴⁰ Neurons do not store glycogen and suppress the enzymatic machinery to synthesize glycogen and therefore rely on the constant flow of oxygen and glucose from the vasculature in order to maintain their high metabolic rate.^{41,42} Dendrites, regions of frequent membrane hyperpolarizations/depolarizations and ion mobility, are the most energetically consuming region of the neuron. Consistent with their great metabolic rate, neurons rely heavily on glucose oxidative phosphorylation (OXPHOS) for ATP production.^{43–45} Glial cells, on the other hand, are highly glycolytic. In addition, glia also have the ability to store glycogen and convert it to glucose 6-phosphate when glucose supply to the retina is diminished.⁴⁴ Glial cells can provide lactate and other metabolites to the neurons of the retina via Monocarboxylate transporters (MCT) when glucose supply is diminished.⁴⁰

Glucose is supplied to the retina through the choroid (located posterior to the RPE) and the retinal capillaries.⁷ Glucose must first traverse the epithelial barrier of the RPE (outer BRB) or the retinal capillary endothelial vasculature (inner BRB) which controls the exchange of metabolites and waste products between the vasculature and neural retina.⁴⁶ The inner BRB supplies nutrients and oxygen to the ONL, INL, and GCL, whereas the outer BRB is responsible for neuronal support of the inner and outer segments of the photoreceptors.⁴⁶ Pericytes surround the endothelial cells and secrete angiopoietin 1 to induce tight junction protein expression.⁴⁷ Both the inner and outer BRB develop junctional complexes between the vascular endothelial/epithelial cells to allow limited movement of ions and solutes into the neural retina.^{46,48} Tight junctions are made up of transmembrane proteins known as, claudins, occludin and the junctional adhesion molecules which are linked to the cytoskeleton of the adjacent cells.⁴⁶ Tight junctions lead to the fusion of two adjacent endothelial or epithelial cells and limit intracellular space to reduce diffusion of ions, pathogens, lipids, proteins, reactive oxygen species (ROS) and foreign substances into the sensitive neuronal space. Epithelial cells of the RPE act as a regulatory interphase between the choroid and outer retina. The basal surface of the RPE contains many invaginations to enhance selective transport and secretion along the vasculature.¹⁵ Due to the continual active transport of substrates across the RPE and vasculature barrier, cells are highly metabolic and subsequently require high levels of ATP to maintain efficient transport. The endothelial/epithelial structure of the BRB is considered to be the first line of defence to protect the neural retina from invasion of pathogens as they can engulf foreign antigens.¹⁵ In addition,

epithelial cells of the RPE can excrete various cytokines to initiate an inflammatory response.⁴⁹ Dysfunctions of the BRB are associated with DR as the dysfunction leads to pathological changes such as chronic inflammation and hypoxia triggering retinopathy progression.^{50,51} In addition, the outer BRB harbours the choroid vasculature and limits neovascularization in the retina. A loss of integrity also leads to AMD and subsequent vision loss.

1.8 Electrophysiological examination can assess health of the visual system in vivo

Electroretinography can be used to assess the integrity of phototransduction in the retina. The electroretinogram (ERG) is a massed, retinal response measured by corneal electrodes following a light stimulation. The major components of the ERG include, the a-wave and the b-wave with high-frequency oscillations known as oscillatory potentials (OP) on the ascending limb of the b-wave (Figure 1.2).¹¹ The initial output wave, known as the a-wave, is associated with photoreceptor hyperpolarization.⁵² The secondary wave, known as the b-wave, is associated with the depolarization of bipolar cells and Müller cells.⁵³ The exact origin of the oscillatory potentials remains unknown, although they are believed to be generated by interactions between amacrine, bipolar and ganglion cells.^{54–} ⁵⁶ Electroretinography can be used to assess the function of both rod- and cone-based systems. "Dark-adaptation" of the subject prior to the assay will result in rod hyperpolarization following a light stimulus.⁵⁷ "Light-adaptation" of the subject prior to the assay will result in hyperpolarization of the cone photoreceptors as the rods will be saturated.⁵⁷ Electroretinography performed under scotopic conditions assesses the functionality of rod photoreceptors and can be used to examine the inner retinal pathways

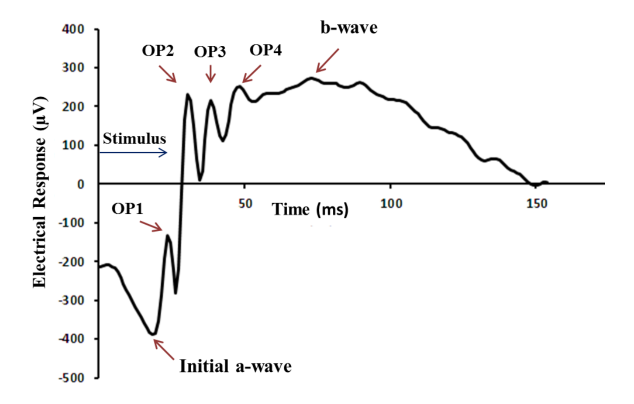


Figure 1.2. A schematic representation of a trace electroretinogram elicited by a healthy, dark-adapted mouse retina following a light stimulus of 30 cd·s/m² (Flash 11). All major characteristic components of the electroretinogram are evident. The a-wave represents the initial hyperpolarization of the photoreceptors and is the initial negative deflection. This is followed by the intermediate oscillatory potentials (OP) peaks 1-4 (OP1, OP2, OP3, OP4) on the ascending limb of the b-wave. It is apparent that early OP peaks are correlated with the function of the photoreceptors whereas late OP peaks are indicative of ganglion and amacrine negative feedback in the outer retina.⁵⁶ The final component, the b-wave, represents the depolarization of the bipolar and Müller cells, and is the cumulative response of the retina.

and mechanisms rods initiate. Since the mouse retina is composed of well over 97% rods ⁵⁸ ERGs performed under scotopic conditions provide a more general, comprehensive analysis of retinal function and health. The ERG provides the only non-invasive, *in vivo*, diagnostic tool that can be used to quantify retinal health by testing functionality of photoreceptors and neural cells of the retina.

1.9 Oscillatory potentials can provide early in vivo examination of INL and GCL function

Oscillatory potentials (OPs) are the low amplitude, high-frequency oscillations that are found on the ascending limb of the b-wave.¹¹ The exact origin of OPs remains unknown and their interpretation remains difficult, however attritional changes are associated with diseases of the retina particularly retinopathy and retinal degeneration. Early OPs are believed to be associated with the hyperpolarization of the photoreceptors and intermediate OPs with action-potential (graded) independent neurons such as horizontal cells and bipolar cells.⁵⁶ Later OPs however, are associated with more proximal retinal responses, that of action-potential dependent mechanisms of amacrine and ganglion cells.⁵⁶ Due to the fact that OPs appear as oscillations on the leading edge of the b-wave, it has been suggested that OPs may be the result of neurons with large lateral mobility in the IPL i.e. amacrine cells.^{54,59} Additionally, the interplexiform cell which spans the OPL and IPL may also generate radial current (feedback) from lateral synapses and may be responsible for the oscillations.^{54,55} OPs have been used previously, to study the function of the inner retina and have been useful in the diagnosis of a variety of progressive retinal eye diseases including; increased intraocular pressure, diabetic

retinopathy, retinal vascular occlusions, and retinopathy of prematurity.^{59–66} OPs have been useful in the diagnosis of DR prior to any structural cell loss or ocular symptoms in both human and animal models.^{59,62,65} Therefore OPs can be used as a versatile, disease tracking, diagnostic tool prior to any structural and noticeable changes.

There is little consensus on the most appropriate method for extracting oscillatory potentials from the ERG. Signal conditioning is the first step that must be performed for isolation of OPs from the ERG trace. Current recommendations set by the International Society for Clinical Electrophysiology of Vision (ISCEV) suggest a simple band-pass filter applied at 75 to 300 Hz to remove any a-wave contribution or high-frequency noise.⁶⁷ In mice, these cut-offs discard most of the rod-driven OPs and are therefore rarely followed.^{68–70} The most common method of OP isolation involves a preconditioning waveform to remove any photoreceptor contribution^{68,71–74} followed by a fifth order Butterworth transformation with a band-pass of 65 to 300 Hz (Figure 2.2).^{68–70,75} The amplitude and latency of each OP wavelet can be analyzed individually or as a sum in the time domain; or through a fast Fourier transform in the frequency domain.

The underlying goal of any OP examination is to surmise physiological origins of the parameters or to identify a clinically relevant, diagnostic parameter that may infer disease mechanisms prior to disease onset.⁷⁶ In the time domain (Figure 2.3), analysis of individual OP amplitudes is advantageous because the amplitudes offer implicit information about associated retinal regions given retinal layers and signal transduction through the retina.⁶⁰ Summed amplitude of all OPs encompasses a full inner retinal response of the INL and GCL. Initial latency is a measure of synapsis timing and periodicity of the INL/GCL retinal response.⁷⁷ Decreases in both OP summed amplitude

and initial latencies have found in retinopathy,^{66,78,79} diabetic retinopathy,^{63,65} and glaucoma.⁶¹ The time domain analysis of OPs contains relevant information in regards to individual components and mechanisms occurring in the retina, however until an exact understanding of OP generation is established only qualitative interpretations can infer disease. The frequency domain (Figure 2.3) analysis of the extracted OP waveform following application of a fast Fourier transform is advantageous because it provides a holistic examination of the OP waveform, particularly when signal-to-noise ratio is low.^{66,76} In addition, the frequency domain analysis is much more objective as it does not require the subjective identification of individual OPs which may become troublesome when signal response is low or noise is high.⁸⁰ Since both time- and frequency-domain analysis provide distinct information about OP parameters and general health of the retina, they were both examined in our investigation to identify an early diagnostic feature of retinal degeneration.

1.10 Nonmetric multidimensional scaling as a method for understanding OP waveform changes

In general, the mammalian ERG contains five OP wavelets existing on the leading edge of the b-wave, each with its own characteristic period (latency) and amplitude.⁵⁴ OP profiles depend on the intensity of the light stimulus, with increasing intensity leading to increased OP amplitude and decreased latency until retinal phototransduction saturation occurs.^{66,74,78} Examination of the OPs occurs following a single high-intensity flash and subsequent comparisons of latencies and amplitudes of each OP between a disease model and a healthy control. In addition, summed OP amplitude is useful for understanding

whole retinal response in the time domain.^{76,78} Examination of OPs at a single flashintensity has limitations since it may not be useful for understanding complete waveform changes across multiple stimuli intensities where disease mechanisms may be evident in retinal neurons only activated under low light intensity. The frequency domain provides a much more holistic approach to whole OP waveform interpretation; however, similar to the time domain, this approach can only be examined at a single flash intensity to examine differences in frequency, power, and energy between a disease model and a healthy control.^{66,80,81}

To understand whole OP waveform changes, a comprehensive nonmetric multidimensional scaling (NMDS) of OP parameters across multiple flash intensities can be applied. NMDS is a monotonic, numeric, ordination technique that involves spatial representation of observed variables to fitted distances.^{82–84} The method involves the assignment of derived coordinates for every sample and mandates that rank order may not be changed but observed proximities can be adjusted.^{82–84} In this way, many variables can be examined for each sample and modelled into a previously selected explicit number of axes, provided that the original rank order is not changed.⁸² Stress is a measure of how well the ordination fits the observed distances among samples and the error associated with modelling.⁸⁵ Stress increases with the number of samples and the number of variables fit into the ordination.

1.11 Mitochondrial dysfunction is a common theme in retinal degeneration

The mitochondrion is the organelle responsible for oxidative phosphorylation (OXPHOS). In highly metabolic tissues, the density of mitochondria tends to be much higher to keep up with the energetic demands of the cell.⁸⁶ The visual pathway is amongst the most energetically demanding processes in the body, as it requires continual movement of ions across membranes for maintenance of resting potentials in addition to the continual recycling of neurotransmitters.^{87,40} Due to the persistent energetic demands of the retina, it has the highest relative oxygen consumption of any tissue in the body, necessary to facilitate OXPHOS.^{88,89} Therefore, adequate and consistent delivery of oxygen is vital to retinal homeostasis and any deviations can lead to mitochondrial dysfunction and subsequent oxidative stress. Mitochondrial dysfunction is a common theme in retinal degenerative diseases and has been implicated in AMD, OAG, and DR due to oxidative damage of photoreceptors.⁹⁰

1.12 The mouse is a valuable model organism for vision research

The use of animal models continues to be vital as an ethical means to investigate fundamental biological principles. Mouse models are amongst the most common due to their short life cycle, quick sexual maturity, large litter size, and similar physiological and genetic composition to humans.⁹¹ In addition, isogenic, inbred strains reduce variation amongst individuals and allow complex diseases to be studied with smaller sample sizes while achieving higher power to detect significant differnces.¹¹ Many similarities exist between the human and mouse eye; including a large cornea and lens' as well as

analogous retinal nerve layer physiologies and cell types.⁹² The mouse retina is quite similar to the human retina with only minor differences in the visual pathway. For example, mice have a higher proportion of rod photoreceptors (over 97%)⁵⁸ which makes their retinas much more effective in scotopic light. Another major difference is the presence of a macula in the human retina. The macula is a dense, cone-rich area in the central region of the retina responsible for central vision acuity.⁷ In mice this is replaced with a central region of highly concentrated rod photoreceptors.⁵⁸ While humans possess three types of cone photoreceptors, mice only have two; green- and blue-sensitive.⁷ This property, although a major difference, assists researchers because mouse "dark-adapted" experiments can be performed in red-light. While subtle differences do exist between the mouse and human retina, common physiologies, genetics and cell types as well as the progression of similar diseases make mice an excellent mimic for human retinal disease mechanisms and subsequently vision research.⁹²

1.13 The harlequin (hq) mouse is a model of mitochondrial dysfunction and neurodegenerative disease

The *harlequin* (*hq*) mouse $(X^{hq}X^{hq}, X^{hq}Y \text{ or } X^{hq}X)$ is a model of mitochondrial dysfunction associated with neurodegenerative disease of the retina and cerebellum.⁹³ The term "*hq*" refers to the *hqY* genotype unless otherwise specified. Characteristic markers of the *hq* phenotype include; low body mass, a patchy coat, ataxia, low subcutaneous fat reserves as well as characteristic retinal and cerebellar degeneration.⁹³ These phenotypic characteristics are the result of a spontaneous proviral insertion into

intron 1 of the X-linked *Apoptosis-inducing factor* (*Aif*) gene which encodes a mitochondrial flavin adenine dinucleotide-dependent oxidoreductase.⁹⁴ Since the proviral insertion is in a suspected regulatory region of intron 1, there is an overall global downregulation of AIF by nearly 80% in most tissues.⁹³

As described previously, *AIF* is located on the human X-chromosome in the Xq25-26 region.⁹⁴ The gene is syntenic in the mouse and is also located on the X-chromosome in the A6 region.⁹⁴ The AIF amino acid sequence is highly conserved in both organisms, sharing 92% sequence homology of the active domain as well as an amino-terminal mitochondrial-localization sequence.⁹⁴ Upon translocation to the intermembraneous space of the mitochondrion, the localization sequence is cleaved leaving a functional 57 kDa flavoprotein.⁹⁴ AIF is ubiquitously expressed across most tissues in mice⁹⁵ with demonstrated embryonic lethality at day 9 in a complete *Aif* mouse knockout.⁹⁶ This indicates AIF's vital role in development as well as cell life and death.

The function of AIF was first characterized as a mitochondrial bound caspaseindependent death effector protein that translocates to the nucleus and aids with chromatin condensation and DNA fragmentation when apoptosis is induced.^{94,97} However, recent investigations have led to the description of multiple functions of AIF. AIF's primary role *in vivo* may be linked to OXPHOS efficiency, particularly that of NADH dehydrogenase (complex I). *Aif*-knockdowns have displayed limited formation of complex I *in vivo* indicating its role in maintenance and facilitation of complex I formation.⁹⁸ *hq* mice have an overall decrease in mitochondrial complex I efficiency, particularly that of the retina (~30% of normal function) and brain (~60% of normal function).⁹⁵ Although *Aif* causes a decrease in complex I function, no changes in the transcriptome or proteome indicate dysfunction of complex I subunits posttranslationally.⁹⁸ Further analysis of AIF indicates that it shares sequence similarities⁹⁴ and similar crystal structure^{99,100} with bacterial oxidoreductases and may function as a mitochondrial scavenger of free-radicals including ROS. *hq* mice demonstrate elevated oxidative stress in dying neurons of the retina and cerebellum.⁹³ However, this function of AIF remains controversial because mitochondria isolated from brain of *hq* mice show no traces of increased ROS production as a result of *Aif* deficiency.¹⁰¹

1.14 Timeline of the hq disease progression

Since *Aif* is mapped to the X-chromosome, affected *hq* males are hemizygous for the proviral insertion. Females can either be homozygous or heterozygous for the proviral insertion. Homozygous females demonstrate a severe, early-onset disease phenotype, similar to that of hemizygous males.⁹³ Heterozygous females however, show delayed onset, moderate disease progression and moderate disease phenotype,^{102,103} likely due to random X-inactivation.

The progression of the hq disease (homozygous and hemizygous) is well characterized in both the retina and the cerebellum. The hq disease is an ageingassociated retinal degeneration with primary structural losses occurring as early as three months of age in amacrine and ganglion cells.⁹³ The progression of the hq disease is quite rapid with histological analysis indicating significant losses in the INL and ONL by four months of age.^{19,93} Examination of the IPL and OPL in the hq retina indicate losses of neurites, particularly that of dendrites.⁹³ Functional deficits precede structural losses in the hq retina with electroretinography showing rapid functional deficits in ERG a-wave amplitudes at three months and b-wave at two months of age.¹⁹ By ten months of age, hqmice elicit no functional response of the retina.⁹³ Physiological changes begin as early as 1.8 months in the hq retina, consistent with a loss of homeostasis and activation of microglial cells in the IPL¹⁰⁴. This is indicative of an inflammatory response and may contribute to the early structural and functional losses observed. Chronic activation of microglial cells continues through hq disease progression and leads to microglial migration into the ONL and GCL by four months of age, consistent with cell losses observed previously.^{93,104}

The degeneration of the hq cerebellum occurs with a much slower progression in comparison to the retina.¹⁰² Early histological analysis indicates apoptosis in the hq cerebellum as early as four months of age in granule cell nuclei leading to smaller total cerebella by seven months.^{93,105} Loss of over 50% of granule cells was observed by twelve months of age.⁹³ Purkinje cell loss in the hq cerebellum occurs later in comparison to granule cell loss initiating at seven months of age.^{93,105} Characteristic markers of microglia activation in the hq cerebellum have yet to be tracked *in situ*.

1.15 Deficiencies in OXPHOS lead to neuronal malfunction and ATP deficiencies

It is well established that OXPHOS is the major source of ATP for all cells, in particular neuronal cells of the brain and retina. Energetic failure of OXPHOS in neuronal cells has been linked with Alzheimer's disease,¹⁰⁶ Parkinson's disease,¹⁰⁷ Huntington's disease,¹⁰⁸ amyotrophic lateral sclerosis¹⁰⁹ and other neurodegenerative disorders. Neuronal cells maintain consistent internal ATP concentrations and minimal energy gradients by axonally transporting mitochondria via microtubules to regions of high energy consumption.¹¹⁰ The presynaptic compartment of the neuron, responsible for neurotransmitter release, is the region of highest energy consumption.^{110,111} High amounts of ATP are required for continuous neurotransmitter recycling as well as ion pumping to maintain membrane potentials.¹¹¹ Deficiencies in OXPHOS particularly affect the presynaptic cleft and other highly metabolic regions leading to neurite loss and synaptic degeneration.^{111,112}

In addition to being "the powerhouse of the cell," the mitochondrion is also an intracellular regulator of calcium. Mitochondrial dysfunction as a result of OXPHOS deficiency causes mitochondrial overload of Ca^{2+} , leading to opening of the mitochondrial permeability transition pore and Ca^{2+} release.^{113,114} This furthers induces ATP deficiency, successive glycolytic ATP dependency, reduced mitochondrial mobility, release of ROS and subsequent activation of the caspase execution cascade leading to apoptosis and cell death in the tissue.^{114–116}

1.16 OXPHOS deficiencies in the hq mouse

The variability associated with the hq disease phenotype is large, with timing of disease onset and progression occurring differently across individuals.⁹⁵ Additionally, the effect of the *Aif* proviral insertion is also not equivalent across tissues of the *hq* mouse. Western blotting reveals that AIF expression in *hq* mice, in comparison to controls, was approximately 20-40% in the cerebellum, spinal cord, cortex, skeletal muscle, and

kidney.⁹⁵ AIF reaches almost 60% expression levels in the liver, with minimal to nil expression in the retina.⁹⁵

In association with decreased AIF expression, compromised OXPHOS efficiency was found in all regions of the brain and retina however no OXPHOS deficiency was evident in the heart or the liver.⁹⁸ A complete knockout of *Aif* in the mouse heart does however lead to reduced OXPHOS activity indicating the buffering threshold of the *Aif* knockout may be organ specific.¹¹⁷ Tissue specific *Aif* knockout for liver, muscle and heart leads to increased glucose uptake (via *glucose transporter 1* and *4*) as well as a reduction in overall ATP levels and high NADH/NAD+ ratios and lactate production. This suggests the switch towards glycolytic metabolism and generation of anaerobic ATP.^{117,118} In addition, *hq* mice demonstrate similar characteristics of increased insulin sensitivity and glucose uptake/tolerance with reduced predisposition to obesity on a high fat diet.¹¹⁸

The complete *Aif* knockout in embryonic stem cells identifies consistent consequences leading to high levels of lactate production.⁹⁸ The addition of 2-D-deoxyglucose, an inhibitor of glycolysis, increases apoptosis in these *Aif*-deficient embryonic stem cells and indicates high dependency on glycolytic ATP and glucose as a primary metabolite.⁹⁸ Taken together, this thorough examination of AIF dysfunction indicates a metabolic change towards anaerobic and lactate ATP production in *Aif*-deficient cells particularly in tissues of mesodermal and ectodermal origin.

The knockout or knockdown of *Aif* predisposes embryonic stem cells, muscle, liver and heart cells to mitochondrial OXPHOS deficiency and reduces overall ATP

generation. Aif-deficient cells must compensate and switch their metabolic processes towards a glycolytic pathway of ATP production and promote high glucose uptake and metabolism. Glycolysis is the conversion of a glucose molecule into pyruvate in a 1:2 ratio with the reduction of two NAD⁺ molecules to NADH and the net overall production of two ATP molecules. The switch to glycolysis allows for ATP production when OXPHOS is an inefficient source of energy production and provides the necessary energy for metabolic need. Unlike other cell types and tissues, glycolysis is intrinsically repressed in neurons and the central nervous system to maintain low oxidative stress.^{45,119} Therefore they do not have the ability to increase glycolysis when OXPHOS integrity is interrupted.⁴⁵ PFKFB3 enzymatically produces fructose-2,6-bisphosphate (a strong glycolytic activator) and is down-regulated in neurons. Its overexpression leads to oxidative stress and cell death *in vivo*.⁴⁵ Since neurons of the central nervous system rely almost completely on OXPHOS-generated ATP, they are highly vulnerable to decreases in ATP generation.¹²⁰ In the hq mice, this is evident due to the severe and irreversible onset of retinal and cerebellar degeneration.

1.17 Parainflammation is a chronic response in the hq retina

Inflammation is the biological response to maintain homeostasis following a negative insult *in vivo*. Inflammation is commonly the product of a tissue injury or infection¹²¹ and solicits the homeostatic response of the innate immune system and the adaptive immune system. The recognition of pathogens by the innate immune system begins with the specific binding of the pathogen-associated molecular patterns (PAMPs) expressed by microbes via Pattern Recognition Receptors (PRP)¹²². Following binding,

immune cells (macrophages) are recruited and the pathogen is destroyed and removed. The adaptive immune response is triggered for the selection of an antibody to better target the pathogen during future insult. In the event of a tissue injury, the innate immune system must respond to cellular damage of an endogenous injured cell and therefore PRPs are not present. The response to these injuries is facilitated by endogenous molecules known as alarmins that are released during non-programmed cell death and recruit the innate immune response.^{123,124} One such example is that of architectural chromatin protein, High Mobility Group Box 1 (HMGB1).¹²⁵ Under apoptotic conditions HMGB1 binds irreversibility to nucleosomes and elicits no immune response, however under necrotic conditions HMGB1 is released into the extracellular space and becomes a potent activator of inflammation.¹²⁵

In addition, alternative endogenous mechanisms beyond necrosis can induce inflammation. The formation of advanced glycation end-products (AGEs) begins under hyperglycemic or oxidative conditions and catalyzes the interaction of glucose with protein amino acids and lipids.¹²⁶ Once formed, they bind with the receptor for advanced glycation end-products (RAGEs) which can induce a pro-inflammatory response through the secretion of Tumor necrosis factor α (TNF- α) and stimulation of T-cells.¹²⁶ Under oxidative conditions; proteins, lipids and DNA can be oxidized. Lipids are oxidized into highly reactive aldehydes which in turn induce macrophage secretion of TNF- α and elicit an inflammatory response.¹²⁷ Finally, inflammatory cytokines released by activated macrophages can further induce inflammatory responses. Chronic cytokine release by macrophages causes further tissue damage and the potential breakdown of blood

barriers.¹²¹ Inflammatory cytokines can alter the functionality of the endothelial/epithelial tight junctions leading to barrier degradation.¹²⁸

Microglial cells are the resident macrophages of the retina and cerebellum that are responsible for the innate immune response during an inflammatory stimulus.¹² Resting. dormant microglia cells generally take on a ramified cellular morphology with many protrusions as a means to monitor their microenvironment as well as clearing any metabolic waste and tissue debris.¹²⁹ Microglial cells become activated during neuronal injury, stress and infection changing their cell morphology to an amoeboid macrophage. The surface membrane, G-coupled purinergic membrane receptor P2Y₁₂ can detect small changes of ADP and ATP concentrations in the extracellular space released by stressed or necrotic neurons and prompts microglial activation.¹³⁰ Activated microglia become highly mobile and exhibit strong phagocytic ability and further induce the release of cytokines to enhance microglia recruitment to mediate tissue repair.¹²¹ During healthy conditions, this is generally a low-grade tissue adaptive response known as basal inflammation. In contrast, chronic activation of microglia cells as a result to neuronal stress can lead to a state of parainflammation. Parainflammation is an intermediate response between overt-inflammation and non-inflammatory removal of apoptotic cells.¹²¹ Chronic microglial activation is associated with the release of tissue damaging ROS, nitric oxide, proteases, and further cytokines causing exaggerated recruitment of microglial cells and retinal degeneration.¹³¹ Parainflammation and chronic microglial activation have previously been observed in the hq retina as early as 1.8 months of age as a result of AIF downregulation. Chronic microglial activation may be the cause of subsequent structural losses in the hq retina. The exact triggers leading to chronic

microglial activation has yet to be defined and may be the result of a continuous endogenous stress (tissue stress) or the breakdown of a retinal barrier allowing chronic influx of exogenous pathogens into the sensitive retinal space (Figure 1.3).

1.18 Central hypothesis

I hypothesize that since the *hq* mouse is hypomorphic for AIF function in oxidative phosphorylation, health of the inner retina is compromised without loss of blood-retinal-barrier integrity and evidence of compensatory ATP production mechanisms exist.

1.19 Experimental Aims

The first experimental aim is to determine if electroretinogram oscillatory potentials can infer early mechanisms of retinal degeneration, particularly operational in the inner retina. The use of OPs as a non-invasive diagnostic feature could infer physiological changes as well as the health of the inner retina prior to any cell losses. The rapid onset of the hemizygous hqY disease leads to functional losses as well as later structural losses. OPs have been used to track disease progression prior to any symptomatic functional losses in a variety of retinal diseases as OP features infer the health of various neural networks in the retina. It is therefore predicted that OPs can be used to diagnose early functional changes in the hq retina prior to any structural losses. To track and examine disease progression, OP from hq mice were analyzed monthly from two months of age to ten months of age. This provides long term disease progression

information and functional retinal documentation at a later age. To complement the examination of OPs in the severe and rapid disease phenotype, the investigation of the heterozygote phenotype can provide valuable information regarding disease progression. Female heterozygotes demonstrate a longer timeframe to observe disease mechanisms prior to cell death and were examined at three months, eleven months and fifteen months of age.

The second experimental aim is to assess RPE (outer BRB) and inner retina endothelial vasculature barrier (inner BRB) integrity to ascertain the potential for exogenous triggers of microglial activation and subsequent retinal degeneration (Figure 1.3). This examination is instrumental in determining if the metabolic deficiency associated with hq neurons compromises the functionality of the highly metabolic RPE and inner retina endothelial vasculature barrier. Metabolic deficiency in the hq retina may result in the degradation of tight junctions and lead to vascular endothelial/epithelial leakage into the sensitive inner retinal space and thereby elicit microglial activation. Alternatively, it can be hypothesized that microglial activation in the hq retina is the consequence of endogenous photoreceptor and inner retinal neuron cell stress due to ATP deficiency triggering chronic microglial activation. To examine the integrity of the BRB, injections of Evans Blue followed by a complete vascular perfusion assesses diffusion of albumin-bound Evans Blue into the inner retina and potential breakdown of the BRB. Two age groups of mice were examined for both wild-type and hq mice: a younger age group (< 4 month) prior to ONL losses, and an older age group (> 7 months) corresponding to chronic microglial activation at 1.8 months. It is hypothesized

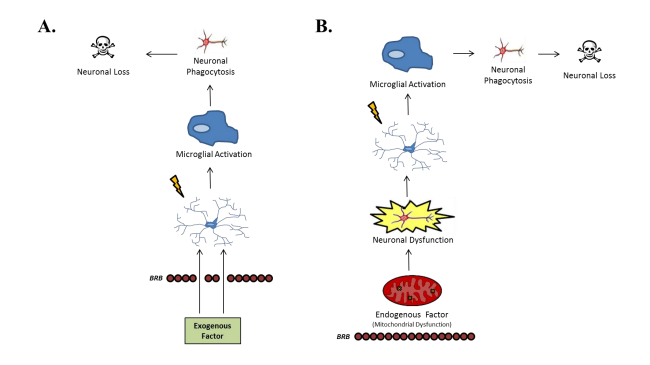


Figure 1.3. Activation of retinal microglia may be the result of blood-retinal-barrier breakdown (exogenous) or inner retinal dysfunction (endogenous). (A) Microglia activation (ramified to amoeboid) may be a consequence of BRB breakdown causing diffusion of harmful exogenous antigens into the sensitive retinal space. Highly metabolic epithelial and endothelial cells of the BRB may malfunction due to intracellular ATP deficiency. (B) Alternatively, microglia activation and ATP deficiency leading to cell stress, apoptotic and necrotic conditions thereby inducing parainflammation and chronic microglia activation.

that microglial activation in the retina is the result of endogenous ATP deficiency leading to cellular stress and apoptotic neurons rather than a compromised BRB inducing microglial activation.

The third and final aim is to confirm retinal transcriptome changes consistent with mitochondrial dysfunction and subsequent metabolic disease during the onset and progression of hq retinal and cerebellar degeneration. The downregulation and reduced expression of AIF leads to alternative metabolism in mesodermal tissues. It was predicted that similar compensating mechanisms were expected to transpire in the hq retina and cerebellum. Although glycolysis is unlikely to be upregulated in hq neurons, alternative pathways of ATP production such as lactate production, glycerol metabolism and glucose uptake are predicted to be upregulated. To confirm transcriptome changes consistent with alternative metabolism three ages were examined corresponding to disease onset and progression (two, three and four months of age). Since the cerebellum is not as ATP demanding as the retina⁴⁰ and histological changes occur later in the hq cerebellum it is expected that metabolic changes in the hq mouse are not as evident by four months of age.

Chapter 2. Materials and Methods

2.1 Animal care and housing

The Canadian Council on Animal Care and The Animal Use Subcommittee of The University Council on Animal Care approved all protocols prior to the commencement of any experimental work (Appendix A). Mice (B6:CBACaA^{w-J}/A-Pcdc8^{hq/J}, 97% CBA/CaJ genetic background) [Jackson Laboratories, Bar Harbor, ME] were housed at a constant temperature of $21 \pm 1^{\circ}$ C with a relative humidity of 44% to 60% and a light/dark cycle of 14/10 hours. The mice received a standard diet [PMI Foods, St. Louis, MO] and water *ad libitum*.

2.2 Genotyping of the hq allele

A tri-primer polymerase chain reaction (PCR) was used to determine the presence of the *Aif hq* allele.⁹³ Tissue samples for genotyping were taken as ear-notches when mice were approximately 12 days old. DNA was amplified using the TerraTM PCR Direct [Clontech, Mountain View, CA] standard protocol. Amplified DNA was subjected to gel electrophoresis through a 1.5% agarose gel stained with SYBR[®] Safe DNA Gel Stain [Life Technologies Inc., Burlington, ON]. The presence of a 725 bp amplicon indicated the *Aif hq* disease allele and the presence of a 537 bp amplicon indicated a wild-type *Aif* allele. Female heterozygous mice show both a 537 bp and a 725 bp amplicon. PCR primer sequences used for *Aif* genotyping are provided in Appendix B (Table B.1). Reference to the *hq* phenotype refers to the *hq*Y genotype unless otherwise specified. Wild-type (WT) mice refer to male (XY) mice with a functional *Aif*.

2.3 Collection of trace electroretinograms

Repeated electroretinogram (ERG) analyses were performed on cohorts of wildtype mice and hqY mice at two, three, four, five to seven and eight to ten months of age to assess retinal function and severe disease progression (Figure 2.1A,B).¹³² Additionally, ERG analyses were performed on female carrier mice (hqX) along with an age-matched wild-type cohort at three, eleven and fifteen months to assess moderate disease progression of the hq carrier disease (Figure 2.1B).^{102,103} Electroretinography was performed with a Colordome Stimulator [Diagnosys, Lowell, MA] on both dark-adapted (scotopic) eyes through a series of eleven flashes of increasing intensities (0.001 – 25 cd•m/s²).

2.4 Processing of trace electroretinogram for waveform modelling

Original exported ERG trace files contained only data corresponding to parameters of the a-wave and b-wave components. A thorough review of the ERG user manual was performed and allowed for export of all raw data corresponding oscillatory potential components. A macro written into Excel [Microsoft, Redmond, VA] known as "MLTRIM" automatically sorted the original trace file into averaged neuronal responses for each eye and presented them as μV across all flash intensities.¹³³ In addition, MLTRIM was used to convert ERG trace data into a format readable by MATLAB [The MathWorks, Natick, MA].

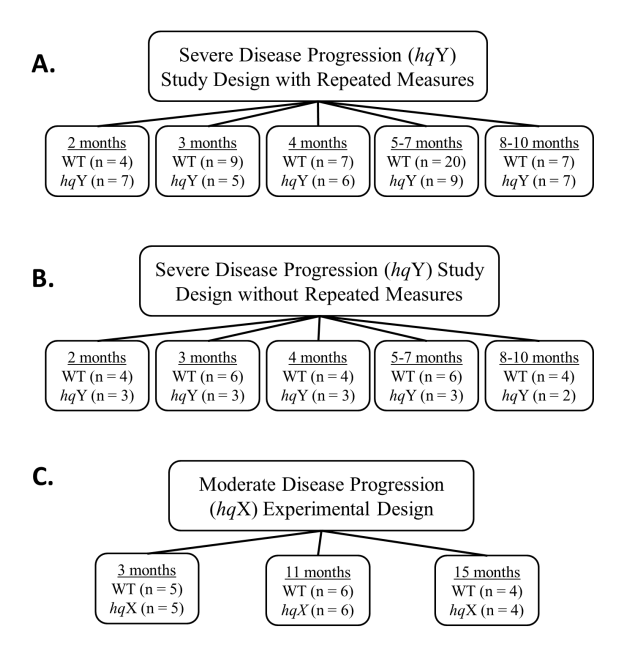


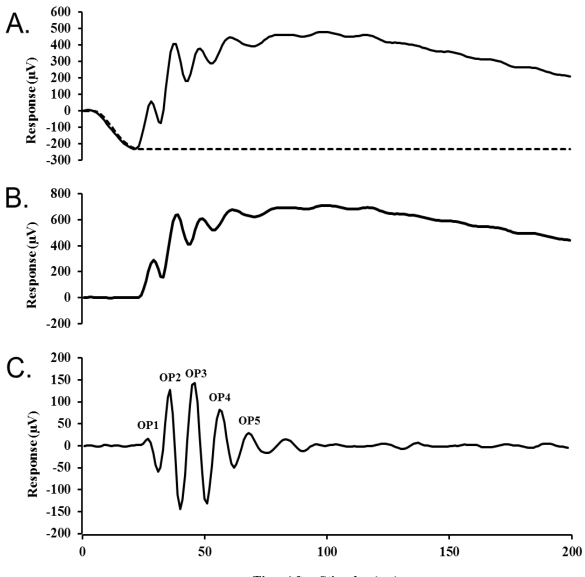
Figure 2.1. Experimental design and cohort sizes for oscillatory potential analysis. (A) The complete experimental design for the severe disease progression hqY analysis.¹³² A single mouse may appear multiple times due to repeated measures at multiple ages. (B) Repeated measures of the severe disease progression experiment were removed randomly to allow study of hqY mice without repetition. (C) The complete experimental design for the moderate disease progression hqX experiment.¹⁰³ No repeated measures were present in this study.

2.5 Signal conditioning and oscillatory potential waveform extraction

Scripts (Appendix C, MATLAB Script A,B) were written into MATLAB [The MathWorks, Natick, MA] to model and extract relevant oscillatory potential waveforms (Figure 2.2).¹³³ To limit contribution of photoreceptor hyperpolarization, the a-wave was digitally modelled and subtracted from the original trace (Figure 2.2A).^{66,68,69,71-74} Additional high frequency and low frequency noise were removed by processing the waveform through a fifth-order Butterworth filter with a band pass of 65-300 Hz (Figure 2.2C).^{69,70} Previous literature suggests a lower band pass limit of 40 Hz for characterization of mouse rod photoreceptor oscillatory potentials, however 65 Hz was chosen to avoid interference by the 60 Hz line noise as previously described.^{69,70} The remaining bandwidth made up the final OP waveform, herein known as the "extracted waveform".

2.6 Parameter analysis of the extracted waveform

The extracted waveform was first analyzed in the time-domain. The amplitude (μV) , initial latency (ms) and interpeak distance were determined for OP2, OP3, OP4, and OP5 (Figure 2.3A). Following waveform extraction, OP1 was typically present, however due to inconsistencies OP1 was not included in further analysis. Amplitude of each OP was defined as the difference between the trough and the peak immediately preceding it (Figure 2.3A). Initial latency was defined as the time from stimulus to the onset of OP2 (Figure 2.3A). Interpeak distance was the measurement of time (ms)



Time After Stimulus (ms)

Figure 2.2 Signal conditioning of oscillatory potentials from the ERG trace waveform. (**A**) A digital subtraction of photoreceptor contribution (dashed line) was performed by a mathematical fitting and subtraction of the a-wave from the initial ERG (solid line). (**B**) The ERG waveform following complete a-wave digital subtraction. (**C**) The final waveform was passed through a fifth order Butterworth transformation (65-300 Hz) to remove any low or high frequency noise resulting in the final "OP extracted waveform" (solid line).

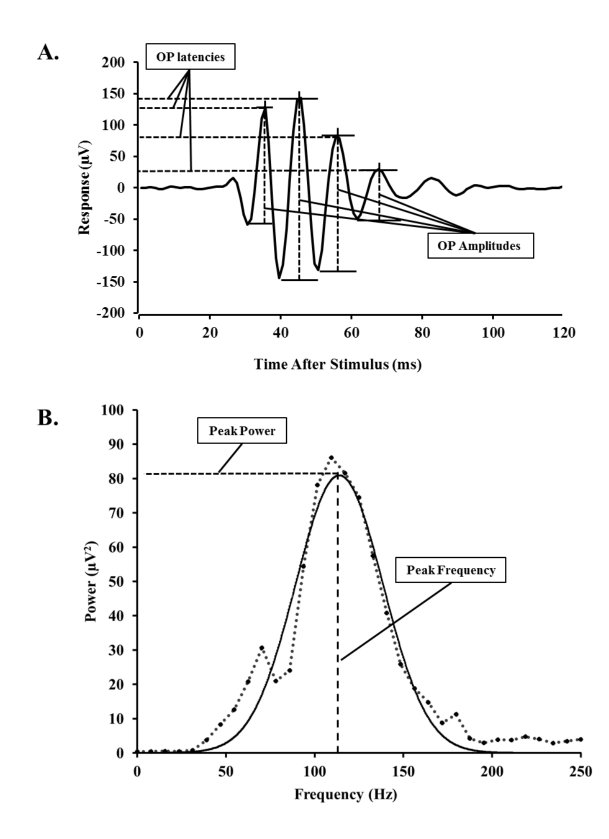


Figure 2.3. Measurement of individual Oscillatory Potential features in the timedomain and frequency-domain. (A) Following waveform extraction, measurements of OP latency (ms) and amplitude (μ V) were taken for OP2, OP3, OP4, and OP5. Amplitude of each OP was defined as the difference between the trough and the peak immediately preceding it. Initial latency was defined as the time from stimulus to the onset of the OP2 peak. Interpeak distance was the measurement of time (ms) between adjacent peaks. OP1 was not measured due to inconsistencies and the potential for awave contamination. (B) A fast Fourier transform (FFT) was applied to the extracted OP waveform and gave a single-sided smoothed frequency power spectrum (dotted Line). The single-sided smoothed frequency power spectrum was fitted to a two-term Gaussian envelope (solid line) and measurements of peak power (P_{peak}), peak frequency (F_{peak}) and total energy (waveform integration) were taken. between adjacent peaks (i.e. OP2 and OP3) (Figure 2.3A). Total OP response in the time domain was measured as the summation of OP2-OP5 amplitudes.

2.7 Fast Fourier Transformation (FFT) of the extracted waveform

The frequency spectrum of the extracted waveform was also analyzed by a Fast Fourier Transformation (FFT) to provide a holistic frequency waveform interpretation.⁵⁴ Scripts (Appendix C, MATLAB Script C) written in MATLAB [The MathWorks, Natick, MA] performed this operation and converted the frequency-power spectra to a single-sided smoothed frequency power spectrum. To avoid "powers of two" problems, only the first 128 samples of the extracted waveform (contained entire OP waveform) were applied to the FFT.⁶⁶ To calibrate the output of the FFT, a sine wave of known periodicity and amplitude was subjected to the algorithm. These produced a single peak indicative of a properly calibrated FFT algorithm. To avoid the problem known as 'overspill' the FFT output was fit to a two-term Gaussian envelope with the following equation: ^{66,75}

$$P(F) = (Ppeak) \left[-\frac{1}{2} \left(\frac{F - Fpeak}{S} \right)^2 \right]$$

where P is the power associated with the frequency response (F, Hz). F_{peak} is the frequency at the peak of the Gaussian envelope, and P_{peak} is the corresponding power (Figure 2.3B). Major frequency (F_{peak}) was determined as the dominant frequency of the Gaussian frequency-power spectrum and is related to OP periodicity. S indicates the width of the time distribution. The area under the original frequency power spectrum (Total energy) was calculated by integration of the original frequency-power spectrum (Figure 2.3B).

2.8 Non-Metric Multidimensional Scaling (NMDS) of oscillatory potential parameters

The longitudinal nature of the *hq*Y study does not allow simple statistical testing (Figure 2.1). Approximately five mice were euthanized at the end of each age cohort age (2, 3, 4, 5-7, 8-10 respectively) and the attrition associated does not allow the application of a simple repeated-measures two-way ANOVA. In addition, OPs contain a wealth of information across multiple flash intensities and a comprehensive analysis was pursued as described in the introduction. A clustering method known as Non-metric Multidimensional Scaling (NMDS) was used in R Statistical Software [R Core Team, Vienna, Austria] (Appendix D, NMDS R-code A) to model every variable from each respective OP peak at all four flash intensities (Flash 8-11). This resulted in a total of 32 variables being analyzed in the time domain and 12 variables in the frequency domain (Figure 2.4). Each variable was first converted into a z-score using the following equation:

$$z - scores = \frac{x - \mu}{\sigma}$$

where x is each value in the dataset, μ is the mean of the dataset and σ is the standard deviation of the dataset. The Euclidean distance between all mice was calculated encompassing all variables and a dissimilarity matrix was computed to demonstrate the absolute distances between mice.⁸² A NMDS was then applied to model data in a twodimensional ordination plot such that rank order matches the order of dissimilarity between mice. Stress is indicative of the best rank order of agreement between the dissimilarities as well as fitting the data to a monotonic regression (Shepard plot). Lowest

Α.

Time-Domain Measurements Taken				
Flash 8 (0.63 cd•s/m ²)	Flash 9 (4 cd•s/m²)	Flash 10 (10 cd•s/m²)	Flash 11 (25 cd•s/m²)	
Latency OP2 (ms)	Latency OP2 (ms)	Latency OP2 (ms)	Latency OP2 (ms)	
Amplitude OP2 (µV)	Amplitude OP2 (μV)	Amplitude OP2(µV)	Amplitude OP2 (μV)	
Latency OP3 (ms)	Latency OP3 (ms)	Latency OP3 (ms)	Latency OP3 (ms)	
Amplitude OP3 (µV)	Amplitude OP3 (μV)	Amplitude OP3 (µV)	Amplitude OP3 (µV)	
Latency OP4 (ms)	Latency OP4 (ms)	Latency OP4 (ms)	Latency OP4 (ms)	
Amplitude OP4 (µV)	Amplitude OP4 (µV)	Amplitude OP4 (µV)	Amplitude OP4 (μV)	
Latency OP5 (ms)	Latency OP5 (ms)	Latency OP5 (ms)	Latency OP5 (ms)	
Amplitude OP5 (μV)	Amplitude OP5 (µV)	Amplitude OP5 (μV)	Amplitude OP5 (μV)	
8 Meaurements	8 Meaurements	8 Meaurements	8 Meaurements	
32 Total Measurements Per Mouse				

Β.

Frequency-Domain Measurements Taken				
Flash 8 (0.63 cd•s/m ²)	Flash 9 (4 cd•s/m ²)	Flash 10 (10 cd•s/m ²)	Flash 11 (25 cd•s/m ²)	
Gaussian Peak	Gaussian Peak	Gaussian Peak	Gaussian Peak	
Frequency (Hz)	Frequency (Hz)	Frequency (Hz)	Frequency (Hz)	
Gaussian Power (µV ²)	Gaussian Power (µV ²)	Gaussian Power (µV ²)	Gaussian Power (µV²)	
Total Energy (μV ² •S)	Total Energy (µV ² •S)	Total Energy (μV ² •S)	Total Energy (μV ² •S)	
3 Measurements	3 Measurements	3 Measurements	3 Measurements	
12 Total Measurements Per Mouse				

Figure 2.4. Total number of measurements taken from a single mouse across four flash intensities. (A) A total of 32 parameters are measured in the time-domain, eight measurements for each flash intensity. (B) A total of 12 parameters are measured in the frequency-domain, three for each flash intensity.

stress was found at the global optima following 100 iterations. Stress below 20% is deemed fair.⁸² For NMDS axis interpretation a Pearson's correlation coefficient was used to interpret the axis in relation to various time domain and frequency domain parameters.

2.9 Experimental design for Blood-Retinal-Barrier (BRB) integrity phenotyping

Male hqY mice and age-matched wild-type controls were selected at two age cohorts of two-to-four months of age and seven months of age. The number of mice in each cohort was between three and five individuals. Evans Blue dye [Sigma Aldrich, St. Louis, MO] dissolved in 1X Phosphate Buffered Saline (PBS) (50µL) [Santa Cruz Biotechnology, Dallas, TX] was intravenously injected over 15 seconds through the tail vein at a concentration of 45 mg/kg.^{134,135} Following injection, the dye was allowed to circulate through the vasculature for a period of 30 minutes. During this period the mouse turned visibly blue particularly in the extremities, confirming the uptake and distribution of the dye.¹³⁶ The mouse was then euthanized via CO₂ inhalation and the chest cavity was opened. A transcardial perfusion with 60 mL 1X PBS was performed until all blood was removed from the vasculature. Tissue was stored at -80°C until further processing.

2.10 Generation of a standard curve to estimate Evans Blue tissue concentrations

A standard curve was generated to calculate concentrations of EB in respective tissues. Serial dilutions of EB in formamide were made from 10000 ng/mL to 250 ng/mL and background-subtracted absorbances (620 nm - 740 nm) measurements were made in triplicate. Standards were prepared on different days by three different individuals.

2.11 Evans Blue Extravasation

Immediately following PBS perfusion, eyes were enucleated and the retinas were carefully microdissected away and rinsed with PBS. Liver, cerebellum and cerebrum tissue were also harvested and rinsed with PBS. Tissues were flash frozen in liquid nitrogen and stored at -80°C until further processing.

The wet weight of tissues was determined and they were then homogenized using a blunt homogenizer. The approximate weight of a mouse retina was 1 milligram and therefore approximately 1 mg of liver tissue was also processed as a technical control. EB was extracted from liver and retinal tissue by incubating in 100 µL of formamide [Sigma Aldrich, St. Louis, MO] for 24 hours at 75°C.¹³⁴ Cellular debris was removed by centrifugation at 4000 rpm for 45 minutes. Approximately 1 µL of supernatant was used for triplicate spectrophotometric measurements using the NanoDrop ND-1000 Spectrophotometer [Thermo Scientific, Wilmington, DE]. A background-subtracted absorbance was determined with measurements at 620 nm and 740 nm, known absorbance maximum and minimums for EB dye.¹³⁴ The concentration of the dye in the extracts was determined using the standard curve and was normalized to liver concentrations from the same animal to account for technical variation such as albumin dye uptake and injection efficiencies. Statistical significance was determined using a two-way ANOVA.

2.12 Examining daily food consumption in the hq mouse

Four-month-old hqY (n = 4) and age-matched, wild-type (n = 4) mice participated in a feeding study to determine average daily food consumption (grams of food consumed/ grams of body mass/ day) differences in hq mice. Food intake (mass) and mouse mass were measured every-other-day over a two-week period to determine daily food consumption. At the end of the two-week period, average daily food consumption for each mouse was determined and divided by the mouse mass. Statistical significance was determined by a repeated-measures ANOVA.

2.13 Microarray pathway analysis

To identify early triggers and mechanisms of retinal degeneration, gene expression microarray [Affymetrix, Mouse Gene 1.0 ST Array, Santa Clara] data were analyzed extensively. The microarray data were obtained by Alex Laliberté with 4-month-old hq and age-matched wild-type mice.¹⁰³ To determine early mechanisms of retinal degeneration selection criterion were set; at a significant change of p < 0.05 and a threshold fold change (mRNA transcript level) of at least ± 1.75. Canonical pathway analysis was performed using KEGG Pathway Analysis [Kyoto University, Kyoto, Japan] and Ingenuity Pathway Analysis [Ingenuity Systems Inc., Redwood City, CA]. A thorough literature review of *Aif* function complemented this pathway analysis and indicated a loss of *Aif* function leads to increased glucose metabolism, lactate metabolism, as well as glycerol and fatty acid metabolism.⁹⁸ Due to the specific nature of retinal degeneration markers and multiple genetic isoforms used in the retina, gene

ontology databases did not provide a thorough review of microarray data. Therefore grouping and annotation of differentially-expressed genes was performed manually and based on extensive literature survey on above pathways in NCBI [National Institute of Health, Bethesda, MD] (Appendix E). Multiple genes (>1.75 fold change, p < 0.05) from each above stated pathway were selected for confirmation of transcriptome changes via quantitative PCR (qPCR).

2.14 Tissue harvest for gene expression assays

Cohorts of hqY mice aged two-, three-, and four-months of age and age-matched wild-type mice (n = 5 per cohort) were euthanized via CO₂ inhalation according to Animal use Protocol (AUP) 2009-033 (Appendix A). Eyes were enucleated and retinas were microdissected and cleaned with 1X PBS in an RNase-free environment. Cerebellum, cerebrum and liver tissues were also harvested. Extracted retinas and cerebella were flash frozen in liquid nitrogen and stored at -80°C. All tools involved in microdissection were baked overnight at 200°C or treated with RNaseZAP [Sigma Aldrich, St. Louis, MI] to ensure an RNase-free environment and RNA integrity.

2.15 Retinal RNA extraction

Retinal samples (~1 mg) were homogenized and lysed using a sand homogenization protocol as previously described.¹³⁷ Ottawa Sand (20-30 mesh) [Fisher Scientific, Fair Lawn, NJ] was rinsed through multiple, serial washes of ddH₂O, 0.5M NaOH and 0.5M HCl and baked overnight at 200°C. Following homogenization, RNA was isolated using the Qiagen[®] RNeasy Mini Spin Columns Kit [Qiagen Canada Inc., Montreal, QC]. This included binding of total RNA to a silica membrane within the spincolumn. A series of subsequent RPE[®] and RW1[®] washes were performed to clean the RNA and increase efficacy of RNA binding to the silica column membrane. An oncolumn Qiagen[®] RNase-Free DNase [Qiagen Canada Inc., Montreal, QC] step was performed to remove genomic DNA (gDNA) contamination. RNA was resuspended in nuclease-free ddH₂O and stored at -80°C.

2.16 Cerebellar RNA extraction

Cerebellar samples (30 mg) were lysed with Qiazol[®] Lysis Buffer [Qiagen Canada Inc., Montreal, QC] and were homogenized by 20 strokes in a 7 mL Type A Glass Dounce Homogenizer [Wheaton Science Products, Millville, NJ]. Following homogenization, total RNA was isolated using the standard protocol for the Qiagen[®] RNeasy Mini Spin Column Kit [Qiagen Canada Inc., Montreal, QC] as described above. An additional on-column RNase-Free DNase step was added and RNA was resuspended in nuclease-free ddH₂O and stored at -80°C.

2.17 Assessment of RNA purity and quality

Retinal and cerebellar RNA quality was first assessed with a NanoDrop ND-1000 Spectrophotometer [Thermo Scientific, Wilmington, DE] to determine A260/A280 ratios (>1.9). Cerebellar samples were then subjected to gel electrophoresis and electrophoresed through a 1.5% agarose gel to determine the presence, integrity and ratios of the 18S and 28S rRNA subunits. This quality assurance stage was not performed with the retinal samples as the total amount of RNA isolated was considerably lower and not sufficient to perform gel electrophoresis. However, to ensure RNA purity, both cerebellar and retinal samples (1 μ L) were assessed with the Agilent 2100 Bioanalyzer [Agilent Technologies Inc., Palo Alto, CA] to determine RNA integrity (RNA Integrity Number - RIN > 8).

2.18 cDNA synthesis

Mouse retinal RNA samples were selected for single-stranded, complementary DNA (cDNA) synthesis from concentrations ranging from 15 ng/µL to 72 ng/µL. Mouse cerebellar samples were selected for cDNA synthesis from concentrations ranging from 84 ng/µL to 354 ng/µL. In each age cohort, the lowest concentration was selected as the rate-limiting sample, indicating the highest total RNA (µg) that could be added to the cDNA synthesis reaction based on the final volume of 16 µL. cDNA synthesis was performed using the Superscript[®] VILOTM cDNA Synthesis Kit [Life Technologies, Burlington, ON] and its associated standard protocol. A negative control did not contain any RNA and ddH₂O was used as a substitute. Following synthesis, cDNA was diluted to 100 µL to increase downstream throughput and stored at -20°C.

2.19 Taqman[®] gene expression experimental design

qPCR was performed using Taqman[®] probes [Life Technologies, Burlington, ON]. Probes that spanned exon junctions were selected to avoid the possibility of false amplification due to gDNA contamination. Genes were assayed in a block design for each age cohort (Appendix B, Figure B.1). Each gene of interest was assayed in three technical replicates including triplicate No-Template Controls (NTC). Biological replicates were assayed in parallel to avoid the possibility of additional technical variation (n = 4). For each reaction; 5µL of Tagman[®] Fast Advanced MasterMix [Life Technologies, Burlington, ON], 0.5 µL of Taqman[®] probes [Life Technologies, Burlington, ON], 2.5 µL of nuclease free water and 2 µL of cDNA were added to a final volume of 10 µL per reaction well. The assay was performed using the ViiATM 7 Real-Time PCR System [Life Technologies, Burlington, ON]. qPCR cycling conditions began with a 2 minute hold at 50°C for Uracil N Glycosylase incubation, a 20 second hold at 95°C for complete cDNA denaturation, followed by 40 cycles of 1 second at 95°C for denaturation and 20 seconds at 60°C for primer annealing and polymerase extension. Two reference genes, 18S Ribosomal RNA (Rn18s) and Glyceraldehyde-3-phosphate dehydrogenase (Gapdh) were selected as normalizer genes due to their documented high and consistent expression levels in all tissues.^{138,139} Rn18s was selected to ensure expression of *Gapdh* was consistently expressed between *hq* and wild-type mice. Upon confirmation of high, consistent uniform expression of *Gapdh* across genotypes, *Gapdh* was selected as the primary normalizer. Aif was assayed across all cohorts to ensure a downregulation in hq mouse retina and cerebellum.

2.20 Statistical analysis of Taqman[®] gene expression changes

The $\Delta\Delta$ Ct relative quantification method was used to determine fold-changes across genotypes.¹⁴⁰ Δ Ct values were determined as the difference between the measured average amplification cycle of the gene of interest and the average amplification cycle of *Gapdh*. Significance was determined with a one-way ANOVA in Excel [Microsoft, Redmond, VA] comparing Δ Ct values between *hq* and wild-type mice in each cohort. To calculate fold-change the *hq* Δ Ct values were subtracted from wild-type Δ Ct values to yield a $\Delta\Delta$ Ct value which was then input in to the following equation to determine fold change:

Fold Change = $2^{-\Delta\Delta Ct}$

Chapter 3. Results

3.1 Mouse phenotyping and Aif PCR genotyping confirmed the presence or absence of an Aif proviral insertion

Characteristic phenotypic hq markers such as a patchy coat, ataxia and low body mass were consistently observed among $X^{hq}X^{hq}$ and $X^{hq}Y$ mice. Genotypes of mice used in this study were confirmed by PCR using the tri-primer method. A 725 bp amplicon indicated the presence of an *Aif* hemizygous males (*hqY*) proviral insertion whereas the presence of a 537 bp amplicon indicated a wild-type *Aif* allele (Figure 3.1). Female heterozygotes (*hqX*) showed amplification of both 537 bp and 725 bp amplicons (Figure 3.1).

3.2 hqY mice show longer latency as early as three months and progressive decreases in OP summed amplitude following a 10 cd \cdot s/m² stimulus

Time-domain analysis of ERG OPs revealed no early (two month) delays of initial OP latency in hq disease mice when compared to the wild-type mice (Figure 3.2A). However, by three months of age hq mice demonstrate a significant delay (p = 0.005, Figure 3.2A) of initial OP latency by 24.6%. This is indicative of neural retinal feedback delay between the photoreceptors, INL and GCL of the retina. The trend of OP delayed response continues at four months and is significantly delayed at five to seven months (p = 0.0115, Figure 3.2A). Summed OP amplitudes demonstrated no significant changes at two or three months of age in the hq mice indicating normal INL and GCL function

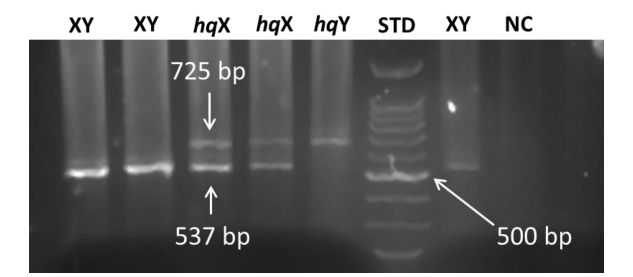


Figure 3.1. PCR amplification confirmed *Aif* genotype. *Aif* genotyping PCR amplicons were electrophoresed through a 1.5% agarose gel and stained with SYBR[®] Safe DNA Gel Stain (Invitrogen, Burlington, ON). A 100 bp DNA ladder (FroggiaBio Inc, Toronto, ON) was used as a size standard (STD) to estimate amplicon size. A single 537 bp amplicon indicated a wild-type *Aif* allele (XY) while a single 725 bp amplicon indicated the presence of an *Aif* proviral insertion.⁹³ Heterozygote, *hq* carriers demonstrated the presence of both 537 bp and 725 bp amplicons. A no DNA control (NC) was created with water prior to amplification and added to the PCR mastermix to eliminate the possibility of a false positive.

(Figure 3.2B). However by four months of age, a significant deterioration of summed OP amplitudes (p = 0.0222) by 26.5% occurred in *hq* mice resulting in reduced INL and GCL function (Figure 3.2B). The largest effect on summed OP amplitudes occurred at five to seven months with a significant reduction (p = 0.003, Figure 3.2B) by 54.4%. No statistical conclusions can be made about initial OP latency or summed OP amplitudes at eight to ten months of age as the *hq* cohort only contains two mice, although general trends appear to continue (Figure 3.2A,B).

3.3 Frequency-domain analysis of ERG OPs show early frequency changes in hqY disease mice followed by later changes in OP power and energy (10 cd·s/m²⁻

stimulus)

*hq*Y OP frequency demonstrated early, two-month changes with a significant reduction by nearly 14% in comparison to the wild type. Significant reductions were also present at three (p = 0.0015, 26.6%) and four months (p = 0.0248, 25.9%) of age (Figure 3.3A). This is indicative of delayed periodicity and reduced neural feedback mechanisms. *hq*Y overall OP power showed no early significant changes at two, three, or four months in comparison to the wild type (Figure 3.3B). This indicated adequate uniform retinal response of the INL and GCL of the *hq* mice. However by five to seven months of age, overall OP power demonstrated a significant reduction (p = 0.0028) by nearly 54.5% in comparison to the wild type (Figure 3.3B). Total OP waveform energy, indicative of total retinal INL, GCL and retinal response across the entire retina, demonstrated no early changes at two or three months of age in *hq*Y mice (Figure 3.3C). A significant decline

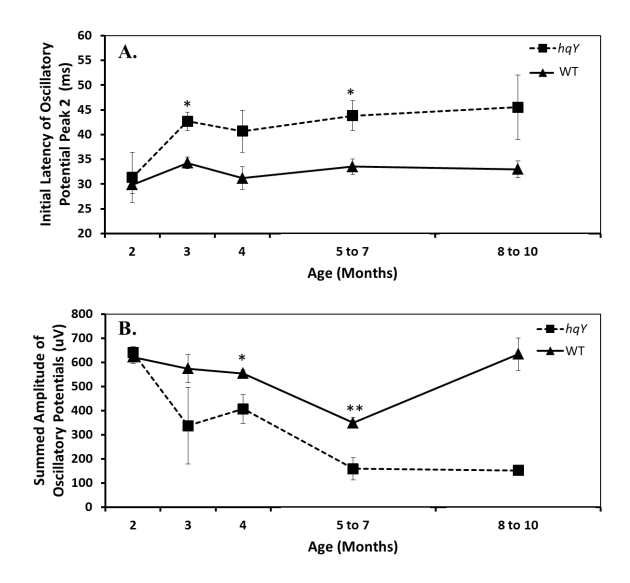


Figure 3.2. Electroretinography demonstrates delayed retinal responses and decreases in initial OP latency followed by subsequent reductions in summed OP amplitudes in the hq retina following a 10 $cd \cdot s/m^2$ stimulus. (A) Initial latency of OP2 indicates the functional response of the INL and GCL cells to photoreceptor hyperpolarization. Delayed initial latency of OP2 is evident at three months in hq mice with a significant delay of nearly 24.6% in comparison to the wild type. The delay in hqinitial OP response is also evident at five to seven months with an overall delay by 30.8%. (B) Summed amplitude is indicative of the overall retinal response and continuity of neural feedback of the INL and GCL. Significant reductions in hq OP summed amplitude are first apparent at four months of age resulting in an overall reduction by nearly 26.5%. In five-to-seven-month-old hq mice, the disease reduces summed OP amplitudes by 54.5%. The effect of age on genotype was also examined for all measured variables (data not shown, Appendix B, Table B.6A) Sample size per cohort can be found in Figure 2.1B. Asterisks indicate statistical significance (* p < 0.05, ** p < 0.005) and error bars represent ± 1 standard error of the mean.

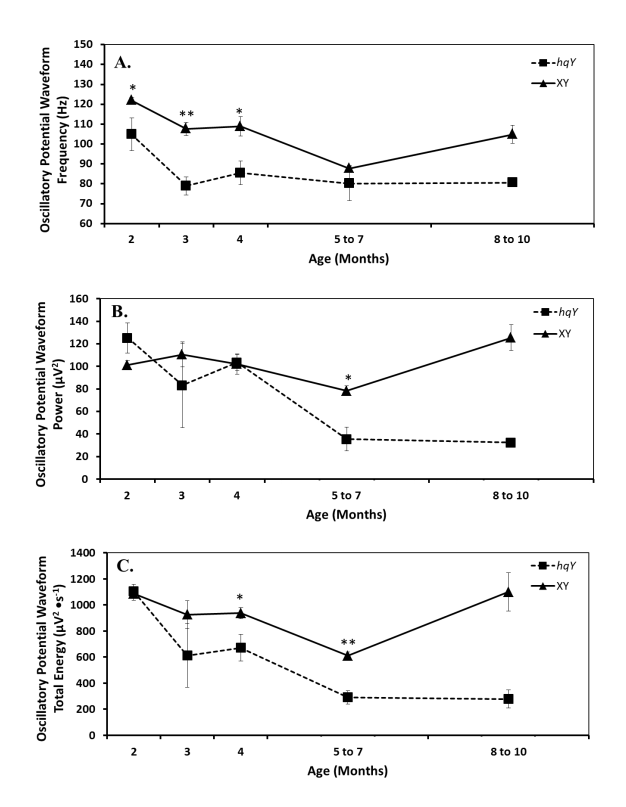


Figure 3.3. hqY mice have changes in OP waveform frequency as early as two months of age, followed by subsequent reductions of OP power and energy following stimulus of 10 cd•s/m². (A) Significant OP frequency reductions were evident immediately at two months of age with a 14% decline in comparison to the wild type. OP frequency continued to decline in hqY mice at 3 (26.6% reduction) and four months (25.9% reduction) of age in comparison to the wild type. This was indicative of a prolonged INL and GCL periodicity. (B) OP power is a measure of uniform retinal response of the INL and GCL. OP power showed a significant reduction of 54.5% by five to seven months in hq mice and continued to decline until eight to ten months. (C) OP energy is the summation of total OP retinal energy and was not significantly different at two or three months of age in hq mice. However, by four months of age hq mice show a significant reduction of OP energy by nearly 28.3% which progresses to a nearly 52.3% reduction by five to seven months of age. This indicates progressive disease in the hqretina particularly in the INL and GCL response. The effect of age on genotype was also examined for all measured variables (data not shown, Appendix B, Table B.6B) Sample size per cohort can be found in Figure 2.1B. Asterisks indicate statistical significance (* p < 0.05, ** p < 0.005) and error bars represent ± 1 standard error of the mean.

(p = 0.029) is evident at four months of age resulting in a 28.3% reduction of OP energy (Figure 3.3C). The decline in OP energy continues as the hq disease progresses and by five to seven months an overall significant reduction (p = 0.0006) by 52.3% is apparent (Figure 3.3C). No statistical conclusions can be made about OP frequency, power or total energy at eight to ten months of age as the hq cohort only contains two mice, although general trends appear to continue (Figure 3.3A, B,C).

3.4 Time-domain analysis of OPs indicate decreased summed amplitude in hq carrier mice as early as eleven months of age following a 10 cd \cdot s/m² stimulus

Initial latency of OP2 revealed no significant changes across all ages of the hqX heterozygote when compared to the wild type (Figure 3.4A). Summed amplitude, a measure of total inhibitory feedback mechanisms of the INL and GCL demonstrated no significant changes at three months when compared to the wild type. In the elevenmonth cohort, a significant reduction of summed OP amplitude was evident (p = 0.011; Figure 3.4B) at nearly 40.6%. Similarly, a significant reduction of 70.9% was observed in summed OP amplitudes at fifteen months (p = 0.0341; Figure 3.4B) indicating overall INL and GCL dysfunction and progression of the hq disease.

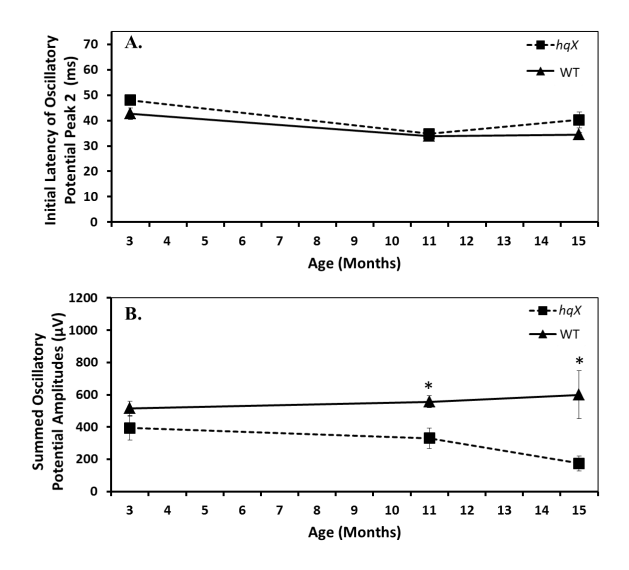


Figure 3.4. OP2-5 summed amplitudes indicates progressive loss of functional feedback mechanisms of the *hq* carrier retina following a 10 cd·s/m² stimulus. (A) Initial latency of OP2 is indicative of the initial neural response between photoreceptors and bipolar and horizontal cells of the INL and GCL. Although not statistically significant at any age it is apparent that *hqX* may have a lengthened initial latency in comparison to the wild type. (B) Summed amplitude of all OPs is indicative of the health of the neural retina and the continuity of feedback mechanisms of the INL and GCL. Significance is evident at eleven months of age with a 40.6% reduction in overall summed amplitude. INL and GCL function continued to decline as demonstrated by a significant reduction of OP summed amplitude by 70.9% at fifteen months. The effect of age on genotype was examined for all measured variables (data not shown, Appendix B, Table B.6C) Sample size per cohort can be found in Figure 2.1C. Asterisks indicate statistical significance (* p < 0.05) and error bars represent ± 1 standard error of the mean.

3.5 OPs of hq carrier mice indicate early functional periodicity deficits and subsequent reductions in retinal response continuity following a 10 cd \cdot s/m² stimulus

OP frequency, a measure of response of periodicity and neural feedback mechanisms across the INL and GCL, showed early functional deficits in hq carrier mice at three months of age (p = 0.049) with an overall reduction of 10.9% (Figure 3.5A). However, at eleven and fifteen months of age hq carrier mice showed no significant difference in overall OP frequency (Figure 3.5A). Overall power, a measure of uniform retinal response, showed no significant changes in hq carrier mice when compared to the wild type at three months of age (Figure 3.5B). By eleven months of age, retinal continuity in hqX mice is significantly reduced (p = 0.013) by 43.7% and progressive decreases lead to a significant (p = 0.0047) 62.9% reduction by fifteen months (Figure 3.5B). OP total energy, a summation of total retinal response and analogous to OP summed amplitude demonstrated no significant differences at both three and eleven months of age in the hq carrier mice (Figure 3.5C). A significant difference was observed in retinal OP total energy at fifteen months with a significant reduction (p = 0.002) of 66.3% (Figure 3.5C).

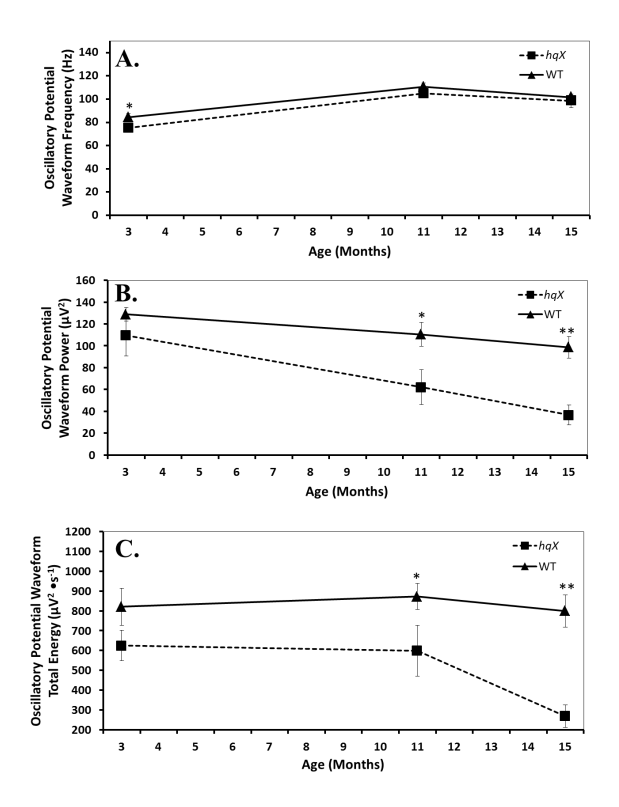
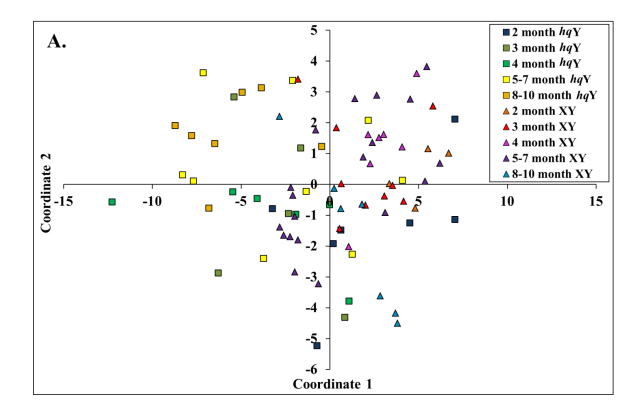


Figure 3.5. Frequency-domain OPs expose early functional differences of periodicity in three-month-old hq carrier mice and subsequent losses of OP power and energy following a stimulus of 10 $cd \cdot s/m^2$. (A) Frequency of OPs is indicative of periodicity of feedback inhibitory mechanisms of the INL and GCL. Significance in OP frequency response changes in the hq carrier mice at three months, with the hqX mice demonstrating a significantly lower OP frequency response. (B) OP Power is a measure of uniform retinal response of the INL and GCL. Statistical significance is evident in hq carrier mice at eleven months of age with an overall reduction of OP power by 43.7%. OP power and INL and GCL function continued to decline in comparison to the wild type leading to an overall reduction by 62.9% by fifteen months of age. (C) Energy of the OP waveform is the summation of the total OP response and was not statistically significant in three- and eleven- month-old hq carrier mice when compared to the wild type. However by 15 months of age, OP energy was significantly reduced by 66.3% in hq carrier mice indicting the progressive nature of the hq disease on INL and GCL function. The effect of age on genotype was also examined for all measured variables (data not shown, Appendix B, Table B.6D) Sample size per cohort can be found in Figure 2.1. Asterisks indicate statistical significance (* p < 0.05, ** p < 0.005) and error bars represent ± 1 standard error of the mean.

3.6 NMDS of time-domain hqY OP parameters indicate functional differences in OP latencies and amplitudes as early as two months of age

The stress associated with modelling all 32 time-domain variables from the severe hqY disease progression study into a two-dimensional axis was 11.31%. Pearson's correlation coefficients were used for axis interpretation following NMDS of hqY OP time-domain parameters (Appendix B, Table B.2). The x-axis (Coordinate 1) is highly correlated to OPs with shorter latencies, particularly initial latencies of OP2 and OP3 at all flash intensities (Figure 3.6). The x-axis is also positively correlated with increasing OP amplitudes particularly amplitudes of OP4 and OP5 at all flash intensities. The y-axis (Coordinate 2) is positively correlated to decreasing OP amplitudes across all flash intensities (Figure 3.6). Young hqY and wild-type mice cluster distinctly at two months of age with hqY mice demonstrating decreased OP amplitudes in comparison to the wild type (Figure 3.6A). Aging of the hqY mice is associated with a sharp decline in OP amplitudes and latencies between two and three months of age indicative of reduced INL and GCL function (Figure 3.6A). The disease appears to be progressive, leading to further reductions of OP latencies and amplitudes as the hqY mice age (Figure 3.6A). Inter-animal variation is much higher in the hqY mice, likely due to variation in the severity of retinal degeneration occurring in the hqY mice (Figure 3.6A). Wild-type mice show little OP amplitude change and slight decreases in OP latencies through time (Figure 3.6A).



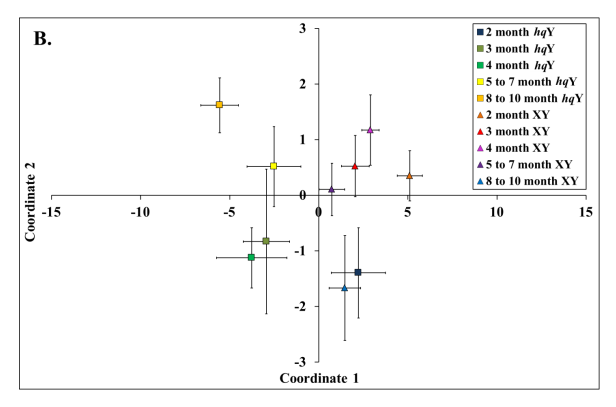
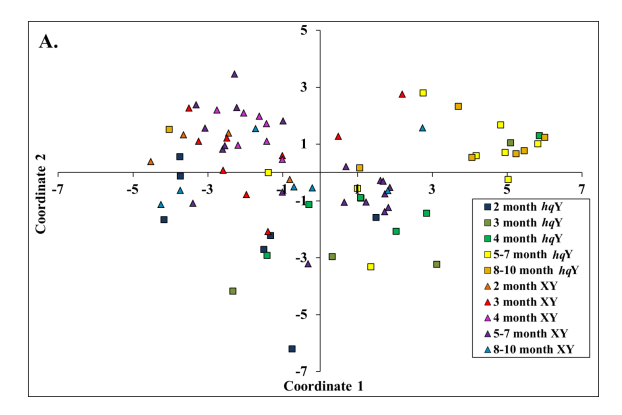


Figure 3.6. hqY time-domain OP parameters indicate early (two months) losses of INL and GCL periodicity and reduced retinal continuity at later ages. (A) Young hqY mice cluster distinctly from young, wild-type mice at two months of age as a result of reduced OP amplitudes and OP latencies. Progression of the hq disease leads to continued loss of OP amplitude (particularly OP4 and OP5) and latencies across all other age groups (months) resulting in severe losses of INL and GCL function. hq retina degeneration shows great inter-animal variation. Young, wild-type mice at two months demonstrate high OP amplitudes and short OP latency responses indicative of healthy retinal responses and have marginal declines from ages two through seven months. By eight to ten months, wild-type retinal function begins to decline with delayed OP latency and decreasing OP amplitudes. (B) The average coordinates for each cohort were plotted to assist with interpretation of NMDS clustering. Error bars represent ± 1 standard error of the mean. Stress is equal to 11.31%.

3.7 NMDS of frequency-domain OP parameters reveal distinct clustering of young hqY mice associated with decreased OP power, energy and frequency

The stress associated with modelling all 12 frequency-domain variables from the severe hqY disease progression study into a two-dimensional scale was 5.67%. Pearson's correlation coefficients were used for axis interpretation following NMDS of hqY OP frequency-domain parameters (Appendix B, Table B.3). Coordinate 1 is highly correlated with decreasing energy and frequency and slightly correlated with decreasing power at all flash intensities (Figure 3.7). Coordinate 2 is highly correlated to decreasing power and highly correlated to increasing OP frequencies at all flash intensities (Figure 3.7). Wild-type and hqY mice cluster distinctly at two months of age with hqY mice demonstrating immediate lower total OP power, energy and frequency (Figure 3.7A). Distinctive hqY clustering continues across all ages with progressive reductions of OP frequency, power and energy (Figure 3.7A). Inter-animal variation is much higher in the hqY mice (Figure 3.7A). Young, wild-type mice cluster with high OP energy, power and frequency and with only marginal reductions in any parameter with age (Figure 3.7A).



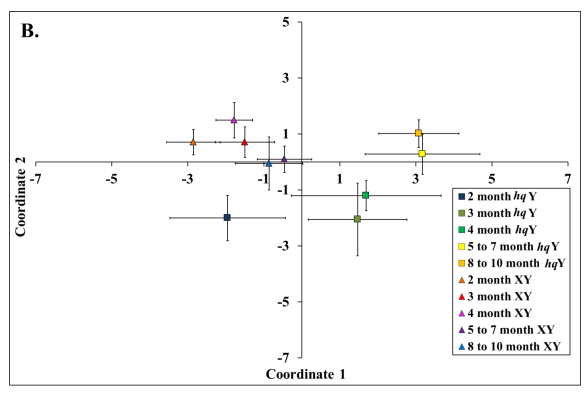


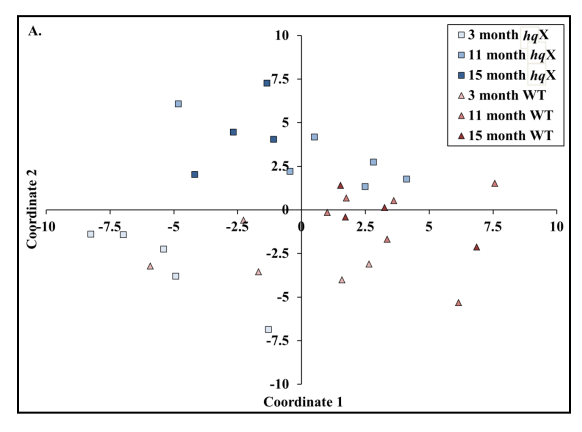
Figure 3.7. hqY frequency-domain OP parameters reveal distinct loss of overall retinal periodicity and continuity of the INL and GCL response as early as two months. (A) Young hqY and wild-type mice cluster distinctly demonstrating an immediate reduction in overall OP frequency, power and total energy and loss of retinal function. Aging of hqY mice (three-ten months) leads to progressive loss of all frequency-domain OP parameters. Severe losses in OP energy and frequency are most prominent between two and three months of age in hqY mice. Young, wild-type mice (two months) cluster with high OP frequency, power and total energy and have slight age related decreases (three to ten months). Progression of aging in wild-type mice shifts NMDS parameters similar to that of young hqY mice (two months). (B) The average coordinates for each cohort were plotted to assist with interpretation of NMDS clustering. Old wild-type mice cluster similarly with young hq disease mice. Error bars represent \pm 1 standard error of the mean. Stress is equal to 5.67%.

3.8 NMDS of hq heterozygote time-domain parameters expose distinct clustering at eleven and fifteen months of age

The stress associated with modelling all 32 time-domain variables from the moderate hqX disease progression study into a two-dimensional scale was 9.84%. Pearson's correlation coefficients were used for axis interpretation following NMDS (Appendix B, Table B.4). The x-axis (Coordinate 1) is highly correlated to short OP latencies, particularly of OP2 and OP3 as well as high OP3 and OP4 amplitudes (Figure 3.8). The y-axis (Coordinate 2) is highly correlated to decreasing OP amplitudes across all flash intensities (Figure 3.8). Young hq carrier and wild-type mice (three months) cluster together with long OP latencies and intermediate amplitudes (Figure 3.8A). However by eleven months, hq mice shift towards decreased OP amplitudes and shorter latencies indicative of enhanced periodicity but loss of continuity of neural feedback response. By fifteen months of age hq carrier mice demonstrate a large delay in OP latencies with small OP amplitudes resulting in overall decreased retinal function (Figure 3.8A). Aging of wild-type mice (eleven and fifteen months) is associated with marginal decreases in OP amplitudes and enhanced OP latencies (Figure 3.8A).

3.9 NMDS of frequency-domain hq carrier mice OP parameters demonstrate progressive loss of OP power and energy at eleven and fifteen months

The stress associated with modelling all 12 frequency-domain variables from the moderate hqX disease progression study into a two-dimensional scale was 3.84%. Pearson's correlation coefficients were used for axis interpretation following NMDS of



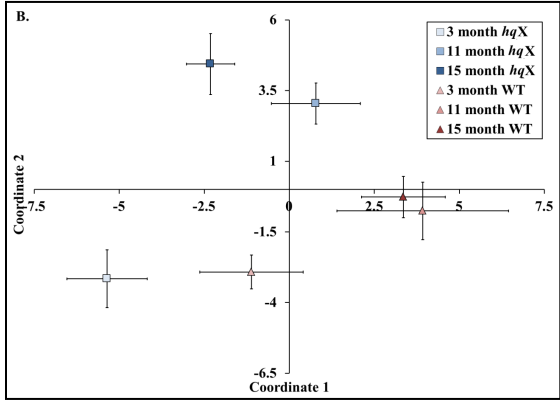
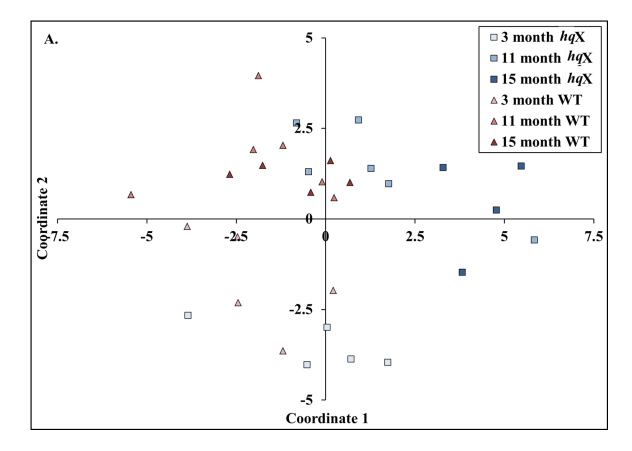


Figure 3.8. hq carrier time-domain OP parameters demonstrates distinct clustering at 11 and fifteen months associated reduced initial INL/GCL response and decreased continuity of overall retinal response. (A) Young hq carrier and wild-type mice cluster similarly with reduced OP initial latency and intermediate amplitudes at three months of age. This is indicative of reduced photoreceptor and INL and GCL periodicity and continuity of INL and GCL function at a young age. By eleven months of age, hq carrier mice demonstrate distinct clustering associated with delayed initial OP latency and a reduction in OP amplitudes when compared to age-matched wild type. As the hq disease progresses, further reductions are evident in both OP initial latency and amplitudes at fifteen months and as a result of incomplete retinal function. Aging of wild-type mice is associated with marginal decreases in OP amplitude and increases in OP latencies and ultimately little change in retinal function. (B) The average coordinates of each age cohort were plotted to assist with interpretation of NMDS clustering. Error bars represent ± 1 standard error of the mean. Stress is equal to 9.84%.



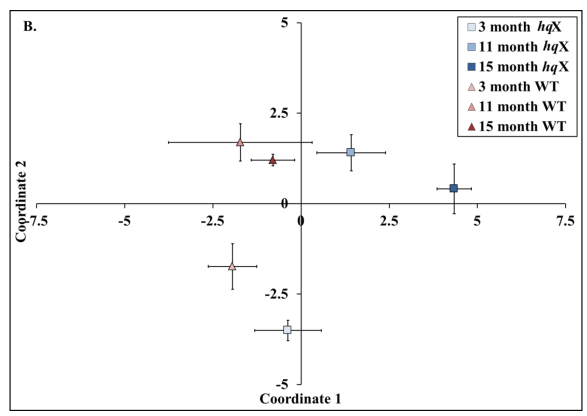


Figure 3.9. Frequency-domain OP parameters of hq carrier mice indicates distinct clustering associated with INL and GCL dysfunction at eleven and fifteen months. (A) Young (three months) hqX and wild-type mice cluster similarly based upon high OP power and energy and low frequency. hq disease progression at eleven months is indicative of large reductions in total OP power and energy as well as increased OP frequency. Further reductions of OP power and energy are evident at fifteen months with additional reduced OP frequency. Aging of wild-type mice is associated with increases in OP frequencies and minimal reductions in overall OP total power and energy. (B) The average coordinates for each cohort were plotted to assist with interpretation of NMDS clustering. Error bars represent ± 1 standard error of the mean. Stress is equal to 3.84%. hqX OP frequency-domain parameters (Appendix B, Table B.5). The x-axis (Coordinate 1) is highly negatively correlated to both OP total energy and power across all flashes while the y-axis (Coordinate 2) is highly positively correlated to frequency across all flashes (Figure 3.9). hq carrier and wild-type mice cluster similarly at three months of age with hqX mice demonstrating slightly lower OP frequency, power and energy (Figure 3.9A). By eleven months of age, hq carrier mice cluster distinctly with higher OP frequency and significantly reduced OP power and energy (Figure 3.9A). This general trend progresses as hq carrier mice progress to older ages with further reductions in OP power and energy indicative of reduced continuity of INL and GCL retinal response (Figure 3.9A). Aging of wild-type mice is associated with increases in OP frequency and marginal decreases in OP power and energy (Figure 3.9A).

3.10 hq mice maintain blood retinal barrier integrity at both a young and old age

To estimate concentrations of the Evans Blue (EB) dye in tissue, a standard curve was created through a series of dilutions of EB in formamide. EB dye dissolved in formamide maintain a linear relationship between 250 to 10000 ng/mL (Figure 3.10A, r^2 = 0.999, y = 7.87 x10⁻⁶x + 0.002). The liver is a region of full vascularization and therefore no blood endothelial barrier is present. To infer sensitivity of the assay a series of tests were performed as described below. Significantly more EB was found in the liver of EB injected wild-type mice than in uninjected controls (n = 4 per cohort, One-way ANOVA, p < 0.001). No significant difference in EB amount was found in the retina of wild-type mice injected with EB and uninjected wild-type mice (n = 4 per cohort, Oneway ANOVA, p > 0.05) indicating an intact BRB in wild-type mice. In addition, wildtype mice show significantly more EB in liver than retina at both a young and old age (Figure 3.10 B, n = 4 per cohort, Two-way ANOVA, p < 0.005) but no significant reduction with age (Figure 3.10B, n=4 per cohort, Two-way ANVOA, p > 0.05). *hq* mice show a similar trend with significantly more EB in the liver than retina at both ages (Figure 3.10 C, Two-way ANOVA, p < 0.005) but no significant reduction with age (Figure 3.10C, Two-way ANOVA, p > 0.05). In order to account for technical variation, retina EB amount was normalized to liver EB amount and no significant difference in retina/liver ratio was found across all cohorts indicating a similar amount of EB is present in the retina across all cohorts. This evidence suggests that there is no degradation of the BRB in *hq* mice (Figure 3.10, Two-way ANOVA, p > 0.05).

3.11 Pathway analysis reveals differential expression of focal adhesion,

extracellular remodeling, cell stress and inflammation in the four-month hq retina

Four-month-old hq retinal gene expression microarray data (n=3 per cohort) previously performed by A. Laliberté (2010),¹⁰³ was filtered using selection criteria set at a 1.75 fold change (mRNA transcript level) and a p-value less than 0.05. This revealed a total of 521 genes that were differentially expressed between retinal tissue from hqY mice and age-matched wild-type mice. A total of 182 genes were downregulated by greater than 1.75 fold, and a total of 339 were upregulated by greater than 1.75 fold. KEGG pathway analysis revealed pathways with significant proportions of differentially expressed genes. Significant pathways involved in extracellular remodelling, ABC transport, immune response, inflammation, cell cycle and p53 signalling were

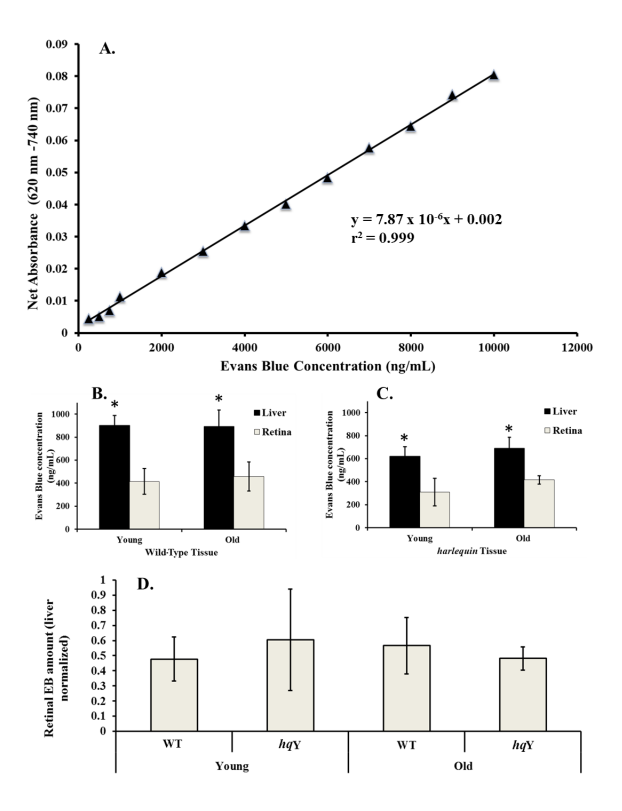


Figure 3.10. hqY mice maintain a blood-retinal-barrier similar to that of agematched wild-type mice. (A) Evans Blue standards in formamide maintain a linear relationship between concentration and background-subtracted absorbance ($r^2 = 0.999$, y = 7.87 x10⁻⁶x + 0.002). Standards were tested by three different people in triplicate and were virtually identical. (B) A significantly higher concentration of Evans Blue dye is present in the liver in comparison to retina in wild-type mice at both ages. (C) Similarly, a significantly higher concentration of Evans Blue dye is present in the liver of hqY mice in comparison to retina across both ages. (D) Liver-normalized retinal Evans Blue amount is not significantly different across any cohort indicating an intact blood-retinal barrier. differentially expressed in the *hq* retina (p < 0.01) (Table 3.1). Additional DNA repair and cancer pathways were also differentially expressed (p < 0.05) (Table 3.1).

KEGG pathway analysis allowed for a comprehensive overview of differential expression in the *hq* mouse retina. However, due to the nature of KEGG gene annotations (not retinal specific) and containing only single genetic isoforms, only 177 gene IDs out of the 521 were represented the software analysis. Therefore, well over 65% of genes differentially expressed in the expression microarray were not used by KEGG pathway software. To account for this bias, a manual annotation of all differentially expressed genes was performed (Appendix E). A thorough review yielded an alternative metabolism theme, common amongst AIF knock-downs including; the upregulation of glucose metabolism, lactate acidosis, and glycerol metabolism. Genes present in these pathways were selected for Taqman[®] qPCR analysis (Table 3.2).

3.12 Cerebellar and retinal RNA quality was assessed prior to transcriptional analysis

Spectrophotometry, gel electrophoresis and Bioanalyzer spectra all indicated the presence of high quality cerebellar RNA. Cerebellar samples were only chosen if they indicated a concentration greater than 140 ng/uL and a high absorbance ratio (A260/A280 > 1.8) (Appendix B, Table B.7). Gel electrophoresis allowed quantification of the 28S and 18S ribosomal RNA (rRNA) subunits as well as the potential for gDNA contamination. Samples were only selected if they demonstrated distinct rRNA subunits with no gDNA contamination (Figure 3.12). Samples were also analyzed via Bioanalyzer spectra and selected if there was clear separation of 28S and 18S peaks (Figure 3.13A)

Pathway	p-value [‡]	Number of Differentially Expressed Genes	Gene Enrichment Percentage (%)
ECM-receptor interaction	6.9 x 10 ⁻⁶	13	2.5
ABC transporters	5.5 x 10 ⁻⁵	9	1.8
Focal Adhesion	1.0 x 10 ⁻⁴	18	3.5
Viral myocarditis	5.5 x 10 ⁻⁴	11	2.1
p53 signaling pathway	4.9 x 10 ⁻³	8	1.6
Cell cycle	5.7 x 10 ⁻³	11	2.1
Complement and coagulation cascades	7.8 x 10 ⁻³	8	1.6
Tight junction	8.2 x 10 ⁻³	11	2.1
Regulation of actin cytoskeleton	0.015	14	2.7
Pathways in cancer	0.02	18	3.5
Small cell lung cancer	0.046	7	1.4

Table 3.1. KEGG pathway analysis based on microarray data of differentially expressed genes from retinal samples of four-month-old hq disease mice.^{*,†}

* Microarray was performed by A. Laliberté (2010).¹⁰³

[†] Pathway analysis was performed with the Database for Annotation, Visualization, and Integrated Discovery (DAVID) [National Cancer Institute at Frederick, Frederick, MD].^{141,142}

‡ Pathway is ordered by increasing p-value.

Gene Name	Symbol	Function†	p-value	Fold Change
Adenylate cyclase 10	Adcy10	Key intracellular regulation of pH and bicarbonate buffering	0.005	+2.858
Glycerol-3- phosphate dehydrogenase 2	Gpd2	Secondary step of glycerol metabolism, feeds into glycolysis	0.04	+2.377
Aldehyde dehydrogenase 2	Aldh2	Primary step for conversion of Acetyl-CoA to acetate	0.045	+2.345
Glycerol Kinase 5	Gk5	Primary step of glycerol metabolism, feeds into glycolysis	0.048	+1.998
TBC1 domain family, member 4	Tbc1d4	Responsible for shuttling <i>Glut4</i> to the plasma membrane from the endoplasmic reticulum	0.03	+1.992
Hexokinase 1	Hk1	Initial step of glycolysis in the retina – drives glycolytic rate	0.037	+1.964
Glucose transporter, member 4 [‡]	Glut4	Rapid, insulin dependent glucose transporter responsible for glucose intake in neurons	0.023	+1.4
Lactate dehydrogenase C^{\ddagger}	Ldhc	Responsible for reverse reaction of L-lactate to pyruvate	0.018	-1.63
Pyruvate dehydrogenase, component X	Pdhx	Structural component of pyruvate dehydrogenase complex	0.016	-2.482
Sodium bicarbonate transporter, member 10	Slc4a10	Bicarbonate transporter to remove excess CO ₂ from mitochondria	0.027	-2.873
Apoptosis- inducing factor 1	Aif	Complex I efficiency and maintenance, apoptosis inducer.	0.0002	-6.012

Table 3.2. Genes of interest from four-month-old hq mice selected from retinal microarray data for Taqman[®] validation.

[†] Functional annotation was based on NCBI, GENE search annotation [National Institute of Health, Bethesda, MD]

[‡] Previous literature suggests increased glucose metabolism and lactate acidosis in hq mice, therefore *Glut4* and *Ldhc* (p < 0.05, < 1.75 fold change) were added for examination.

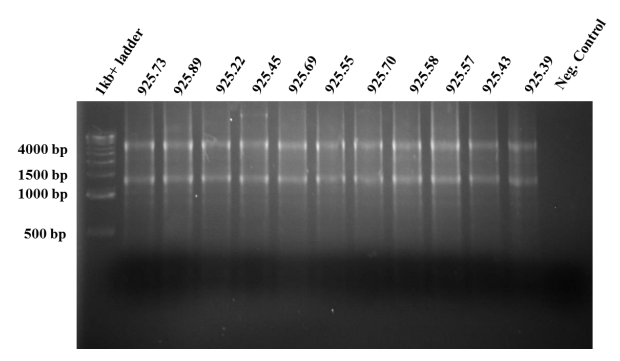
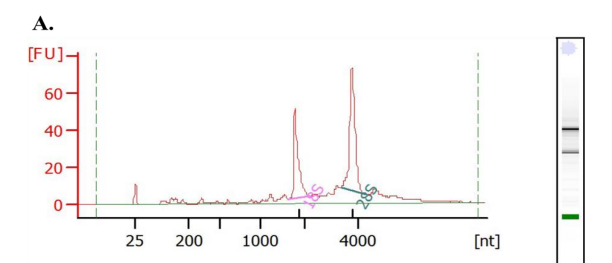
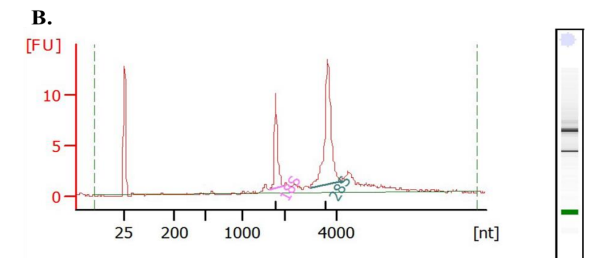


Figure 3.11. Agarose gel electrophoresis demonstrates high quality cerebellar **RNA samples with distinct 28S and 18S ribosomal subunits.** A representative gel of 11 samples indicates the presence of high quality cerebellar RNA (distinct 28S and 18S bands) electrophoresed with a 1 kb+ DNA ladder [Life Technologies, Burlington, ON]. Sample 925.45 (lane 5) presented contamination with high molecular weight gDNA. A negative control of water was electrophoresed in the final lane (lane 13).





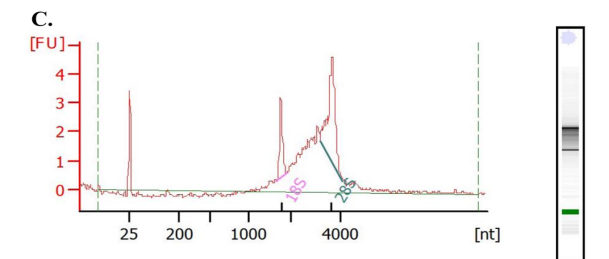


Figure 3.12. Electropherograms indicate high RNA quality of both cerebellar and retinal samples. (A) A representative electropherogram of intact, high quality cerebellar RNA using the Agilent 2100 Bioanalyzer [Agilent Technologies Inc., Palo Alto, CA]. RNA quality is a measure of fluorescence (FU) and nucleotide (nt) size. An initial fluorescence peak at 25 nt represents a size marker, followed by a clear distinction of the 18S (~1500 nt) and 28S rRNA subunits (~4000 nt). A representative agarose gel image is created (right hand side) based on fluorescence of rRNA peaks. (B) A representative electropherogram of intact, high quality retinal RNA. Fluorescence is much lower due to a lower concentration, however a clear separation of the 18S and 28S rRNA subunits is still evident. (C) A retinal RNA electropherogram with gDNA contamination. Sheared gDNA and degradation products show similar properties to the 28S rRNA subunit and lead to skewing of the rRNA peak.

as well as an RNA integrity number (RIN) greater than 8. Retinal samples were subject to the same spectrophotometric (concentration > 16 ng/uL) and Bioanalyzer assessment (Figure 3.13B) (Appendix B, Table B.7). Gel electrophoresis was not performed with retinal samples as the RNA concentrations were too low. All retinal RNA samples used for transcriptome analysis were of high quality.

3.13 Taqman[®] quantitative PCR confirms metabolic and cell regulation changes in the hq mice retina and cerebellum

Aif was significantly downregulated (p < 0.001, fold change < 11.312) in the hq retina at four months, three months, and two months (Table 3.3) of age as well as significantly (p < 0.001, fold change = -8.8183) downregulated in the four-month hq cerebellum (Table 3.4). Cyclin-dependent kinase inhibitor 1A (Cdkn1a), responsible for inhibiting cell cycle, was significantly (p < 0.05) upregulated in the hq retina at four months, three months and two months of age (Table 3.3) as well as the four-month cerebellum (Table 3.4). Genes associated with metabolism were upregulated in the fourmonth hq retina (Table 3.3) including; Adcy10 (p = 0.0013, fold change = +1.7995), Slc4a10 (p = 0.0024, fold change = 1.8102), Gk5 (p = 0.0021, fold change = +1.7995), Aldh2 (p = 0.017, fold change = +1.660), Gpd2 (p = 0.023, fold change = +1.274) and Pdhx (p = 0.027, fold change = +1.250). No genes involved in metabolism were differentially regulated in the three-month hq retina (Table 3.3). However, Gk5 (p = 0.022, fold change = +1.2688) and Aldh2 (p = 0.0465, fold change = +1.289) were found to be differentially expressed early in the two- month hq retina (Table 3.3). One metabolism gene was found to be differentially expressed in the four-month ha cerebellum (Table 3.4); Adcy10 (p = 0.0265, fold change = +2.2422).

Gene	Two months	Three months	Four months	
Name	Fold change [‡]	Fold change ‡	Fold change [‡]	
	$(\mathbf{p}$ -value [†])	$(\mathbf{p}$ -value [†])	(p-value [†])	
Aifm1	- 12.315 (<0.001)	- 17.301 (<0.001)	- 11.312 (<0.001)	
Adcy10	+ 1.7285 (0.0878)	+ 1.5042 (0.6607)	+ 1.7995 (0.0013)	
Gk5	+ 1.2688 (0.0219)	+ 1.0664 (0.60)	+ 1.8102 (0.0021)	
Slc4a10	+ 1.1164 (0.1655)	+ 1.0728 (0.4052)	+ 1.4990 (0.0024)	
Cdkn1a	+ 2.2391 (0.0334)	+ 1.6654 (0.0053)	+ 3.2214 (0.0063)	
Aldh2	+ 1.2889 (0.0465)	- 1.0963 (0.5967)	+ 1.660 (0.017)	
Gpd2	+ 1.2586 (0.1036)	- 1.106 (0.6446)	+ 1.274 (0.023)	
Pdhx	+1.1222 (0.1319)	+1.0285 (0.6297)	+1.250 (0.027)	
Hk1	+ 1.1442 (0.1780)	- 1.007 (0.997)	+ 1.286 (0.060)	
Rn18s	+ 1.4432 (0.3058)	+ 1.0531(0.7299)	+ 1.361 (0.270)	
Tbc1d4	+ 1.2331 (0.2442)	n/a	+ 1.091 (0.344)	
Glut4	+ 3.2467 (0.1107)	- 2.8596 (0.3161)	- 2.898 (0.832)	
Ldhc	n/a	n/a	n/a	

Table 3.3. Differential gene expression of selected genes in the *hq* retina at two, three and four months of age based on Tagman[®] RT-PCR.^{*}

* Gene order was established by ascending four month p-value, bold genes indicate significance.

 \dagger An ANOVA was used in Excel [Microsoft, Redmond, VA] to determine significance of the Δ Ct values.

 \ddagger Fold change was determined with DataAssistTM software [Life Technologies, Burlington, ON] using the comparative Ct ($\Delta\Delta$ Ct) method.¹⁴⁰

n/a - indicates late amplification.

Gene Name	p-Value [†]	Fold Change [‡]
Aifm1	<0.001	- 8.8183
Cdkn1a	0.0251	+ 1.6291
Adcy10	0.0265	+ 2.2422
Slc4a10	0.1409	+ 1.4275
Tbc1d4	0.1409	+ 1.2278
Glut4	0.1624	+ 1.6189
Gk5	0.1792	+ 1.4212
Pdhx	0.2217	+ 1.1763
Rn18s	0.2776	+ 1.3111
Hk1	0.4030	+ 1.1589
Gpd2	0.4065	+ 1.1127
Aldh2	0.4159	+ 1.1105
Ldhc	n/a	n/a

Table 3.4. Differential gene expression of selected genes in the *hq* cerebellum at four months of age based on Taqman[®] RT-PCR.^{*}

* Gene order was established on ascending p-value, bolded genes indicate significance.

 \dagger An ANOVA was used in Excel [Microsoft, Redmond, VA] to determine significance of the Δ Ct values.

‡ Fold change was determined with DataAssistTM software [Life Technologies, Burlington, ON] using the comparative Ct ($\Delta\Delta$ Ct) method.¹⁴⁰

n/a - indicates late amplification.

3.14 hqY mice have a decreased total body mass but display hyperphagia at four months

The mean body mass of hqY diseased mice at four months (22.422 ± 0.9993g) was significantly lower (p < 0.001) than age-matched wild-type mice (29.512 ± 0.5362g) (Figure 3.13A) with a 24% reduction in total body mass. Although hqY mice exhibit smaller mean body mass, they display hyperphagia at four months with a significant increase (p < 0.001) in mean food consumption (0.1991 ± 0.01 grams of food/gram of body mass/day) in comparison to the wild type. This overall increase is nearly 34.4% (0.1481 ± 0.002 grams of food/gram of body mass/day) in hqY mice (Figure 3.13B).

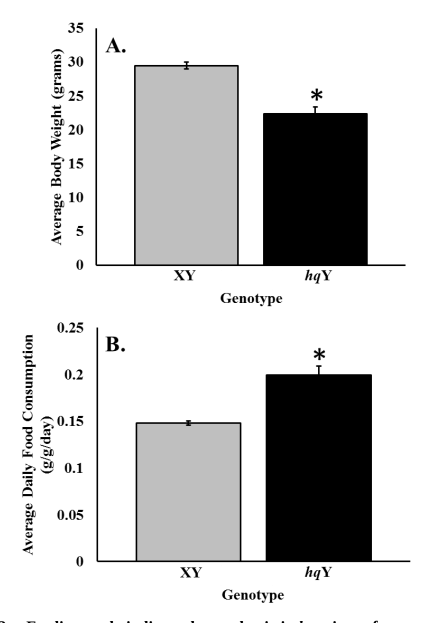


Figure 3.13. Feeding study indicates hyperphagia in hq mice at four months of age. (A) hq mice demonstrate a significantly lower mean body mass in comparison to agematched wild-type mice. (B) Mean daily food consumption was significantly higher in hqdisease mice indicating hyperphagia. Sample size was four per cohort, asterisks indicate significance (*p < 0.001) and error bars represent ± 1 standard error of the mean.

Chapter 4. Discussion

4.1 Early hq retinal dysfunction can be tracked to the inner retina through the study of oscillatory potentials

Previously, it was shown that OXPHOS dysfunction in the hq retina is associated with parainflammatory disease as early as 1.8 months of age, resulting in microglial migration to the ONL and subsequent ONL cells losses.^{103,104} To date, early mechanisms prompting retinal degeneration in the hq mouse have yet to be elucidated. It was predicted that oscillatory potentials could be used to track hq disease progression because of their association with hq hallmarks of the retinal disease including degeneration of the ONL, IPL, OPL as well as ganglion and amacrine cells. Herein, at the retinal physiological level, *in vivo* diagnosis of electrophysiological response in the hq retina was able to identify early, diagnostic candidates of retinal malfunction including OP frequency and initial latency. NMDS provided a thorough visualization of OP waveforms across multiple flash intensities and identified changes across all measured parameters of OP profiles.

Due to the highly metabolic nature (continuous transport) of the BRB, it was unknown whether the severe neurodegeneration in the hq retina was the result of BRB degradation or endogenous retinal stress caused by reduced AIF function. Loss of BRB integrity leading to exogenous antigens larger than the Evans Blue albumin complex traversing the BRB was rejected as a proposed mechanism for chronic microglial activation in the hq retina. Thus, triggers of chronic microglial activation are hypothesized to be the result of an endogenous cellular stress mechanism. Early transcriptome changes in the hq retina are consistent with partial compensating metabolic mechanisms in an attempt to maintain production of ATP *in vivo*. Such as, glycerol metabolism to generate $FADH_2$ which enters OXPHOS distal to NADH dehydrogenase. This study provides the first, *in vivo*, preliminary evidence for alternative metabolism in the retina of the *hq* mouse and is the basis for subsequent future examinations of *hq* retinal metabolism.

4.2 Oscillatory potentials are relevant for the study of hq retinal degeneration

Oscillatory potentials (OPs) are the intermediate peaks on the ascending limb of the ERG b-wave. OPs are believed to correspond to inner retinal function of the ON-, OFFand feedback mechanisms of the INL and GCL. OP subgroups, beginning with intermediate OPs represent the proximal action-potential independent (bipolar and horizontal cell graded potentials) and action-potential dependent mechanisms of ganglion cells and amacrine cells of the inner retina (Figure 4.1).⁵⁶ Additionally, the interplexiform cell which spans the OPL and IPL may generate radial current flow (feedback) from lateral synapses of the OPL and IPL and may be responsible for oscillations.^{54,55} This is particularly beneficial for the study of *hq* retinal degeneration. Neurites of the OPL and IPL contain the synapse between dendrites and axons of adjacent neurons relaying electrochemical signal from the OS. They are the most energetic region of the neurons and hence *hq* mice exhibit considerable thinning of the neurites⁹³ as well as upregulation of neurite remodelling and maintenance genes.¹⁰³ These genes include; *Sodium channel beta 4 subunit (Scn4b)*, and *Sterile alpha motif domain containing 4 (Samd4)* both of

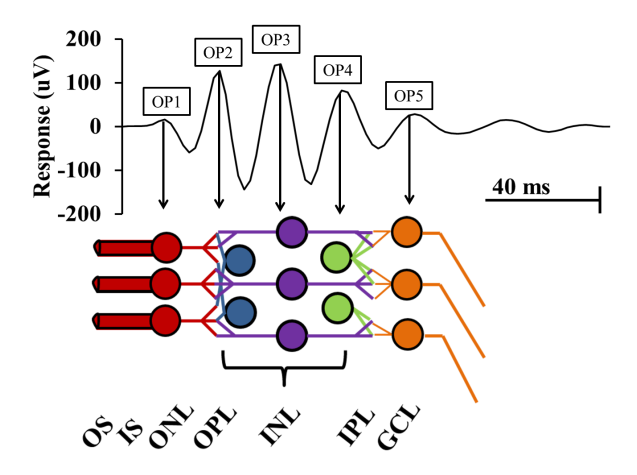


Figure 4.1. Isolated oscillatory potentials (OP1-OP5) may correspond to retinal layer functionality. OP1 is associated with photoreceptor hyperpolarization (red). Intermediate peaks OP2-OP3 are believed to be the response of second-order horizontal (blue) and bipolar (purple) action-potential independent (graded) neurons. Subsequent late OP peaks are believed to be associated with action-potential dependent amacrine (green) and ganglion (orange) cells. Intermediate OP peaks are thereby generally associated with INL function and later OP peaks with GCL function. Radial electrochemical cycling is the result of feedback mechanism occurring across the IPL of the retina.

which are involved in maintenance of neurite function and synaptic health.¹⁰³ In addition to neurite loss, the first histological hallmarks of retinal degeneration begin at three months of age, with retinal thinning of the ganglion and amacrine cells.⁹³ Oscillatory potentials measure retinal health and signal transduction through the retina and can therefore assess the health of both the neurites and the ganglion cells prior to structural losses. OPs are capable of detecting early changes as a result of mitochondrial dysfunction prior to any functional losses in the *hq* retina as diagnosed by the ERG awave and b-wave.

4.3 The hqY mouse shows early neuronal functional deficits

In the longitudinal study of ERG OPs through the hq disease, OP profiles changed in both the time and frequency domain. The initial latency of OPs indicated that functional changes began as early as three months of age in hq mice. Simultaneously, changes in the major frequency of the OPs began at two months of age in the hq mouse and were followed by a steep reduction in OP frequency occurring between two and three months of age. Major frequency and initial latency are informative of periodicity and likely a response of radial current cycling through the inner retina and therefore are indicative of the speed of the graded potential changes in the INL and GCL as a result of photoreceptor synapsis leading to depolarization of the bipolar cells. Analysis in the frequency domain is advantageous when examining periodicity of OPs because it identifies significant changes across hq retina function prior to the time domain. Significant decreases in OP summed amplitude began at four months of age in the hq mouse and continued into later ages with diminished total OP amplitude. Analogous changes in the frequency domain demonstrate decreases in total OP energy occurring at four months of age. Both the summed amplitude and total energy are indicative of total energetic response of the OPs, and therefore the functionality and health of all cells in the inner retina. Decreases of both hq OP summed amplitude and total energy are likely due to neuronal cell loss of the ONL, INL and GCL as decreasing OP amplitudes have been implicated in structural retinal loss.⁵⁶ Summed OP amplitude can therefore be used as a measure of overall inner retinal health as previously described in other models.^{66,78,79}

In the hq mouse, inner retinal functional deficits as a result of changes in OP frequency and latency occur prior to any documented structural losses at four months.^{19,93,102} These functional differences occur as early as two months of age and are consistent with a neuronal synaptic malfunction and microglial activation at complementing ages in the hq retina. The activation of microglial cells and OP functional deficits of the inner retina coincide at two months and indicate the loss of homeostasis and neuronal synaptic malfunction.

4.4 The hqX mouse demonstrates slower disease progression of the inner retina

The hq carrier mouse phenotype displays a moderate, extended disease phenotype in comparison to the hemizygous or homozygous disease mice.^{93,102,103} The functionality of the inner retina, as measured by OPs of the ERG, demonstrate consistent moderate phenotypic results. In fact, no OP initial latency or frequency changes were observed in the hq carrier mice at any age. This is indicative of no periodicity change in the retina of the hq carrier mice and sufficient responses of the synapsis between ONL: INL and INL:GCL neurons and overall vitality of the ONL photoreceptors. However, decreases in both amplitude and analogous energy and power indicate inner retinal cell losses and decreases in INL and GCL function at eleven months of age. Retinal thinning in the hqcarrier retina is not evident until approximately fifteen months when structural losses of the INL and ONL occur.¹⁰³ Therefore OPs identify physiological changes of the INL (bipolar and amacrine cells) prior to any structural cell loss and reveal a diagnostic parameter to assess retinal health before cell death has occurred.

4.5 hqY OP functional losses mimic retinopathy and may help reveal the origin of OPs

Mammalian retinopathy begins with photoreceptor loss in the ONL. Retinopathy leads to hyperoxia in the inner retina caused by oxidative challenges, neovascularization, and inflammation.^{143,144} Recent studies, of both human and rat scotopic ERGs have demonstrated the functional disease progression of retinopathy and its effect on OPs. Retinopathy leads to decreases in all OP amplitudes, particularly that of early OPs, indicative of persistent dysfunction of photoreceptors.^{66,78} Deterioration of OP amplitudes and total OP energy in *hq* disease mice indicates similar disease mechanisms to that of retinopathy as health of photoreceptors is diminished.

Aging has been associated with nuclear losses in the INL and GCL leading to compromised retinal function with senescence.^{66,145,146} Aging leads to the progressive

loss in OP periodicity (initial latency and frequency) in wild-type rats and humans,^{66,146} analogous to periodicity changes in the hq retina. The deficits in hq OPs demonstrate persistent effects of hypomorphic AIF on retinal function leading to mitochondrial dysfunction, photoreceptor cell death, retinal degeneration and an advanced aging phenotype. A thorough histological characterization of hq retina neuron losses indicates OP are generated in the INL and GCL and are helpful in the investigation of the source of OP production in the retina.^{102,132}

4.6 Non-metric multidimensional scaling offers a comprehensive visualization of inner retinal function with genotype and age

The measurement of each OP amplitude and latency in the time domain provided a total of eight measurements for each flash intensity (excluding OP1 due to inconsistencies and photoreceptor contamination). Four flash intensities (0.63, 4, 10 and 25 cd \cdot s/m²) were examined as they all contained strong and measureable OP responses. Across the four flash intensities 32 measurements were taken in the time domain and 12 measurements in the frequency domain for each mouse.

NMDS computationally-derived coordinates fitted to 32 time-domain variables into two dimensions and 12 frequency-domain variables into two dimensions. The stress associated with modelling multiple parameters into two dimensions was low across all four analyses and demonstrates a valid, comprehensive evaluation of OPs across multiple stimulus intensities. NMDS analysis provided a valuable interpretable aid in examining disease progression and provides translational relevance in determining origins of disease and potentially identifying drug targets. Pearson's correlation coefficients were used to examine the correlation of each axis with each of the fitted variables to determine which variables participate most into axis determination. For example, flash eight OP2 amplitude may be highly positively correlated with the x-axis and therefore any sample with high flash eight OP2 amplitude will receive a high ordination on the x-axis. NMDS of OP parameters both in the time domain and frequency domain provided a visual aid in examination of disease progress as well as an interpretation of OP parameters and their impact on disease.

4.7 NMDS of hqY mice reveals distinct functional retinal changes by two months of age

NMDS of time-domain OP parameters reveals that young, hqY mice (two months of age) cluster distinctly from wild-type mice and demonstrate early characteristics of retinal dysfunction. Through axis interpretation with Pearson's correlation coefficients it is evident that young, hqY mice (two months of age) cluster in regions of low, early OP amplitudes (OP2-OP3) as well as decreasing initial latencies across all flash intensities. Age-matched wild-type mice cluster in regions of high, late OP amplitudes (OP4-OP5) at all flash intensities as well as short initial latencies across all flash intensities. It is therefore apparent that mechanisms of the hq disease lead to early OP latency increases and amplitude loss followed by both latency increases and amplitude reductions across all OPs later in life. This is consistent with previous findings indicating that hq mice experience ONL losses at four months of age.^{19,93} However, this evidence suggests that functional changes may occur as early as two months in the hq photoreceptors and INL cells. An interesting observation is that ten-month-old wild-type mice cluster similarly to that of two-month-old hq mice indicating that young hq mice show retinal functional losses similar to that of middle-age wild-type mice. Ageing has been shown to affect rod photoreceptor function leading to a 40% reduction in mouse rods.¹⁴⁷ This suggests that hq mice exhibit ERG features similar to an advanced aging phenotype leading to photoreceptor losses and decreased INL and GCL function.

In the frequency domain, young two-month-old hqY mice cluster distinctly with reductions in power and frequency in comparison to wild-type mice across all flash intensities. Wild-type mice cluster in regions of high energy, power and frequency across all ages with minimal OP changes with age. Aging of the hq mouse demonstrates dramatic reductions in OP frequency, power and energy, particularly between two and three months of age. This is indicative of reduced radial current feedback in the hq retina due to reductions in periodicity and overall INL and GCL function resulting in reduced neural retina response and OP generation.

4.8 NMDS of hqX mice reveals distinct retinal changes at eleven months of age

NMDS of time-domain parameters indicates that middle-aged (eleven months of age) hqX mice cluster distinctly with reductions in both early and later OP amplitudes (OP2-OP5) as well as shorter overall latencies. Age-matched wild-type mice indicate slight reductions in all OP amplitudes but shortened overall latencies across all flash intensities. These functional decreases would indicate that the hqX mice lose the functionality of the overall retinal response and likely experience neurite losses in both

the OPL and IPL. Cell loss and subsequent retinal thinning does not occur in the INL and GCL until fifteen months of age in the hqX mice,¹⁰³ however functional changes are evident early through NMDS of time-domain OPs.

In the frequency domain, middle-aged hqX mice (eleven months of age) cluster distinctly with large overall reductions in OP power and energy across all flash intensities and increases in OP frequency with age. Wild-type mice of similar ages cluster in regions of sustained OP power and energy with increased frequency. Functional losses in total OP energy and power indicate a delayed and stunted response of the radial current feedback of the INL and GCL. This is prior to any structural losses and indicates an early functional diagnosis of the hqX retinal disease.

4.9 The hq mouse retina maintains an intact inner and outer blood-retinal barrier

Assessments of the blood-retinal barrier via Evans Blue circulation indicate no increase in EB concentrations in the retina of hq mice when compared to age-matched wild-type mice. Although endothelial cells of the inner and epithelial cells of the outer retina (RPE) blood barrier are highly metabolic, they retain functional tight junctions with limited diffusion of exogenous EB from the vasculature into the sensitive inner retinal space. This is consistent with the alternate hypothesis of activation of hq retinal microglial cells from an endogenous source due to diminished AIF function leading to cell stress and loss of retinal neurons. Although retinal endothelial and epithelial vasculature are rich in mitochondria and highly metabolic, a probable explanation of integrity is due to maintenance by glia.¹⁴⁸ Glial cells rely heavily on glycolytic ATP as

well as lactate and therefore a deficiency in OXPHOS will not compromise glial function.¹¹⁹ Glial cells have been shown to enhance blood-retinal barrier properties by increasing tight junction protein expression, endothelial-1,¹⁴⁹ as well as providing neuronal support¹⁵⁰ for both the outer¹⁵¹ and inner³³ retina. It would be expected that a reduction in complex I functionality and ATP production accompanying *hq* mice leads to a blood-retinal barrier malfunction but continued support from aerobic glial cells promotes the continued formation of tight junctions and maintenance of the blood-retinal barrier.

4.10 Transcriptome changes demonstrate Aif downregulation

Gene expression analysis of hq Aif is consistent with previous findings suggesting a global downregulation of Aif mRNA transcript by over 80%.⁹³ Contrary to previous findings downregulation of Aif in the retina surpasses an 80% reduction in the retina of four-, three-, and two-month-old mice to a downregulation by 91.16%, 94.22% and 91.88% respectively. This finding is consistent with quantified protein expression in the hq retina indicating minimal to nil endogenous AIF expression and complex I efficiency in the hq retina.^{95,98} Regulation of hq Aif gene expression occurs prior to transcription and specifies that the regulatory region of intron 1 is more vital in the retina than other tissues. The significant downregulation of Aif in the hq retina may explain the rapid onset of retinal degeneration preceding other tissue malfunctions. Functional retinal deficits begin at two months of age and an early reduction of Aif mRNA expression may be the catalyst to early retinal degeneration in the hq mouse.

4.11 The hq retinal transcriptome suggests alternative metabolism

As functional losses occur in the ONL of the hq retina, it is important to characterize endogenous activation of microglial cells and changes that lead to cell loss. The loss of *Aif* efficacy leads to reduced complex I function and expression resulting in reduced ATP yield across a variety of tissues.^{95,98,118} As predicted, the hq retina at two-, three- and four-months of age are unable to upregulate early, key enzymatic reactions of the glycolytic cycle as demonstrated by *Hexokinase I* (*Hk1*). *Hk1* catalyzes the first regulatory, enzymatic reaction of glycolysis in the cytoplasm by phosphorylating glucose to glucose 6-phosphate (Figure 4.2A).¹⁵² Overexpression of *Hk1* drives high rates of aerobic glycolysis, largely in cancers.¹⁵³ *Hk1* demonstrated no change in transcript levels in the *hq* retina even after microarray data suggested it may be upregulated.

Previous studies suggest the role of increased glucose uptake in hq mesodermal/endodermal tissues as a result of OXPHOS deficiency and a reliance on glycolytic ATP.^{98,118} Glycolytic flux is believed to be primarily controlled at the level of glucose uptake and phosphorylation of glucose by Hkl.¹⁵⁴ In agreement with no glycolytic changes in the hq retina, the expression of *Glucose transporter 4* (*Glut4*), a glucose transporter in the retina,¹⁵⁵ demonstrates no significant changes in mRNA expression. *Glut4* is a rapid, time-scale glucose transporter that translocates to the plasma membrane from the endoplasmic reticulum when insulin binds its receptors on the cell surface (Figure 4.2A).¹⁵⁶ This translocation is mediated by *TBC1 Domain Family*, *Member 4* (*TBC1D4*)¹⁵⁷ which also showed no regulation changes in the hq retina at any age. The regulation of *Glut* and *Hk1* exhibit the high control over glycolytic flux¹⁵⁴ and

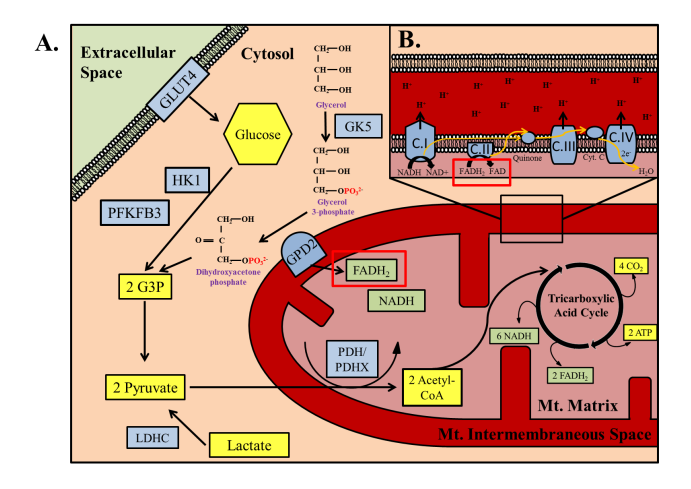


Figure 4.2. Proposed mechanisms of altered metabolism in the hq mouse based on significant alterations in the hq retinal transcriptome. Boxes coloured: blue indicate enzymes, yellow indicate substrates/metabolites and green indicate reducing equivalents. (A) Glucose metabolism in the hq retina, beginning with the uptake of glucose from the extracellular space by GLUT4 and followed by regulatory steps of glycolysis (HK1, PFKFP3) remain essentially unchanged. Metabolism of glycerol in the hq retina is differentially upregulated beginning with the conversion of glycerol to glycerol 3phosphate by GK5. Mitochondrial bound GPD2 facilitates the conversion of glycerol 3phosphate to dihydroxyacetone phosphate by reducing FAD to FADH2. The glycerol shuttle shuttles FADH2 into the mitochondria. In addition, the formation of dihydroxyacetone phosphate from glycerol metabolism feeds into glycolysis and through the formation of glyceraldehyde 3-phosphate (G3P). The dysfunction of complex I in the hq retina causes an accumulation of NADH and decreased ATP production. (B) FADH2, produced by the glycerol shuttle, can donate electrons distal to complex I and allow improved functionality of OXPHOS in the hq retina.

show no differential expression in the hq retina (Figure 4.2A). These observations support no upregulation of transcript levels of glycolytic compensatory mechanisms for neuronal ATP deficits cause by AIF dysfunction. This further confirms that neurons may be unable to upregulate glycolysis as increased glycolytic rate leads to oxidative stress and apoptotic cell death.⁴⁵

Neuronal and glial cells do not preferentially metabolize triglyceride components under healthy conditions.¹⁵⁸ β -oxidation of fatty acids is not favoured in neurons because; 1) it requires high levels of oxygen thereby increasing risk of becoming hypoxic 2) it leads to generation of superoxides and 3) has a slower ATP turnover *in vivo*.¹⁵⁸ Glycerol however, can traverse the blood-brain-barrier¹⁵⁹ and be utilized as a neuronal metabolite.¹⁶⁰ A suspected similar mechanism may occur in the retina during periods of ATP depletion and metabolic need. Glycerol metabolism is highly energetic as it fuels ATP formation as well the ability to participate in the glycerol-3-phosphate (G3P) shuttle by transferring reducing agents into the mitochondrion and leading to the formation of FADH₂ (Figure 4.2AB).¹⁶⁰ The oxidative metabolism of glycerol leads to the formation of one FADH₂ reducing molecule before forming 3-carbon, dihydroxyacetone phosphate which can enter the glycolytic pathway (Figure 4.2A). *Glycerol-3-phosphate dehydrogenase* (*Gpd2*) and *Glycerol kinase* 5 (*Gk5*) encode the primary and

secondary enzymatic mediators leading to and facilitating glycerol metabolism in the brain (Figure 4.2A).¹⁶⁰ Both *Gpd2* and *Gk5* transcript levels were upregulated in the *hq* retina at four months of age inferring neuronal metabolism of glycerol as a metabolic means for ATP and FADH₂ production. GK5 facilitates the primary step of glycerol metabolism by phosphorylating glycerol to L-glycerol 3-phosphate and was upregulated

at two-months of age and suggests a potential link between early hq disease and glycerol metabolism.

Since OXPHOS of the hq mouse becomes dysfunctional, alternative mechanisms for generation of ATP must occur in order for neurons of the retina to remain viable. Excess glycerol fed into the later stages of glycolysis would lead to a bottleneck of pyruvate oxidation (pyruvate dehydrogenase) and as expected the *Pyruvate* dehydrogenase complex component X (Pdhx) of pyruvate dehydrogenase is upregulated at four months. *Pdhx* is a structural component of the pyruvate dehydrogenase complex that links the catalytic and regulatory subunits of the functional complex.¹⁶¹ Pyruvate dehydrogenase complex is the metabolic link between glycolysis and the Tricarboxylic (TCA) acid cycle by decarboxylating pyruvate to form acetyl-CoA (Figure 4.2A). Deficiencies in Pdhx cause pyruvate dehydrogenase malfunction and have been associated with neurological dysfunction and lactate acidosis.¹⁶¹ Although the directionality of *Pdhx* expression based on Tagman[®] assay and microarray for *Pdhx* do not agree, dysregulation infers an intracellular mechanism that may lead to lactate acidosis in the hq retina and/or enhanced production of acetyl-CoA as a result of increased glycerol metabolism. In vivo examination of PDHX activity may help resolve its exact function in the hq retina.

OXPHOS complex I reduction in the hq mouse⁹⁸ likely results in the inadequate processing and utilization of reducing agents (NADH and FADH₂) produced by the mitochondrial TCA Cycle (Tricarboxylic Acid Cycle) (Figure 4.2). This is of particular importance for NADH, which is primarily oxidized by the NADH dehydrogenase complex (complex I) of the electron transport chain. A dysfunction of NADH dehydrogenase may disrupt the NAD+ pool, increasing the NADH/NAD+ ratio thereby altering the redox state of the cell.¹⁶² In 80% of patients with complex I deficiency, high levels of lactate/pyruvate were evident with severe increases in NADH/NAD+ ratios.¹⁶³ Since it is likely that *hq* OXPHOS has compromised utilization of NADH, the mitochondria may compensate by decreasing the throughput of the TCA Cycle (high NADH concentrations decrease the activity of malate dehydrogenase, the primary step of the TCA cycle¹⁶⁴) and increase glycerol metabolism as a means to increase FADH₂ production. FADH₂ electrons enter the electron transport chain distal to complex I and therefore can be used as an alternative electron source for driving the electron transport chain in order to maintain its function.¹⁶⁵

Severe increases in NADH/NAD+ are also associated with ketoacidosis and lactate acidosis as a result of acetyl-CoA being converted into acetoacetic acid and beta-hydroxybutyric acid or lactate.^{166,167} Neuronal excitability is affected by intracellular acid-base homeostasis,¹⁶⁸ where a rise in pH is associated with increased neuronal excitability, and a drop in pH reducing neuronal excitability.^{169,170} Acidic buffering to maintain pH, decreases intracellular concentration of HCO_3^- and further reduces buffering capabilities of the cell.¹⁷¹ Consequently, two key intracellular pH regulators and bicarbonate transporters are transcriptionally upregulated in the *hq* retina at four-months. *Solute carrier family 4, sodium bicarbonate transporter, member 10 (Slc4a10)* is a sodium-coupled bicarbonate transporter that exploits the transmembrane Na⁺ gradient in neurons to uptake HCO_3^- and reduces acid load.¹⁷² Adenylate cyclase 10 (Adcy10) is believed to be a universal, evolutionarily conserved bicarbonate sensor.¹⁷³ Unlike other intracellular pH regulators, *Adcy10* functions as a bicarbonate sensor rather than a proton

sensor which allows a more direct regulation of bicarbonate to maintain pH homeostasis.¹⁷⁴ Almost every cellular protein is sensitive to fluctuations in pH, however ADCY10 exhibits a unique ability to remain functional during extreme pH fluctuations.¹⁷³ In addition, ADCY10 exhibits the ability to regulate the TCA cycle. Discrepancy between generation and utilization of reducing agents (NADH and FADH₂) produced mainly by the TCA cycle, leads to ROS production and becomes harmful for the cell.¹⁷⁴ Adcy10 uses CO₂ produced by the TCA cycle to serve as a metabolic signal for TCA cycle turnover and can reduce the intra-mitochondrial cAMP concentration leading to reduced output of the TCA cycle.¹⁷⁴ Adcy10 is upregulated in both the four-month hqretina and the four-month hq cerebellum inferring an increased cellular need for pH homeostasis and TCA cycle regulation. High NADH/NAD+ ratios will also push increased lactate production which has been observed in Aif-deficient cells and tissues of the hq mouse.^{95,98,118} Expression analysis of Lactate dehydrogenase C (Ldhc), which catalyzes the reverse reaction of lactate to pyruvate, was not examined due to low transcript levels of the gene (Figure 4.2A

Previous transcriptome studies of the hq-mouse retina performed by our laboratory are consistent with ammonia toxicity and protein degradation leading to amino acid metabolism as a source for ATP generation *in vivo*.¹⁷⁵ The catabolism of amino acids is associated with the production of aldehyde by-products which can lead to glycation of amino-acids,^{176,177} and subsequent inflammatory responses. Thereby, *Aldehyde dehydrogenase 2* (*Aldh2*) is likely transcriptionally upregulated in the hq retina to eliminate toxic aldehydes as a consequence of increased amino acid catabolism.¹⁷⁸

4.12 The limitations of transcriptome analyses and future in vivo examinations

The transcriptome of the hq retina identifies many useful preliminary therapeutic targets and potential early disease mechanisms, however many limitations exist with transcriptome investigation. Transcriptional analysis occurs prior to post-transcriptional processing, translation and post-translational regulation and therefore does not reflect the quantity or functionality of the enzyme/protein in vivo. In addition, other interacting targets may be responsible for the phenotype observed. Further examinations at the level of the proteome are important for characterizing changes and complete interactions as a result of the Aif downregulation in vivo. One recommendation is the examination of key enzymatic factors of the glycerol pathway, GK5 and GPD2, to determine if increased neuronal glycerol metabolism is a feasible mechanism for FADH₂ production. In addition to proteome analysis, gas-chromatography mass spectrophotometry could be used to determine the *in vivo* concentrations of metabolites such as NADH, NAD+, lactate, pyruvate, dihydroxyacetone phosphate, and FADH₂ in comparison to the wild type. This will give us better insight into the mechanisms that occur in the hq retina to cope with NADH dehydrogenase dysfunction and reduced ATP production.

4.13 Preliminary suggestions for hq mitochondrial dysfunction therapy

In addition to transcriptome quantification, a preliminary study of metabolic rate suggests that hq mice are hyperphagic and require increased food consumption. This is likely the result of reduced OXPHOS function causing a reduced ratio of ATP produced per molecule of glucose. hq mice are resistant to progressive weight gain and lipid

accumulation when fed a high fat diet inferring potential reliance and utilization of triglycerides as predicted.¹¹⁸ This is likely due to the fact that fatty acids and glycerol yield a much higher FADH₂ and NADH than glucose and are commonly used for treatment of complex I deficiencies.¹⁶²

From a therapeutic point of view, treatment of hq mice must take a direct approach to correcting mitochondrial dysfunction. Increased glycerol metabolism in the hq mouse predicts an increased reliance on FADH₂ as the primary reducing agent in the mitochondria. The benefits' of a high fat diet are likely to reduce the severity of the ha phenotype, although the benefits are unlikely to provide continued treatment due to NADH accumulation in the cell. Thereby, a probable treatment may be in the form of Riboflavin (vitamin B₂). Riboflavin is a flavoprotein precursor and is the central component of FAD. Treatment with riboflavin will allow the cell to produce more FADH₂ and facilitate the enzymatic reactions of the TCA cycle.¹⁷⁹ Complex I and II deficiencies show betterment of symptoms and delayed disease progression following treatment with riboflavin.^{180–182} In addition, riboflavin has been used to treat human patients with OXPHOS deficiency caused by an Aif amino-acid deletion (R201 del).¹⁸³ Riboflavin treatment improves neurological symptoms and increases the functionality of OXPHOS in these patients.¹⁸³ Although the hq disease is associated with a severe downregulation of Aif as opposed to a deletion of an amino acid, riboflavin supplementation demonstrates a promising preliminary therapeutic treatment for improving the *hq* disease.

4.14 Conclusions

Retinal degeneration in the hq mouse is characterized by mitochondrial dysfunction leading to parainflammation and closely mimics aging and mitochondrial-associated retinal degeneration. Early disease mechanisms resulting in degeneration of the hq retina are associated with early neurite losses and photoreceptor cell losses that can be characterized by diagnostic oscillatory potentials. This study has developed a novel framework for the analysis of the entire OP profile across multiple stimuli, allowing functional examination of all retinal cell types and their health when combined with awave and b-wave analyse. We have identified diagnostic ERG markers associated with the hq disease as well as a pipeline for identifying diagnostic ERG markers in other models of retinal disease. Parainflammation in the hq retina is more likely the result of endogenous retinal AIF deficiency and transcriptome alterations are consistent with compensatory mechanisms of ATP production such as glycerol metabolism. Studies aimed at a thorough characterization of glycerol metabolism as well as the accumulation of specific primary metabolites will help elucidate the role of AIF and bring us closer to understanding mitochondrial-associated retinal degeneration.

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Appendix A. The Canadian Council on Animal Care and The Animal Use Subcommittee of The University Council on Animal Care approval for animal use in research.

This appendix contains a copy of the 2012 annual approval (**A**) for animal use by the Hill laboratory as well as the 2013 annual approval (**B**) for animal use by the Hill laboratory. Approved by The Canadian Council on Animal Care and The Animal Use Subcommittee of The University Council on Animal Care 2009-033::3:

AUP Number: 2009-033 AUP Title: Mutational Mechanisms: Relevance and Role of Oxidative Stress

Approval Date: 08/18/2009

The YEARLY RENEWAL to Animal Use Protocol (AUP) 2009-033 has been approved.

- 1. This AUP number must be indicated when ordering animals for this project.
- 2. Animals for other projects may not be ordered under this AUP number.
- Purchases of animals other than through this system must be cleared through the ACVS office. Health certificates will be required.

REQUIREMENTS/COMMENTS

Please ensure that individual(s) performing procedures on live animals, as described in this protocol, are familiar with the contents of this document.

The holder of this Animal Use Protocol is responsible to ensure that all associated safety components (biosafety, radiation safety, general laboratory safety) comply with institutional safety standards and have received all necessary approvals. Please consult directly with your institutional safety officers.

А.

AUP Number: 2009-033

AUP Title: Mutational Mechanisms

Approval Date: 09/04/2013

Official Notice of Animal Use Subcommittee (AUS) Approval: Your new Animal Use Protocol (AUP) entitled "Mutational Mechanisms

* has been APPROVED by the Animal Use Subcommittee of the University Council on Animal Care. This approval, although valid for four years, and is subject to annual Protocol Renewal. 2009-033::5

- 1. This AUP number must be indicated when ordering animals for this project.
- 2. Animals for other projects may not be ordered under this AUP number.
- 3. Purchases of animals other than through this system must be cleared through the ACVS office. Health certificates will be required.

The holder of this Animal Use Protocol is responsible to ensure that all associated safety components (biosafety, radiation safety, general laboratory safety) comply with institutional safety standards and have received all necessary approvals. Please consult directly with your institutional safety officers.

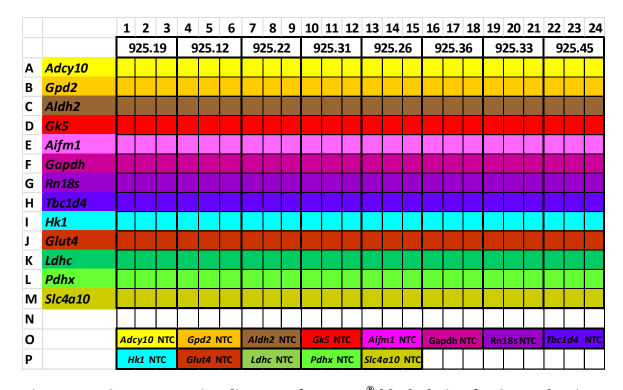
B.

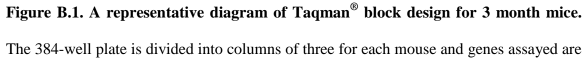
Appendix B. Supplementary Figures and Tables

This appendix contains figures and tables to assist with understanding of presented material.

Primer Position	Sequence (5' to 3')	Amplicon Length (bp)
Forward	AGTGTCCAGTCAAAGTACCGGG	537 (WT)
Proviral Forward	GAACAAGGAAGTACAGAGAGAGGC	725 (hq)
Reverse	CTATGCCCTTCTCCATGTAGTT	

Table B.1. PCR Primers used for Aif Genotyping





represented by rows. A total of 13 genes were assayed per mouse in triplicate. A notemplate control (NTC) for each gene was assayed in rows O & P to ensure no false amplification and was also assayed in triplicate.

Table B.2. Pearson's correlation coefficients from nonmetric multidimensional scaling analysis of hqY time-domain OP parameters. Tables are ordered by ascending Coordinate 1 value (left chart) and ascending Coordinate 2 value (right chart) to assist with axis interpretation.

Measurement	Coordinate 1	Coordinate 2	Measurement	Coordinate 1	Coordinate 2
F11 Latency OP2	-0.859006334	-0.151377761	F9 Amplitude OP2	0.560034669	-0.66660931
F11 Latency OP3	-0.853657492	-0.15753476	F10 Amplitude OP2	0.543878698	-0.644420042
F9 Latency OP3	-0.846274465	-0.169707181	F9 Amplitude OP3	0.729042222	-0.615488338
F10 Latency OP2	-0.841057093	-0.193566945	F11 Latency OP4	-0.638909514	-0.589130854
F8 Latency OP3	-0.830526046	-0.065715862	F8 Amplitude OP3	0.599566532	-0.557316795
F10 Latency OP3	-0.822981718	-0.208552355	F8 Amplitude OP4	0.684464179	-0.552811113
F9 Latency OP2	-0.82194351	-0.125478715	F11 Latency OP5	-0.468400077	-0.541219708
F9 Latency OP4	-0.80694756	-0.298008836	F10 Latency OP4	-0.684617696	-0.529856532
F8 Latency OP4	-0.799695375	-0.207521146	F8 Amplitude OP2	0.618144453	-0.461141832
F8 Latency OP2	-0.742677191	-0.101943569	F10 Amplitude OP3	0.812086192	-0.448484634
F9 Latency OP5	-0.733660057	-0.250762361	F11 Amplitude OP2	0.424307269	-0.412374408
F10 Latency OP5	-0.685039837	-0.356267863	F8 Amplitude OP5	0.672790759	-0.40557432
F10 Latency OP4	-0.684617696	-0.529856532	F10 Latency OP5	-0.685039837	-0.356267863
F11 Latency OP4	-0.638909514	-0.589130854	F9 Amplitude OP4	0.728464308	-0.343319214
F11 Latency OP5	-0.468400077	-0.541219708	F9 Latency OP4	-0.80694756	-0.298008836
F8 Latency OP5	-0.344040045	-0.265368694	F8 Latency OP5	-0.344040045	-0.265368694
F11 Amplitude OP2	0.424307269	-0.412374408	F9 Latency OP5	-0.733660057	-0.250762361
F10 Amplitude OP5	0.481785605	0.164778075	F10 Latency OP3	-0.822981718	-0.208552355
F10 Amplitude OP2	0.543878698	-0.644420042	F8 Latency OP4	-0.799695375	-0.207521146
F9 Amplitude OP2	0.560034669	-0.66660931	F10 Latency OP2	-0.841057093	-0.193566945
F11 Amplitude OP5	0.578306152	0.186810407	F11 Amplitude OP3	0.870065089	-0.178484534
F8 Amplitude OP3	0.599566532	-0.557316795	F9 Latency OP3	-0.846274465	-0.169707181
F8 Amplitude OP2	0.618144453	-0.461141832	F11 Latency OP3	-0.853657492	-0.15753476
F9 Amplitude OP5	0.65045626	-0.111703757	F11 Latency OP2	-0.859006334	-0.151377761
F8 Amplitude OP5	0.672790759	-0.40557432	F9 Latency OP2	-0.82194351	-0.125478715
F8 Amplitude OP4	0.684464179	-0.552811113	F9 Amplitude OP5	0.65045626	-0.111703757
F9 Amplitude OP4	0.728464308	-0.343319214	F8 Latency OP2	-0.742677191	-0.101943569
F9 Amplitude OP3	0.729042222	-0.615488338	F8 Latency OP3	-0.830526046	-0.065715862
F10 Amplitude OP4	0.748886709	0.043278448	F10 Amplitude OP4	0.748886709	0.043278448
F11 Amplitude OP4	0.799065572	0.162761807	F11 Amplitude OP4	0.799065572	0.162761807
F10 Amplitude OP3	0.812086192	-0.448484634	F10 Amplitude OP5	0.481785605	0.164778075
F11 Amplitude OP3	0.870065089	-0.178484534 _.	F11 Amplitude OP5	0.578306152	0.186810407

Table B.3. Pearson's correlation coefficients from nonmetric multidimensional scaling analysis of hqY frequency-domain OP parameters. Tables are ordered by ascending Coordinate 1 value (left chart) and ascending Coordinate 2 value (right chart) to assist with axis interpretation.

Measurement	Coordinate 1	Coordinate 2	Measurement	Coordinate 1	Coordinate 2
F10 Total Energy	-0.97963505	-0.080079919	F9 Gaussian Power	-0.590242553	-0.770721391
F11 Total Energy	-0.968656764	0.088457664	F8 Gaussian Power	-0.439755859	-0.745030923
F9 Total Energy	-0.938426703	-0.236289954	F10 Gaussian Power	-0.735730899	-0.563367817
F11 Gaussian Power	-0.837221842	-0.260841987	F8 Total Energy	-0.817230902	-0.292346888
F10 Gaussian			F11 Gaussian Power	-0.837221842	-0.260841987
	-0.821513771	0.543230455	F9 Total Energy	-0.938426703	-0.236289954
Frequency			F10 Total Energy	-0.97963505	-0.080079919
F8 Total Energy	-0.817230902	-0.292346888	F11 Total Energy	-0.968656764	0.088457664
F9 Gaussian			F10 Gaussian		
Frequency	-0.810590977	0.546777937	Frequency	-0.821513771	0.543230455
F8 Gaussian			F9 Gaussian		
Frequency	-0.784149192	0.563671382	Frequency	-0.810590977	0.546777937
Frequency	-0.776138181	0.594753436	F8 Gaussian		
F10 Gaussian Power	-0.735730899	-0.563367817	Frequency	-0.784149192	0.563671382
F9 Gaussian Power	-0.590242553	-0.770721391	F11 Gaussian		
F8 Gaussian Power	-0.439755859	-0.745030923	Frequency	-0.776138181	0.594753436

Table B.4. Pearson's correlation coefficients from nonmetric multidimensional scaling analysis of hqX frequency-domain OP parameters. Tables are ordered by ascending Coordinate 1 value (left chart) and ascending Coordinate 2 value (right chart) to assist with axis interpretation.

Measurement	Coordinate 1	Coordinate 2	Measurement	Coordinate 1	Coordinate 2
F10 Latency OP2	-0.8465227	-0.33263146	F9 Amplitude OP4	0.37176222	-0.87729291
F9 Latency OP2	-0.84122833	-0.26833499	F10 Amplitude OP4	0.3800024	-0.8572363
F10 Latency OP3	-0.8286985	-0.28526331	F8 Amplitude OP4	0.42295653	-0.74131346
F11 Latency OP3	-0.7839018	-0.39646655	F8 Amplitude OP3	0.61054609	-0.72685793
F9 Latency OP4	-0.78382521	-0.21714434	F9 Amplitude OP5	0.33357947	-0.71742454
F10 Latency OP5	-0.77789174	-0.07591251	F10 Amplitude OP5	0.12469163	-0.71649367
F8 Latency OP2	-0.75190675	-0.47672993	F11 Amplitude OP5	0.1437659	-0.68752732
F8 Amplitude OP3	-0.74317829	-0.22739861	F11 Amplitude OP4	0.48735872	-0.64154307
F11 Latency OP2	-0.73761625	-0.47558814	F11 Amplitude OP3	0.76840952	-0.57446596
F8 Amplitude OP4	-0.73351111	-0.04430713	F11 Latency OP4	-0.63576058	-0.55236009
F9 Latency OP5	-0.72513157	-0.46055913	F8 Amplitude OP5	0.50542379	-0.54592269
F10 Latency OP4	-0.67806932	-0.06781597	F9 Amplitude OP2	0.68950648	-0.52237812
F11 Latency OP4	-0.63576058	-0.55236009	F10 Amplitude OP3	0.64286454	-0.50677443
F9 Latency OP3	-0.61737348	-0.35646637	F8 Latency OP2	-0.75190675	-0.47672993
F8 Amplitude OP5	-0.59553985	-0.2987983	F11 Latency OP2	-0.73761625	-0.47558814
F11 Latency OP5	-0.27739875	-0.23479761	F9 Latency OP5	-0.72513157	-0.46055913
F10 Amplitude OP5	0.12469163	-0.71649367	F11 Amplitude OP2	0.78171173	-0.41812742
F11 Amplitude OP5	0.1437659	-0.68752732	F11 Latency OP3	-0.7839018	-0.39646655
F9 Amplitude OP5	0.33357947	-0.71742454	F10 Amplitude OP2	0.76133209	-0.36759832
F9 Amplitude OP4	0.37176222	-0.87729291	F9 Amplitude OP3	0.59040284	-0.36471528
F10 Amplitude OP4	0.3800024	-0.8572363	F9 Latency OP3	-0.61737348	-0.35646637
F8 Amplitude OP4	0.42295653	-0.74131346	F8 Amplitude OP2	0.65078245	-0.34984908
F11 Amplitude OP4	0.48735872	-0.64154307	F10 Latency OP2	-0.8465227	-0.33263146
F8 Amplitude OP5	0.50542379	-0.54592269	F8 Amplitude OP5	-0.59553985	-0.2987983
F9 Amplitude OP3	0.59040284	-0.36471528	F10 Latency OP3	-0.8286985	-0.28526331
F8 Amplitude OP3	0.61054609	-0.72685793	F9 Latency OP2	-0.84122833	-0.26833499
F10 Amplitude OP3	0.64286454	-0.50677443	F11 Latency OP5	-0.27739875	-0.23479761
F8 Amplitude OP2	0.65078245	-0.34984908	F8 Amplitude OP3	-0.74317829	-0.22739861
F9 Amplitude OP2	0.68950648	-0.52237812	F9 Latency OP4	-0.78382521	-0.21714434
F10 Amplitude OP2	0.76133209	-0.36759832	F10 Latency OP5	-0.77789174	-0.07591251
F11 Amplitude OP3	0.76840952	-0.57446596	F10 Latency OP4	-0.67806932	-0.06781597
F11 Amplitude OP2	0.78171173	-0.41812742	F8 Amplitude OP4	-0.73351111	-0.04430713

Table B.5. Pearson's correlation coefficients from nonmetric multidimensional scaling analysis of hqX frequency-domain OP parameters. Tables are ordered by ascending Coordinate 1 value (left chart) and ascending Coordinate 2 value (right chart) to assist with axis interpretation.

Measurement	Coordinate 1	Coordinate 2	Measurement	Coordinate 1	Coordinate 2
F9 Total Energy	-0.969372022	0.053267539	F8 Gaussian Power	-0.778672152	-0.477976579
F10 Total Energy	-0.96440625	0.209308386	F9 Gaussian Power	-0.852374157	-0.462860557
F11 Gaussian Power	-0.947405504	0.019228962	F10 Gaussian Power	-0.936559689	-0.249515924
F10 Gaussian Power	-0.936559689	-0.249515924	F11 Gaussian Power	-0.947405504	0.019228962
F8 Total Energy	-0.925313065	0.134905804	F9 Total Energy	-0.969372022	0.053267539
F11 Total Energy	-0.898996534	0.364182853	F8 Total Energy	-0.925313065	0.134905804
F9 Gaussian Power	-0.852374157	-0.462860557	F10 Total Energy	-0.96440625	0.209308386
F8 Gaussian Power	-0.778672152	-0.477976579	F11 Total Energy	-0.898996534	0.364182853
F8 Gaussian			F8 Gaussian		
Frequency	-0.103790027	0.939471085	Frequency	-0.103790027	0.939471085
F9 Gaussian			F9 Gaussian		
Frequency	-0.088637167	0.951935284	Frequency	-0.088637167	0.951935284
F10 Gaussian			F11 Gaussian		
Frequency	-0.062301485	0.980003901	Frequency	-0.045683178	0.969829489
F11 Gaussian			F10 Gaussian		
Frequency	-0.045683178	0.969829489	Frequency	-0.062301485	0.980003901

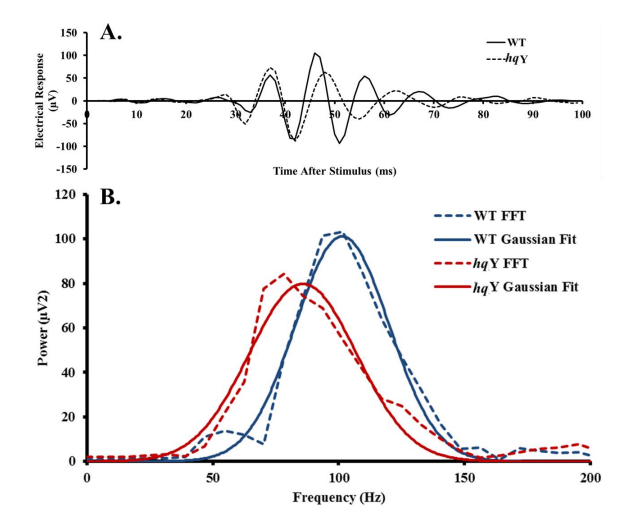


Figure B.2. A schematic representation of OP parameters elicited by a three month hqY and wild-type mouse (10 cd·s/m² stimulus). (A) Time-domain representation of ERG OP parameters indicate functional changes as early as three months of age with changes in OP periodicity. (B) Frequency-domain representation of ERG OP parameters at three months indicate functional changes in OP frequency. It is evident that power, and total energy are decreased in this schematic representation.

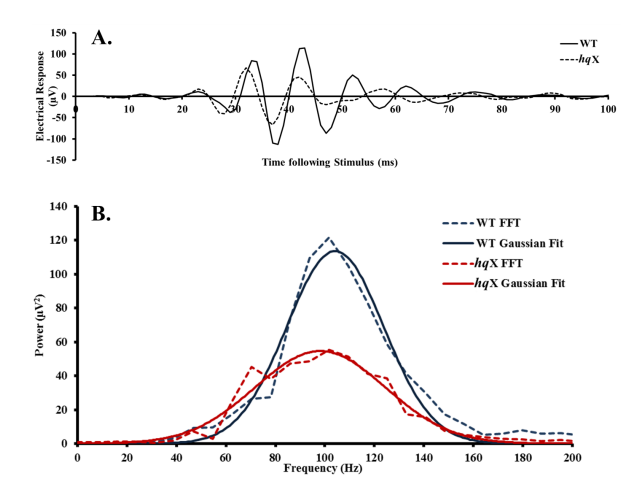


Figure B.3. A schematic representation of OP parameters elicited by an eleven month hqX and wild-type mouse (10 cd·s/m² stimulus). (A) Time-domain representation of ERG OP parameters indicate functional changes at eleven months of age with changes in OP amplitude. (B) Frequency-domain representation of ERG OP parameters at eleven months indicate functional changes in OP total energy and power. It is evident that OP frequency remains unchanged in this schematic representation.

Table B.6. Two-way ANOVA's performed in Excel [Microsoft, Redmond, VA] on oscillatory potentials parameters both in the time and frequency domain to explore the effect of age on genotype. Significance is equal to p < 0.05.

Α.	
T T •	

hqY Time-Domain Parameters					
Initial Latency	Summed Amplitude				
Genotype (p < 0.001)	Genotype (p < 0.001)				
Age (p = 0.07337)	Age = $(p < 0.001)$				
Interaction $(p = 0.212)$	Interaction (p < 0.01)				

B.

hqY Frequency-Domain Parameters							
Peak Frequency Peak Power Total Energy							
Genotype (p < 0.001)	Genotype (p < 0.05)	Genotype (p < 0.01)					
Age (p < 0.001)	Age (p < 0.01)	Age (p < 0.01)					
Interaction (p < 0.05)	Interaction (p < 0.01)	Interaction (p < 0.01)					

С.

hqX Time-Dom	hqX Time-Domain Parameters						
Initial Latency	Summed Amplitude						
Genotype (p < 0.01)	Genotype (p < 0.001)						
Age (p < 0.001)	Age $(p = 0.63994)$						
Interaction $(p = 0.212)$	Interaction $(p = 0.15089)$						

D.

hqX Frequency-Domain Parameters								
Peak Frequency Peak Power Total Energy								
Genotype (p < 0.01)	Genotype (p < 0.001)	Genotype (p < 0.001)						
Age (p < 0.001)	Age (p < 0.01)	Age $(p = 0.7796)$						
Interaction $(p = 0.88369)$	Interaction ($p = 0.16913$)	Interaction ($p = 0.13802$)						

cerebella			Retina				Cerebel	lum	
		Sample ID	Concentration (ng/µL)	A260/A280	RIN	Sample ID	Concentration (ng/µL)	A260/A280	RIN
		*924.3	33.9	2.06	9.6	*924.3	257.2	2.028	9.3
		*925.16	41.7	2.04	9.6	*925.16	185.6	2.053	9.2
		*925.77	44.9	2.03	9.5	*925.77	232	2.064	8.8
	hq	*925.83	72.2	2.21	8.9	925.83	287.6	2.043	8.7
		925.89	n/a	n/a	n/a	*925.89	231.2	2.141	8.8
		925.90	24	2.1	9.1	925.90	n/a	n/a	n/a
		*925.93	72.3	2.13	9.2	925.93	n/a	n/a	n/a
4 Month		924.4	n/a	n/a	n/a	*924.4	273.2	1.957	9.3
		*924.5	16.3	2.43	8.9	924.5	n/a	n/a	n/a
		*924.11	23.7	2.01	9.3	924.11	n/a	n/a	n/a
	wт	*925.14	34.1	2.02	9.2	*925.14	140	1.977	8.8
	VV I	925.72	n/a	n/a	n/a	925.72	146	2.086	9.1
		*925.73	29.1	2.18	8.8	*925.73	140	2.077	8.9
		925.75	12.1	2.59	8.4	*925.75	210	2.315	9.2
		*925.95	41.1	2.11	9.1	925.95	n/a	n/a	n/a
		*925.19	32.7	2.15	8.8	*925.19	212	2.008	8
		*925.22	16.2	1.85	8.2	*925.22	210	2.05	9.1
	hq	*925.26	32	2.04	9.6	925.26	216	2.050	7.5
		925.27	35.6	2.05	9	*925.27	240	2.040	8.2
3 Month		*925.33	33.6	2.02	9.8	*925.33	154	1.873	9.1
5 Monun		*925.12	43.9	2.05	8.9	*925.12	353.6	2.007	7.7
		925.13	12.8	2.23	None	925.13	240	1.911	7.6
	WT	*925.31	53.4	2.03	9.8	*925.31	84	1.812	8.8
		*925.36	29.6	2.1	9.4	*925.36	214	2.030	8.3
		*925.45	47.1	2.18	9.9	*925.45	85	2.095	9.2
		*925.53	26.6	2.12	9.5	*925.53	215	2.011	8.8
		925.54	8.2	2.21	9.2	*925.54	182	1.97	8.8
	hq	*925.55	52.9	2.11	9.6	925.55	199.6	1.965	9.3
	1	*925.69	27	2.13	9.2	*925.69	221.6	1.965	8.9
		*925.70	50.2	2.1	9.6	*925.70	167.2	2.133	9.1
2 Month		*925.39	44.4	2.1	9.8	925.39	110	2.202	8.9
		*925.43	31	2.18	9.6	*925.43	310	2.102	8.5
	WT	*925.51	32.3	2.03	9.4	*925.51	194	2.008	9
		*925.58	36.1	2.15	9.1	*925.58	188	1.983	8.6
		925.57	17	2.15	8.6	*925.57	209.2	2.135	9
	L	743.31	1/	2.30	0.0	- 743,31	209.2	2.133	7

 Table B.7. Concentrations and quality assurance measures for extracted retinal and

 cerebellar RNA samples

*Indicates sample that was chosen for transcriptome analysis. In addition, light shadings indicate that the sample was not used.

Appendix C. MATLAB functions and Scripts

This appendix contains MATLAB scripts for the isolation of oscillatory potentials from the original ERG waveform and fast Fourier transformation. MATLAB Script A. "P3fit" function written into MATLAB for modelling photoreceptor contribution to the ERG.

```
function err = p3fit(rktd)
global datatofit timestofit
options.MaxFunEvals=1000000;
i = find(timestofit<rktd(3));
model(i) = 0;
j = find(timestofit>=rktd(3));
model(j) = rktd(1)*(1-exp(-rktd(2)*(timestofit(j)-rktd(3)).*(timestofit(j)-rktd(3)));
err = sum((model' - datatofit).^2);
```

MATLAB Script B. "P3driver" function written into MATLAB for modelling photoreceptor contribution to the ERG, followed by a fifth order Butterworth transformation (65-300Hz) for oscillatory potential waveform extraction.

```
global datatofit timestofit
options.MaxFunEvals=1000000;
for icol = 16:19
clear model
%modelling the p3
%details
minval = min(data(:,icol));
imin = find(data(:,icol) == minval);
iremain = data(imin+1:200,icol);
timestofit = data(1:imin,1);
datatofit = data(1:imin,icol);
datatofit = datatofit - datatofit(1);
initialguess = [ min(datatofit) 0.5 0];
rktd = fminsearch(@p3fit,initialguess);
i = find(timestofit<rktd(3));</pre>
model(i) = 0;
j = find(timestofit>=rktd(3));
model(j)
          =
                     rktd(1)*(1-exp(-rktd(2)*(timestofit(j)-
rktd(3)).*(timestofit(j)-rktd(3))));
k = (find(iremain)+(imin));
model(k) = min(model);
```

```
timestoplot(i) = i;
timestoplot(j) = j;
timestoplot(k) = k;
%Subtracting model from data to form p2
p2(i) = ((data(i,icol)-data(1,icol)) - model(i)');
p2(j) = ((data(j,icol)-data(1,icol)) - model(j)');
p2(k) = ((data(k, icol) - data(1, icol)) - model(k)');
%Butterworth filtering the p2
Wn = [65 \ 300] / 500;
[B,A] = butter(5,Wn);
fildata = filter(B, A, p2);
%Plots to see how things are going
figure
subplot(2,1,1),hold on,
plot(data(:,1), data(:,icol)-data(1,icol));
plot(timestoplot,model,'r');
plot(timestoplot,p2,'g');
plot(timestoplot,fildata,'black');
subplot(2,1,2),plot(timestoplot,fildata,'black');
```

end

MATLAB Script C. "P3driverFFT" function written into MATLAB for modelling photoreceptor contribution to the ERG, followed by a fifth order Butterworth transformation (65-300Hz) and fast Fourier transformation (Gaussian).

```
global datatofit timestofit
options.MaxFunEvals=1000000;
for icol = 16;
clear model
%modelling the p3
%details
minval = min(data(:,icol));
imin = find(data(:,icol) == minval);
iremain = data(imin+1:200,icol);
timestofit = data(1:imin,1);
datatofit = data(1:imin,icol);
datatofit = datatofit - datatofit(1);
initialguess = [ min(datatofit) 0.5 0];
rktd = fminsearch(@p3fit,initialguess);
i = find(timestofit<rktd(3));</pre>
model(i) = 0;
j = find(timestofit>=rktd(3));
                      rktd(1)*(1-exp(-rktd(2)*(timestofit(j)-
model(j)
             =
rktd(3)).*(timestofit(j)-rktd(3))));
k = (find(iremain) + (imin));
model(k) = min(model);
timestoplot(i) = i;
timestoplot(j) = j;
timestoplot(k) = k;
Subtracting model from data to form p2
p2(i) = ((data(i,icol)-data(1,icol)) - model(i)');
p2(j) = ((data(j,icol)-data(1,icol)) - model(j)');
p2(k) = ((data(k, icol) - data(1, icol)) - model(k)');
%Butterworth filtering the p2
Wn = [65 \ 300] / 500;
[B,A] = butter(5,Wn);
fildata = filter(B, A, p2);
```

```
fs = 1000;
                                    % Sample frequency (Hz)
t = 0:1/fs:0.2-1/fs;
                                    % 10 sec sample
x = (fildata);
b = fildata(1:128);
                            % Window length
m = length(b);
n = pow2(nextpow2(m)); % Transform length
y = fft(x, n);
                       % DFT
f = (0:n-1)*(fs/n); % Frequency range
p = sqrt(y.*conj(y)/n); % Power of the DFT
plot(f(1:floor(n/2)), p(1:floor(n/2)))
xlabel('Frequency (Hz)')
ylabel('Power')
title('{\bf Periodogram}')
A=((p(1)/2 + sum (p(2:end-1) + p(end)/2))/2) % area under
the curve
%Plots to see how things are going
%figure
%subplot(2,1,1),hold on,
%plot(data(:,1), data(:,icol)-data(1,icol));
%lot(timestoplot,model,'r');
%plot(timestoplot,p2,'g');
%plot(timestoplot,fildata,'black');
%subplot(2,1,2),plot(timestoplot,fildata,'black');
g = fit(f.', p.', 'gauss2')
plot(g)
end
```

Appendix D. Non-metric multidimensional scaling R code.

This appendix contains R code for performing non-metric multidimensional scaling on oscillatory potential parameters in both the time and frequency domain.

NMDS R-code A. Non-metric multidimensional scaling performed by calculating Euclidean distances between samples followed by subsequent correlation analysis.

```
> getwd()
# This will list the location that R will read files from
> list.files()
#List of all files in the working directory
> data <- read.csv("ZSCORES HQY&XY FFT DATA.csv")</pre>
# Information in file will become a variable named "data"
> data
# View "data" variable
> (D <- dist(data))
  This will calculate Euclidean distance between each
#
sample
        based
                 on
                      input
                              z-scores.
                                          Presented as
                                                            а
#dissimilarity matrix, assigned as variable D.
> summary (D)
# Displays dissimilarity matrix
> library ("MASS")
# Reads in library package to perform NMDS
# Install the HSAUR2 package
> data mds <- isoMDS(D)</pre>
# This will perform NMDS on the dissimilarity matrix (D)
based on 50 iterations. Assigns NMDS as #data mds
> x <- data mds$Points[,1]</pre>
# Assigns the first coordinate of the NMDS variables to the
variable x
> y <- data mds$Points [,2]</pre>
# Assigns the first coordinate of the NMDS variables to the
variable y
# Variables plotted in excel
# NMDS coordinates put into a new z-score file and read
into excel for correlation analysis
> x <- read.csv("ZSCORES (Coorelation) HQY&XY FFT DATA</p>
(ERIC DOLINAR).csv")
```

```
# Assigns variable x to z-score file with NMDS coordinates
> cor(x, use = "complete.obs", method = "pearson")
# Performs a Pearson's correlation on the variable x
# Shepard plots
> MDS.sh <- Shepard (D, data_mds$points)
> plot(MDS.sh, pch =".")
> plot(MDS.sh, pch =".", xlab = "Dissimilarity", ylab =
"Distance")
> lines(MDS.sh$x, MDS.sh$yf, type="S")
```

Appendix E. Functional annotation of differentially expressed retinal genes

This appendix contains functional annotations based on NCBI Gene searches¹⁸⁴ of 521 retinal genes differentially expressed in hq mice at four months of age. Differential expression is based on a retinal gene expression microarray [Affymetrix, Mouse Gene 1.0 ST Array, Santa Clara] previously performed by the Hill laboratory.¹⁰²

Gene Symbol	p-value‡	Fold Change	Annotation/Function†
Aifm1	0.000163	-6.01196	Mitochondrial intermembrane flavoprotein involved in OXPHOS and apoptosis
Cyp3a25	0.0307866	-4.68949	Cytochrome P450 variant involved in metal binding and oxidoreductase activity
Нс	0.0116071	-3.90775	Complement activation (C5) and innate immune response
Ttc18	0.0024609	-3.43071	Extracellular vesicular exosome important to ciliated cells
Trim36	0.0275573	-3.41135	Ubiquitin ligase activity with zinc-binding protein activity
Abca6	0.0450953	-3.09029	ATP-binding cassette involved in macrophage lipid metabolism and homeostasis
Iqgap2	0.0440039	-3.01678	Regulates cell morphology and mobility through interaction with the cytoskeleton
Myh6	0.0058866	-2.90362	Encodes heavy chain of myosin, an intracellular motor protein
Slc4a10	0.0266117	-2.87259	Neuronal intracellular bicarbonate transporter and pH regulator
Robo3	0.0008035	-2.81071	Controls neurite outgrowth and axonal navigation
Olfm4	0.015151	-2.77354	Antiapoptotic protein and facilitates cell adhesion
Haus6	0.0362302	-2.70602	Microtubule and kinetochore assembly
Atm	0.0358792	-2.69275	Cell cycle checkpoint kinase protein
Nbr1	0.0227154	-2.64043	Peroxisome autophagic degradation
BC030307	0.0060405	-2.62532	Unknown
Slc26a4	0.0160909	-2.60606	Sulfate and chloride transporter
Dpy19l1	0.0274067	-2.54374	Unknown
Ppp1r9a	0.0149737	-2.53199	Phosphatase I involved in cytoskeleton organization
4930519F16Rik	0.0061143	-2.51977	Unknown
Wdr60	0.0071325	-2.51121	Cell cycle regulator and formation of cilia
D19Ertd652e	0.0298091	-2.50222	Unknown
Pdhx	0.0161061	-2.48204	Structural component of pyruvate dehydrogenase, binds subunits together
Aqp4	0.0018312	-2.47148	Neuronal aquaporin to allow water-selective transport
Pkd1l2	0.0066857	-2.46823	Cation channel pore, cation transport
Siglece	0.0321588	-2.46084	Sialic acid binding protein involved in immune cell response
BC055324	0.0449079	-2.40619	Unknown
Garnl1	0.0495052	-2.38863	Activation of Ral GTPase
Smc1b	0.0466303	-2.3795	Mitotic and meiotic DNA recombination and chromatid formation
Klra5	0.0094125	-2.36588	Membrane protein involved in cellular adhesion during immune response
Pear1	0.0419941	-2.35637	Responsible for platelet aggregation

Table E.1. Functional annotation of 521 differentially expressed retinal genes based on microarray data from four-month-old hq disease mice.^{*}

Trio	0.0159855	-2.345	Promotes leukocyte transendotheial migration
Adal	0.0032844	-2.3368	Purine, drug and small molecule catabolism
Kmo	0.0428267	-2.32506	Mitochondrial protein involved in NADH oxidase activity and FAD binding
4922505G16Rik	0.0380333	-2.30153	Unknown
Mansc1	0.03965	-2.28514	Plasma membrane component
Rbms3	0.0109571	-2.27729	Cytoplasmic RNA metabolism
Golga3	0.0392264	-2.27699	Golgi apparatus transport of proteins and lipids to nucleus during mitosis
Carl	0.0257676	-2.2678	Catalyzes to reversible hydration of carbon dioxide
Hdc	0.0008771	-2.23775	Conversion of L-histidine to histamine involved in neurotransmission
Ankfy1	0.0015828	-2.23773	Cytoplasmic protein and vesicle transport
Msh3	0.0048097	-2.2069	Initiates post-replication DNA mismatch repair
Dnahc2	0.0235939	-2.20618	Involved in axonemal microtubule formation
Apafl	0.0367202	-2.19996	Cytoplasmic initiator of apoptosis through formation of apoptosome
Col7a1	0.0174913	-2.19387	Component of collagen and extracellular remodelling
Sall2	0.0137244	-2.18272	Involved in transcriptional regulation during growth and development of neurons
Katnal2	0.0307008	-2.1754	Microtubule severing through ATPase activity
Slc38a11	0.0438682	-2.17502	Plasma membrane amino acid and sodium transporter
Col4a2	0.02681	-2.16956	Component of collagen involved in inhibiting angiogenesis
Npy2r	0.0213178	-2.16915	Neuropeptide signalling activity
Rad50	0.0452278	-2.16269	DNA double-stranded break repair and cell cycle regulation
Nmbr	0.039171	-2.15422	Binds neuromedin B to promote mitosis
Tada3l	0.0106471	-2.15409	Promotes DNA-binding of transcriptional factors
Nlrp3	0.0341535	-2.15371	NF-kappa B signalling to regulated inflammation and immune response
Dhx29	0.0398495	-2.15276	Involved in mRNA translation initiation
Col9a2	0.0063438	-2.14271	Component of collagen involved in extracellular remodelling and axon guidance
Abca16	0.0127381	-2.13578	ATP-binding cassette, function unknown
Duox2	0.0052049	-2.11419	NADH oxidase activity and cytokine release
Wrn	0.0050367	-2.1116	RNA and DNA helicase
Pard3b	0.0249889	-2.10666	Involved in cell cycle and tight junction formation
Ptpru	0.0008582	-2.10598	Protein binding regulation of cell growth, differentiation and mitosis
Clasp2	0.048169	-2.08765	Microtubule end binding and anchoring
Grin2b	0.0422529	-2.0848	Glutamate receptor for prominent neurotransmitter in the retina
Dnahc11	0.0020171	-2.06752	Ciliary dynein protein from microtubule motor activity
4921528I07Rik	0.0376126	-2.06285	Unknown
Ms4a4c	0.0019349	-2.06017	Unknown

Tg	0.0232324	-2.06013	Thyroglobin (thyroid hormone)
Nudt13	0.0313719	-2.05882	Unknown
Oca2	0.0226697	-2.05639	Integral membrane protein - tyrosine transport
Itgal	0.013379	-2.05613	Integrin expressed in leukocytes for intracellular adhesions
Scn5a	0.0192309	-2.04663	Sodium channel, voltage-gated
Spna1	0.0108579	-2.04024	Actin cytoskeleton organization
Dpp10	0.0499214	-2.03872	Serine protease and binds K+ channels and alters functionality
Coq7	0.0001781	-2.03198	Coenzyme Q involved in ubiquinone biosynthesis
Rfc1	0.0063545	-2.02704	DNA replication factor C for DNA replication
Telo2	0.008154	-2.00621	S-phase checkpoint protein
Nlrc5	0.0164919	-2.0028	Inhibits NF-kappa-B activation and interferon signalling
Brca1	0.021531	-2.00024	Tumour suppressor acts as a DNA damage sensor and mediates repairs
Casc5	0.0081067	-1.99927	Microtubule-kinetochore attachments and chromosome separation
L3mbtl	0.0497075	-1.99779	Chromatin modification and cell cycle regulation
2310033E01Rik	0.0307551	-1.98669	Phospholipidase inhibitor
Lctl	0.00774	-1.96934	Glycosidase
Tcp10b	0.0271985	-1.96595	Unknown
Mfn1	0.0067646	-1.96589	Mediates mitochondrial fusion
Pfkfb3	0.0191326	-1.96108	Activator of the glycolysis pathway through early primary steps
2610034M16Rik	0.0169822	-1.96036	Unknown
Lars2	0.0062127	-1.95613	Aminoacyl-tRNA synthetase
Aox4	0.0361653	-1.94431	Aldehyde oxidase involved in biosynthesis of retinoic acid
Cdh23	0.0006493	-1.94368	Intracellular adhesion glycoprotein
Ptbp2	0.0088319	-1.94299	Neuronal RNA regulatory protein
Pla2g4d	0.0325627	-1.94056	Phospholipidase, liberates fatty acids
Wdr52	0.0464484	-1.93782	mRNA processing and/or cytoskeleton assembly
Csmd1	0.0052928	-1.93496	Integral membrane protein
Txndc3	0.0474444	-1.93491	Unknown
BC059842	0.0456172	-1.93344	Unknown, potential for CNS development
Scnn1b	0.0456916	-1.93172	Sodium channel, voltage-gated control of electrolytes across epithelia
C1qtnf5	0.001318	-1.92401	Basement membrane and cell adhesion
Ift140	0.0364322	-1.91511	Microtubule formation of cilia
Cdx4	0.0260372	-1.9096	RNA polymerase core promoter binding protein
Hephl1	0.006727	-1.90675	Copper ion transport
Slc5a4a	0.0281977	-1.90241	Unknown, transporter protein
9030420J04Rik	0.0062036	-1.90156	Unknown
Nphp4	0.0382606	-1.89934	Microtubule development and function

Kif2c	0.0299502	-1.89863
Hkdc1	0.0242806	-1.8967
Nbeal2	0.0373284	-1.89554
Pole	0.019369	-1.89426
Gucy2c	0.0005898	-1.89361
Acot12	0.0272659	-1.89312
Lhx8	0.0443032	-1.89031
1700040L02Rik	0.0025767	-1.88678
Abcc5	0.0410045	-1.88581
Cep152	0.0360524	-1.88357
Gm6190	0.0055903	-1.87934
Lrp2	0.0249245	-1.8778
Stag3	0.0005808	-1.87163
4930555F03Rik	0.0406336	-1.8666
Prkdc	0.0292855	-1.86298
Vps13c	0.0425298	-1.85981
Optc	0.0157026	-1.85973
Col27a1	0.0486116	-1.85899
Enoph1	0.0344511	-1.85735
Cd2ap	0.0110717	-1.85617
Mcm5	0.0444294	-1.85324
Usp33	0.0228026	-1.85306
Lcp2	0.0018684	-1.85165
Senp6	0.0247478	-1.85165
Hsd17b3	0.0295071	-1.84551
2010204N08Rik	0.0058221	-1.84151
Setdb1	0.0326697	-1.83627
Lef1	0.0049875	-1.8353
Kndc1	0.0267347	-1.8341
Gm6986	0.0120064	-1.82782
Col6a2	0.020632	-1.8273
Ippk	0.0300244	-1.82475
Timeless	0.0430905	-1.82091
C730048C13Rik	0.0126639	-1.82063
Slc9a10	0.0313326	-1.82026
Enah	0.011655	-1.81313

Kinesin-like microtubule motor protein, mitotic
chromosome segregation Hexokinase activity and glucose metabolism
Regulates neurotransmitter receptor trafficking
DNA polymerase subunit
Enterotoxin receptor
Acetyl-CoA hydrolase activity
DNA binding transcription factor
Unknown, important for cilia function
ATP-binding cassette involved in export of cyclic nucleotides
Microtubule centrosome function
Unknown
Uptake of lipoproteins, sterols, hormones and vitamins from extracellular space
Cohesion of sister chromatids during mitosis
Unknown
Protein kinase mediates double-stranded break DNA repair Unknown
Binds collagen fibrils and regulations fibril formation and morphology
Component of structural extracellular matrix
Unknown, amino-acid metabolism
Scaffolding protein involved in organization of actin cytoskeleton
Facilitates cell cycle and mitosis initiation
Deubiquinating enzyme
T-cell receptor and immune response
Ubiquitin-like molecule
Conversion of androstenedione to testosterone
High carbohydrates induces histone acetylation
Mediates gene silencing and transcriptional repression
Transcription factor for T-cell receptors
Regulates Ras GTPase activity and dendrite morphogenesis
Unknown
Component of collagen binds extracellular matrix proteins
Mediates DNA repair and endocytosis through inositol- phosphate signalling
Involved in DNA polymerase activity, telomere
maintenance and cell morphology
Unknown, solute carrier
Na+ transporting carboxylic acid decarboxylase
Axon guidance and T-cell receptor signalling

Zfp597	0.0215991	-1.81127	Unknown
Mbd2	0.0030594	-1.81122	Binds methylated DNA and represses transcription
4430402118Rik	0.0030179	-1.80948	Unknown
B3galtl	0.0208824	-1.80685	Beta 1,3-galactosyltransferase
Vps13b	0.0355605	-1.80646	Intracellular vesicle protein transport of CNS
Lgr5	0.0418022	-1.8045	G-protein-coupled receptor
Chd11	0.039933	-1.80225	DNA helicase for DNA repair
Rnf10	0.0066076	-1.80152	Ring finger protein DNA binding and transcription regulation
Hmcn1	0.0369989	-1.79986	Immunoglobin, involved in anchorage of neurons to epidermis
Wdr90	0.0235019	-1.79898	Unknown
Ccdc33	0.032919	-1.79699	Unknown
Pcsk5	0.0191332	-1.79594	Protein trafficking through the endoplasmic reticulum
Htr3b	0.049761	-1.79425	Ion channel for fast, depolarizing response in neurons
Myh7b	0.000595	-1.79347	Heavy chain of myosin (transport)
F11	0.0400904	-1.79324	Factor 11 in blood coagulation
Col4a3	0.0341158	-1.79188	Component of collagen
Zbtb17	0.0187941	-1.78929	Regulation of c-myc (transcription factor)
Srrt	0.031086	-1.78731	Regulation of transcription
Chkb	0.0003588	-1.78714	Formation of phosohocholine or phosphoethanolamine by kinase activity
Tecrl	0.0019138	-1.7857	Unknown
Alg13	0.0429268	-1.78543	Lipid oligosaccharide biosynthesis
Ank3	0.0474462	-1.78453	Ankyrin, cytoskeleton anchoring and cell mobility
Hnfla	0.0046702	-1.78376	Liver transcription factor
Clic6	0.0270728	-1.78355	Chloride channel, intracellular transport
Myo7a	0.01778	-1.78178	Myosin component, compromised in retinal degeneration
Ubr2	0.0197904	-1.78082	Ubiquitin protein ligase, targets proteins for degradation
Clcn1	0.0254486	-1.77847	Chloride channel, voltage-sensitive (muscle)
Cdkl4	0.0059487	-1.777	Cyclin dependent kinase, cell cycle regulation
Hdac7	0.0285512	-1.77682	Histone deacetylase, transcription and cell cycle regulation
Eme2	0.0158106	-1.77538	Endonuclease involved in DNA repair
Myh15	0.0186594	-1.77466	Myosin component, motor activity
Myo15	0.0261537	-1.77444	Myosin component, motor activity
Lypd6b	0.0207377	-1.77414	Unknown
Reg3d	0.0443465	-1.77411	Enhances glucose tolerance, unknown function
Vtila	0.0191915	-1.76861	Vesicle transporting, SNARE interactions
Tyr	0.035644	-1.76852	Facilitates conversion of tyrosine to melanin
Morc1	0.0141049	-1.76743	Potential in regulating apoptosis
Cizl	0.0417301	-1.76299	Zinc-finger protein for CDKN1A interaction
Itgb2l	0.0265953	-1.76047	Integrin, intracellular adhesions
C8a	0.0459502	-1.76019	C8 of complement system

Cenpk	0.0417527	-1.75707	Kinetochore assembly and mitosis progression
Ahctf1	0.0486254	-1.75615	Transcription factor for mitotic gene transcription
Ltbp2	0.0023953	-1.75599	Growth factor, involved in cell adhesion
Dnahc12	0.0032213	-1.75459	Dynein, microtubule motor activity
Chd4	0.0391803	1.75029	DNA helicase, histone modifications
Plg	0.0028306	1.75042	Inhibitor of angiogenesis
Fras1	0.0203383	1.75069	Extracellular matrix protein for cell adhesion
Rbl2	0.0395343	1.75129	Retinoblasma-like involved in cell cycle regulation
Steap3	0.0263896	1.75146	Ferrireductase
Ldlr	0.0340576	1.75369	Low density lipoprotein receptor, endocytosis, cholesterol metabolism
Ostf1	0.0028003	1.75443	Cell adhesion, unknown function
Nusap1	0.0061925	1.75444	Nucleus spindle microtubule formation and integrity
Esyt1	0.0333224	1.75465	Unknown
Cdh19	0.0447529	1.75584	Cadherin, glycoprotein involved in cell adhesion
Ptpn14	0.0377965	1.75649	Mitosis, microtubule associated
Plbd2	0.0444493	1.75824	Phospholipase B, lipid metabolism
Trim37	0.0419797	1.75946	Protein-protein interactions
Mov1011	0.0119256	1.76008	RNA helicase (testis)
Asap3	0.0050632	1.76035	Promotes differentiation, cell morphology and mobility
Plcb2	0.0048519	1.76224	Phospholipase C, lipid metabolism, synaptic function
Rictor	0.0231411	1.76235	Component of MTOR, growth factor
Arfgap3	0.0098453	1.76245	GTPase activity, promotes vesicle transport
Col5a3	0.0262117	1.76265	Component of collagen
Scn11a	0.0163679	1.76309	Sodium channel, voltage gated, ion regulation
Lbp	0.0303646	1.76368	Immune response, binds lipopolysaccharides
Aire	0.0118822	1.76399	Binds CREB, regulates expression of immunity genes
Dkc1	0.0101854	1.76501	DNA damage repair
Hepacam2	0.0296601	1.76551	Promotes mitosis and metaphase
Mmp14	0.0165032	1.76592	Extracellular matrix remodelling
Nln	0.02361	1.76613	Formation of neurotensin
Plcb3	0.0120545	1.76731	Phospholipase C, lipid metabolism, signal transduction
Ubash3b	0.0103796	1.7674	Ubiquitin domain inhibits endocytosis of growth factor
Psat1	0.0206356	1.77061	Phosphoserine aminotransferase involved in schizophrenia
Skap2	0.0081311	1.771	Responsible for src signalling pathway
Strc	0.0479517	1.77119	Stereocilin involved in stereocilia formation
Atf5	0.0004549	1.77199	Transcription factor inhibiting astrocyte differentiation
Fam83b	0.0452632	1.77223	MAPK activation and epithelial cell transformation
Vwf	0.0212629	1.77274	Glycoprotein involved in blood coagulation
Irak3	0.0261513	1.77409	Interleukin receptor, inhibits Toll-like receptors in glial cells
Cchcr1	0.0388618	1.77477	Cellular organization and proliferation
Pls3	0.0403178	1.77554	Plastin involved in actin-binding

Patzl0.03845991.77933DNA binding transcription factor and chromatin remodellingSiae0.02638211.78277Deacetylation of sialic acidsQk0.02438011.78342RNA binding protein for oligodendrocyte differentiationCmna30.00312981.7871Intercellular interactions, cell adhesionPlin30.03927881.78726Mannose receptor for transport of lysosmal hydrolase to endosomeCol3a10.00508131.78733Component of collagenDock90.02279071.78862Mediator of cytokinesisCdb10.00206941.7907Cell checkpoint cell, mediates DNA damage repairCdb30.01659281.79195Junction protein, can inhibit apotosis processes in vivoMmr120.01741331.79408Lipid phosphataseTrpv50.04107511.79854Calcium channel involved in ion regulationUnc93b10.02046581.80234Heterochromatin protein to maintain chromatin structureRegulates Toll-receptors trafficking from endoplasmic reticulum, immune responseChiorid channel, intracellular pH and transportNsm20.03484471.80258Extracellular organizationTrpm50.0107071.80627Cation channel, ion homeostasisTrmm20.01646321.80926Bicarbonate transported for intracellular pH maintenanceSlc23a00.02167911.81083Small molecule inner mitchondrial matrix transporterSlc24a40.0340311.80922Bicarbonate transported for inflammatory cytokinesSlca4a40.0340	Ifih1	0.0230229	1.77892	RNA helicase to remove secondary structure, ribosomal formation
Ok0.02438011.78342RNA binding protein for oligodendrocyte differentiation $Okan0.00312981.7871Intercellular interactions, cell adhesionPlin30.03927881.78726Mannose receptor for transport of lysosomal hydrolaseto endosomeCol3a10.00508131.78733Component of collagenDock90.02279071.78862Mediator of cytokinesisCdkn1a0.02006941.7907Cell checkpoint cell, mediates DNA damage repairCd530.01659281.7919Cell checkpoint cell, mediates DNA damage repairCd530.01659281.7919Junction protein, can inhibit apoptosis processes in vivoMtmr120.0141731.79408Lipid phosphataseTrpv50.04107511.79854Calcium channel involved in ion regulationCbci0.00954261.80234Heterochromatin protein to maintain chromatin structureUnc93b10.02046581.80258Chloride channel, intracellular pH and transportNsun20.03481471.80296mRNA translation, ensures codon/anti-codon matchingTrpm50.01907371.80627Cation channel, ion homeostasisTmem20.01646321.80853Uracil and thymidine metabolism, rate-limitingpyrimidine metabolismSlc25a300.02167911.8108Small molecule inner mitochondrial matrix transporterSpc/20.0117151.81131UhknownSlc30010N08Rik0.02211121.81575UnknownRbH10.04648951.8154Interacts wi$	Patz1	0.0384599	1.77933	
L_{mad} 1.7871 Intercellular interactions, cell adhesion $Plin3$ 0.0392788 1.78716 Mannose receptor for transport of lysosomal hydrolase to endosome $Collaal$ 0.0050813 1.78733 Component of collagen $Dock9$ 0.0227907 1.78862 Mediator of cytokinesis $Cdkn1a$ 0.000694 1.7907 Cell checkpoint cell, mediates DNA damage repair $Cd53$ 0.0165928 1.7919 Junction protein, can inhibit apoptosis processes in vivo $Mmr12$ 0.0174133 1.79408 Lipid phosphatase $Trpv5$ 0.0410751 1.79854 Calcium channel involved in ion regulation $Cbx1$ 0.0095426 1.80234 Heterochromatin protein to maintain chromatin structure $We93b1$ 0.0204658 1.80241 Regulates Toll-receptors trafficking from endoplasmic reticulum, immune response $Clic4$ 0.038684 1.80258 Chloride channel, intracellular pH and transport mRNA translation, ensures codon/anti-codon matching Dyrd $Nsun2$ 0.0164632 1.80853 Extracellular organization $Dyyd$ 0.017076 1.80853 Uracil and thymidine metabolism pyrimidine metabolism $Slc25a30$ 0.0216791 1.81068 Small molecule inner mitochondrial matrix transport $Msun2$ $Slam/7$ 0.0089023 1.81231 Inhibits production of inflammatory cytokines $Rap1gap$ $Rap1gap$ 0.0216791 1.81068 Small molecule inner mitochondrial matrix transport $Msun2$ $Slc25a30$ 0.0216791 1.81068 <td< td=""><td>Siae</td><td>0.0263821</td><td>1.78277</td><td>Deacetylation of sialic acids</td></td<>	Siae	0.0263821	1.78277	Deacetylation of sialic acids
Plinal0.03927881.78726Mannose receptor for transport of lysosomal hydrolase to endosome $Plinal$ 0.03927881.78733Component of collagen Dock9 $Dock9$ 0.02279071.78862Mediator of cytokinesis $Cdk1a$ 0.02006941.7907Cell checkpoint cell, mediates DNA damage repair Cell growth and mobility, recruits T cells and macrophages $Cdk3$ 0.01659281.7919Junction protein, can inhibit apoptosis processes in vivo Lipid phosphatase $Fam129b$ 0.04321931.79408Calcium channel involved in ion regulation reticulum, immune response $Chr1$ 0.00954261.80234Heterochromatin protein to maintain chromatin structure Regulates Toll-receptors trafficking from endoplasmic reticulum, immune response $Clic4$ 0.0386841.80258Chloride channel, intracellular pH and transport mRNA translation, ensures codon/anti-codon matching Dyrd $Nsun2$ 0.03481471.80296Itaracellular organization $Dydd$ 0.0170761.80883Uracil and thymidine metabolism, pyrimidine metabolism $Slc24a4$ 0.03420311.80922Bicarbonate transported for intracellular pH maintenance Syleg $Slc25a30$ 0.02167911.81131Unknown $Slamf7$ 0.00890231.81231Inhibits production of inflammatory cytokines Gap1gap $Rap1gap$ 0.02313721.81154Unknown $Slamf7$ 0.03046131.82246Anoctamic alcuium-cholied channel Apfsg1 0.0464895 1.81564Mitochondrial ATP synthase for proton transport during <br< td=""><td>Qk</td><td>0.0243801</td><td>1.78342</td><td>RNA binding protein for oligodendrocyte differentiation</td></br<>	Qk	0.0243801	1.78342	RNA binding protein for oligodendrocyte differentiation
Pun3 $0.0392/88$ $1.78/26$ Col3a1 0.0050813 1.78733 Component of collagenDock9 0.0227907 1.78862 Mediator of cytokinesisCdkn1a 0.0200694 1.7907 Cell checkpoint cell, mediates DNA damage repairCd53 0.0165928 1.7919 Cell checkpoint cell, mediates DNA damage repairCds1 0.00174133 1.79195 Junction protein, can inhibit apoptosis processes <i>in vivo</i> Mtmr12 0.0174133 1.79408 Lipid phosphataseTrpv5 0.0407511 1.79854 Calcium channel involved in ion regulationCbx1 0.0095426 1.80234 Heterochromatin protein to maintain chromatin structureRegulates Toll-receptors trafficking from endoplasmic reticulum, immune responseChloride channel, intracellular pH and transportNsun2 0.0348147 1.80256 Extracellular organizationNsun2 0.017076 1.80833 Uracil and thymidine metabolism, rate-limiting pyrimidine metabolism,	Ctnna3	0.0031298	1.7871	Intercellular interactions, cell adhesion
Dock90.02279071.78862Mediator of cytokinesis $Cdkn1a$ 0.02006941.7907Cell checkpoint cell, mediates DNA damage repair $Cd53$ 0.01659281.7919Cell checkpoint cell, mediates DNA damage repair $Cd53$ 0.01659281.7919Cell growth and mobility, recruits T cells and macrophages $Fam129b$ 0.04321931.79195Junction protein, can inhibit apoptosis processes <i>in vivo</i> $Mimr12$ 0.01741331.79408Lipid phosphatase $Trpv5$ 0.04107511.79854Calcium channel involved in ion regulation $Ckx1$ 0.00954261.80234Heterochromatin protein to maintain chromatin structure $Unc93b1$ 0.02046581.80241Regulates Toll-receptors trafficking from endoplasmic reticulum, immune response $Clic4$ 0.0386841.80258Chloride channel, intracellular pH and transport mRNA translation, ensures codon/anti-codon matching Dryd $Trpm5$ 0.01907371.80627Cation channel, ion homeostasis $Tmem2$ 0.01646321.80883Extracellular organization $Dpyd$ 0.0170761.80883Uracil and thymidine metabolism $Slc4a4$ 0.03420311.80922Bicarbonate transported for intracellular pH maintenance $Slc25a30$ 0.02167911.81068Small molecule inner mitochondrial matrix transporter $Spef2$ 0.0117151.81131Unknown $Anp5g1$ 0.04648951.81564Mitochondrial ATP synthase for proton transport during OXPHOS $Brip1$ 0.03236281.81949Inte	Plin3	0.0392788	1.78726	
Calkn la0.02006941.7907Cell checkpoint cell, mediates DNA damage repairCd530.01659281.7919Cell growth and mobility, recruits T cells and macrophagesFam129b0.04321931.79195Junction protein, can inhibit apoptosis processes <i>in vivo</i> Mtmr120.0171131.79408Lipid phosphataseTrpv50.04107511.79854Calcium channel involved in ior regulationCbx10.00954261.80234Heterochromatin protein to maintain chromatin structureUnc93b10.02046581.80241Regulates Toll-receptors trafficking from endoplasmic reticulum, immune responseClic40.0386841.80258Chloride channel, intracellular pH and transportNsun20.03481471.80296mRNA translation, ensures codon/anti-codon matching pyrimidine metabolismTrpm50.01907371.80627Cation channel, ion homeostasisSlc4a40.03420311.80922Bicarbonate transported for intracellular pH maintenanceSlc2a300.02167911.81068Small molecule inner mitochondrial matrix transporterSlc4a40.03231721.81131UnknownAmp70.00890231.81231Inhibits production of inflammatory cytokinesRap1gap0.02211121.81575UnknownRbl10.04172791.81876Retinoblasma-like involved in cell cycle regulationInteracts with BRAC1 to mediate double-stranded DNA break repairMitochondrial ATP synthase for proton transport during OXPHOSGbp100.03236281.81246Anoctamin calcuiu	Col3a1	0.0050813	1.78733	Component of collagen
Cd530.01659281.7919Cell growth and mobility, recruits T cells and macrophages $Fam129b$ 0.04321931.79195Junction protein, can inhibit apoptosis processes <i>in vivo</i> $Mimr12$ 0.01741331.79408Lipid phosphatase $Trpv5$ 0.04107511.79854Calcium channel involved in ion regulation Ckl 0.00954261.80234Heterochromatin protein to maintain chromatin structure $Unc93b1$ 0.02046581.80241Regulates Toll-receptors trafficking from endoplasmic reticulum, immune response $Clic4$ 0.0386841.80256Chloride channel, intracellular pH and transport $Nsun2$ 0.01646321.80833Extracellular organization $Dpyd$ 0.0170761.80883Uracil and thymidine metabolism, pyrimidine metabolism $Slc4a4$ 0.03420311.80922Bicarbonate transported for intracellular pH maintenance $Slc25a30$ 0.02167911.81068Small molecule inner mitochondrial matrix transporter $Spef2$ 0.0117151.81131Unknown $Slamf7$ 0.00890231.81231Inhibits production of inflammatory cytokines $Rap1gap$ 0.02211121.81575Unknown $Rbl1$ 0.04172791.81876Retinoblasma-like involved in cell cycle regulation $Brip1$ 0.03236281.81240Interacts with BRAC1 to mediate double-stranded DNA break repair $Gbp10$ 0.0375261.82236GTPase activation $Ano5$ 0.03046131.82236ATPase, unknown function $Ijlti1$ 0.046469 </td <td>Dock9</td> <td>0.0227907</td> <td>1.78862</td> <td>Mediator of cytokinesis</td>	Dock9	0.0227907	1.78862	Mediator of cytokinesis
Call0.01039281.7919macrophages $Fam129b$ 0.04321931.79195Junction protein, can inhibit apoptosis processes in vivo $Mnmr12$ 0.01741331.79408Lipid phosphatase $Trpv5$ 0.04107511.79854Calcium channel involved in ion regulation $Cbx1$ 0.00954261.80234Heterochromatin protein to maintain chromatin structure $Unc93b1$ 0.02046581.80241Regulates Toll-receptors trafficking from endoplasmic reticulum, immune response $Clic4$ 0.0386841.80258Chloride channel, intracellular pH and transport $Nsun2$ 0.03481471.80296mRNA translation, ensures codon/anti-codon matching $Trpm5$ 0.01907371.80627Cation channel, ion homeostasis $Tmem2$ 0.01646321.80853Extracellular organization $Dpyd$ 0.0170761.80883Uracil and thymidine metabolism $Slc4a4$ 0.03420311.80922Bicarbonate transported for intracellular pH maintenance $Slc25a30$ 0.02167911.81068Small molecule inner mitochondrial matrix transporter $Spef2$ 0.0117151.81131Unknown $Slamf7$ 0.00890231.81231Inhibits production of inflammatory cytokines $Rap1gap$ 0.02211221.81564GTPase activity $Atp5g1$ 0.04648951.81564Mitochondrial ATP synthase for proton transport during OXPHOS $E030010N08Rik$ 0.02211121.8176Retinoblasma-like involved in cell cycle regulation $Rip11$ 0.0456491.82246Anoctami	Cdkn1a	0.0200694	1.7907	Cell checkpoint cell, mediates DNA damage repair
Mtmr120.01741331.79408Lipid phosphatase $Trpv5$ 0.04107511.79854Calcium channel involved in ion regulation $Cbx1$ 0.00954261.80234Heterochromatin protein to maintain chromatin structure $Unc93b1$ 0.02046581.80241Regulates Toll-receptors trafficking from endoplasmic reticulum, immune response $Clic4$ 0.0386841.80258Chloride channel, intracellular pH and transport mRNA translation, ensures codon/anti-codon matching Cation channel, ion homeostasis $Nsun2$ 0.01646321.80853Extracellular organization $Dpyd$ 0.0170761.80883Uracil and thymidine metabolism, rate-limiting pyrimidine metabolism $Slc4a4$ 0.03420311.80922Bicarbonate transported for intracellular pH maintenance Small molecule inner mitochondrial matrix transporter $Sle72$ 0.0117151.81131Unknown $Slamf7$ 0.00890231.81231Inhibits production of inflammatory cytokines $Rap1gap$ 0.02211121.81564Mitochondrial ATP synthase for proton transport during OXPHOS $E030010N08Rik$ 0.02211121.81575Unknown $Rbl1$ 0.04172791.81876Retinoblasma-like involved in cell cycle regulation $Brip1$ 0.03236281.81242Immune response via interferon activation $Erbb2$ 0.04147821.82236Epidermal growth factor involved in mitosis activation $Ano5$ 0.03046131.82246Anoctamin calcuium-cholride channel $Ano5$ 0.03046131.8238Unknown $Inte$	Cd53	0.0165928	1.7919	macrophages
Trpv50.04107511.79854Calcium channel involved in ion regulation $Trpv5$ 0.00954261.80234Heterochromatin protein to maintain chromatin structure $Unc93b1$ 0.02046581.80241Regulates Toll-receptors trafficking from endoplasmic reticulum, immune response $Clic4$ 0.0386841.80258Chloride channel, intracellular pH and transport mRNA translation, ensures codon/anti-codon matching Trpm5 $Nsun2$ 0.03481471.80296mRNA translation, ensures codon/anti-codon matching Uracil and thymidine metabolism $Dpyd$ 0.0170761.80883Uracil and thymidine metabolism $Slc4a4$ 0.03420311.80922Bicarbonate transported for intracellular pH maintenance $Slc25a30$ 0.02167911.81068Small molecule inner mitochondrial matrix transporter $Spef2$ 0.0117151.81131Unknown $Slantf7$ 0.00890231.81231Inhibits production of inflammatory cytokines $Rap1gap$ 0.02211121.81575Unknown $Rbl1$ 0.04648951.81564Mitochondrial ATP synthase for proton transport during OXPHOS $Brip1$ 0.03236281.81949Interacts with BAC1 to mediate double-stranded DNA break repair $Gbp10$ 0.00795261.82236Epidermal growth factor involved in mitosis activation $Ano5$ 0.03046131.82246Anoctamin calcuium-cholride channel $Anp13a5$ 0.0251781.82328ATPase, unknown function $Ifltd1$ 0.0465691.8238Unknown $Depdc1b$ 0.02045871.	Fam129b	0.0432193	1.79195	
Cbx10.00954261.80234Heterochromatin protein to maintain chromatin structure $Unc93b1$ 0.02046581.80241Heterochromatin protein to maintain chromatin structure $Unc93b1$ 0.02046581.80241Regulates Toll-receptors trafficking from endoplasmic reticulum, immune response $Clic4$ 0.0386841.80258Chloride channel, intracellular pH and transport mRNA translation, ensures codon/anti-codon matching Trpm5 $Nsun2$ 0.01907371.80627Cation channel, ion homeostasis $Tmem2$ 0.01646321.80853Extracellular organization $Dpyd$ 0.0170761.80883Uracil and thymidine metabolism, rate-limiting pyrimidine metabolism $Slc4a4$ 0.03420311.80922Bicarbonate transported for intracellular pH maintenance Small molecule inner mitochondrial matrix transporter $Spef2$ 0.0117151.81131Unknown $Slamf7$ 0.00890231.81231Inhibits production of inflammatory cytokines $Rap1gap$ 0.0211121.81564Mitochondrial ATP synthase for proton transport during OXPHOS $E030010N08Rik$ 0.02211121.8175Unknown $Rbl1$ 0.04172791.81876Retinoblasma-like involved in cell cycle regulation $Brip1$ 0.03236281.82162Immune response via interferon activation $Artp 53$ 0.0251781.82236Epidermal growth factor involved in mitosis activation $Ano5$ 0.03046131.82246Anoctamin calcuium-cholride channel $Atp 13a5$ 0.0251781.82328ATPase, unknown function<	Mtmr12	0.0174133	1.79408	Lipid phosphatase
Unc 93b10.02046581.80241Regulates Toll-receptors trafficking from endoplasmic reticulum, immune responseClic40.0386841.80258Chloride channel, intracellular pH and transport mRNA translation, ensures codon/anti-codon matching Trpm5Nsun20.03481471.80296mRNA translation, ensures codon/anti-codon matching Trpm5Trpm50.01907371.80627Cation channel, ion homeostasisTmem20.01646321.80853Extracellular organizationDpyd0.0170761.80883Uracil and thymidine metabolism, rate-limiting pyrimidine metabolismSlc4a40.03420311.80922Bicarbonate transported for intracellular pH maintenanceSlc25a300.02167911.81068Small molecule inner mitochondrial matrix transporterSpef20.0117151.81131UnknownSlamf70.00890231.81231Inhibits production of inflammatory cytokinesRap1gap0.02211121.8155UnknownRbl10.04172791.81876Retinoblasma-like involved in cell cycle regulationInteracts with BRAC1 to mediate double-stranded DNA break repairEpidermal growth factor involved in mitosis activationAno50.03046131.82246Anoctamin calcuium-cholride channelAno50.03046131.8238MINue Pidermal growth factor involved in mitosis activationAno50.0251781.8238ATPase, unknown functionIftid10.0465691.8238GTPase activationAno50.02045871.82578GTPase activation <td< td=""><td>Trpv5</td><td>0.0410751</td><td>1.79854</td><td>Calcium channel involved in ion regulation</td></td<>	Trpv5	0.0410751	1.79854	Calcium channel involved in ion regulation
Unc93b1 0.0204658 1.80241 $Clic4$ 0.038684 1.80258 Chloride channel, intracellular pH and transport $Nsun2$ 0.0348147 1.80296 mRNA translation, ensures codon/anti-codon matching $Trpm5$ 0.0190737 1.80627 Cation channel, ion homeostasis $Tmem2$ 0.0164632 1.80853 Extracellular organization $Dpyd$ 0.017076 1.80883 Uracil and thymidine metabolism, rate-limiting pyrimidine metabolism $Slc4a4$ 0.0342031 1.80922 Bicarbonate transported for intracellular pH maintenance $Slc25a30$ 0.0216791 1.81068 Small molecule inner mitochondrial matrix transporter $Spef2$ 0.011715 1.81131 Unknown $Slamf7$ 0.0089023 1.81231 Inhibits production of inflammatory cytokines $Rap1gap$ 0.0221112 1.81575 Unknown $Rbl1$ 0.044895 1.81564 Mitochondrial ATP synthase for proton transport during $OXPHOS$ $E030010N08Rik$ 0.0221112 1.81575 Unknown $Rbl1$ 0.0079526 1.82162 Immune response via interferon activation $Erbb2$ 0.0414782 1.82236 Epidermal growth factor involved in mitosis activation $Ano5$ 0.0304613 1.82246 Anoctamin calcuium-cholride channel $Ano5$ 0.025178 1.8238 Unknown $InteractsunknownfunctionIftd10.0465691.82578GTPase activationRnf213$	Cbx1	0.0095426	1.80234	Heterochromatin protein to maintain chromatin structure
Nsun2 0.0348147 1.80296 mRNA translation, ensures codon/anti-codon matching $Trpm5$ 0.0190737 1.80627 Cation channel, ion homeostasis $Tmem2$ 0.0164632 1.80853 Extracellular organization $Dpyd$ 0.017076 1.80883 Uracil and thymidine metabolism, rate-limiting pyrimidine metabolism $Slc4a4$ 0.0342031 1.80922 Bicarbonate transported for intracellular pH maintenance $Slc25a30$ 0.0216791 1.81068 Small molecule inner mitochondrial matrix transporter $Spef2$ 0.011715 1.81131 Unknown $Slamf7$ 0.0089023 1.81231 Inhibits production of inflammatory cytokines $Rap1gap$ 0.0221122 1.81564 Mitochondrial ATP synthase for proton transport during OXPHOS $E030010N08Rik$ 0.0221112 1.81575 Unknown $Rbl1$ 0.0417279 1.81876 Retinoblasma-like involved in cell cycle regulation $Brip1$ 0.0323628 1.81949 Interacts with BRAC1 to mediate double-stranded DNA break repair $Gbp10$ 0.0079526 1.82162 Immune response via interferon activation $Arb53$ 0.025178 1.82236 Epidermal growth factor involved in mitosis activation $Arb13a5$ 0.025178 1.8238 ATPase, unknown function $Ifttd1$ 0.046549 1.8238 Unknown $Depdc1b$ 0.0204587 1.82578 GTPase activation $Rnf213$ 0.0016575 1.82609 Zinc-finger protein for inter-protein interaction <td>Unc93b1</td> <td>0.0204658</td> <td>1.80241</td> <td>• • • •</td>	Unc93b1	0.0204658	1.80241	• • • •
Trpm50.01907371.80627Cation channel, ion homeostasis $Tmem2$ 0.01646321.80853Extracellular organization $Dpyd$ 0.0170761.80883Uracil and thymidine metabolism $Dpyd$ 0.0170761.80883Uracil and thymidine metabolism $Slc4a4$ 0.03420311.80922Bicarbonate transported for intracellular pH maintenance $Slc25a30$ 0.02167911.81068Small molecule inner mitochondrial matrix transporter $Spef2$ 0.0117151.81131Unknown $Slamf7$ 0.00890231.81231Inhibits production of inflammatory cytokines $Rap1gap$ 0.02211721.81564Mitochondrial ATP synthase for porton transport during OXPHOS $E030010N08Rik$ 0.02211121.81876Retinoblasma-like involved in cell cycle regulation $Brip1$ 0.03236281.81949Intracts with BRAC1 to mediate double-stranded DNA break repair $Gbp10$ 0.00795261.82162Immune response via interferon activation $Ano5$ 0.03046131.82236Epidermal growth factor involved in mitosis activation $Ano5$ 0.0251781.82328ATPase, unknown function $Ifltd1$ 0.04656491.8238Unknown $Depdc1b$ 0.02045871.82578GTPase activation $Rnf213$ 0.00165751.82609Zinc-finger protein for inter-protein interaction	Clic4	0.038684	1.80258	Chloride channel, intracellular pH and transport
TypeInstantExtracellular organization $Tmem2$ 0.01646321.80853Extracellular organization $Dpyd$ 0.0170761.80883Uracil and thymidine metabolism, rate-limiting pyrimidine metabolism $Slc4a4$ 0.03420311.80922Bicarbonate transported for intracellular pH maintenance $Slc25a30$ 0.02167911.81068Small molecule inner mitochondrial matrix transporter $Spef2$ 0.0117151.81131Unknown $Slamf7$ 0.00890231.81231Inhibits production of inflammatory cytokines $Rap1gap$ 0.02313721.81419GTPase activity $Atp5g1$ 0.04648951.81564Mitochondrial ATP synthase for proton transport during OXPHOS $E030010N08Rik$ 0.02211121.81575Unknown $Rbl1$ 0.04172791.81876Retinoblasma-like involved in cell cycle regulation Interacts with BRAC1 to mediate double-stranded DNA break repair $Gbp10$ 0.00795261.82162Immune response via interferon activation $Ano5$ 0.03046131.82236Epidermal growth factor involved in mitosis activation $Ano5$ 0.0251781.8238ATPase, unknown function $Ifltd1$ 0.04656491.8238GTPase activation $Depdc1b$ 0.02045871.82578GTPase activation	Nsun2	0.0348147	1.80296	mRNA translation, ensures codon/anti-codon matching
Dpyd0.0170761.80883Uracil and thymidine metabolism, rate-limiting pyrimidine metabolismSlc4a40.03420311.80922Bicarbonate transported for intracellular pH maintenanceSlc25a300.02167911.81068Small molecule inner mitochondrial matrix transporterSpef20.0117151.81131UnknownSlamf70.00890231.81231Inhibits production of inflammatory cytokinesRap1gap0.02313721.81419GTPase activityAtp5g10.04648951.81564Mitochondrial ATP synthase for proton transport during OXPHOSE030010N08Rik0.02211121.81575UnknownRbl10.0472791.81876Retinoblasma-like involved in cell cycle regulation Interacts with BRAC1 to mediate double-stranded DNA break repairGbp100.00795261.82162Immune response via interferon activation Erabs2Ano50.03046131.82246Anoctamin calcuium-cholride channelAtp13a50.0251781.8238UnknownIfItd10.04656491.8238UnknownDepdc1b0.02045871.82578GTPase activationRnf2130.00165751.82609Zinc-finger protein for inter-protein interaction	Trpm5	0.0190737	1.80627	Cation channel, ion homeostasis
Dpyd $0.01/076$ 1.80883 pyrimidine metabolismSlc4a4 0.0342031 1.80922 Bicarbonate transported for intracellular pH maintenanceSlc25a30 0.0216791 1.81068 Small molecule inner mitochondrial matrix transporterSpef2 0.011715 1.81131 UnknownSlamf7 0.0089023 1.81231 Inhibits production of inflammatory cytokinesRap1gap 0.0231372 1.81419 GTPase activityAtp5g1 0.0464895 1.81564 Mitochondrial ATP synthase for proton transport during OXPHOSE030010N08Rik 0.0221112 1.81575 UnknownBrip1 0.0323628 1.81949 Interacts with BRAC1 to mediate double-stranded DNA break repairGbp10 0.0079526 1.82162 Immune response via interferon activationAno5 0.0304613 1.82246 Anoctamin calcuium-cholride channelAtp13a5 0.025178 1.8238 UnknownIftld1 0.046549 1.8238 GTPase activationDepdc1b 0.0204587 1.82578 GTPase activation	Tmem2	0.0164632	1.80853	Extracellular organization
Slc25a30 0.0216791 1.81068 Small molecule inner mitochondrial matrix transporter $Spef2$ 0.011715 1.81131 Unknown $Slamf7$ 0.0089023 1.81231 Inhibits production of inflammatory cytokines $Rap1gap$ 0.0231372 1.81419 GTPase activity $Atp5g1$ 0.0464895 1.81564 Mitochondrial ATP synthase for proton transport during OXPHOS $E030010N08Rik$ 0.0221112 1.81575 Unknown $Rbl1$ 0.0417279 1.81876 Retinoblasma-like involved in cell cycle regulation $Brip1$ 0.0323628 1.81949 Interacts with BRAC1 to mediate double-stranded DNA break repair $Gbp10$ 0.0079526 1.82162 Immune response via interferon activation $Ano5$ 0.0304613 1.82246 Anoctamin calcuium-cholride channel $Atp13a5$ 0.025178 1.82328 ATPase, unknown function $Ifltd1$ 0.0465649 1.82578 GTPase activation $Depdc1b$ 0.0204587 1.82609 Zinc-finger protein for inter-protein interaction	Dpyd	0.017076	1.80883	
Spef20.0117151.81131UnknownSlant70.00890231.81231Inhibits production of inflammatory cytokinesRap1gap0.02313721.81419GTPase activityAtp5g10.04648951.81564Mitochondrial ATP synthase for proton transport during OXPHOSE030010N08Rik0.02211121.81575UnknownRbl10.04172791.81876Retinoblasma-like involved in cell cycle regulation Interacts with BRAC1 to mediate double-stranded DNA break repairBrip10.03236281.81949Immune response via interferon activationErbb20.04147821.82236Epidermal growth factor involved in mitosis activation Ano5Atp13a50.0251781.82328ATPase, unknown function Infltd1Jfltd10.04656491.8238UnknownDepdc1b0.02045871.82578GTPase activation Finger protein for inter-protein interaction	Slc4a4	0.0342031	1.80922	Bicarbonate transported for intracellular pH maintenance
SpecialStant/F (0.011116) (1.01131) Slamt/7 (0.0089023) 1.81231 Inhibits production of inflammatory cytokinesRap1gap (0.0231372) 1.81419 GTPase activityAtp5g1 (0.0464895) 1.81564 Mitochondrial ATP synthase for proton transport during OXPHOSE030010N08Rik (0.0221112) 1.81575 UnknownRb11 (0.0417279) 1.81876 Retinoblasma-like involved in cell cycle regulationBrip1 (0.0323628) 1.81949 Interacts with BRAC1 to mediate double-stranded DNA break repairGbp10 (0.0079526) 1.82162 Immune response via interferon activationErbb2 (0.0414782) 1.82236 Epidermal growth factor involved in mitosis activationAno5 (0.0304613) 1.82246 Anoctamin calcuium-cholride channelAtp13a5 (0.0204587) 1.8238 UnknownDepdc1b (0.0204587) 1.82578 GTPase activationRnf213 (0.0016575) 1.82609 Zinc-finger protein for inter-protein interaction	Slc25a30	0.0216791	1.81068	Small molecule inner mitochondrial matrix transporter
Rap1gap0.02313721.81419GTPase activityAtp5g10.04648951.81564Mitochondrial ATP synthase for proton transport during OXPHOSE030010N08Rik0.02211121.81575UnknownRbl10.04172791.81876Retinoblasma-like involved in cell cycle regulationBrip10.03236281.81949Interacts with BRAC1 to mediate double-stranded DNA break repairGbp100.00795261.82162Immune response via interferon activationErbb20.04147821.82236Epidermal growth factor involved in mitosis activationAno50.03046131.82246Anoctamin calcuium-cholride channelAtp13a50.0251781.8238ATPase, unknown functionIfltd10.04656491.82578GTPase activationDepdc1b0.02045871.82609Zinc-finger protein for inter-protein interaction	Spef2	0.011715	1.81131	Unknown
Atp5g10.04648951.81564Mitochondrial ATP synthase for proton transport during OXPHOSE030010N08Rik0.02211121.81575UnknownRbl10.04172791.81876Retinoblasma-like involved in cell cycle regulationBrip10.03236281.81949Interacts with BRAC1 to mediate double-stranded DNA break repairGbp100.00795261.82162Immune response via interferon activationErbb20.04147821.82236Epidermal growth factor involved in mitosis activationAno50.03046131.82246Anoctamin calcuium-cholride channelAtp13a50.0251781.8238UnknownDepdc1b0.02045871.82578GTPase activationRnf2130.00165751.82609Zinc-finger protein for inter-protein interaction	Slamf7	0.0089023	1.81231	Inhibits production of inflammatory cytokines
Atp3g10.04648951.81564OXPHOSE030010N08Rik0.02211121.81575UnknownRbl10.04172791.81876Retinoblasma-like involved in cell cycle regulationBrip10.03236281.81949Interacts with BRAC1 to mediate double-stranded DNA break repairGbp100.00795261.82162Immune response via interferon activationErbb20.04147821.82236Epidermal growth factor involved in mitosis activationAno50.03046131.82246Anoctamin calcuium-cholride channelAtp13a50.0251781.8238UnknownIfltd10.04656491.8238UnknownDepdc1b0.02045871.82578GTPase activationRnf2130.00165751.82609Zinc-finger protein for inter-protein interaction	Rap1gap	0.0231372	1.81419	GTPase activity
Rb110.04172791.81876Retinoblasma-like involved in cell cycle regulationBrip10.03236281.81949Interacts with BRAC1 to mediate double-stranded DNA break repairGbp100.00795261.82162Immune response via interferon activationErbb20.04147821.82236Epidermal growth factor involved in mitosis activationAno50.03046131.82246Anoctamin calcuium-cholride channelAtp13a50.0251781.82328ATPase, unknown functionIfltd10.04656491.8238GTPase activationDepdc1b0.02045871.82578GTPase activationRnf2130.00165751.82609Zinc-finger protein for inter-protein interaction	Atp5g1	0.0464895	1.81564	
Brip10.03236281.81949Interacts with BRAC1 to mediate double-stranded DNA break repairGbp100.00795261.82162Immune response via interferon activationErbb20.04147821.82236Epidermal growth factor involved in mitosis activationAno50.03046131.82246Anoctamin calcuium-cholride channelAtp13a50.0251781.82328ATPase, unknown functionIfltd10.04656491.8238UnknownDepdc1b0.02045871.82578GTPase activationRnf2130.00165751.82609Zinc-finger protein for inter-protein interaction	E030010N08Rik	0.0221112	1.81575	Unknown
Brip10.03236281.81949break repairGbp100.00795261.82162Immune response via interferon activationErbb20.04147821.82236Epidermal growth factor involved in mitosis activationAno50.03046131.82246Anoctamin calcuium-cholride channelAtp13a50.0251781.82328ATPase, unknown functionIfltd10.04656491.8238UnknownDepdc1b0.02045871.82578GTPase activationRnf2130.00165751.82609Zinc-finger protein for inter-protein interaction	Rbl1	0.0417279	1.81876	Retinoblasma-like involved in cell cycle regulation
Erbb20.04147821.82236Epidermal growth factor involved in mitosis activationAno50.03046131.82246Anoctamin calcuium-cholride channelAtp13a50.0251781.82328ATPase, unknown functionIfltd10.04656491.8238UnknownDepdc1b0.02045871.82578GTPase activationRnf2130.00165751.82609Zinc-finger protein for inter-protein interaction	Brip1	0.0323628	1.81949	
Ano5 0.0304613 1.82246 Anoctamin calcuium-cholride channel Atp13a5 0.025178 1.82328 ATPase, unknown function Ifltd1 0.0465649 1.8238 Unknown Depdc1b 0.0204587 1.82578 GTPase activation Rnf213 0.0016575 1.82609 Zinc-finger protein for inter-protein interaction	Gbp10	0.0079526	1.82162	Immune response via interferon activation
Atp13a5 0.025178 1.82328 ATPase, unknown function Ifltd1 0.0465649 1.8238 Unknown Depdc1b 0.0204587 1.82578 GTPase activation Rnf213 0.0016575 1.82609 Zinc-finger protein for inter-protein interaction	Erbb2	0.0414782	1.82236	Epidermal growth factor involved in mitosis activation
IfItd1 0.0465649 1.8238 Unknown Depdc1b 0.0204587 1.82578 GTPase activation Rnf213 0.0016575 1.82609 Zinc-finger protein for inter-protein interaction	Ano5	0.0304613	1.82246	Anoctamin calcuium-cholride channel
Depdc1b0.02045871.82578GTPase activationRnf2130.00165751.82609Zinc-finger protein for inter-protein interaction	Atp13a5	0.025178	1.82328	ATPase, unknown function
Rnf2130.00165751.82609Zinc-finger protein for inter-protein interaction	Ifltd1	0.0465649	1.8238	Unknown
5	Depdc1b	0.0204587	1.82578	GTPase activation
<i>Lrp2bp</i> 0.0483587 1.82778 Unknown	Rnf213	0.0016575	1.82609	Zinc-finger protein for inter-protein interaction
	Lrp2bp	0.0483587	1.82778	Unknown

Klhl4	0.037315	1.82864	Actin binding, unknown function
Dapk2	0.0441918	1.82947	Regulator of pre-programmed cell death
Ddx60	0.0002651	1.83023	RNA helicase to remove secondary structure
Fyttd1	0.0002602	1.83227	Unknown
Lcp1	0.0305862	1.83464	B-cell and T-cell activation and cell migration
Sh3pxd2a	0.0016584	1.8374	Regulates invadopodia (extracellular protrusions)
Ccnb3	0.0017382	1.838	Cyclin involved in cell cycle regulation
Cfb	0.0018594	1.83836	Complement factor B for complement activation and leukocyte recruitment
Tmc3	0.0438492	1.84376	Ion transport, unknown function
Nmt2	0.0001749	1.84398	N-myristoyltransferase 2
Naip5	0.0364918	1.84443	Apoptosis inhibitory protein, inhibits proteins involved in apoptosis
Plcg2	0.0092755	1.84448	Phospholipase C, signal transduction and cellular messaging
Dock1	0.0133594	1.84479	Dedicator of cytokinesis, induces cell migration and phagocytosis
Gbp5	0.0440096	1.84668	Stimulates inflammation response to foreign antigens
Cyp27a1	0.0372999	1.84918	Cytochrome P450 variant, drug metabolism
Zfp330	0.0358616	1.85066	Zinc finger protein, unknown function
Ttc12	0.0098723	1.85072	Unknown
Wdr64	0.0307521	1.85187	Unknown
Cadps2	0.0171056	1.85342	Calcium secretion, synaptic function in neurons
Itga8	0.0114879	1.85524	Integrin, intracellular adhesions, cytoskeleton formation
Atp1a4	0.0212119	1.85636	ATPase, Na+/K+ transporter, neuron electrochemical gradient maintenance
Tmem77	0.007951	1.85869	Cellular apoptosis inducing
Slc25a13	0.0030538	1.86195	Mitochondrial glutamate/aspartate transfer
Parp12	0.018547	1.86243	mRNA post-translational modifications, ADP ribose
Hydin	0.0313929	1.86592	Cilia formation
Oprm1	0.0181287	1.87057	Opioid receptor, neuronal G-protein
Rps6ka1	0.0374657	1.87268	Ribosomal protein kinase, controls cell differentiation
Gpr64	0.025621	1.87376	G protein-coupled receptor, transmembrane
Mthfd2	0.0019253	1.87397	Nuclear mitochondrial NAD sequestering
Nid2	0.0172362	1.87464	Binds collagen, involved in cell adhesion
Lsp1	0.0468358	1.8747	Immune cell expression, involved in cell adhesion and extracellular remodelling
Ppfia4	0.0098483	1.87761	Protein tyrosine phosphatase, receptor type, tyrosine phosphorylation
1700003E16Rik	0.0106796	1.88475	Unknown
Btk	0.0300985	1.88622	Formation and functionality of B-cells
Maged1	0.0174606	1.88731	Programmed cell death, cancer antigen presentation
Slc22a4	0.0382704	1.89398	Solute carrier, cations and zwitterion transport
Egfr	0.0279557	1.89959	Epidermal growth factor receptor, cell proliferation
Fanci	0.0060002	1.90404	Fanconi anemia, complementation group I activation

Muc5b	0.0041903	1.90453	Mucin 5B, glycosylated protein involved in mucus formation
Sympk	0.0036757	1.90828	Post-transcriptional processing, involved in polyadenylation
Meox1	0.019515	1.9114	Developmental signalling
Atp2c2	0.0266884	1.91293	ATPase, cation transporter, neuron function
Ano1	0.0409161	1.91376	Anoctamin 1, calcium activated chloride channel, epithelial repair
Padi2	0.0202878	1.91377	Post-translational modifications mediated by calcium, CNS
Apoe	0.0235635	1.9157	Apolipoprotein E, metabolism of triglycerides (liver)
Ncf2	0.0177262	1.91758	Oxidoreductase activity, neutrophil specific
Map3k6	0.0247556	1.91763	Mitotic activated protein kinase
Dock11	0.0036392	1.91909	Dedicator of cytokinesis 11, extracellular remodelling
BC057079	0.0142697	1.92342	Focadhesin, cell adhesion, unknown function
Col6a3	0.0133423	1.92357	Component of collagen, extracellular matrix binding
Ppp1r12c	0.0066925	1.92469	Protein phosphatase 1, cytoskeleton formation and organization
Elmo3	0.0392572	1.92727	Engulfment of apoptotic cells, inflammation
Ccdc88c	0.0496785	1.93007	Developmental interactions, wnt signalling
Abca8b	0.0061563	1.9319	ATP-binding cassette, unknown function
Dnahc17	0.0288871	1.93335	Dynein, microtubule motor activity
Lrrfip1	0.026914	1.9367	DNA, RNA binding, regulation of transcription
Slc7a7	0.0483406	1.94032	Solute carrier, amino acid transporter (cationic and neutral)
4833423E24Rik	0.0361036	1.94104	Unknown
Sytl1	0.0258357	1.94714	GTPase activating protein
Rtkn	0.0205158	1.9493	Rhotekin scaffold protein inhibits GTPase Rho
Orc6l	0.0257836	1.95019	Initiates DNA replication in the nucleus
Sec24d	0.045669	1.95193	Vesicle transporting in vivo
Pnldc1	0.0114914	1.95203	Binds poly-A tail of mRNA, unknown function
Ppp2r1b	0.0226536	1.95427	Protein phosphatase, regulates protein function <i>in vivo</i> , cancer associated
Abca14	0.0374404	1.95828	ATP-binding cassette, unknown function
Grhl3	0.0425524	1.96065	Developmental transcription factor, endothelial cell migration
Sorbs2	0.0308871	1.96262	Cytoskeleton modifications
Hk1	0.0367158	1.96385	Hexokinase, glucose metabolism and glycolysis regulation
Itpr3	0.0090388	1.96463	Inositol 1,4,5-trisphosphate receptor, regulates calcium homeostasis
Pyroxd2	0.0302111	1.96623	Unknown
Soat1	0.0179412	1.96706	Formation of fatty acids and cholesterol
Eng	0.043215	1.97004	Vascular endothelial glycoprotein receptor
Slc7a1	0.0123391	1.97223	Solute carrier, cationic amino acids
Trim6	0.0210848	1.9726	Unknown

Fbxl13	0.0324725	1.97696	Cleave ubiquitin from ubiquitination products
Spata6	0.0324723	1.97090	Spermatogenesis, cell differentiation, unknown function
Lrrc9	0.0356599	1.9848	Unknown
Myh3	0.0268801	1.98504	Myosin component, motor activity, cell transport
Zfp599	0.0370838	1.9856	Zinc finger protein, unknown function
2310008H04Rik	0.0400248	1.98643	Unknown
Cacna1h	0.0118042	1.98808	Calcium channel, voltage-dependent, maintenance of membrane potentials
Tbc1d4	0.0315198	1.99166	Mediates transport of GLUT4 from endoplasmic reticulum to plasma membrane
Dhx40	0.0420537	1.99427	RNA metabolism, helicase
Cep250	0.0243675	1.99597	Centrosome formation and function during mitosis
Gk5	0.0480136	1.99773	Primary, regulatory step of glycerol metabolism
Slc12a1	0.0416418	1.99777	Solute carrier, unknown function
Abca9	0.0438495	1.99907	ATP-binding cassette, unknown function
Klk1b16	0.0251047	2.00083	Unknown
Lama2	0.01227	2.0026	Laminin component, extracellular component for remodelling and organization
Atp8b1	0.0192593	2.00292	Aminophospholipid transporter
Baiap3	0.0003945	2.00342	Angiogenesis regulator in the brain, involved in synaptic function
Igfbp3	0.0358018	2.00685	Insulin-like growth factor binding protein
Lmna	0.0164179	2.00747	Nuclear lamin, nucleus stability and chromatin structure
Ephx1	0.0429007	2.00765	Conversion of epoxides to trans-dihydrodoils for excretion
Scn4b	0.0418185	2.0079	Sodium channel, voltage-gated
Samd4	0.0318405	2.00926	RNA-binding, postranscriptional regulation
Cyfip1	0.0210452	2.0136	Regulates cytoskeleton and axon growth/formation
Fbln2	0.0057021	2.01365	Extracellular, neuronal structure development
Ankrd26	0.0291523	2.02058	Ankyrin, protein-protein interactions
Eya4	0.0268545	2.02241	Eye development, transcription activation
Mx2	0.0307161	2.02348	Dynamin and GTPase activity, heterochromatin formation
Ifnar2	0.0370614	2.02687	Interferon receptor, activates Janus kinases
Wdr62	0.0108659	2.03409	Cerebral development, function unknown
Kif14	0.0261951	2.03591	Kinesin, microtubule motor protein for transport
Aldh111	0.0205617	2.03926	Aldehyde dehydrogenase for removal of cytotoxic aldehydes
Syk	0.0130348	2.04443	Tyrosine kinase involved in cell proliferation, differentiation and phagocytosis
Ddb2	0.0142656	2.04721	DNA damage repair from UV light damage
Flnc	0.0458499	2.04823	Filamin involved in anchoring of actin cytoskeleton
4930452B06Rik	0.0425945	2.05001	Unknown
Kntc1	0.0116476	2.05073	Mitotic chromosome separation during anaphase
Arhgef10	0.0173248	2.0533	Rho GTPase, stimulates G-proteins

Lamc2	0.0297158	2.05539	Laminin component, extracellular component for remodelling and organization
Stab1	0.0215188	2.05942	Stabilin, receptor that endocytoses AGES, bacteria and lipoproteins
Notch1	0.0027652	2.06196	Intracellular signalling pathway mediator
BC006779	0.0150289	2.06524	DNA helicase, transcriptional activation
Klf4	0.0404938	2.06701	Transcription regulation and activation
Sfmbt2	0.0316905	2.07112	Regulation of transcription
Rgs22	0.0449781	2.0731	Regulator of G-protein signalling
Atp8b4	0.0050661	2.07605	Cation ATPase channel for phospholipid transport
Pde2a	0.0162072	2.07675	Phosphodiesterase, regulates endothelial cell proliferation and angiogenesis
Col25a1	0.0282262	2.07705	Component of collagen, brain specific
Tmbim1	0.0386395	2.07758	Inhibits apoptosis by reducing FAS trafficking
Abcc3	0.0025065	2.07795	ATP-binding cassette, transport of anions
Msn	0.0492673	2.07988	Moesin, links plasma membrane to actin as well as filopodia formation
Sult5a1	0.0181227	2.08116	Unknown
Baiap211	0.0451011	2.08365	Remodelling of the plasma membrane and cytoskeleton pedestal formation
Col19a1	0.0015986	2.08366	Component of collagen, maintenance of extracellular matrix
Pnpla7	0.0413459	2.08459	Lipid metabolism, function unknown
Col5a2	0.0343231	2.08489	Component of collagen
Tap2	0.0128675	2.08507	ATP-binding cassette, transport of peptides to endoplasmic reticulum
Fgfr2	0.0317797	2.08935	Fibroblast growth factor receptor, mitosis and differentiation
Fblim1	0.0248279	2.10192	Cell-cell adhesion modifications
Ecm2	0.0480193	2.10255	Extracellular matrix protein, function unknown
Dock2	0.0211604	2.10412	Cytoskeleton modifications for lymphocyte mobility
Taf1d	0.0085981	2.11135	RNA transcription regulation
Col1a2	0.0138499	2.11752	Component of collagen
Magt1	0.023233	2.12737	Magnesium transporter, N-glycosylation associated
Nckap11	0.0115969	2.12884	Cell morphology regulation (hematopoietic)
Ccdc150	0.0115571	2.12925	Unknown
Cd68	0.0202543	2.13446	Glycoprotein in microglial cells, promotes activation of macrophages
Svep1	0.0457553	2.13766	Unknown
Frem2	0.0232823	2.13927	Basement membrane epithelia protein
Vwa1	0.008256	2.14086	Extracellular matrix protein
Pebp4	0.0195098	2.14353	Lipid binding, unknown
Lrrk1	0.0116593	2.14701	Unknown
Gbp4	0.0277205	2.15078	Guanylate binding protein, interferon induced
Plcl2	0.009802	2.152	Phospholipase C, modifications to GABA receptors
Sgsm1	0.0143354	2.15698	G-protein signalling, GTPase activity

Ccl24	0.0205018	2.15907	Cytokine CC, activates T-lymphocytes
Kitl	0.0309238	2.16085	Tyrosine-kinase receptor, neural cell development, cell mobility
Kif20a	0.0312462	2.16108	Kinesin, microtubule motor protein for transport
Dnahc1	0.0089948	2.17058	Dynein, microtubule motor activity
Calcrl	0.0195045	2.17483	Calctonin receptor
Abcc4	0.0495595	2.18036	ATP-binding cassette involved in anion cycling
Abca13	0.0068074	2.18061	ATP-binding cassette, transport (unknown function)
H2-D1	0.0232616	2.1838	Major histocompatibility component
Hpse	0.0361428	2.18872	Heparanase, component of extracellular matrix and cell mobility
Nt5dc1	0.0014142	2.19853	5'-nucleotidase protein
Abca12	0.0210858	2.20527	ATP-binding cassette, small molecule transport
Ecm1	0.0133792	2.20762	Cell structure interaction for extracellular remodelling
Pth1r	0.0414317	2.21192	Parathyroid hormone receptor
Mertk	0.0157256	2.21711	MER/AXL/TYRO3 receptor kinase, involved in RPE phagocytosis regulation
Myom2	0.0104973	2.21919	Myomesin 2, muscle cell component
Clca1	0.0381724	2.22377	Chloride channel accessory protein
Pappa2	0.0093243	2.22752	Regulates intracellular insulin-like growth factor availability
AW551984	0.0204154	2.23252	Unknown
Nrap	0.0457788	2.23485	Nebulin-related anchoring protein, actin binding
Myo5c	0.0035317	2.23524	Myosin component, actin motor activity
Pld2	0.0463916	2.23549	Hydrolysis of phosphatidylcholine, cell cycle and transcription regulation
Serpina3a	0.0458079	2.23824	Serine peptidase inhibitor
Iqgap3	0.0152522	2.23908	GTPase activation
Ermp1	0.026493	2.24009	Endoplasmic reticulum metallopeptidase
Tnxb	0.0431861	2.24326	Tenascin, reduces cell adhesion
Fut8	0.0406937	2.24542	Transfers fucose to glycopeptides
4933428G20Rik	0.0347269	2.24603	Unknown
Mpa2l	0.0362691	2.24733	Macrophage activation like, GTPase activity
Trpm4	0.0428844	2.25314	Calcium ion channel, non-selective
Myof	0.0059642	2.27105	Membrane vesicle formation, membrane fusion
Pkhd111	0.0083783	2.2745	Potential role in immune response
Poln	0.0388691	2.2817	DNA polymerase subunit
Fn1	0.0160342	2.2906	Fibronectin glycoprotein, cell mobility
Sfrp4	0.0167618	2.29162	Regulates Wnt signalling, receptor protein
Myh11	0.02687	2.30494	Myosin, microtubule motor activity
Plin1	0.0405267	2.30902	Lipid storage, lipolysis regulation and inhibition
Ccdc46	0.0344049	2.31067	Unknown
Efcab7	0.0026008	2.32595	Unknown
Scml2	0.033619	2.33493	Transcription regulation (inhibition), polycomb protein

Myo15b	0.0067032	2.33608	Myosin, microtubule motor activity	
Cd93	0.0376391	2.33913	Immune response to apoptotic cells, cellular migration	
Aldh2	0.0450249	2.34462	Aldehyde dehydrogenase for removal of cytotoxic aldehydes	
Kynu	0.0160715	2.34696	Amino acid metabolism of tryptophan, NAD regeneration	
C920025E04Rik	0.0365824	2.34911	Major histocompatibility binding protein	
Galnt3	0.0292916	2.35444	Polypeptide N-acetylgalactosaminyltransferase	
Dse	0.0420728	2.36637	Induces immune response	
Gpd2	0.0399365	2.37661	Secondary step of glycerol metabolism, glycerol shuttle	
Pdzrn4	0.0249827	2.39325	Zinc-ring finger protein, unknown function	
Adhfe1	0.0358737	2.39843	Alcohol dehydrogenase	
Vipr1	0.0160948	2.40598	Activation of adenylate cyclase	
Fxyd5	0.0435462	2.40968	Ion transport regulator, functions during high chemokine production	
B2m	0.0238666	2.42012	Major histocompatibility complex heavy chain associated, microglial activation	
Renbp	0.0156324	2.42411	N-acetylglucosamine metabolism	
Dlec1	0.0277845	2.4261	Deleted in cancer progression, unknown function	
Itpr2	0.039317	2.43982	Inositol 1,4,5-trisphosphate receptor, cell signalling	
Slc44a5	0.0399251	2.44105	Solute carrier, unknown function	
Igtp	0.0366583	2.44422	Interferon gamma subunit	
Fas	0.0160064	2.45018	TNF receptor for cell apoptosis	
Mvp	0.0233917	2.45571	Regulates MAP kinase for drug metabolism	
Ncapd2	0.0265338	2.46031	Binds histones during mitosis	
Sec16b	0.0451595	2.46535	Endoplasmic reticulum organizational transport of mature proteins	
Tspan12	0.0455787	2.47621	Cell surface tetraspanin for cell growth and mobility	
Slc15a1	0.0488859	2.48534	Oligopeptide transporter for protein metabolism	
Osmr	0.0118207	2.49015	Interleukin receptor for inflammation response	
Gm7298	0.0104095	2.49301	Unknown	
Thsd7b	0.0234948	2.51864	Unknown	
1700011F14Rik	0.0159698	2.53491	Unknown	
Clic5	0.0454731	2.54845	Chloride intracellular channel	
Diap1	0.038375	2.54863	Axon guidance and elongation	
Nup133	0.0421358	2.54899	Formation of nuclear envelope during mitosis	
Mmel1	0.0193461	2.56265	Endopeptidase for phosphate homeostasis	
Atp10d	0.0486888	2.57034	ATPase ion transport across concentration and electrochemical gradients	
Obfc2a	0.0140833	2.57695	Single-stranded DNA binding protein for replication, repair and recombination	
Nos2	0.0316639	2.58763	Nitric oxide synthase for regulation of nitric oxide	
F3	0.0378579	2.58869	Factor 3 in blood coagulation	
Dnahc8	0.0312568	2.60645	Dynein, microtubule motor activity	
Sdk2	0.0422683	2.60932	Immunoglobin, synaptic maintenance	

Pola1	0.0082495	2.61239	Subunit of DNA polymerase involved in initiation of DNA replication	
Myole	0.0024885	2.61773	Non-muscle myosin for actin motor activity	
Dysf	0.0130688	2.64452	Membrane repair and regeneration	
Zfp748	0.0486775	2.67676	Unknown, zinc-finger protein	
Csmd2	0.00496	2.68896	Unknown	
Mllt4	0.0447334	2.70289	Organization of cell junctions and extracellular interactions	
Cast	0.0018883	2.71118	Calpain inhibitor which is involved in neuronal vesicle trafficking	
Aox311	0.0153934	2.71173	Retinol and amino-acid metabolism	
Asah2	0.0132518	2.71609	Production of sphingosine, induce apoptosis	
Fndc3c1	0.0074993	2.72731	Fibronectin	
Lama3	0.0032464	2.78592	Extracellular remodelling and migration	
Rassf2	0.0486192	2.78738	Tumor suppressor gene, Ras associated	
Mug1	0.0151463	2.80096	Unknown	
Cps1	0.0242031	2.81088	Primary, regulatory step of urea cycle and ammonia removal	
Il18r1	0.0389371	2.81156	Interleukin receptor for microglial activation and cytokine release	
Myh9	0.0255247	2.82263	Non-muscle myosin for cell mobility and morphology	
B230120H23Rik	0.0352319	2.82642	Unknown	
Utp20	0.0292205	2.84246	Small nucleolar RNA for 18S assembly	
Abca15	0.0133014	2.84975	ATP-binding cassette	
Adcy10	0.0050507	2.85833	Intracellular bicarbonate sensor and pH regulator	
Dock10	0.0090131	2.89996	Directs cytokinesis	
Hhip	0.0086926	2.91551	Involved in hedgehog recruitment for development	
Nrk	0.0136839	2.94234	Activates JNK pathway	
Clspn	0.0035071	2.94334	Cell cycle checkpoint protein, arrests cell cycle	
Gmnn	0.0215492	2.99088	Inhibits DNA replication, cell cycle checkpoint protein	
Cd97	0.0149192	2.99546	Intercellular interactions, involved in leukocyte activation and recruitment	
Xdh	0.0064959	3.00028	Metabolism of purines, involved in microglial activation	
Myo1c	0.0374662	3.00731	Myosin component for actin motor transport	
Als2cl	0.0282106	3.0433	Protein localization and transport	
Myolf	0.0001664	3.04358	Myosin component for microtubule transport	
H2-K1	0.0371741	3.06774	Major histocompatibility component	
Nfkbil2	0.0214434	3.14832	Inhibits NF-kappa-B activation	
F13a1	0.0234937	3.21301	Factor 13 in blood coagulation	
Plec1	0.0253006	3.22776	Unknown	
Kif4	0.0073226	3.3724	Kinesin microtubule motor activity	
Trpmб	0.0435942	3.47521	Ion channel for magnesium transport in epithelial cells	
Csmd3	0.0211546	3.53038	Unknown	
Bub1b	0.0050054	3.78249	Spindle checkpoint to inhibit cell cycle until proper chromosome separation	

Dpp4	0.0179589	4.02866	T-cell activation, involved in immune response
Muc5ac	0.0183852	4.03264	Extracellular remodelling

* Microarray was performed by A. Laliberté (2010).¹⁰³ Cut-offs set at: Fold change > \pm 1.75, p < 0.05.

[†] Annotation based on NCBI Gene¹⁸⁴ [National Center for Biotechnology Information, U.S. National Library of Medicine, Bethesda MD]

‡ Pathway is ordered by increasing p-value.

Curriculum vitae:

EDUCATION

Eric Dolinar

Department of Biology, The University of Western Ontario 1151 Richmond St. North, London, Ontario

EDUCATION	
Western University	London, Canada
Masters of Science in Biology Candidate – Cell and Molecular Biology	Expected: August 2014
• Supervisor: Dr. Kathleen Hill	
 Proposed Thesis Title: Early Mechanisms of Retinal De 	generation in the harlequin Mouse
Western University	London, Canada
Bachelor of Science (Honors) – Honors Specialization in Genetics	2012
Université de Moncton	Moncton, Canada

Completed the Explore 5-week Summer Immersion French Program

SCHOLARSHIPS & ACADEMIC HONORS

Best Graduate Presentation Travel Award – Ophthalmology Research Day (\$1000)		
Ontario Graduate Scholarship (\$15K)		
Western Science Graduate Entrance Scholarship (\$2500)		
Dean's Honor List, Faculty of Science, Western University		
Laurene Paterson Estate Scholarship (\$2000)	2011-2012	
James F. Birch Bursary (\$650)		
William F.M. and Helen M. Haight 125th Anniversary Alumni Bursary (\$1000)		
Western Scholarship of Excellence (\$2000)		
Queen Elizabeth Aiming for the Top Scholarship (\$3500)		
Governor General Academic Medal (Bronze)	2008	

PRESENTATIONS AND ABSTRACTS

Dolinar EA, MacPherson TC, Laliberté AM, Van Gaalen J, Hutnik CML, Hill KA. Non-metric Multidimensional Scaling of Electroretinogram Oscillatory Potential Data achieves Early Diagnosis of Retinal Degeneration in a Mouse Model of Mitochondrial Dysfunction. The Association for Research in Vision and Ophthalmology 2014, *poster presentation*.

Dolinar EA, Van Gaalen J, MacPherson TC, Hutnik CML, Hill KA. Electroretinogram Oscillatory Potentials provide a Non-invasive Diagnostic Analysis of Retinal Cell Function in a Mouse Model of Mitochondrial Dysfunction and Retinal Degeneration. Ophthalmology Research Day 2013 - Ivey Eye Institute, London Ontario, *oral presentation*

Li A, **Dolinar EA**, Van Gaalen J, MacPherson TC, Hutnik CML, Hill KA. Memantine Treatment does not shift the Aberrant Electroretinogram Oscillatory Potential Profiles of the Harlequin Mouse Model of Retinal Degeneration due to Mitochondrial Dysfunction. Ophthalmology Research Day 2013 - Ivey Eye Institute, London Ontario, *oral presentation*

Van Gaalen J, Bedore, J, McDonald I, **Dolinar EA**, Li A, Martyns A, Prado V, Prado M. Impaired Retinal Function is Associated with a Lack of Acetylcholine-Mediated Retinal Waves during Development. Ophthalmology Research Day 2013 - Ivey Eye Institute, London Ontario, *oral presentation*

Summer 2008

Faraz, MA, Li A, **Dolinar, EA**, Hill KA. Assessment of the Integrity of the Blood Retina Barrier in a Mouse Mimic of Oculodentodigital Dysplasia (ODDD)-Related Glaucoma. Ophthalmology Research Day 2013 - Ivey Eye Institute, London Ontario, *oral presentation*

Dolinar EA, Wagner J, Faraz MA, Cadesky A, Hill KA. An Eye to the Future: Identifying Early Disease Mechanisms in a Mouse Model of Retinal Degeneration. Biology Graduate Research Forum 2013 – Department of Biology, London, Ontario, *oral presentation*

Faraz, MA, Li A, Mayers J, **Dolinar EA**, Hill KA. Early mechanisms of retinal degeneration in a mouse mimic of oculodentodigital dysplasia (ODDD)-related glaucoma. International Conference on Clinical & Experimental Ophthalmology 2014, *oral presentation*

RESEARCH & TEACHING EXPERIENCE

Western University

Graduate Student – Department of Biology 2012-2013

- Managed the Experimental Eye Research Facility and animal colony for the duration of my project
- Technical Molecular biology skills include: PCR, DNA and RNA isolation, qPCR, histology, cloning, fully body mouse dissection including brain and retina, injections, and fluorescent microscopy
- Developed protocols for mathematically modelling (MATLAB) retinal waveforms using electroretinography
- Responsible for continual organization of all molecular biology-based projects for five undergraduate students including NSERC scholarship winners and fourth-year honors thesis students

Western University

Teaching Assistant – Gene Regulation (3597), Advanced Genetics (3595) and Organismal Physiology (2601) 2012-2013

- Organized tutorial discussions for a class of 150 students on current topics in gene regulation
- Lectured concepts, theories and novel ideas of gene regulation and advanced genetic techniques
- Graded final exams, lab reports, tutorial assignments and term papers

London, Canada

London, Canada