ASSESSMENT OF CEREBELLAR AND HIPPOCAMPAL MORPHOLOGY
AND BIOCHEMICAL PARAMETERS IN THE COMPOUND
HETEROZYGOUS, TOTTERING/LEANER MOUSE

A Thesis
by
EMILY MARY MURAWSKI

Submitted to the Office of Graduate Studies of
Texas A&M University
in partial fulfillment of the requirements for the degree of
MASTER OF SCIENCE

December 2009

Major Subject: Biomedical Sciences
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Approved by:

Chair of Committee, Louise C. Abbott
Committee Members, C. Jane Welsh
George Stoica
Head of Department, Evelyn Tiffany-Castiglioni

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ABSTRACT

Assessment of Cerebellar and Hippocampal Morphology and Biochemical Parameters in the Compound Heterozygous, Tottering/leaner Mouse. (December 2009)

Emily Mary Murawski, B.S., Texas A&M University

Chair of Advisory Committee: Dr. Louise C. Abbott

Due to two different mutations in the gene that encodes the $\alpha_{1A}$ subunit of voltage-activated CaV 2.1 calcium ion channels, the compound heterozygous tottering/leaner ($tg/tg^{la}$) mouse exhibits numerous neurological deficits. Human disorders that arise from mutations in this voltage dependent calcium channel are familial hemiplegic migraine, episodic ataxia-2, and spinocerebellar ataxia 6. The $tg/tg^{la}$ mouse exhibits ataxia, movement disorders and memory impairment, suggesting that both the cerebellum and hippocampus are affected. To gain greater understanding of the many neurological abnormalities that are exhibited by the 90-120 day old $tg/tg^{la}$ mouse the following aspects were investigated: 1) the morphology of the cerebellum and hippocampus, 2) proliferation and death in cells of the hippocampal dentate gyrus and 3) changes in basic biochemical parameters in granule cells of the cerebellum and hippocampus.

This study revealed no volume abnormalities within the hippocampus of the mutant mice, but a decrease in cell density with the pyramidal layer of CA3 and the hilus of the dentate gyrus. Cell size in the CA3 region was unaffected, but cell size in the
hilus of the dentate gyrus did not exhibit the gender difference seen in the wild type mouse. The cerebellum showed a decrease in volume without any decrease in cerebellar cellular density. Cell proliferation and differentiation in the subgranular zone of the hippocampal dentate gyrus remained normal. This region also revealed a decrease in cell death in the \textit{tg/tg}^{la} mice.

Basal intracellular calcium levels in granule cells show no difference within the hippocampus, but an increase in the \textit{tg/tg}^{la} male cerebellum compared to the wild type male cerebellum. There was no significant difference in granule cell mitochondrial membrane potential within the wild type and mutant animals in either the hippocampus or cerebellum. The rate of reactive oxygen species (ROS) production in granule cells revealed no variation within the hippocampus or cerebellum. The amount of ROS was decreased in cerebellar granule cells, but not granule cells of the hippocampus. Inducing ROS showed no alteration in production or amount of ROS produced in the hippocampus, but did show a ceiling in the amount of ROS produced, but not rate of production, in the cerebellum.
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<table>
<thead>
<tr>
<th><strong>NOMENCLATURE</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Ca$^{2+}$</td>
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<td>CGC</td>
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<td>Cacna1a&lt;sup&gt;rok&lt;/sup&gt;</td>
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<td>EA-2</td>
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<td>&lt;i&gt;tg/tg&lt;/i&gt;&lt;sup&gt;la&lt;/sup&gt;</td>
</tr>
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<td>TRH</td>
</tr>
<tr>
<td>SCA-6</td>
</tr>
<tr>
<td>VDCC</td>
</tr>
</tbody>
</table>
TABLE OF CONTENTS

<table>
<thead>
<tr>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABSTRACT .............................................................................................................. iii</td>
</tr>
<tr>
<td>ACKNOWLEDGEMENTS....................................................................................... v</td>
</tr>
<tr>
<td>NOMENCLATURE .......................................................................................... vi</td>
</tr>
<tr>
<td>TABLE OF CONTENTS ........................................................................................ vii</td>
</tr>
<tr>
<td>LIST OF FIGURES .......................................................................................... ix</td>
</tr>
<tr>
<td>CHAPTER</td>
</tr>
<tr>
<td>I INTRODUCTION ................................................................................ 1</td>
</tr>
<tr>
<td>Calcium Channel Mutant Mice ................................................................. 1</td>
</tr>
<tr>
<td>Voltage Dependent Calcium Channels .................................................. 16</td>
</tr>
<tr>
<td>Cerebellum ................................................................................................. 21</td>
</tr>
<tr>
<td>Hippocampus and Neurogenesis ............................................................. 23</td>
</tr>
<tr>
<td>Cellular Environment ............................................................................... 27</td>
</tr>
<tr>
<td>Objective of This Thesis .......................................................................... 28</td>
</tr>
<tr>
<td>II MORPHOLOGY OF THE CEREBELLUM AND HIPPOCAMPUS OF TOTTERING/LEANER MICE ............................................................................ 31</td>
</tr>
<tr>
<td>Introduction ................................................................................................. 31</td>
</tr>
<tr>
<td>Experimental Procedures ........................................................................ 32</td>
</tr>
<tr>
<td>Results ........................................................................................................ 34</td>
</tr>
<tr>
<td>Discussion ................................................................................................. 49</td>
</tr>
<tr>
<td>III PROLIFERATION AND DEATH IN THE DENTATE GYRUS OF TOTTERING/LEANER HIPPOCAMPUS ................................................................. 51</td>
</tr>
<tr>
<td>Introduction ................................................................................................. 51</td>
</tr>
<tr>
<td>Experimental Procedures ........................................................................ 54</td>
</tr>
<tr>
<td>Results ........................................................................................................ 59</td>
</tr>
<tr>
<td>Discussion ................................................................................................. 71</td>
</tr>
</tbody>
</table>
# LIST OF FIGURES

<table>
<thead>
<tr>
<th>FIGURE</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.1 Comparison of phenotypic onset and severity of mutant genotype disorders.</td>
<td>15</td>
</tr>
<tr>
<td>1.2 Diagram of α1A VDCC subunit</td>
<td>20</td>
</tr>
<tr>
<td>1.3 Representative photo of the mouse hippocampus</td>
<td>26</td>
</tr>
<tr>
<td>2.1 Quantitative comparison of volume within brain regions</td>
<td>36</td>
</tr>
<tr>
<td>2.2 Cellular density in the hippocampus</td>
<td>38</td>
</tr>
<tr>
<td>2.3 Cellular density in the cerebellum</td>
<td>42</td>
</tr>
<tr>
<td>2.4 Cell size in hippocampal areas</td>
<td>48</td>
</tr>
<tr>
<td>3.1 Representative image of neuronal death in the subgranular zone of the dentate gyrus as revealed by Fluoro-Jade stain</td>
<td>61</td>
</tr>
<tr>
<td>3.2 Image and graph of BrdU immunohistochemistry</td>
<td>63</td>
</tr>
<tr>
<td>3.3 Representative images of PCNA immunohistochemistry</td>
<td>64</td>
</tr>
<tr>
<td>3.4 Graphs of comparison of 3-month-old $tg/tg^{la}$ and wild type PCNA labeled regions</td>
<td>65</td>
</tr>
<tr>
<td>3.5 Images of NeuN/BrdU immunohistochemistry</td>
<td>68</td>
</tr>
<tr>
<td>3.6 Images of GFAP/BrdU immunohistochemistry</td>
<td>69</td>
</tr>
<tr>
<td>3.7 Graphs of percent cells labeled GFAP/BrdU and NeuN/GFAP</td>
<td>70</td>
</tr>
<tr>
<td>4.1 Image of DAPI/NeuN labeled granule cells</td>
<td>79</td>
</tr>
<tr>
<td>4.2 Image of CaG-1 and bright field granule cells</td>
<td>80</td>
</tr>
<tr>
<td>4.3 Graphs of CaG-1 percent control of fluorescence</td>
<td>81</td>
</tr>
<tr>
<td>4.4 Image and graph of TMRM percent control fluorescence</td>
<td>84</td>
</tr>
<tr>
<td>4.5 Graph of amount and rate of ROS production</td>
<td>87</td>
</tr>
</tbody>
</table>
4.6 Comparison of results from thesis experiments for $tg/tg^{ja}$ mouse as compared to known observations in $tg$ and $tg^{ja}$ mice.
CHAPTER I
INTRODUCTION

CALCIUM CHANNEL MUTANT MICE

Mutation Characterization

A number of spontaneously occurring mutations and genetically engineered alleles of the $\alpha_{1A}$ subunit of P/Q type voltage dependent calcium channels (VDCC) has allowed for greater understanding of this complex structure. The CaV 2.1 channel is highly expressed throughout the mouse cerebral cortex, hippocampus, olfactory bulb, tectum, and hindbrain (Fletcher et al., 1996). The pore-forming subunit of the Cav2.1 voltage gated calcium channel (VGCC), alternatively known as the P/Q-type VGCC, is highly expressed in cerebellar granule and Purkinje cells (Westenbroek et al., 1995).

Both the tottering ($tg$) and leaner ($tg^{la}$) allelic mutations reside in chromosome 8 (Green and Sidman, 1962) and in the gene that encodes the alpha1A subunit of the voltage-activated CaV 2.1 calcium ion channels (Fletcher et al., 1996). This $tg$ gene is prevalent throughout the mouse central nervous system in areas such as the cerebral cortex, hippocampus, olfactory bulb, tectum, hindbrain and cerebellar granule and Purkinje cells (Fletcher et al., 1996). Several mouse mutants that display different calcium channel mutations are of great interest because of their relationship to a variety of devastating human neurological disorders. Extensive research into autosomal

This thesis follows the style of Brain Research.
recessive mutations at the tg locus, including the leaner and homozygous tottering mutations, has proven valuable in understanding human diseases such as familial hemiplegic migraine (FHM), episodic ataxia-2 (EA-2) (Ophoff et al., 1996), and spinocerebellar ataxia 6.

**Human Disorders**

FHM, an autosomal dominant mutation on chromosome 19 (Joutel et al., 1993), is an inherited form of migraine with some rare sporadic, noninherited, cases. FHM is characterized by visual perceptual disturbances, weakness in half of the body, motor and sensory losses (Blau and Whitty, 1955; Whitty, 1953) and in some cases, ocular motor defecit (Elliott et al., 1996), cerebellar atrophy (Terwindt et al., 1996), ataxia, nausea, vomiting (Blau and Whitty, 1955), acute paranoid psychosis with anxiety and visual hallucinations (Spranger et al., 1999). Rare documented cases have experienced fever, seizures and coma (Ducros, 2000). These preliminary aura symptoms are known to occur for 10 minutes to hours before onset of migraine headache which can last anywhere from two hours to seven days (Ahmed et al., 1996; Ducros, 2000). Within affected families and even within each patient, symptoms and severity of aura and migraine can vary (Ducros, 2000). This, in addition with incomplete penetrance within families, suggest that the symptoms of FHM could be a combination of genetic and environmental factors (Ducros, 2000).

Onset of symptoms occur in most often in youth, but can commonly begin in those between the ages of 4 and 30 (Ahmed et al., 1996). FHM is relatively rare, with
only over 47 families affected worldwide (Ahmed et al., 1996). Both stress and minor head trauma are known triggers of this migraine in roughly 2/3 of patients (Ophoff et al., 1996; Terwindt et al., 1996). Because this is an allelic mutation, patient treatment is often aimed at the symptoms of migraine and promising treatments include acetazolamide as a preventative (Battistini et al., 1999) and intranasal ketamine to shorten length of attacks (Kaube et al., 2000).

There is much clinical overlap with FHM and episodic ataxia type 2 (EA 2). EA 2, an autosomal dominant paroxysmal cerebral disorder, lies on the same chromosome and interval as the FHM locus (Kramer et al., 1995; Teh et al., 1995) and is also responsive to acetazolamide. Characteristics of EA 2 include vertigo, visual disturbances, dysarthria, ataxia and in some cases nyastagnus and other cerebellar dysfunctions much less severe than those of patients with SCA 6 (Jodice et al., 1997).

Many argue that SCA 6 and EA 2 are the same disorder with great phenotypic variability (Frontali, 2001; Geschwind et al., 1997; Jodice et al., 1997), while others stand by their belief that these as genetically and phenotypically distinct disorders (Zhuchenko et al., 1997). Predominant hallmarks that distinguish these mutations phenotypically are the presence of extracerebellar signs and a larger proportion of those with ataxia in those with SCA 6 (Frontali, 2001). It is possible that this discrepancy could be derived from different ages of examination or pathological differences between the disorders (Frontali, 2001).

SCA 6 results from a polyglutamine track created by small expansions in the CAG repeat in the coding region of CACNA1A gene(Zhuchenko et al., 1997). Typical
symptoms of SCA 6 include gait disturbances, slurred speech, eye movement disorder, paroxysmal vertigo and migraine (Craig et al., 2004). The characteristic cerebellar atrophy and functional deficit progress in severity over time and in some cases lead to permanent ataxia (Calandriello et al., 1997; Geschwind et al., 1997).

It is interesting to note that this mutation appears to increase calcium channel current and increased expression of \( \alpha_{1A} \) at the cell surface, which could allow calcium ions (\( \text{Ca}^{2+} \)) to enter the cell at toxic levels (Piedras-Renteria et al., 2001). This discovery contradicts other \( \alpha_{1A} \) mutations in which there is a decrease of P/Q type calcium channel current (Dove et al., 1998; Lorenzon et al., 1998; Wakamori et al., 1998).

Like FHM and EA 2, in some reported cases, SCA 6 episodes are responsive to treatment with acetazolamide (Calandriello et al., 1997). In addition, it is proposed that the drugs gabapentin and pregabalin help to decrease ataxia in patients with SCA 6 (Gazulla and Benavente, 2007; Gazulla and Tintore, 2007). SCA 6 is more prevalent in populations than the previously presented human disorders, with an estimated 1 in 19,210 people with SCA 6 or at risk for SCA 6 in the northeast of England alone (Craig et al., 2004).

**Leaner Mutation**

Of these mutations, the leaner mutation (\( \text{tg}^{la} \)) is considered the most severe. Homozygous leaner mice experience early onset of disorders, at postnatal days (P)8-10, which include debilitating ataxia, absence epilepsy and paroxysmal dyskinesia (Herrup and Wilczynski, 1982). These disorders in \( \text{tg}^{la} \) mice result from a splice site mutation in
the 5’ end of the α1A gene that truncates the carboxy terminus of the α1A protein, which likely creates two α1A splice variants (Doyle et al., 1997; Fletcher et al., 1996).

The tg^{la} mouse is known to exhibit cerebellar granule cell (CGC) loss that begins soon after P10, with the greatest amount of death at P20. These mice still P40 still experience a marked CGC loss as compared to age matched wild type mice at P30 and P40 (Lau et al., 2004). More specifically, there was an increase in CGC death in the anterior lobe as compared to the posterior lobe of the leaner cerebellum (Frank et al., 2003; Lau et al., 2004). This increase in CGC death is by way of apoptosis and continues into adulthood (Bawa and Abbott, 2008). In addition, the pattern of CGC death follows the same spatial pattern in adulthood (Bawa and Abbott, 2008) as initially expressed during postnatal development (Lau et al., 2004).

While there is no apparent change in basal intracellular calcium homeostasis in leaner Purkinje cells (Dove et al., 2000), there is an absolute decrease in the levels of calretinin expression in leaner CGC (Nahm et al., 2002) as well as a marked decrease in the number of CGCs (Bawa and Abbott, 2008). Mitochondrial membrane potential is also decreased in CGCs of the adult leaner mouse as compared to both wild type and tottering mice (Bawa and Abbott, 2008). Interestingly, there is no significant alteration in generation of reactive oxygen species in CGCs of leaner mice as compared to wild type or tottering mice (Bawa and Abbott, 2008). It is hypothesized that CGC death in the leaner mouse may be more dependent upon an apparent mitochondrial dysfunction rather than decreased basal intracellular calcium levels (Bawa and Abbott, 2008).
The complexity of calcium homeostasis in leaner Purkinje cells has been revealed as investigations have ventured into calcium buffering mechanisms within the cell. An additional decrease in calcium current amplitude in the cerebellum possibly creates the noticeable decrease in calbindin and parvalbumin expression in Purkinje cells (Dove et al., 2000). Despite no alteration in leaner Purkinje cell resting somatic Ca\textsuperscript{2+} concentration, there is a reduction in Ca\textsuperscript{2+} buffering (Dove et al., 2000). There is not only diminished Ca\textsuperscript{2+} uptake by the endoplasmic reticulum in leaner mice, but reductions in mRNA levels of Ca\textsuperscript{2+} binding proteins which are a proposed result of the P/Q type mutation that creates this severely affected phenotype (Dove et al., 2000).

As a possible result of altered calcium homeostasis, the adult leaner mouse exhibits decreased neuronal nitric oxide synthase (NOS) mRNA expression and NADPH-diaphorase histochemical staining, with no alteration in levels of neuronal NOS protein as compared to age matched wild type mice (Rhyu et al., 2003). It has been proposed that these decreases may be attributed to their severe neuronal deficits in which nitric oxide acting as a trophic factor is unable to prevent granule and Purkinje cell death in the leaner mouse cerebellum (Rhyu et al., 2003).

Purkinje cells have been closely examined in the leaner mouse as these cells are now known to be highly concentrated in P/Q type VDCC and have a 60% reduction in P/Q type VDCC calcium current and one third the channel open probability of that in the wild type animal (Dove et al., 1998; Lorenzon et al., 1998; Wakamori et al., 1998). This reduction in Ca\textsuperscript{2+} current is not a result of \(\alpha_{1A}\) abundance, as the amounts of mRNA and protein for \(\alpha_{1A}\) subunits are unaltered (Lau et al., 1998). The leaner cerebellar cortex
and inferior olive are marked with neuronal loss (Heckroth and Abbott, 1994). Purkinje cell death in the cerebellum is increased in the leaner mouse compared to wild type mice where it begins at P25, peaks in the vermis at P40, in the hemispheres at P50 and continues to show low levels of cell death at P80 (Frank et al., 2003). In addition, the leaner cerebellum expresses decreased serum and cerebellar insulin-like growth factor-I at P30 compared to age matched wild type mice (Nahm et al., 2003). Exogenous administration of insulin-like growth factor-I is shown to improve cerebellar function in leaner mice, but remains futile in arresting Purkinje cell death (Nahm et al., 2003).

The leaner mouse, in addition to tottering and compound heterozygous tottering/leaner, exhibits enlarged parallel fiber varicosities that synapse on multiple Purkinje cells dendritic spines in both juvenile and adult animals (Rhyu et al., 1999). With closer examination, the leaner mouse has a greater amount of these synapses to the Purkinje cells dendritic spines than the tottering and compound heterozygous tottering/leaner mouse which may indicate greater morphological abnormality linked to a more severe phenotype (Rhyu et al., 1999).

Because this malformation precedes the expression of the disorder in the tottering mouse, it is possible that it is not a result of the pathogenic change (Rhyu et al., 1999). On the other hand, Purkinje cell and parallel fiber synapse alterations are more apparent in the adult animal and this increase could be a result of prolonged interaction and compensation of this mutation (Rhyu et al., 1999). It is also noted that $tg$, $tg^{la}$ and $tg/tg^{la}$ juvenile and adult genotypes all display axonal swellings which allude to Purkinje cell
damage that could cause impaired axonal transport (Abbott et al., 1996; Rhyu et al., 1999).

**Tottering Mutation**

The tottering (tg) mutation remains less severe with a later phenotypic onset at P21-28, and clinical signs that include: mild ataxia, paroxysmal dyskinesia and absence seizures (Noebels and Sidman, 1979). The tottering deficits are a result of a point mutation residing in the extracellular region linking the IIS5 and IIS6 domains in the α1A protein, which is responsible for a proline-to-leucine amino acid substitution (Fletcher et al., 1996).

Various cerebellar morphological deficits exist in the adult tottering mouse compared to age-matched wild type mice. Overall cerebellar volume is decreased in the mutant, along with a decrease in the volume of the Purkinje cells in the cerebellar molecular layer of 3-month-old tottering mice, with specific reduction in the posterior cerebellar region (Isaacs and Abbott, 1995). Molecular layer Purkinje cell density remains unchanged in the mutant cerebellum, without evidence of excessive Purkinje cell loss (Isaacs and Abbott, 1995). Forebrain, cerebellum and hindbrain weight are significantly less in the adult tottering mouse as compared to age matched wild type, although malnutrition is not thought to be responsible for this weight decrease (Isaacs and Abbott, 1995). The ratio of cerebellum to body weight remains greater in adult tottering mice than in age matched wild type mice which rules out the hypothesis that
smaller brain weight is merely a result of overall body weight loss (Isaacs and Abbott, 1995).

Early postnatal tottering mice do not exhibit the expected increase in P-Type VDCC current density in Purkinje cells and at P8 continue to show a 25% decrease as compared to age matched wild type mice (Erickson et al., 2007). Because tottering mice at P15 begin to display normal levels of Purkinje cell P-Type VDCC current density, it is believed that there is an obvious developmental delay that precedes any visible behavioral abnormalities (Erickson et al., 2007).

While the P/Q type VDCC are likely to be the primary responsibility for the observed neurological deficits, it is proposed that L-type VDCC may play a small role in episodes of paroxysmal dyskinesia (Campbell and Hess, 1999). L-Type VDCC blocking agents have shown to reduce this specific type of motor deficit in the tottering mouse by presumably acting on the central nervous system rather than skeletal or cardiac muscle (Campbell and Hess, 1999). In addition, tottering mouse Purkinje cells exhibit mRNA upregulation of the L-type VDCC α1C subunit (Campbell and Hess, 1999), and experience a two fold increase in calcium current density at P15 through these channels as compared to wild type animals (Erickson et al., 2007). Such an increase would be consistent with an upregulation of this VDCC in response to P-type VDCC dysfunction (Erickson et al., 2007). Further analyses indicate compensation of Cav1 channels in basal forebrain neurons (Etheredge et al., 2007) and N-type channels in the Schaffer collateral cells neurotransmitter release (Qian and Noebels, 2000) in response to the Cav2.1 channel mutation.
Tottering Purkinje cells show axonal damage beginning at 3 to 5 weeks of age and persist through adulthood, encouraging the interpretation that postsynaptic activities may be altered in cerebellar nuclei and help to create their ataxic phenotype (Hoebeek et al., 2008).

The tottering mutation is likely the culprit for an observed decrease in CGC basal intracellular calcium homeostasis (Bawa and Abbott, 2008). In addition, there is a noticeable decrease in the tottering CGC mRNA expression of calretinin, calcium binding protein, which could be correlated with impaired calcium homeostasis and motor coordination (Cicale et al., 2002). The tottering Purkinje cell layer in the cerebellar flocculus has shown a decrease in the mRNA expression of ryanodine receptor type 1, a calcium channel in the endoplasmic reticulum, which may alter synaptic plasticity and be attributed to their ataxic phenotype (Cicale et al., 2002).

Tyrosine hydroxylase (TH) expression in the tottering and compound heterozygous tottering/leaner cerebellum also is altered (Abbott et al., 1996; Austin et al., 1992; Hess and Wilson, 1991), while expression of Zebrin II remains like that of normal mice and exhibits areas positive for TH immunoreactive for Zebrin II and those negative for TH displaying no immunoreactivity for Zebrin II (Abbott et al., 1996).

In contrast to the leaner mouse, there is a significant increase in NOS mRNA and protein in the cerebellum and NADPH-diaphorase histochemical stain in the cerebellar molecular and granule cell layer of adult tottering and rolling Nagoya mice as compared to age-matched wild type mice (Rhyu et al., 2003). This is presumed to be a result of
their VDCC malformation and could be a potential byproduct of some mechanism that prevents their cell death (Rhyu et al., 2003).

Epileptic seizures of the tottering mouse are postulated to be correlated with excitability defects in the CA3 region of the hippocampus (Helekar and Noebels, 1991). Closer examination reveals prolonged paroxysmal depolarizing shifts in hippocampal CA3 pyramidal neurons after exposure to high potassium levels in vitro during the developmental stage of the tottering mouse (Helekar and Noebels, 1992). As this abnormality is most prominent in young tottering mice, it appears to possibly be a part of a genetically signaled neuronal synchronization mutation rather than an effect of a life of seizures (Helekar and Noebels, 1992).

**Compound Heterozygous, Tottering/Leaner Mutation**

The compound heterozygous, tottering/leaner (tg/tg\(^{la}\)) mouse carries one tottering allele and one leaner allele. Much less is known about the tg/tg\(^{la}\) mouse, which also exhibits several neurological deficits to a lesser degree than the leaner mouse, but to a greater degree than the tottering mouse. The characteristic phenotype of tg/tg\(^{la}\) mutant mice is noticeable around P20-22 with a moderate ataxia (Green and Sidman, 1962; Herrup and Wilczynski, 1982).

While the tg/tg\(^{la}\) mouse appears to be a “compromise” between the leaner and tottering mutations, the morphological and biochemical characteristics remain unique, seemingly mimicking neither mouse. The tottering and leaner mutations result in ataxia and movement disorders, suggesting that the cerebellum is affected. Previous studies of
cerebellar structure in mice carrying these mutations show atypical functioning of Purkinje cells that are likely related to the ataxic phenotype (Rhyu et al., 1999). It is understood that the leaner mouse but not tottering mice, experience significant cerebellar granule, Golgi and Purkinje cell death, but little is known regarding potential excessive neuron cell death the tg/tg<sup>la</sup> genotype (Herrup and Wilczynski, 1982; Isaacs and Abbott, 1995). In addition, because some mice that carry mutations in this gene exhibit atypical spatial and memory learning, the hippocampus is also a possible site of abnormality.

Preliminary investigation of tg/tg<sup>la</sup> mice has revealed that the weight of the forebrain, cerebellum and hindbrain are significantly decreased in the adult tg/tg<sup>la</sup> mouse as compared to age matched wild type mice (Isaacs and Abbott, 1995). Malnutrition is not thought to be responsible for the significant decrease in brain weights (Isaacs and Abbott, 1995). Like the tottering mouse, overall cerebellar volume is decreased in the tg/tg<sup>la</sup> mouse, along with a decrease in the volume of the Purkinje cells in the molecular layer of the 3 month old tottering cerebellum (Isaacs and Abbott, 1995). Purkinje cell density per unit of cerebellar molecular layer remains unchanged in the tg/tg<sup>la</sup> cerebellum, without evidence of excessive Purkinje cell loss (Isaacs and Abbott, 1995). Adult tg/tg<sup>la</sup> mice show a ratio of cerebellum to body weight that is greater than that of age-matched wild type mice, which rules out the hypothesis that smaller brain weight is merely a result of overall body weight loss (Isaacs and Abbott, 1995).

As reported previously, the tg/tg<sup>la</sup> mouse exhibits enlarged parallel fiber varicosities that synapse on multiple Purkinje cells dendritic spines in both juvenile and adult animals (Rhyu et al., 1999). Purkinje cells in the tg/tg<sup>la</sup> cerebellum show
significantly higher mRNA expression of tyrosine hydroxylase in both juvenile and adult pre seizure animals as compare to age matched wild type controls, indicating that the occurrence of these neurological abnormalities is likely to be independent of seizure onset (Austin et al., 1992).

Additional mutations affecting the $\alpha_{1A}$ subunit include the Nagoya rolling (Cacna1a$^{rol}$) and rocker (Cacna1a$^{rkr}$) mutations. The Nagoya (rolling) mouse, which is a Cacna1a mutant that originated from Nagoya University in Japan, has an autosomal recessive mutation, ($tg^{rol}$) on chromosome 8 that demonstrates an ataxic phenotype with $\alpha$ motor disturbances in its hind legs, much like the tottering and leaner mouse in which this mutation shares a locus (Oda, 1981; Tsuji and Meier, 1971). An ataxic gait and difficulty balancing beginning at P10-14 is a characteristic phenotype of these mice (Oda, 1981). Similarities with the leaner or tottering mutant mouse include loss of granule cells in the anterior cerebellum (Nishimura, 1975; Suh et al., 2002) presumably by way of apoptosis (Suh et al., 2002), altered Purkinje cell synapses (Rhyu et al., 1999), altered ryanodine receptor expression (Sawada et al., 2008) and persistent tyrosine hydroxylase expression in the cerebellum (Muramoto et al., 1981). Cerebellar tyrosine hydroxylase expression in Nagoya rolling mice is also correlated with increased numbers of corticotrophin-releasing factor-positive climbing fibers, but not mossy fibers, which could alter the function of connecting Purkinje cells (Sawada et al., 2001).

The cerebellum, parietal cortex, ventral tegmental area and basal ganglia of the rolling mouse is understood to have low levels of local cerebral glucose utilization (Kato et al., 1982; Kinoshita et al., 1995; Yamaguchi et al., 1992), a neuronal metabolic index
for neuronal activity (Sokoloff et al., 1977; Sokoloff, 1981). Dysfunction in the basal
ganglia is thought to lie predominantly in the striatal projection neurons (Taniwaki et al.,
1996). Thyrotropin-releasing hormone (TRH: L-pyroglutamyl-L-histidyl-L-
prolinamide) and its analog TA-0910 ((-)N-[S]-hexahydro-L-methyl-2,6-dioxo-4-
pyrimidinylcarbonyl]-L-histidyl-L-prolinamide tetrahydrate) are observed to reduce
ataxia in the rolling mouse (Kinoshita et al., 1995) possibly by way of activating
thryotrophin-releasing hormone receptors in the ventral tegmental field directly and
through receptors in the cuneiform nucleus indirectly (Kinoshita et al., 1997).

A more recent discovery of a Cacna1a allelic mutant mouse is rocker
(Cacna1a<sup>kr</sup>), which stems from the fourth generation offspring of a male mouse
mutagenized with ethylnitrosourea (Bode, 1984). A proline to leucine substitution at
position 1310 on chromosome 8 affects the alteration of polarity of the extracellular
membrane near the pore-forming region of P/Q type VDCC (Zwingman et al., 2001).

Characteristics of the rocker mouse appear very similar to that of the tottering
mouse (Figure 1.1), with an ataxic gait initiated by an action tremor (Zwingman et al.,
2001). Onset of behavioral conditions is around P21-28 (Zwingman et al., 2001).
Although the rocker mouse is phenotypically reminiscent of the tottering mouse, its
brain shows quite different patterns of neurological abnormalities. Cerebellar
morphology and cytoarchitecture remains normal along with no Purkinje cell loss or
abnormal staining patterns with various neuronal or non-neuronal antibodies (Zwingman
et al., 2001). There are abnormalities in dendritic structure in cerebellar Purkinje cells as
rocker mice age, with normal appearance in 6-month-old rocker mice, but reduced
dendritic mass and secondary growth of distal dendrite tips in 1-year-old rocker mice (Zwingman et al., 2001). In addition, Purkinje cells exhibited no axonal swellings, ectopic spines on distal branches or aberrant TH expression in adult rocker mice (Zwingman et al., 2001).

Rocker mice show identical morphological and temporal patterns of cortical burst activity and behavioral seizure arrest to that observed for tottering mice (Zwingman et al., 2001). Rocker mice display moderately reduced calcium current density in cerebellar Purkinje cells with normal gating and voltage dependence (Kodama et al., 2006). While presynaptic parallel fibers to Purkinje cell synapses appear normal, postsynaptic function is impaired (Kodama et al., 2006). Decreased number and density of postsynaptic a-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors may suggest that even a mild calcium mutation, such as rocker, can have profound neurological effects (Kodama et al., 2006).

<table>
<thead>
<tr>
<th></th>
<th>tottering</th>
<th>tottering/leaner</th>
<th>leaner</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cerebellar ataxia</strong></td>
<td>Mild ~P20</td>
<td>Moderate ~P16</td>
<td>Severe ~P10</td>
</tr>
<tr>
<td><strong>Absence epilepsy</strong></td>
<td>Present (1-3 sec.) ~P24</td>
<td>Present (2-5 sec.) ~P20</td>
<td>Present (4-7 sec.) ~P14</td>
</tr>
<tr>
<td><strong>Paroxysmal dyskinesia</strong></td>
<td>Present (20-30 min.) ~P25</td>
<td>Present (20-40+ min.) ~P22</td>
<td>Present (short) ~P16 Disappear after ~P40-60</td>
</tr>
</tbody>
</table>

Figure 1.1
Comparison of phenotypic onset and severity of mutant genotype disorders.
VOLTAGE DEPENDENT CALCIUM CHANNELS

Calcium homeostasis and neuronal calcium ions in neurotransmission are critical components for cell function and survival. Many experts in the field hypothesized that multiple types of VDCC existed, but not until the late 1980’s was clear progress made to actually describe them (Bean, 1989). VDCC are classified into two distinct groups based on the principal of voltage-activation threshold. These two groups or categories are high-voltage (HVA) channels and low-voltage activated (LVA) channels (Carbone and Lux, 1984; Fedulova et al., 1985). LVA, or T type channels, exhibit a voltage activation threshold of less than -50mV and inactivate quickly in contrast to HVA, that function at a voltage activation threshold of less than -20mV and inactivate more slowly (Carbone and Lux, 1984; Nowycky et al., 1985).

HVA channels are further divided into several different channels (L, N, P, Q, O and R types), based upon specific properties of each channel with respect to physiological and pharmacological parameters (Bean, 1989; Hess, 1990; Tsien et al., 1988). Of the HVA channels, the L type is sensitive to dihydropyridine (DHP) and several peptide toxins, while N, P, Q, O and R are all insensitive DHP. Second to be discovered was the N type channel, which is a class of calcium ion channel with greater heterogeneity and therefore is best characterized by its blockage by ω-conotoxin GVIA (McCleskey et al., 1987; Nowycky et al., 1985; Tsien et al., 1988). Further analysis of peptide toxins revealed the R type channel, which is believed to have multiple subtypes (Tottene et al., 1996) and whose complete blockage is still to be elucidated (Randall and Tsien, 1995).
Of greatest importance for this study are the P and Q type channels. The P type channels exhibit similar conductance as L and T type channels and voltage dependence between N and T type channels. P type channels were first discovered in Purkinje cells (Llinas et al., 1989; Regan, 1991) and were noted for their resistance to L and T type channel blocking agents such as DHP and w-conotoxin. One particular toxin has proven to block P type channels is the venom of the funnel-web spider, *agelenopsis aperta* and more specifically, its toxic peptide component, ω-Aga IVA (Llinas et al., 1992; Mintz et al., 1992; Takahashi and Momiyama, 1993). Another toxin, ω-CgTx MVIIC, derived from the marine cone snail, *Conus magus*, similarly inhibits P type channels in addition to Q- and O- channel conductance and N- type channels (Hillyard et al., 1992; Takahashi and Momiyama, 1993). The P type channel persists throughout the mammalian nervous system, with particularly high expression in cerebellar granule cells, (Randall and Tsien, 1995) neuroendocrine cells and the cell soma and dendrites of Purkinje cells (Usowicz et al., 1992), and P type channels are thought to have critical function in neuronal integration and death (Llinas et al., 1992).

While the Q type channel is fully blocked by ω-CgTx MVIIC (Hillyard et al., 1992), it’s partial resistance to ω-Aga IVA at concentrations known to block that of the P type channel is its novel characteristic (Randall and Tsien, 1995; Zhang et al., 1993). The Q type is the most highly represented of the channels in cerebellar granule cells (Randall and Tsien, 1995). Because Q type channel malformation that is so highly present in CGCs, this type of cell begs for closer examination regarding its role in the \(tg/tg^{la}\) phenotype.
VDCC consist of a complex structure that allows for specific channel functions. There are four to five subunits, β, γ, α2-δ and α1, involved in forming the complete channel. The β subunit is intracellular with α helices (Ruth et al., 1989). The γ subunit is a glycosylated hydrophobic protein with four transmembrane domains and two N-linked glycosylation sites (Jay et al., 1990). A single gene for the α2-δ subunit is post-translationally cleaved to allow for a disulfide link between α2 and δ proteins (De Jongh et al., 1990; Ellis et al., 1988; Jay et al., 1991). The α2-δ subunit is a transmembrane domain which mediates subunit interactions and is attached to a glycosylated extracellular domain that enhances current amplitude (Gurnett et al., 1996).

While each subunit plays an integral part, the diversity of calcium channels is owed to the α1 subunit (Hofmann et al., 1994). The α1 subunit was discovered to act in all voltage dependent calcium channels and is understood to act as the conduction pore, voltage sensor and gating apparatus and a common site of second messenger, drugs and toxins channel regulation (Sather et al., 1993; Smith et al., 1996; Stea et al., 1994). In mammals, the α1 subunit is controlled by at least ten genes. It consists of four homologous mostly hydrophobic domains I, II, III and IV and is believed to have six membrane spanning subunits (S1-6) within each domain. In the α1 subunit, the S4 subunit is believed to act as specifically as part of the voltage-sensing mechanism for the channel whereas the channel pore proper is formed between the hydrophobic domains of S5 and S6.

Using cDNA analysis and Northern blotting, the P/Q type calcium channel was found to have the α1A subunit, also referred to as Cav2.1. This subunit is prevalent
throughout the nervous system, heart and pituitary, with high concentration in the cerebellum. It accounts for the largest number of VDCC subunits within the entire nervous system (Mori et al., 1991) and can be found in cell bodies, dendrites and presynaptic terminals (Westenbroek et al., 1995). It is noted that CaV 2.1 channels account for about 45% of calcium channels in cerebellar granule cells and over 90% of calcium channels on the Purkinje cell somata of mammals (McDonough et al., 1997; Mintz et al., 1995; Randall and Tsien, 1995).

The $\alpha_{1A}$ subunit (Figure 1.2) is thought to express various electrophysiological properties according to the type of $\beta$ subunit coexpressed (Stea et al., 1994). The coexpression of $\alpha_{1A}$ with a $\beta 2\alpha$ subunit appears more similar to the P type VDCC current waveform, while $\beta 1b$ or $\beta 3$ lend themselves to mimic the Q type VDCC current waveform (Stea et al., 1994). The $\alpha_{1A}$ subunit is most convincingly present in the P type VDCC because of the high localization of $\alpha 1\alpha$ transcripts and common immunoreactivity between the two (Westenbroek et al., 1995).

Recent discoveries highlight the important role of signal transducing molecules on modulating Cav2.1 channels. In the mouse Purkinje cells, immunohistochemical and electrophysiological evidence has shown that this subunit is also responsible for GABAergic inhibitory transmission (Stephens et al., 2001). It also colocalizes tightly with GAD, indicating that GABA is likely produced in this region thought to comprise mostly of basket cell terminals (Stephens et al., 2001).

Calcium modulating protein, calmodulin (CaM) is known to mediate $\text{Ca}^{2+}$-dependent facilitation and inactivation as it binds to local calcium after a voltage-gated
calcium influx through Cav2.1 channels and induces conformational changes (DeMaria et al., 2001; Lee et al., 2003). An intricate mechanism allows for positive regulation of Cav2.1 channels in the instances of local \( \text{Ca}^{2+} \) increases and negative regulation in instances of global \( \text{Ca}^{2+} \) increases (Lee et al., 2003). CaM is also believed to interact with neuronal \( \text{Ca}^{2+} \)-binding protein 1 (CaBP1) in Cav2.1 channels to inhibit neuronal calcium influx (Lee et al., 2002). Additional calcium sensors such as neuronal calcium sensor 1 (NCS-1) are recently hypothesized to alter P/Q-type VDCC presynaptic calcium channel current (Tsujimoto et al., 2002).

![Diagram of α1A VDCC subunit. Spontaneous mutations responsible for the tottering, leaner, rocker and rolling phenotypes indicated.](image)

**Figure 1.2**
Diagram of α1A VDCC subunit. Spontaneous mutations responsible for the tottering, leaner, rocker and rolling phenotypes indicated.
(Adapted from Pietroban 2002)
CEREBELLUM

The cerebellum is a brain region marked with diverse postulations regarding its complicated and intriguing functions. It is linked to roles in motor coordination (Bastian, 2006; Thach et al., 1992), processing in cognition (Booth et al., 2007; Gordon, 2007; Thach, 2007), sensation (Andreasen et al., 1998; Gao et al., 1996), emotion (Schmahmann and Sherman, 1998; Schmahmann, 2004; Schmahmann and Caplan, 2006), and behavioral learning (Cheron, 1990; Miles and Lisberger, 1981; Raymond et al., 1996).

The cerebellum is divided into 10 distinct lobules (I-X), which can be combined or further divided into sublobules depending on the species (Larsell, 1970). Layers of the cerebellum include an inner white matter core composed of axons surrounded by a granule cell layer. The granule cell layer, in addition to numerous granule cells is also populated with Golgi cell somata, Lugaro cell interneurons and unipolar brush cells (Mugnaini et al., 1997). The Purkinje cell layer lies on top of the granule cell layer and is composed primarily of Purkinje cell somata with a sparse scattering of Bergmann glia cell bodies (Voogd and Glickstein, 1998) and candelabrum cells throughout (Laine and Axelrad, 1994). The outermost layer, called the molecular layer, is comprised of Purkinje cell dendrites, climbing fibers from inferior olive neurons and granule cell axons along with a thinly dispersed number of stellate, basket cell interneurons, as well as rows of Bergmann glia fibers (Voogd and Glickstein, 1998).

Sensory nerves innervating the cerebellum include mossy fibers that synapse in the granule cells layer and climbing fibers that synapse in the molecular cell layer.
Mossy fibers originate from many areas, but primarily innervate the cerebellum from their origin of the basilar pontine nuclei, external cuneate nucleus, vestibular nuclei and lateral reticular nuclei (Harvey and Napper, 1991). They also are known to project from all levels of the spinal cord in rodents and along with projections from the brain, end in the cerebellum by entering through the superior, middle and inferior cerebellar peduncles.

Each climbing fiber emanates from a neuronal cell body found in the inferior olive and forms multiple synapses with one Purkinje cell dendritic tree in the cerebellar molecular layer (Eccles et al., 1966; Thach, 1968). In addition, climbing fibers are also able to make multiple synaptic contacts with NG2-immunoreactive glia cells (Lin et al., 2005).

Noradrenergic neurons originating from the locus coeruleus project to the cerebral cortex, cerebellum, brain stem and spinal cord (Abbott and Sotelo, 2000) and this pathway allows one to respond and avoid dangerous situations brought upon by unexpected environmental stimuli. Serotonergic neurons also project to many areas in the brain, including the cerebellar cortex and cerebellar nuclei from their origin in the raphe nuclei (Strahlendorf and Hubbard, 1983). Cholinergic neurons extending from the pedunculopontine tegmental nucleus to the cerebellum, lateral paragigantocellular nucleus and various raphe nuclei to the cerebellum (Jaarsma et al., 1997) and are thought to possibly aid in regulation of sleep and arousal (McCormick, 1989).

The cerebellum can be further divided into interwoven transverse zones in the cerebellar vermis and hemispheres: anterior zone, posterior zone and nodular zone (Ahn
et al., 1994; Hawkes and Eisenman, 1997). These zones are revealed with patterns of zebrin II expression in Purkinje cells (Brochu et al., 1990; Ozol et al., 1999) a transgene expression in Purkinje cells such as L7/pcp2-LacZ transgene 4 and an olfactory marker protein (OMP)-LacZ transgene (Nunzi et al., 1999), and even some types of cerebellar neurodegeneration (Sarna and Hawkes, 2003). Cerebellar long term depression at the cerebellar parallel fiber-Purkinje cell synapse may also be contained within this striping pattern as many proteins involved with glutamatergic neurotransmission display parasagittal stripes in cerebellar Purkinje cells (Ekerot and Kano, 1985; Ito and Kano, 1982; Ito et al., 1982).

**HIPPOCAMPUS AND NEUROGENESIS**

The hippocampus is a well-studied brain region responsible for various aspects of memory storage. The circuitry of the hippocampus includes stellate cells from the entorhinal cortex giving input through the perforant pathway to synapse on granule cells. Mossy fibers, granule cell axons, then synapse onto CA3 pyramidal cells that synapse on CA1 pyramidal cells, ending with their axons leaving the hippocampus proper. It is now understood that neurogenesis occurs in the adult hippocampus (Altman, 1963; Altman and Das, 1965b) and olfactory system (Altman and Das, 1965a; Altman, 1969). As hippocampal neurogenesis is most pertinent to these studies, it will be given greater attention.

The role of hippocampal neurogenesis is still relatively unclear, but various studies using techniques to halt neurogenesis are proving promising in elucidating this
mystery. Hippocampal neurogenesis is thought to possibly play a role in timing or task
difficulty in learning such as trace eye blink conditioning (Bruel-Jungerman et al., 2007;
Shors et al., 2001). Adult hippocampal neurogenesis may also be responsible for long
term spatial memory storage (Snyder et al., 2005) and spatial memory, but no alteration
in contextual fear conditioning (Dupret et al., 2008). It is possible that neurogenesis
improves these tasks because of the activity-dependent plasticity of newly formed
neurons in the DG (Bruel-Jungerman et al., 2007). Hippocampal dependent learning is
believed to increase the amount of survival (Gould et al., 1999; Leuner et al., 2007) and
possibly proliferation (Dobrossy et al., 2003; Lemaire et al., 2000) in adult hippocampal
neurogenesis. In addition, exercise and environmental enrichment may affect
neurogenesis (Brown et al., 2003; Bruel-Jungerman et al., 2007; van Praag et al., 1999).

The path of neurogenesis migration in the hippocampus begins with the
proliferation of the precursor cells, quiescent progenitor cells, in the sub granular zone
(SGZ) of the dentate gyrus. Precursor cells are composed of both horizontal astrocytes
and radial astrocytes, multipotent cells resembling radial glia (Seri et al., 2004). These
cells express various markers such as nestin, precursor cell marker protein and glial
fibrillary acidic protein (GFAP).

Daughter cells of quiescent progenitor cells become the amplifying neural
progenitors (Encinas et al., 2006). These cells express nestin and Sox2, but not GFAP,
can be labeled with BrdU and predominantly cluster along the SGZ (Encinas et al.,
2006). Amplifying neural progenitor cells gives rise to the intermediate precursors, D2
which then mature to D3 cells (Seri et al., 2004) and express a variety of different
markers not found in its parent cells (Encinas and Enikolopov, 2008; Rao and Shetty, 2004; Seki and Arai, 1993; Seri et al., 2004). D3 cells look more like immature granule cells rather than glia and extend processes through the granule cell layer and hilus of the dentate gyrus (Seri et al., 2004).

Once D3 cells are in the dentate gyrus they begin to mature and make neuronal connections with mature granule cells and transiently express doublecortin, calretinin, and neuronal markers NeuN and calbindin (Kempermann, 2005). Within 4-10 days of differentiation, the maturing cells are able to make axonal connections to neurons in CA3 (Markakis and Gage, 1999). These cells continue to send axons called mossy fibers that project to the CA3 region to synapse with pyramidal cells and the polymorphic layer (Patestas, 2006). Polymorphic-layer neurons send fiber projections and are able to modify information by with synapses to the dendrites and cell bodies of granule cells (Patestas, 2006). Pyramidal cells of the CA3 region (Figure 1.3) are able to emit Schaefer collaterals that project within itself, the CA1 region and septal nuclei (Patestas, 2006). CA3 axons are also able to join the band of white matter in the medial hippocamous, the fimbria (Patestas, 2006). At seven weeks, these neurons are fully functional and posses the ability to produce action potentials and complete integration within the hippocampal network (van Praag et al., 1999). There is an excess of newly proliferated cells and a portion of them will die in an apoptotic manner before reaching maturity (Biebl et al., 2000).
Figure 1.3
Representative photo of the mouse hippocampus. The following specific regions are labeled: pyramidal cell layer of CA3 (CA3) and CA1 (CA1), oriens layer (OL), lacunosum molecular layer (LML), hilus (H) and granular cell layer (G) of the dentate gyrus.
CELLULAR ENVIRONMENT

The dynamics related to mitochondrial responses to Ca$^{2+}$ are physiologically important. Mitochondria are primarily involved in the production of ATP by oxidative phosphorylation, but also are linked to cellular functions such as buffering cytosolic [Ca$^{2+}$], apoptosis (Kroemer et al., 1997; Kroemer, 1999), the urea cycle (Lehninger, 1970), beta oxidation of fatty acids (Lehninger, 1970) and metabolic synthesis.

Mitochondria employ methods of efflux and influx as well as mitochondrial permeability transition to control calcium ion transport. Calcium influx in mitochondria is mediated by the mitochondrial calcium uniporter located in the organelles inner membrane (Deluca and Engstrom, 1961; Kirichok et al., 2004). Other methods of Ca$^{2+}$ influx are driven by rapid mode or RaM and the mitochondrial ryanodine receptor (Beutner et al., 2001). Methods regulating Ca$^{2+}$ efflux include the Na$^+$ dependent and Na$^+$ independent mechanisms (Puskin et al., 1976).

While mitochondria have the highest Ca$^{2+}$ capacity of all cellular components, the endoplasmic reticulum and nuclear envelope are also known sources.

Reactive oxygen species (ROS) are byproducts of the mitochondrial respiration chain in the inner mitochondrial membrane that creates ATP. There are multiple forms of ROS such as hydrogen peroxide (H$_2$O$_2$), superoxide anions (O$_2^-$), hypochlorous acid (HOCl), singlet oxygen (O$_2$) and hydroxyl radical (OH). O$_2^-$ is the most prevalent form of ROS that is primarily produced by complex I and II of the electron transport chain. While ROS is believed to play a role in normal cell signaling (Brookes et al., 2002), an overabundance of ROS in a cell can overwhelm antioxidant machinery and prove
deadly. These deadly levels of ROS, oxidative stress, are believed to play a role in various neurodegenerative diseases, cancer and aging (Cross et al., 1987; Halliwell et al., 1992).

There are several antioxidant enzymes in the cell that act to degrade ROS such as superoxide dismutase (SOD), catalase and glutathione peroxidase. As mentioned previously, oxidative stress can arise because of an overabundance of ROS, but can also be a response to a lack of sufficient antioxidants. Mechanisms in the cellular matrix work to detoxify superoxide anions in the cell through reactions of manganese superoxide dismutase (MnSOD) that create the much less deleterious H$_2$O$_2$ (Melov, 2000). The most abundant form of SOD, inter-membrane space copper zinc superoxide dismutase (CuZn-SOD), creates H$_2$O$_2$ and oxygen forms superoxide anions within the intramembrane. In addition to mitochondria, ROS is produced in other cellular compartments such as the nuclear membrane.

**OBJECTIVE OF THIS THESIS**

In light of knowing that the cerebellum is responsible for many of the deficits associated with several mutations at the tottering locus and the strong possibility that the hippocampus also could be affected, it becomes interesting to investigate these regions in the $tg/tg^{la}$ mouse. Interesting differences observed in the cellular environment, and neuronal proliferation in the leaner and tottering mouse are cause for more detailed study of the less known $tg/tg^{la}$ mouse. This thesis seeks to determine if and how the $tg/tg^{la}$ mouse is morphologically and biochemically different from wild type mice with respect
to the cerebellum and hippocampus. In this thesis, cerebellar and hippocampal morphology and cellular biochemistry of neurons in those two brain regions in adult male and female tottering/leaner mice that are 90-120 days of age have been assessed.

**Overall Hypothesis**

*The cerebellum and hippocampus of adult tottering/leaner mice are morphologically and biochemically different from age- and sex-matched wild type mice.*

*This hypothesis has been tested through completion of three experimental aims:*

1) *examination of the morphology of the cerebellum and hippocampus of tottering/leaner mice* 2) *assessment of proliferation and death in the dentate gyrus of tottering/leaner hippocampus* 3) *investigation of changes in basic biochemical parameters in granule cells of the cerebellum and hippocampus in tottering/leaner mice.*

**Experimental Protocols**

I examined both male and female mice. All experiments examined adult mice at ages 90-120 days. Six animals of each genotype and gender were used in each experiment, except when including fewer animals in each group resulted in observing significant differences. For histological analyses both frozen, unfixed brains and fixed brains were used. For volume analyses all mice will be anesthetized with isoflurane (Abbott Laboratories, North Chicago, IL, USA) and decapitated. To obtain fixed tissue, mice were deeply anesthetized with intraperitoneal injections of xylazine (Vedco, St.
Joseph, MO, USA) and ketamine (Vedco, St. Joseph, MO, USA; 0.5mg xylazine plus 3.0 mg ketamine per 20 gm body weight) and then perfused.
INTRODUCTION

Determine Hippocampal and Cerebellar Morphology

Previous work from this lab has revealed decreased forebrain and hindbrain weights in mice that are homozygous for the tottering and leaner mutations as well as the compound heterozygous tottering/leaner mice, but the volume of the cerebellum and hippocampus has not been previously published (Huang et al., 2007). To gain greater understanding of the brain regions potentially responsible for the many neurological abnormalities exhibited by these mice, volumes of both the hippocampus and cerebellum of $tg/tg^{la}$ and age- and gender-matched wild type mice have been measured.

It was hypothesized that cerebellar volume of $tg/tg^{la}$ mice would be significantly less than that of age-matched wild type mice, as it is an area that is known to be affected by both the tottering and leaner mutations. Another hypothesis was that there would be no significant difference in the hippocampal volume of the $tg/tg^{la}$ mouse, as this brain region does not exhibit as many overt alterations as those seen in the cerebella of leaner and tottering mice.

Potential differences in volume of these structures could result from a change in cell density in either region. For these experiments, density is referred to as numbers of cells per unit area. Any potential change in cell density could reveal further deficit in
terms of altered neuronal death or proliferation. An absence of density change could imply synapse loss without significant loss of neuronal cell bodies or processes or increased numbers of glial cells.

In the assessment of cell density, the original hypothesis was that granule cell density in the cerebellum would be decreased in all lobules investigated, while the hippocampus will have fewer cells per unit area in the \( tg/tg^{la} \) mouse as compared to age- and gender-matched wild type mice.

Volume or density differences could also be the result of a change in the size of neurons in various brain regions within the hippocampus. Various diseases such as Alzheimer’s disease (van de Nes et al., 2008) and epilepsy (Andres et al., 2005; Bothwell et al., 2001) show abnormalities in neuronal size in some brain regions of affected patients. The hypothesis regarding neuronal size was that average cell size in specific regions of the hippocampus of the \( tg/tg^{la} \) mouse would be significantly smaller when compared to the same specific regions in age- and gender-matched wild type mice.

EXPERIMENTAL PROCEDURES

Hippocampal and Cerebellar Volume

For volume studies, every fifth, coronal, 25 micrometer thick frozen section of unfixed mouse brain was thaw-mounted onto microscope slides, stained with 0.1% thionin (Thermo Fisher Scientific Inc., Fair Lawn, NJ, USA) in 0.1M acetate buffer, dried and coverslipped using Permount (Thermo Fisher Scientific Inc.). The slides were coded to blind the investigator as to sex and genotype during measurements. A 2X
objective on a Nikon E400 microscope with a Nikon DXM1200 video camera and ACT1 software was used to capture images for measurement. After outlining the images of sections of the cerebellum and hippocampus with Image J software (Rasband, 1997-2008), the area was computed and then multiplied by section thickness and number of sections skipped to determine total volume.

Cell Density in the Hippocampus and Cerebellum

Cellular density in six different regions of the hippocampus was examined: pyramidal cell layer of CA1 and CA3, oriens layer, lacunosum molecular layer, granule cell layer of the dentate gyrus, and hilus of the dentate gyrus. In the cerebellar vermis, rostral lobe II and caudal lobes VI and X were examined for density of cells in the granule layer, Purkinje cell layer and molecular layer.

Coronal, 5 µm thick paraffin sections of each region were stained with hematoxylin (Sigma Aldrich, Milwaukee, WI, USA) and eosin (Sigma Aldrich, Milwaukee, WI, USA), coverslipped and coded to blind the investigator as to sex and genotype during scoring. Two sections were counted for each brain region for each animal. A 40X objective on a Nikon E400 microscope with a Nikon DXM1200 video camera and ACT1 software was used to capture images for counting. A uniform area of 100 x 50 µm was randomly selected twice for each region to count numbers of cells in each area. Only cells completely within the specified area that contained an obvious nucleus were counted.
Hippocampal Cell Size

Coronal sections through the rostral hippocampus from sections that were stained with Thionin were used to assess cell density were used to assess neuronal area. In regions where significant differences in density were observed, 10 cells were randomly chosen and images captured with a 40X objective on a Nikon E400 microscope with a DXM1200 video camera and ACT1 software. After outlining individual neurons with Image J software, the areas were measured. To ensure that neuronal area was measured in similar planes through each cell, each measured cell was required to fulfill the criteria of having a nucleus present, a nucleolus visible within the nucleus and a distinct plasma membrane.

Data were analyzed using SPSS Version 12.0.1 Multivariate Analysis of Variance (ANOVA) at $\alpha = 0.05$ to test gender differences. A Univariate Analysis of Variance (ANOVA) at $\alpha = 0.05$ two way ANOVA was used if the Multivariate ANOVA showed that genders could be combined. Tukey’s honest significant difference (HSD) post hoc test was used when significant differences within region and genotypes were assessed.

RESULTS

Hippocampal and Cerebellar Volume

Hippocampal volume showed no significant difference between gender or genotype ($P = 0.841$). This implies that although the forebrain weight in the mutant mouse is significantly less than that of the wild type mouse, there is no apparent volume
variation that is observed in the hippocampus. It is possible that other affected forebrain regions could account for the observed forebrain weight differences.

There was a significant difference in cerebellar volume (Figure 2.1) in the mutant mice compared to age-matched wild type mice (P = 0.021). It is logical to believe that the decreased cerebellar volume in the tottering/leaner mouse could be partially responsible for their decrease in hindbrain weight. As the cerebellum is an area known to be severely affected in mice with similar mutations and noting the gravity of the movement disorders of the tottering/leaner mouse, it is no surprise that the cerebellum exhibits such abnormal architecture.

**Cellular Density in the Hippocampus and Cerebellum**

Even though the hippocampus showed no change in hippocampal volume when the two genotypes were compared, there was a variation in cellular density within some regions of the tottering/leaner hippocampus. The pyramidal cell layer of CA3 (P = 0.001) and the hilus of the dentate gyrus showed a significant decrease in cellular density (P = 0.009). It is possible that there are increases in cellular processes or glial cells that might compensate for the decrease in cellular density loss that allows for no alteration in hippocampal volume.
Figure 2.1: Quantitative comparison of volume within brain regions. Volume of tottering/leaner and wild type mice at 90-120 days of age is determined by measuring area of region in every fifth section, multiplied by section thickness and sections skipped. Comparison of hippocampal volume (A) and cerebellar volume (B). A) No significant difference between gender or genotypes observed. B) Significant difference between genotypes denoted by single asterisk (*), where tottering leaner cerebellar volume is significantly less than wild type cerebellar volume (P = 0.021).
One explanation for the decrease in cellular density (Figure 2.2) of the CA3 region in the \( \text{tg/tg}^{la} \) mouse may be explained by the defects the homozygous tottering mouse shows in this region. Epileptic seizures exhibited by the tottering mouse are postulated to be correlated with excitability defects in the CA3 region of the hippocampus (Helekar and Noebels, 1991). Closer examination reveal prolonged paroxysmal depolarizing shifts in hippocampal CA3 pyramidal neurons after exposure to high potassium levels in vitro during the developmental stage of the tottering mouse (Helekar and Noebels, 1992). As this abnormality appears most prominently in young tottering mice, it appears that this is possibly a genetically signaled neuronal synchronization mutation rather than an effect of a life of seizures (Helekar and Noebels, 1992).

The \( \text{tg/tg}^{la} \) mouse similarly exhibits seizures like the homozygous tottering mouse and could perhaps also have this abnormal CA3 excitability defect. There could be less neurons in the \( \text{tg/tg}^{la} \) mouse hippocampal CA3 region because the abnormal neuronal synchronization may cause them to die. It could also be hypothesized that the decrease in cell density in this region is partly responsible for any abnormal excitability in this region with some underlying cause of neuronal loss waiting to be elucidated.
Figure 2.2: Cellular density in the hippocampus. Cellular density of 90-120 day old tottering/leaner and wild type mice defined as cells per unit area. Six regions of the hippocampus are examined: pyramidal cell layer of CA1 (A) and CA3 (B), granule cell layer of the dentate gyrus (C), and hilus of the dentate gyrus (D), lacunosum molecular layer (E) and oriens layer (F). A), C), E), F) No significant difference between genotypes observed. B) Single asterisk (*) indicates significantly decreased cellular density significantly tottering/leaner mice as compared to wild type (P = .001). D) Double asterisk (**) indicates significantly decreased cellular density in tottering/leaner mice as compared to wild type (P = .009).
C

Granule Cell Layer of the Dentate Gyrus

![Bar graph showing the number of cells per unit area in the Granule Cell Layer of the Dentate Gyrus between Tottering/Leaner and Wild type.]

D

Hilus of the Dentate Gyrus

![Bar graph showing the number of cells per unit area in the Hilus of the Dentate Gyrus between Tottering/Leaner and Wild type.]

Figure 2.2 continued
E  

**Lacunosum Molecular Layer**

- **Tottering/Leaner**
- **Wild type**

F  

**Oriens Layer**

Figure 2.2 continued
In the investigation of cerebellar density (Figure 2.3), there was an observed trend towards decreased density in the anterior and posterior lobes combined in the cerebellum of the $tg/tg^{la}$ mouse in comparison to the wild type mouse ($P = 0.053$). On closer inspection, this discrepancy was found to be a result of a significant decrease specific to the anterior lobe of the tottering/leaner mouse as compared to the posterior lobe of the wild type mouse ($P = 0.016$). There was also a significant decrease in the $tg/tg^{la}$ mouse cellular density in the granule cell layer anterior and posterior lobes combined as compared to the wild type mouse ($P = 0.036$). It appears that there is enough of a subtle difference, but not a significant difference, in that the anterior lobe is slightly decreased in comparison to the posterior lobe in the $tg/tg^{la}$ mouse. The subtle increase in the posterior lobe of the wild type mouse and slight decrease in the anterior lobe of the $tg/tg^{la}$ mouse presumably leads to an overall significance between the wild type mouse posterior lobe and $tg/tg^{la}$ mouse anterior lobe. It is likely that this decrease in cellular density in the $tg/tg^{la}$ mouse comes from the cerebellar granule cell layer.

There was no significant difference within genotypes when comparing lobule 10 or the layers of lobule 10. In addition, there were no other significant differences in the anterior or posterior lobules other than that previously stated. The decrease in cerebellar volume, but no decrease in cerebellar density leads one to believe that there possibly is a decrease in cellular processes.
Figure 2.3: Cellular density in the cerebellum. Graphs depict anterior (lobule 2) and posterior (lobule 6) lobes combined (A) of 90-120 day old $tg/tg^{la}$ and wild type mice and the examination distinct layers in the combined anterior and posterior lobes combined: molecular layer (B), granule layer (C) and Purkinje cell layer (D). Further analyses includes granule cell layer of anterior lobe compared to posterior lobe (E). Lobule 10 is described separately with layers combined (F) and in distinct layers: molecular layer (G), Purkinje cell layer (H) and granule cell layer (I). B), D), F) -I) No significant difference between genotypes observed. B) Number sign (#) indicates no significant difference, but a trend (P=0.053) where tottering/leaner is cellular density is decreased in anterior and posterior lobes combined as compared to wild type anterior and posterior lobes combined. C) Single asterisk (*) indicates significant decrease in $tg/tg^{la}$ cellular density in anterior and posterior lobes combined in granule cell layer as compared to wild type cellular density in corresponding regions (P=0.036) E) Ampersand symbol (&) indicates significant decrease in tottering/leaner anterior lobe as compared to wild type posterior lobe in the granule cell layer (P = 0.016).
Figure 2.3 continued

B

Molecular Layer
Anterior and Posterior Lobes Combined

Cells per unit area

C

Granule Cell Layer
Anterior and Posterior Lobes Combined

Cells per unit area

Tottering/Leaner
Wild type

*
Figure 2.3 continued
Figure 2.3 continued
Figure 2.3 continued
Hippocampal Cell Size

As there was a significant difference in cellular density in the hippocampus in the hilus of the dentate gyrus and in the pyramidal cell layer of CA3, it was necessary to investigate the neuronal size as indicated by area within these two regions. Results showed that there was no difference in the pyramidal cell layer of CA3 between $tg/tg^{la}$ mice and wild type mice (Figure 2.4). There was a significant difference within the hilus of the dentate gyrus with the female wild type cell size being larger than the cell size of all other genders and genotypes ($P = .000$ to $0.001$). This shows that the $tg/tg^{la}$ mouse fails to exhibit the normal gender difference in cell size within hilus of the dentate gyrus when the wild type mouse displays this difference. Alterations in $tg/tg^{la}$ hormonal expression may lie behind this discrepancy.
Figure 2.4: Cell size in hippocampal areas. This includes the pyramidal cell layer of CA3 (A) and the hilus of the dentate gyrus (B) in tottering/leaner and wild type mice at age 90-120 days. 

**A)** No significant difference observed within genotypes ($P = 0.841$). 

**B)** Single asterisk (*) indicates significant increase in female wild type cell size as compared to all other genders and genotypes ($P = 0.000$ to $0.001$).
DISCUSSION

Results revealed no decrease in hippocampal volume in \textit{tg/tg}^{la} mice while cellular density in the CA3 and hilus of the dentate gyrus was decreased and hippocampal cell size remained normal. This indicates that there could be a compensation of cellular processes in the CA3 and hilus of the dentate gyrus to account for the cellular density decrease in these regions as increased hippocampal cell size is ruled out as a possible mechanism.

The \textit{tg/tg}^{la} cerebellum displayed a significant decrease in cerebellar volume, which confirms previous results from our lab that further describe a 24% decrease in the amount of cerebellar molecular layer per Purkinje cell layer without Purkinje cell loss as compared to the wild type mouse (Isaacs and Abbott, 1995). In line with previous assumptions, it is likely that the cerebellar volume loss in our study could be due to one or a combination of the following: glial cell loss, loss or shrinkage of parallel fibers, or a decrease in Purkinje cell dendritic arborizations in the cerebellar molecular layer (Isaacs and Abbott, 1995).

Alternatively, it could be suggested that the slight, but not significant decrease in the \textit{tg/tg}^{la} anterior lobe as compared to the posterior lobe and the slight but not significant increase in the wild type posterior lobe as compared to the anterior lobe are suggestive of an actual biological difference. The homozygous leaner (\textit{tg}^{la}/\textit{tg}^{la}) mouse is known to exhibit a greater decrease in cerebellar granule cells in the anterior lobe when compared to the posterior lobe (Frank et al., 2003; Lau et al., 2004) by way of apoptosis (Bawa and Abbott, 2008), which leads to the possibility that the \textit{tg/tg}^{la} mouse is
following a similar trend that is not as pronounced as that observed for the \( tg^{la}/tg^{la} \) mouse. It would be necessary to measure the volume of the granule cell layer in the \( tg/tg^{la} \) mouse and stain markers of cellular death such as fluoro-jade to investigate this aspect further.
CHAPTER III
PROLIFERATION AND DEATH IN THE DENTATE GYRUS OF TOTTERING/LEANER HIPPOCAMPUS

INTRODUCTION

It was necessary to determine whether changes in the number of neurons observed within the hippocampus as overall neuronal loss could result from increased cell death, decreased proliferation or interactions between these two processes.

Neuronal death has been examined in the subgranular zone of the dentate gyrus by means of Fluoro-Jade staining. Fluoro-Jade is an anionic tribasic fluorescein known to stain cell bodies, axons and axon terminals of degenerating neurons without staining healthy neurons, myelin, vascular elements or neuropil (Schmued et al., 1997). Fluoro-jade has proven successful for staining in a genetic model of cell death in the homozygous leaner mouse cerebellum (Frank et al., 2003). It is a fast and reliable method that does not have a clearly understood, but it is thought to function histologically based upon its chemical properties of high acidity that may attract a degenerating neurons strongly basic molecule (Schmued et al., 1997). As Fluoro-Jade has an emission peak of 550 nm and excitation peaks at 362 and 390 nm, appropriate fluorescence microscope visualization can be obtained with a FITC filter system.

The original hypothesis regarding this experiment was that the number of dying cells stained by Fluoro-jade in the $tg/tg^{la}$ subgranular zone (SGZ) of the hippocampal dentate gyrus (DG) would be significantly higher than the number of Fluoro-jade positive cells observed in age- and gender-matched wild type mice.
In addition to characterizing cell death, cell proliferation and differentiation in the SGZ of the hippocampal DG must be examined using the immunohistochemical analyses. As early as 1965, postnatal neurogenesis in the SGZ of the DG in the hippocampus was characterized in rats using thymidine-\textsuperscript{3}H to label S-phase cells (Altman and Das, 1965b). As research progressed, the immunohistochemical method of detecting of 5-bromo-2'-deoxyuridine (BrdU) incorporated into nuclear DNA was shown to appropriately label the nuclei of dividing cells that are in S-phase (Gratzner, 1982; Miller and Nowakowski, 1988; Miller and Nowakowski, 1991; Nowakowski et al., 1989). Leaner and tottering mice have been shown to exhibit decreased cell proliferation in the SGZ of the DG (Wills, 2006).

Proliferating cell nuclear antigen (PCNA) is an endogenous cell cycle protein that is localized within the cell nucleolus during the S phase of cell replication (Bravo and Celis, 1980; Bravo and Macdonald-Bravo, 1987; Takasaki et al., 1981). Using antibody to PCNA in conjunction with BrdU immunohistochemistry is a powerful tool to aid in cellular birth dating and cell cycle kinetics (Mandyam et al., 2007).

Prior to experimentation, it was hypothesized that there would be a decrease in cell proliferation in the dentate gyrus of \textit{tg/tg} \textit{la} mice as compared to age- and gender-matched wild type mice as revealed using BrdU immunohistochemical staining. PCNA immunohistochemical staining would mirror the results of BrdU immunohistochemical staining and show a decrease in the amount of cell proliferation in the dentate gyrus of the \textit{tg/tg} \textit{la} mice as compared to wild type mice.
While BrdU immunohistochemistry will show cell proliferation, it is not able to give information regarding differentiation of newly formed cells. It was necessary to determine the fate of these proliferating cells to have a more complete understanding of adult neurogenesis in \( tg/tg^{la} \) mice. To identify mature neurons, primary antibody to Neuronal nuclear protein (NeuN) was used to double label the sections along with BrdU. NeuN is neuronal cell-type specific protein that is expressed and will label neurons as early as one week after cell division and differentiation and peaks at four weeks (Mullen et al., 1992).

In addition to neurons, the proliferating cells are also capable of differentiating into astrocytes. Glial fibrillary acidic protein (GFAP) was used to characterize mature cells of astrocytic lineage (Bignami et al., 1972; Bignami and Dahl, 1973). Specifically, mature radial astrocytes and horizontal astrocytes are the two cell types that can be immunostained in the subgranular zone of the dentate gyrus (Seri et al., 2004).

Unpublished data from the Abbott lab has shown no significant difference in percent of cell differentiation of proliferating neurons as mature astrocytes or neurons between leaner, tottering and wild type mice. These previous findings lead to the hypothesis that the \( tg/tg^{la} \) mouse would reveal no significant difference in the percent of cell differentiation of proliferating neurons as mature neurons or astrocytes as compared to age- and gender-matched wild type mice.
EXPERIMENTAL PROCEDURES

Neuronal Death with Fluoro-Jade Stain

Fixed frozen 25µm thick coronal sections of the hippocampal DG were cut on a cryostat and thaw mounted onto gelatin-coated slides. Desired slides of the hippocampal DG were selected and taken through Fluoro-Jade staining according to a standard protocol (Schmued et al., 1997).

According to procedure, slides were thawed and dried for 3 hours in a 50°C incubator to assist adherence of sections to slides. An additional section of fixed frozen mouse testes were taken through this procedure to serve as a positive control. After cooling to room temperature, slides were taken through a series of alcohol immersions; 3 minutes in 100% ethyl alcohol, 1 minute in 70% ethyl alcohol and 1 minute in distilled water.

To help minimize background stain, slides were then placed in a 0.06% potassium permanganate solution (Thermo Fisher Scientific Inc., Fair Lawn, NJ, USA) for 15 minutes, while being gently agitated on a rotating platform. Two rinses in distilled water for 1 minute each is then followed by the immersion of slides into 0.001% fluoro-jade staining solution (Chemicon International, Temecula, CA, USA) while being gently agitated on a rotating platform for 30 minutes. One slide of mouse DG did not enter the fluorojade stain to serve as a negative control. Lastly, slides are rinsed in distilled water three times for 1 minute each, excess water is drained from slides and they are rapidly dried with a hot air gun. After slides are completely dry, they are
immersed in xylene for about 5 minutes, coverslipped with permount and allowed to dry for 24 hours prior to imaging.

Images of the fluoro jade stained SGZ of the DG will be captured using a FITC fluorescent filter on a Zeiss Axioplan Microscope with Zeiss Axiocam. Once coded to eliminate investigator bias, images will be scored according to their bright green fluorescence. Only positive fluorescing cells along the SGZ of the DG and those of appropriate cell size and shape will be counted to ensure proper scoring.

Degree of Cellular Proliferation Using BrdU Labeling and PCNA

Immunohistochemistry

The first experiment examined neuronal proliferation in the SGZ using BrdU immunohistochemical detection. Mice were dosed with 100mg/mL BrdU (Sigma Aldrich, Milwaukee, WI, USA) given in sterile lactated Ringer’s solution in volumes of 0.01mL/g body weight. Injections occurred twice daily with 8-12 hours between each injection for three days and one injection on the final fourth day, for a total of seven injections. One hour after the final injection, mice were perfused with Tyrode’s saline (50mL; pH 7.2-7.4) and 4% phosphate buffered paraformaldehyde (J.T Baker Inc., Phillipsburg, NJ, USA). Brains were stored in 4% phosphate buffered paraformaldehyde for 12-24 hours followed by cryoprotection in 20% sucrose solution (EMD Chemicals Inc., Gibbstown, NJ, USA) in PBS after which they were rapidly frozen using powdered dry ice and stored at -70°C until used.
A cryostat coronally cut brains at 15µm throughout the desired brain regions and the sections were thaw-mounted onto gelatin coated microscope slides. Cut sections were stored at -70°C until they use. Sections containing the rostral DG were stained with a standard BrdU immunostaining protocol (Kempermann et al., 1997).

Slides were initially selected for desired sections of the DG and allowed to thaw and dry at room temperature for 15 minutes. After drying, slides were placed in HCl for 1 hour in a 40°C incubator to open up the cell membrane. Three rinses of sodium borate (Fisher Scientific Co., Fair Lawn, NJ, USA) for 5 minutes neutralized the acid. Slides were then rinsed three times for 10 minutes each in NIH-PBS and then immersed in a detergent, Triton-X (Sigma Aldrich, St. Louis, MO, USA) for one hour. Another three rinses for 10 minutes each in NIH-PBS followed before slides were placed in 5% normal goat serum (Invitrogen, Eugene, OR, USA) for 1 hour at room temperature. Next, slides were immersed in a primary antibody of 1:1000 anti-BrdU (Sigma, St. Louis, MO, USA) in 2% normal goat serum overnight in the refrigerator. One slide was not placed in the primary antibody and used as a negative control, instead, it set in 2% normal goat serum overnight in the refrigerator.

The following day, slides were removed from the primary antibody and rinsed three times in NIH-PBS for 10 minutes each. Slides were then are kept in a dark environment and placed in a solution of 1:1,000 Alexa-fluor 594 (Invitrogen, Eugene, OR, USA), a red fluorescence dye that binds to the anti-BrdU, in 1% bovine serum albumin (EMD Chemicals Inc., Gibbstown, NJ, USA) for six hours. Following this step, slides were rinsed three times for 10 minutes each in NIH-PBS. Moisture was carefully
wicked away from the slides with a kimwipe, avoiding any brain tissue, and then coverslipped in slo-fade gold (Invitrogen, Eugene, OR, USA). Coverslipped slides set for 24 hours and then were sealed along the edges with nail polish to prevent any dehydration. A period of at least 30 minutes took place before slides were imaged.

A Zeiss Axioplan Microscope with Zeiss Axiocam was used to capture fluorescence images of at least four sections of the DG with a gap of at least 75µm between sections so not to count the same nuclei more than once. The investigator counted the number of fluorescent nuclei along the SGZ of the coded images. Size and shape of fluorescent nuclei was examined to ensure that no background or other neuronal processes are counted.

Paraffin sections were used for PCNA staining. Immersion fixed brains in 4% phosphate buffered paraformaldehyde were paraffin embedded and cut coronally in 5 µm thick sections. Slides were heated for 30 minutes in a 40°C oven to ensure adhesion of the sections to the microscope slides. After cooling, the sections underwent deparaffinization in xylene and decreasing concentrations of alcohol. Slides then entered a basic immunohistochemistry procedure with peroxidase quenching (Mandyam et al., 2007). PCNA anti-mouse (Sigma, St. Louis, MO, USA) at a dilution of 1:10,000 in PBS served as the primary antibody. Diaminobenzidine (DAB; Curtin Matheson Scientific, Houston, TX, USA) stain labeled PCNA reactive cells for counting. The investigator counted the number of cells with DAB stained nuclei in the hilus, granular layer and along the SGZ of the DG. At least four sections each separated by at least 25 µm were counted from each animal. Sections of PCNA stained mouse intestines were
used as a positive control, while sections of hippocampus stained without primary antibody were used as a negative control. Positive cell counts per area were averaged for each gender and genotype.

**Fate of Surviving New Cells Using NeuN and GFAP Immunohistochemistry**

After injecting animals with BrdU as described previously, a period of four weeks passed before euthanasia. Sections were cut from paraformaldehyde fixed, cryoprotected brains sectioned at a thickness of 15 µm on a cryostat and thaw-mounted onto gelatin coated microscope slides. The protocol followed in the same manner as previously described for BrdU immunohistochemistry with the exception of adding a second label of immunostaining of NeuN (Chemicon International, Temecula, CA, USA) at concentrations of (1:1000) in bovine serum albumin and fluorescent marker Alexa-fluor 488 (Invitrogen, Eugene, OR, USA). fluorescent nuclei labeled with BrdU were imaged as described previously, but those nuclei stained with Alexa-fluor 488 were captured with a FITC filter. Nuclei were counted in same manner as those stained with BrdU alone, but additional NeuN positive cells were counted in the event of evidence of staining overlap with BrdU labeled nuclei. This ensured that proliferating cells of neuronal lineage were counted.

Using an additional set of slides from the rostral DG, the immunohistochemical staining protocol followed in the same manner as in the BrdU / NeuN double labeling with the exception of the second label that will allow for immunostaining for GFAP (Sigma, St. Louis, MO, USA) at concentrations of (1:2000) and fluorescent marker
Alexa-fluor 488 (Invitrogen, Eugene, OR, USA). Fluorescent cells labeled with BrdU were imaged as described previously, but those cells stained with Alexa-fluor 488 were captured with a FITC filter. Sections of mouse intestine were stained as a positive control, while hippocampal sections were stained without a primary antibody for negative controls. Cells were counted in same manner as those stained with BrdU alone, but additional GFAP-positive cells were counted if there was evident staining overlap with BrdU labeled cells. This ensured only proliferating cells of astrocyte and radial glial cell lineage were counted.

All data were analyzed using SPSS Version 12.0.1 Multivariate Analysis of Variance (ANOVA) at α = 0.05 to test gender differences. A Univariate Analysis of Variance (ANOVA) at α = 0.05 two way ANOVA was used if the Multivariate ANOVA showed that genders could be combined. Tukey’s honest significant difference (HSD) post hoc test was used when significant differences within region and genotypes were assessed.

RESULTS

Neuronal Death with Fluoro-Jade Stain

Fluoro-Jade staining with ANOVA analysis (Figure 3.1) revealed no significant gender difference, but a significant decrease in cell death within the tottering/leaner mouse hippocampus subgranular zone as compared to the wild type mouse (P = 0.039). Positive and negative controls both revealed that this procedure appropriately labels dying cells. An unexpected neuroprotective effect is revealed from this mutant mouse.
While some hippocampal regions experienced decreased density in the tottering/leaner mouse, the subgranular zone is not one believed to have an increase in neuronal loss. As cells in this area are continuing to proliferate, these results give reasonable belief that in the 3 month old tottering/leaner mouse a greater amount of newly formed cells remain viable than in the wild type animal.

Despite the results that there was a decrease in neuronal death in this area, it is yet to be determined whether the cells that are living are fully functional and integrated into the circuitry. It is possible those in the wild type subgranular zone a number of cells die for the benefit of this brain region and that the decrease in neuronal death in the mutant mouse proves unbeneeficial and potentially harmful to hippocampal performance. Death signals may go unnoticed and not brought to completion in the cells of the subgranular zone of the DG in this unique mouse.
Figure 3.1 Representative image of neuronal death in the subgranular zone of the dentate gyrus as revealed by Fluoro-Jade stain. Positive control testes (A), negative control (B) and stained wild type male dentate gyrus (C). Arrows indicate positive cells. Graph depicts amount of fluoro-jade positive cells as averaged per section (D) A single asterisk (*) indicates no significant difference in gender, but a significant decrease in neuronal death in the tottering/leaner mouse as compared to the wild type mouse (P = 0.039).
Degree of Cellular Proliferation Using BrdU Labeling and PCNA

Immunohistochemistry

BrdU labeling (Figure 3.2) revealed no significant difference between gender or genotypes regarding neuronal proliferation in the subgranular zone of the dentate gyrus (P = 0.166), conflicting with my previous hypothesis. This revealed that the decrease in cellular density in some regions of the hippocampus was not due to any changes in neuronal proliferation. It also helps to describe the interaction of proliferation and cell death in that the decrease in cell death is not a result of less new cells being formed.

PCNA further confirmed BrdU labeling results and revealed no significant difference in cell proliferation between genotypes (Figure 3.3; Figure 3.4). There was a slight trend of increased $tg/tg^a$ PCNA labeling in the hilus of the dentate gyrus and the subgranular zone (P = .081 and .087 respectively).
Figure 3.2 Image and graph of BrdU immunohistochemistry. Representative of BrdU labeled wild type 3-month-old dentate gyrus (A). Arrows indicate BrdU positive cells. Cellular proliferation in the subgranular zone of the hippocampal dentate gyrus in 3 month old animals as labeled by BrdU (B). BrdU labeled cells were averaged within each animal examined as number of positive cells per section. B) No significant difference in neuronal proliferation between gender or genotype.
Figure 3.3 Representative images of PCNA immunohistochemistry. Negative control with non-specific staining (A), Positive intestine control (B) and PCNA labeled wild type hippocampus (C). Arrows indicate PCNA positive cells.
Figure 3.4
Graphs of comparison of 3-month-old \(tg/tg^{la}\) and wild type PCNA labeled regions. Granular cell layer of the dentate gyrus (A), hilus of the dentate gyrus (B), and sub granular zone of the dentate gyrus (C). PCNA labeled cells were averaged within each animal examined as number of positive cells per section. (A-C) No significant difference between genotypes. (B) Trend of increased \(tg/tg^{la}\) PCNA positive cells with \(P = .081\). (C) Trend of increased \(tg/tg^{la}\) PCNA positive cells with \(P = 0.087\).
Figure 3.4 continued
Fate of Surviving New Cells Using NeuN and GFAP Immunohistochemistry

While basic cellular proliferation and death allow for a greater understanding of the process of new cell formation, the percent of cells in the subgranular zone of the dentate gyrus double labeled with BrdU and GFAP showed how many of the newly formed cells differentiated into astrocytes. The percentage of cells double labeled with BrdU and GFAP showed no significant difference between gender or genotype (P = 0.706). In line with the number of single labeled BrdU positive cells per section, double labeled cells showed no significant difference when gender or genotype were compared, giving more confidence to the results. The result of this experiment revealed that newly proliferated cells showed no abnormal differentiation to astrocytes (Figure 3.6).

When comparing gender and genotype (P = 0.772) of percent of BrdU/NeuN double labeled cells, no significant difference was found (Figure 3.5). Number of BrdU positive cells per section in BrdU/NeuN double labeled immunohistochemistry showed no significant difference in the same manner as single labeled BrdU results. While no significant difference was shown in percent of newly proliferating cells differentiating into neurons or astrocytes, these results remain in the expected range of normal cellular differentiation observed by our lab in this region (Wills, 2006).
Figure 3.5 Images of NeuN/BrdU immunohistochemistry. Representative images of BrdU labeled (A), NeuN labeled (B) and BrdU/NeuN double labeled (C) 3 month old tg/tgla subgranular zone of the dentate gyrus. Arrows indicate positive cells.
Figure 3.6  Images of GFAP/BrdU immunohistochemistry. Representative images of GFAP labeled (A), BrdU labeled (B) and BrdU/GFAP double labeled (C) 3 month old $tg/tg^{la}$ subgranular zone of the dentate gyrus. Arrows indicate positive cells.
Figure 3.7  Graphs of percent cells labeled GFAP/BrdU and NeuN/GFAP.  Graph comparing percent of 3-month-old $tg/tg$ and wild type positive GFAP/BrdU (A) and NeuN/BrdU (B) double labeled cells per section of subgranular zone of the dentate gyrus.  (A,B) No significant difference between genotypes.
DISCUSSION

The Fluoro-jade, BrdU and PCNA results indicated that the decrease in cell death in the $tg/tg^{la}$ mouse was not due to an increase in cellular proliferation (Figure 3.7). In addition, newly proliferating cells differentiate into neurons and glia in the same percentages as wild type animals. Although the proliferating cells have differentiated into normal lineages, it is unknown whether they are fully functional or integrated into their environment. The $tg/tg^{la}$ mutation appears to have deleterious effect to the brain region as cells that normally would die in a wild type animal are not allowed to enter normal cell death. It is possible that the cells that are not dying as intended, and are not properly functioning which could be adding to this dysfunction.

Unpublished research from our lab reports an increase in neuronal proliferation in the leaner mutant mouse at P42-50, but then a decrease in neuronal proliferation as compared to wild type mice at P100-150 (Wills, 2006). While the $tg/tg^{la}$ mouse shows no variation at P90-120, it is possible that it may follow proliferation fluctuations as the leaner mouse displays. It cannot be ruled out that there are changes in proliferation in the dentate gyrus in $tg/tg^{la}$ juvenile or adult mice aged above the age range this research investigated. As the leaner mutation is more phenotypically severe than the $tg/tg^{la}$ mouse (Green and Sidman, 1962; Herrup and Wilczynski, 1982), it could be that neurogenesis is in fact never altered in these mice.
CHAPTER IV

CHANGES IN BASIC BIOCHEMICAL PARAMETERS IN GRANULE CELLS
OF THE CEREBELLUM AND HIPPOCAMPUS IN TOTTRING/LEANER MICE

INTRODUCTION

Changes in several intracellular biochemical parameters have been observed in
granule cells of homozygous leaner and tottering mice (Bawa and Abbott, 2008).
Previously it was unknown whether any alterations in cellular physiology exist in
granule cells from tg/tg<sup>la</sup> mice.

As the characteristic mutation of these mice results in malformations of the
Cav2.1 calcium channel, it is likely that calcium homeostasis may be altered. It has been
shown that basal free intracellular calcium is reduced by 20% in cerebellar granule cells
from the tottering and leaner mouse as compared to age- and gender-matched wild type
mice (Bawa and Abbott, 2008).

It was previously hypothesized that basal intracellular calcium levels would be
reduced in cerebellar and hippocampal granule cells in the tg/tg<sup>la</sup> mouse as compared to
age- and gender-matched wild type mice.

Calcium homeostasis along with cellular metabolism and bioenergetics are held
in a delicate balance with mitochondrial activity being a critical component. Ca<sup>2+</sup>
accumulation by mitochondria can affect ATP synthesis along with alterations in ATP
synthesis effecting the removal of Ca<sup>2+</sup> from the cytoplasm by ion pumps. Alterations in
mitochondrial membrane potential, an indicator of mitochondrial activity, may
ultimately result in a cascade of cellular events ending in cell death (Nicholls and Ward, 2000). Previous research revealed a reduction in the mitochondrial membrane potential in leaner mouse cerebellar granule cells as compared to wild type cerebellar granule cells, but no significant difference was observed between cerebellar granule cells from tottering and wild type mice (Bawa and Abbott, 2008).

As the \( tg/tg^{la} \) mouse appears to more closely resemble the tottering mouse rather than the leaner mouse with respect to several morphological and biochemical features, the initial hypothesis was based upon the plausibility that there will be no difference in mitochondrial membrane potential, and therefore, mitochondrial activity, in this mutant mouse as compared to the wild type mouse.

With the goal of further characterizing these mutations’ possible effects on cell functioning and viability, examination of several aspects of reactive oxygen species (ROS) production in the cell took place. Reactive oxygen species refers to partially reduced metabolites of oxygen that are more reactive than molecular oxygen and include: superoxide anions, hydrogen peroxide, and hydroxyl radicals (Thannickal and Fanburg, 2000). ROS can result from a variety of processes, and are often found as an abundant byproduct in the mitochondrial respiratory chain. Fluctuations in ROS can serve the cell positively be aiding in intracellular signaling and regulatory functions, but an excess of ROS can cause serious damage to DNA, lipids and protein, possibly leading to cell death (Droge, 2002; Stadtman and Levine, 2000).

This experiment assessed the activity of ROS by using 5-(and-6)-chloromethyl-2',7'-dichlorodihydrofluorescein diacetate, acetyl ester (CM-H\(_2\)DCFDA). This dye
fluoresces in response to removal of acetate groups by intracellular esterases and oxidation within the cell. It is specific for the presence of H$_2$O$_2$, but can also identify H$_2$O$_2$ as a product of superoxide. Research data on generation of reactive oxygen species show no difference between tottering, leaner or wild type mice (Bawa and Abbott, 2008).

In regards to reactive oxygen species, it was hypothesized that the tg/tg$^{la}$ mouse would show no difference in the amount or rate of change in ROS production compared to age- and gender-matched wild type mice.

**EXPERIMENTAL PROCEDURES**

**Levels of Basal Intracellular Calcium Ion Homeostasis**

Calcium homeostasis was assessed by measuring basal levels of free intracellular calcium [Ca$^{2+}$] in granule cells from the cerebellum and hippocampus of 90-120 day old tg/tg$^{la}$ and wild type animals using calcium green-1 fluorescent dye. One tg/tg$^{la}$ and one wild type animal of the same gender were assessed for each session of data collection. Each mouse was anesthetized with isoflurane and decapitated, followed by immediate dissection of the cerebellum and hippocampus. Tissue from each brain region was kept separate, sliced into small pieces and placed in Minimum Essential Medium Eagle (MEM; Sigma Aldrich, St. Louis, MO, USA). A series of proteolytic and mechanical disruptions took place, ending with a final centrifugation. The resulting pellet consisted of whole cells (primarily granule cells) from a specific brain region of origin and were reconstituted in MEM and plated on gelatin and poly-d-lysine (Sigma,
St. Louis, MO, USA) coated coverslips to ensure cells adhered securely. Coverslips were incubated in MEM media for 30 minutes in at 37°C, 95% CO₂ to allow the cells to adhere more tightly to the coverslip and to keep the cells viable. After incubation, media was removed and calcium green 1-AM (CaG-1; Invitrogen, Eugene, OR, USA), a dye that fluoresces intensely when bound to calcium, added at a 1µM concentration in Hanks Balanced Salt Solution (HBSS; Sigma Aldrich, St. Louis, MO, USA) for 20 minutes. CG was then removed and replaced with HBSS for imaging.

Ten images per coverslip were captured at 40X on Zeiss Axiovert 200M microscope using SlideBook 4.1.0 software from 3I, FITC Excitation: BP 470/20, dichroic FT 493, emission BP 505-530nm; or Cy3 Excitation: BP 560/40, dichroic FT 585, emission: BP 630/75 nm. Simple PCI and Imaging System Software were used to measure fluorescence from a minimum of 10 individual granule cells per brain region for each animal. Preliminary investigation of cell types adherent to coverslips and stained with DAPI (Invitrogen, Eugene, OR, USA) and NeuN (Chemicon International, Temecula, CA, USA) ensured that granule cells were the predominate cell found in the resulting cultures. A coverslip without adherent cells was imaged in HBSS alone as a negative control. During data collection, granule cells were identified by size and shape. Fluorescence data per cell will be averaged for each animal and percent control was calculated per experiment and statistically analyzed as previously stated.
Levels of Mitochondrial Activity

Mitochondria membrane potential was measured by employing identical cell isolation and imaging techniques as described for the previous experiment with the exception of using tetramethyl rhodamine methylester perchlorate (TMRM; Invitrogen, Eugene, OR, USA), a lipophilic potentiometric mitochondrial dye, at concentrations of 150nM in HBSS as the imaging dye. Fluorescent measurements and data analysis were done in the same manner as presented earlier, except that a minimum of 10 individual mitochondria in each granule cell were measured for fluorescence intensity. Cells not exposed to TMRM and adherent to a coverslip were imaged in HBSS as a negative control.

Data for levels of basal intracellular calcium ion homeostasis and mitochondrial activity were analyzed using SPSS Version 12.0.1 Multivariate Analysis of Variance (ANOVA) at $\alpha = 0.05$ to test gender differences. A Univariate Analysis of Variance (ANOVA) at $\alpha = 0.05$ two way ANOVA was used if the Multivariate ANOVA showed that genders could be combined. Tukey’s honest significant difference (HSD) post hoc test was used when significant differences within region and genotypes were assessed.

Amount and Rate of Change in Reactive Oxygen Species Production

Cerebellar and hippocampal cells were isolated as previously described with adjustments made to plating of cells, dye and imaging procedure. After centrifugation, individual collections of cells from different brain regions and genotypes were reconstituted in 2mL MEM with 0.5mL placed in each of three wells of a 48 well cell
culture plate. Following incubation as described previously, media were removed from wells and replaced with CM-H$_2$DCFDA (Invitrogen, Eugene, OR, USA) in HBSS. A BioTek Synergy 4 Microplate Reader with Gen5 software read the plate fluorescence with excitation: 485/20, emission 528/20, optics position: top 50% and sensitivity: 75. Readings occurred every 2.3 minutes for 45 minutes ending with a total of 19 observations.

After the initial readings were completed, cells entered a process of staining with Janus green (Sigma Aldrich, Milwaukee, WI, USA), a colorimetric dye detecting protein content (Raspotnig et al., 1999). The plate reader was set at absorbance: 630. Fluorescence values given at each time point for each brain region and genotype in their respective three wells were divided by their Janus green value to get an adjusted ROS value. The three adjusted ROS values for each region and genotype at each specific time point were then averaged. Each reading was then divided by the initial reading value from the same brain region from the wild type animal. Data was analyzed using SPSS Version 12.0.1a Mixed Model Analysis of Repeated Measures with p = 0.05 to detect any significant changes in amount and rate of ROS production. Preliminary investigation of cell types existing in well plates with staining of DAPI and NeuN ensured that granule cells were highly present in resulting cultures.

The above procedure was also repeated with the addition of 50µM tert-butyl hydrogen peroxide (TBHP; Sigma Aldrich, Milwaukee, WI, USA) to CM-H$_2$DCFDA as a way to identify if cells from either genotype are reaching a ceiling of ROS production or the cells in both genotypes are able to produce similar rate or amounts of ROS.
Fluorescence reading of cells in a well plate with HBSS dye alone was used as a negative control.

RESULTS

Levels of Basal Intracellular Calcium Ion Homeostasis

Mice with VDCC mutations exhibit abnormal calcium influx and specifically the leaner mouse displayed decreased levels of basal intracellular calcium ion homeostasis in the cerebellar granule cells (Bawa and Abbott, 2008). Therefore, the level of calcium ion homeostasis the $tg/tg^{la}$ mouse hippocampal and cerebelar granule cells should be investigated.

Basal intracellular calcium ion homeostasis remained normal in the $tg/tg^{la}$ mouse hippocampus as compared to the wild type animal (Figure 4.1; Figure 4.2). As there have been few abnormalities within the mutant hippocampus throughout this study, it comes at no surprise that this region remained normal. Alternatively, the male $tg/tg^{la}$ cerebellar granule cells levels of basal intracellular calcium homeostasis were significantly greater than in the wild type animal ($P = 0.028$). This discrepancy is not found in the female mice. Male $tg/tg^{la}$ cerebellar granule cell levels of basal intracellular calcium homeostasis were decreased as compared to that of the mutant hippocampus ($P = 0.001$).

While wild type male and female mice could not be compared with each other as the data were presented in percent control, examining raw fluorescent data can suggest whether there are gender differences within either region of the wild type animals.
When the range and standard error of the mean of the wild type animals in both genders were examined, it suggested that there was not a real gender difference between wild type animals (data not shown). It is more likely that the gender difference where the \( \text{tg/tg}\text{la} \) male cerebellum exhibited increased basal intracellular calcium ion levels as compared to the wild type animal is due to the mutation’s effect (Figure 4.3).

![Image of DAPI/NeuN labeled granule cells. Representative images of isolated cells stained with DAPI (A), NeuN (B), and double labeled with DAPI/NeuN to confirm isolation of granule cells.](image)

Figure 4.1 Image of DAPI/NeuN labeled granule cells. Representative images of isolated cells stained with DAPI (A), NeuN (B), and double labeled with DAPI/NeuN to confirm isolation of granule cells.
Figure 4.2 Image of CaG-1 and bright field granule cells. Representative images of CAG-1 fluorescent granule cells (green) and bright field (red). 3-month-old male tg/tg la male hippocampus (A) and cerebellum (B).
Figure 4.3 Graphs of CaG-1 percent control of fluorescence. Graphs comparing 3-month-old \( tg/tg^{la} \) and wild type animals percent control of CaG-1 fluorescent granule cells in the male and female cerebellum (A,B), male and female hippocampus (C,D) and as a comparison within the cerebellum and hippocampus of the female (E) and male (F) mutant mouse. (A) Indicates a significant increase of CaG-1 fluorescence in the \( tg/tg^{la} \) male cerebellar granule cells as compared to the wild type (\( P = 0.028 \)). (B-E) No significant difference. (F) \( tg/tg^{la} \) male cerebellar CaG-1 fluorescence significantly decreased as compared to the hippocampus (\( P = 0.001 \)).
C

Male Hippocampus

D

Female Hippocampus

Figure 4.3 continued
Figure 4.3 continued
Figure 4.4  Image and graph of TMRM percent control fluorescence. Mitochondrial membrane potential (MMP) in isolated cerebellar granule cells in wild type and \( tg/tg^{la} \) mice. (A) Representative image of isolated 3-month-old male \( tg/tg^{la} \) isolated cerebellar granule cells, bright field (green) and TMRM fluorescence (red). Graphs depict of the percent control of TMRM fluorescence in the wild type and \( tg/tg^{la} \) mice with genders combined. B) Hippocampus C) Cerebellum. (A,B) No significant difference between genotypes.
Figure 4.4 continued
Levels of Mitochondrial Membrane Potential

It was important to investigate the level of mitochondrial functioning in the \(tg/tg^{la}\) mice as alterations in their functioning could be affected by the levels of calcium ions entering into the cells through P/Q type calcium ion channels. Variations in the mutant’s MMP could be related to dysfunction in the cell’s ability to maintain normal calcium ion levels and could send affected cells into a death cascade such as the apoptosis leaner cerebellar granule cells exhibit. However, both the \(tg/tg^{la}\) male and female mice did not show any evidence of unusual MMP in cerebellar or hippocampal granule cells (Figure 4.4).

Amount and rate of change in reactive oxygen species (ROS) production

The rate and amount of ROS production in the hippocampus remained normal relative to levels observed for wild type mice (Figure 4.5). When ROS production was artificially enhanced by exposure to 50µM TBHP, no variation remained between both genotypes.

The cerebellum revealed more intriguing results as its granule cells show a significant decrease in the amount of ROS produced (P = 0.000). The effect of addition of TB on the amount of ROS produced also revealed a ceiling in the \(tg/tg^{la}\) where the wild type animal continued to increase, but the mutant was significantly decreased and unable to reach normal levels despite a slight increase of its own (P = .002).
Figure 4.5  Graph of amount and rate of ROS production. The amount and rate of ROS production in the hippocampus (A) and cerebellum (C) of 3-month-old $tg/tg^{la}$ and wild type animals. Amount of ROS and rate of ROS production with 50µM TBHP in hippocampus (B) and cerebellum (D) of 3-month-old $tg/tg^{la}$ and wild type animals. (A, B) No significant difference between genotypes. (C) Significant decrease in amount of ROS production in $tg/tg^{la}$ mice as compared to wild type mice ($p = .000$), no significant difference in rate of ROS production. (D) ) Significant decrease in amount of ROS production in $tg/tg^{la}$ mice compared to wild type mice ($p = .002$ ), no significant difference in rate of ROS production.
Cerebellum

Amount and Rate of Production of ROS

Cerebellum + 50µM TBHP

% Change of Fluorescence

Read

D

Amount and Rate of Production of ROS

Cerebellum + 50µM TBHP

% Change of Fluorescence

Read

Figure 4.5 continued
DISCUSSION

The gender difference observed in basal intracellular calcium ion homeostasis could be the illustration of the \( \text{tg}^{\text{la}}/\text{tg}^{\text{la}} \) mouse having abnormal hormone activity as suggested by previous research in the Abbott lab (Serpedin, 2003). Female leaner mice spend a longer amount of time in estrous and nonestrous as compared to age matched wild type females as a result from abnormal hypothalamus activity (Serpedin, 2003). In addition, other reproductive abnormalities exist in the leaner mouse such as diminished reproductive success, along with an increase in plugs and implants as compensation (Serpedin, 2003). Lower amounts of estradiol in the leaner mouse also imply room for a larger gender discrepancy (Serpedin, 2003).

Despite high testosterone levels, estrogen deficiency in males plays a critical role especially in skeletal growth and maturation (Grumbach, 2000). It is possible that while estrogen is less implicated in the male, a decrease in estrogen could have prominent effects on the brain. In light of the alterations in hormones of the female \( \text{tg}^{\text{la}}/\text{tg}^{\text{la}} \) mouse, male \( \text{tg}^{\text{la}}/\text{tg}^{\text{la}} \) mice may operate with estrogen levels so low that it may greatly affect what normal calcium levels cerebellar cells function at. Estradiol is known to play a significant role in cerebellar development (Gottfried-Blackmore et al., 2007), which could suggest that the \( \text{tg}/\text{tg}^{\text{la}} \) cerebellum is severely affected by hormonal aberrations even prior to the phenotypic onset of abnormalities.

Investigating basal intracellular calcium ion levels does not show the whole picture of calcium in granule cells as this method does not allow for computation of
calcium influx, calcium stores in the endoplasmic reticulum and intracellular stores or calcium binding proteins.

In contrast to the \(tg/tg^{\text{la}}\) results, both leaner and tottering mice displayed at least a 20% reduction in basal intracellular calcium ion homeostasis (Bawa and Abbott, 2008). This gives more support to the hypothesis that the \(tg/tg^{\text{la}}\) mutation is biologically very different than the homozygous \(tg/tg\) or \(tg^{\text{la}}/tg^{\text{la}}\) mice.

The decrease in ROS could be a direct result of the mutation or indirectly caused by adjustments in the cellular physiology due to coping with the mutation. A study of postnatal and juvenile animals could shed light on this matter with the absence or presence of altered ROS levels seen prior or during development of the mutant phenotype. It would be interesting to measure the levels of ROS prior to the phenotypic onset of this disorder.

Altered calcium homeostasis likely found in \(tg/tg^{\text{la}}\) mice because of the presence of altered calcium homeostasis in \(tg/tg\) and \(tg^{\text{la}}/tg^{\text{la}}\) mice. Despite no alteration in basal intracellular calcium levels and in mitochondrial membrane potential, the presumed deviation in calcium homeostasis most likely is related to decreased ROS levels. Other calcium channel mutant mice exhibit abnormal calcium binding proteins, which in turn could lead to decreased ROS. There is not only diminished \(\text{Ca}^{2+}\) uptake by the endoplasmic reticulum in these mice, but reductions in mRNA levels of \(\text{Ca}^{2+}\) binding proteins (Dove et al., 2000). In addition, there is a noticeable decrease in the tottering cerebellar granule cell layer mRNA expression of calretinin, calcium binding protein,
which could be correlated with impaired calcium homeostasis and motor coordination (Cicale et al., 2002).

It is possible that the mitochondria, while functional, are not working at levels comparable to the wild type mouse and therefore produce less ROS byproduct. A decrease in amount of ROS could signify increased antioxidants such as scavengers or SODs. This explanation appears less probable as an increase in these antioxidants are linked to positive characteristics not found in these mice such as hippocamal long term potentiation, learning and memory (Hu et al., 2007; Klann, 1998; Levin et al., 1998).

ROS from the mitochondria may be produced in amounts normal to that of the wild type mouse mitochondria, but that the ROS deficits are coming from other areas of production such as the endoplasmic reticulum and nuclear membrane. In light of the ultrastuctural observations, the smooth endoplasmic reticulum is shown to exhibit crystalline-like array in Purkinje cell axons of the three mutant mice, *tg*\(^{la}/tg^{la}\), *tg/tg*, and *tg/tg*\(^{la}\) (Rhyu et al., 1999). In addition, there is diminished Ca\(^{2+}\) uptake (Dove et al., 2000) and decrease in the mRNA expression of ryanodine receptor type 1 in the endoplasmic reticulum of granule cells and Purkinje cells, respectively, in the *tg/tg* mouse (Cicale et al., 2002).
<table>
<thead>
<tr>
<th></th>
<th>tottering</th>
<th>tottering/leaner</th>
<th>leaner</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cerebellum</strong></td>
<td>Decreased volume &amp; weight</td>
<td>Decreased volume &amp; weight</td>
<td>-----</td>
</tr>
<tr>
<td><strong>Hippocampus</strong></td>
<td>----</td>
<td>No change in volume</td>
<td>----</td>
</tr>
<tr>
<td>Cellular Proliferation in the SGZ of the DG</td>
<td>Decreased</td>
<td>No change</td>
<td>Decreased</td>
</tr>
<tr>
<td>Calcium Ion Homeostasis in CGC</td>
<td>Decreased</td>
<td>Female: no change Male: increase</td>
<td>Decreased</td>
</tr>
<tr>
<td>Mitochondrial Membrane Potential in CGC</td>
<td>No change</td>
<td>No change</td>
<td>Decreased</td>
</tr>
<tr>
<td>Amount of ROS</td>
<td>No change</td>
<td>Decrease in normal and TB induced conditions</td>
<td>No change</td>
</tr>
<tr>
<td>Rate of ROS Production</td>
<td>No change</td>
<td>No change</td>
<td>No change</td>
</tr>
</tbody>
</table>

Figure 4.6: Comparison of results from thesis experiments for $tg/tg^{la}$ mouse as compared to known observations in $tg$ and $tg^{la}$ mice.
CHAPTER V

CONCLUSIONS

The \(tg/tg^{la}\) mouse is the least understood of the mice carrying mutations at the tottering locus. While somewhat neglected by previous research, this mouse appears to have potential to act as a new model for investigating the critical and delicate role of calcium in the hippocampus and cerebellum.

The studies presented in this thesis shed light on basic morphological and physiological abnormalities in the \(tg/tg^{la}\) mouse which often differ from known phenomena in the \(tg/tg\) and \(tg^{la}/tg^{la}\) mouse (Figure 4.6). While it could be expected that this mouse may mimic both or either of the heterozygous tottering or leaner mouse model, it remains quite unique.

Overall, the \(tg/tg^{la}\) hippocampus shows the least dysfunction. The only hippocampal difference discovered in this study lies in the increase in cellular density in the pyramidal layer of CA3 and the hilus of the dentate gyrus. It is possible that this increased density could highlight abnormal connectivity throughout the hippocampus resulting directly from the mutation or the mutation’s effect on estrogen levels or due to the occurrence of epileptiform seizures that occur in these mice. While the cellular architecture and environment in this region show little variation, it suggests that functionality of the hippocampus should be investigated further.

The \(tg/tg^{la}\) cerebellum not only reveals a decrease in volume, but also altered basal intracellular calcium ion homeostasis in male \(tg/tg^{la}\) mice and greatly decreased
amounts of ROS production in $tg/tg^{la}$ mice of both genders. This calcium channel mutation is able to greatly affect both architecture and physiology in the cerebellum. It could be hypothesized that the cerebellar abnormalities described in this thesis give rise to the $tg/tg^{la}$ ataxic phenotype. The larger question remains as to whether some of these cerebellar abnormalities are compensations that have occurred in order to maintain life. Future research could investigate the cerebellum morphology at ages prior to the characteristic $tg/tg^{la}$ phenotypic onset, whether there is any increased incidence of neuronal cell death within the cerebellum, modes of calcium storage and removal and functionality of synaptic connections.


Dobrossy, M.D., Drapeau, E., Aurousseau, C., Le Moal, M., Piazza, P.V., Abrous, D.N., 2003. Differential effects of learning on neurogenesis: learning increases or decreases the number of newly born cells depending on their birth date. Mol Psychiatry. 8, 974-82.


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## APPENDIX

### Volume of the Cerebellum

<table>
<thead>
<tr>
<th>genotype</th>
<th>Mean</th>
<th>Std. Error</th>
<th>95% Confidence Interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>50.396</td>
<td>2.165</td>
<td>45.499-55.293</td>
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<tr>
<td>$tg/tg^{la}$</td>
<td>40.711</td>
<td>2.314</td>
<td>35.476-45.946</td>
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### Volume of the Hippocampus

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<tr>
<td>Wild type</td>
<td>13.100</td>
<td>.843</td>
<td>11.403-15.290</td>
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<tr>
<td>$tg/tg^{la}$</td>
<td>13.347</td>
<td>.843</td>
<td>11.157-15.044</td>
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### Hippocampal Density

#### Pyramidal Cell Layer of CA1

<table>
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<th>Mean</th>
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<tr>
<td>$tg/tg^{la}$</td>
<td>22.594</td>
<td>.631</td>
<td>21.281-23.906</td>
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### Pyramidal Cell Layer of CA3

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### Granule Cell Layer of the Dentate Gyrus

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<td>Wild type</td>
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<td>0.608</td>
<td>45.315</td>
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<td></td>
<td>47.896</td>
<td>0.582</td>
<td>46.685</td>
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### Hilus of the Dentate Gyrus

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<td>0.341</td>
<td>9.563</td>
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<td>8.917</td>
<td>0.327</td>
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### Lacunosum Molecular Layer

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<td>.145</td>
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<td>$tg/tg^{ia}$</td>
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<td>.132</td>
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### Oriens Layer

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<td>Wild type</td>
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<tr>
<td>$tg/tg^{ia}$</td>
<td>2.469</td>
<td>.141</td>
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### Cerebellar Density

**Anterior and Posterior Lobes and Layers Combined**

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<td>.827</td>
<td>26.444 - 29.781</td>
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<td>24.190 - 27.439</td>
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### Molecular Layer, Anterior and Posterior Lobes Combined

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### Granule Cell Layer, Anterior and Posterior Lobes Combined

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<td>10.713</td>
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### Purkinje Cell Layer, Anterior and Posterior Lobes Combined

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<td>$tg/tg^a$</td>
<td>4.802</td>
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Granule Cell Layer, Anterior vs. Posterior Lobe

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<th>Lobe</th>
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Lobule 10, Layers Combined

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<th>Std. Error</th>
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Molecular Layer, Lobule 10

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### Purkinje Cell Layer, Lobule 10

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### Granule Cell Layer, Lobule 10

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### Hippocampal Cell Size

### Pyramidal Cell Layer of CA3

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<tr>
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### Hilus of the Dentate Gyrus

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<tr>
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### BrdU

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PCNA

Granular Cell Layer of the Dentate Gyrus

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Hilus of the Dentate Gyrus

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Subgranular Zone of the Dentate Gyrus

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### GFAP/BrdU Double Labeling

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### NeuN/BrdU Double Labeling

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### Basal Intracellular Calcium Ion Homeostasis

#### Cerebellum

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### Hippocampus

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### Mitochondrial Membrane Potential

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Amount and Rate of ROS Production

**Cerebellum**

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**Cerebellum + 50um TB**

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### Hippocampus

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### Hippocampus + 50uM TB

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</table>
VITA

Emily Mary Murawski received her Bachelor of Science degree in psychology from Texas A&M University in 2007. She entered into the Biomedical Science program with an emphasis in neuroscience research at Texas A&M University College of Veterinary Medicine in August 2007 and received her Master of Science degree in December 2009. Her research interests include neurodegeneration in the hippocampus and cerebellum. She entered the College of Veterinary Medicine at Texas A&M University in August of 2009 in pursuit of a DVM degree. She anticipates practicing with her DVM in a mixed animal clinic.

Ms. Murawski may be reached at 4458 TAMU, Texas A&M University College of Veterinary Medicine, College Station, TX 77843-4458

Her email is Emurawski@cvm.tamu.edu