MAGNETIC RESONANCE IMAGING AND HISTOLOGICAL ANALYSIS OF BETA – AMYLOID PLAQUES IN HUMAN ALZHEIMER’S DISEASE AND APP/PS1 TRANSGENIC MICE

A Dissertation in Neuroscience by Mark David Meadowcroft

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Submitted in Partial Fulfillment of the Requirements for the Degree of Doctor of Philosophy

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ABSTRACT

Alzheimer’s Disease (AD) is a progressive neurological disorder characterized by the hallmark beta-amyloid (Aβ) plaques and neurofibrillary tangles (NFT) found in the neocortex of brain tissue samples. Other associated pathological changes in the brain consist of degeneration of brain tissue via loss of neurons and neuronal changes associated with loss of synapses and dendritic branch dearborization (1). The disease consists of a general progression of cognitive and memory impairment leading to an insidious dementia like state followed by eventual patient fatality. Above the age of 65 years old the incidence rate of Alzheimer’s increases exponentially with age. It is believed that one in ten individuals older than the age of 65 and approximately half of those older than 85 years old have been diagnosed with the disease (2). Approximately 5 million Americans have AD at the present time and with the increased survival age of the world’s inhabitants it is therefore poised to become a greater public health threat as our population ages.

Histopathological investigation currently is the only approach that can accurately diagnose an individual with Alzheimer’s disease; however these methods can only be completed post mortem. Methods to noninvasively examine tissue in vivo have been of great interest in the study of AD and other neurological disorders. Magnetic resonance imaging offers the unique ability to perform detailed in vivo evaluation of neural tissue to aid in this endeavor. Imaging of beta-amyloid plaques in human Alzheimer’s disease and transgenic models that mimic plaque production has been challenging for AD research. The
development of MR imaging technology capable of visualizing and quantifying Aβ plaques in the Alzheimer’s brain is critically important for translational and clinical investigations. Understanding the transverse MR relaxation associated with the neurodegenerative processes in AD would provide insight into future technologies that will aid in clinical studies. Currently, the histo-pathological basis of image contrast and the relaxation mechanism associated with Aβ plaques is not well understood. Analysis and comparison between histology of a tissue sample and MRI in a one-to-one fashion is of importance for validating new imaging methods and contrasts, especially in molecular imaging research. To help in this endeavor, our development of a histological RF coil has been realized and implemented. Formaldehyde fixed tissue samples are cut on a cryostat, placed between two glass coverslips for positioning inside of the histological coil for MR imaging, then following the scanning protocols are carefully removed and histologically stained. The experimental data collected with the histological coil demonstrate the feasibility of imaging thin slices of tissue samples and unambiguously co-registering these MR images to various histological staining results. With the aid of the coil, T$_2$*-weighted images and parametric maps were directly compared to histology stains acquired form a set of Alzheimer’s disease and transgenic APP/PS1 tissue slices. For AD tissue samples, transverse T$_2$* relaxation due to Aβ plaques is shown to be directly associated with the gradation of iron concentration and to a minor degree with plaque size. For the transgenic APP/PS1 model, which generates Aβ plaques and aims to mimic pathology, plaque morphology and size were associated with
image contrast and transverse relaxation. It has been assumed within the MR community that MR contrast associated with beta-amyloid plaques is due to iron concentration alone. However, the contrast due to transgenic plaques with considerably less iron load are equally conspicuous as the human AD plaques in the MR images. These data suggest a duality in the relaxation mechanism were both high focal iron concentration and highly compact fibrillar beta-amyloid masses cause rapid proton transverse magnetization decay. For human AD tissue, the former mechanism is likely the dominant source of $T_2^*$ relaxation; for APP/PS1 animals, the latter is likely the major cause of increased transverse proton relaxation rate in beta-amyloid plaques. Histological evaluation on the APP plaques and AD plaques demonstrate important differences in iron load and plaque morphology that are hypothesized to be related to differences in transverse relaxation mechanisms. This insight is essential for understanding the histo-pathological underpinning of MRI measurements associated with Aβ plaques.

Transgenic mouse models that reproduce neuropathology are important in understanding disease etiology and testing new clinical procedures. Utilization of the transgenic APP/PS1 model brought into the light important dissimilarities between the naturally occurring Aβ plaques found in the AD brain and those engineered to amass in the transgenic model. Understanding the benefits and disadvantages of the APP/PS1 mouse model is important in considering the outcome from studies involving their usage and eventually transitioning these treatments to the in vivo AD brain. Using various histological stains and
microscopy methods aimed at viewing plaques morphology, iron management and inflammatory response we show that there are distinct differences between the naturally occurring human AD plaques and those found in the APP/PS1 model. Clear morphological differences in plaque composition and ultra-structure allude to differences in amyloidogenesis and component fibril formation. Transgenic animal plaques are morphologically distinct from Alzheimer’s plaques as seen in both low magnification traditional microscopy and ultra-high magnification transmission electron microscopy. Microglial and astrocyte staining shows a differing pattern of inflammatory response being much more pronounced in the AD neural tissue compared to the transgenic tissue. Iron mismanagement and focal iron deposition is found ubiquitously throughout the AD tissue cortex, with plaques showing a close relationship to iron accumulation within them, and is not seen in the transgenic model. The data presented here emphasizes pronounced histological grounds for MR relaxation differences between naturally occurring Aβ plaques in the AD brain and those instituted to produce in the APP/PS1 transgenic mouse model. The data suggest that utilization of these transgenic models for various magnetic resonance, histological and pharmacological studies of Alzheimer’s disease should be done with the consideration of these observations.

The evolution and translation of the MR approaches outlined in this work into the clinical domain to accurately diagnose AD and quantify \textit{in vivo} beta-amyloid plaques would be advantageous for future medical diagnosis. The data presented here represents a foundation for understanding the transverse
relaxation mechanisms associated with beta-amyloid plaques in Alzheimer’s disease. Translating these results to an in vivo clinical setting presents a challenge to the MR community. Currently the methods presented here are beyond the resolution limitations and time constraints of low field clinical scanners. However, with advancements in future MR technology there is potential for expanding this research into a valid system.
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<td>Å</td>
<td>angstrom</td>
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<tr>
<td>AA</td>
<td>amino acid</td>
</tr>
<tr>
<td>Aβ</td>
<td>beta-amyloid</td>
</tr>
<tr>
<td>Aβ_{40}</td>
<td>beta-amyloid – 40 amino acids in length</td>
</tr>
<tr>
<td>Aβ_{42}</td>
<td>beta-amyloid – 42 amino acids in length</td>
</tr>
<tr>
<td>AD</td>
<td>Alzheimer's disease</td>
</tr>
<tr>
<td>Al</td>
<td>aluminum</td>
</tr>
<tr>
<td>APP</td>
<td>amyloid precursor protein</td>
</tr>
<tr>
<td>APP/PS1</td>
<td>amyloid precursor protein / presenilin 1</td>
</tr>
<tr>
<td>ApoE</td>
<td>apolipoprotein E</td>
</tr>
<tr>
<td>ApoE4</td>
<td>apolipoprotein E4</td>
</tr>
<tr>
<td>B₀</td>
<td>external static magnetic field</td>
</tr>
<tr>
<td>B₁</td>
<td>RF magnetic field</td>
</tr>
<tr>
<td>C</td>
<td>capacitance</td>
</tr>
<tr>
<td>^{12}C</td>
<td>carbon</td>
</tr>
<tr>
<td>°C</td>
<td>Celcius</td>
</tr>
<tr>
<td>C57BL/6</td>
<td>laboratory strain of black 6 mice</td>
</tr>
<tr>
<td>Ca</td>
<td>calcium</td>
</tr>
<tr>
<td>cDNA</td>
<td>circular deoxyribonucleic acid</td>
</tr>
<tr>
<td>cm</td>
<td>centimeter</td>
</tr>
<tr>
<td>Co</td>
<td>copper</td>
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<tr>
<td>dB</td>
<td>decibel</td>
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dH₂O     distilled Water
DAB      3,3’ – diaminobenzine
DNA      deoxyribonucleic acid
EC       extracellular
FAD      familial Alzheimer’s disease
Fe       iron
Fe²⁺     ferrous iron
Fe³⁺     ferric iron
FeCl₃    ferric chloride
FITC     fluorescein isothiocyanate
g        gram
GFAP     glial fibrillary acidic protein
H        hydrogen
¹H       hydrogen proton
H₂O₂     hydrogen peroxide
H-ferritin heavy chain subunit of ferritin
HCl      hydrochloric acid
IBA1     ionized calcium-binding adaptor molecule 1
IC       intracellular
IgG      immunoglobulin G
kb        kilobase pair
kDa      kilodalton
kg        kilogram
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<td>L</td>
<td>inductance</td>
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<tr>
<td>L-ferritin</td>
<td>light chain subunit of ferritin</td>
</tr>
<tr>
<td>MHz</td>
<td>megahertz</td>
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<td>mg</td>
<td>milligram</td>
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<tr>
<td>MGE</td>
<td>multi gradient echo</td>
</tr>
<tr>
<td>ml</td>
<td>milliliter</td>
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<tr>
<td>mm</td>
<td>millimeter</td>
</tr>
<tr>
<td>mM</td>
<td>millimolar</td>
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<tr>
<td>MR</td>
<td>magnetic resonance</td>
</tr>
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<td>MRI</td>
<td>magnetic resonance imaging</td>
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<td>MRM</td>
<td>magnetic resonance microscopy</td>
</tr>
<tr>
<td>ms</td>
<td>millisecond</td>
</tr>
<tr>
<td>n</td>
<td>number</td>
</tr>
<tr>
<td>N</td>
<td>nitrogen</td>
</tr>
<tr>
<td>NaCl</td>
<td>sodium chloride</td>
</tr>
<tr>
<td>NEX</td>
<td>number of excitations (averages)</td>
</tr>
<tr>
<td>nm</td>
<td>nanometer</td>
</tr>
<tr>
<td>NFT</td>
<td>neurofibrillary tangle</td>
</tr>
<tr>
<td>P</td>
<td>phosphorus</td>
</tr>
<tr>
<td>PBS</td>
<td>phosphate buffered saline</td>
</tr>
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<td>PBST</td>
<td>phosphate buffered saline with tween</td>
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<tr>
<td>OH(^-)</td>
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</tr>
<tr>
<td>PS1</td>
<td>presenilin 1</td>
</tr>
<tr>
<td>Acronym</td>
<td>Definition</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
</tr>
<tr>
<td>PS2</td>
<td>presenilin 2</td>
</tr>
<tr>
<td>RER</td>
<td>rough endoplasmic reticulum</td>
</tr>
<tr>
<td>$R_2$</td>
<td>$1/T_2$ – transverse relaxation</td>
</tr>
<tr>
<td>$R_2^*$</td>
<td>$1/T_2^*$ – rapid loss in transverse magnetization</td>
</tr>
<tr>
<td>RF</td>
<td>radio frequency</td>
</tr>
<tr>
<td>ROI</td>
<td>region of interest</td>
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<tr>
<td>ROS</td>
<td>reactive oxygen species</td>
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<tr>
<td>s</td>
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</tr>
<tr>
<td>SNR</td>
<td>signal-to-noise ratio</td>
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<tr>
<td>SOD</td>
<td>superoxide dismutase</td>
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<tr>
<td>T</td>
<td>tesla</td>
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<tr>
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<td>longitudinal relaxation</td>
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<td>$T_2$</td>
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<tr>
<td>$T_2^*$</td>
<td>rapid loss in transverse magnetization</td>
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<tr>
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<td>transmembrane</td>
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<tr>
<td>TMHF</td>
<td>3,5,5 – trimethylhexanoyl-ferrocene</td>
</tr>
<tr>
<td>TE</td>
<td>echo-time</td>
</tr>
<tr>
<td>TR</td>
<td>time to repeat</td>
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<tr>
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<td>volt</td>
</tr>
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ACKNOWLEDGEMENTS

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Chapter 1
Introduction

1.1 – Alzheimer’s disease

Alzheimer’s disease (AD) is a progressive neurological disorder with increasing prevalence in modern society and currently is the sixth leading cause of death in the United States. As many as five million American’s are currently living with Alzheimer’s disease, which is expected to sharply rise over the next century due to the ever increasing human lifespan leading to increasing amounts of elderly individuals in our society. Alzheimer’s disease is the most common form of dementia and is generally described as a loss of memory, neuronal function and intellectual abilities that interfere with daily life. It was first described in 1906 by the German physician Alois Alzheimer, but is considered to have been prevalent undiagnosed in human society for thousands of years prior to its modern description (3). Classical pathological changes seen in AD include the formation and presence of beta-amyloid (Aβ) senile plaques and neural fibrillary tangles (NFT) in the human neocortex. The pathogenesis of Aβ plaques in the AD brain has been found to start in the entorhinal cortex (archicortical regions such as the hippocampal formation) of the human brain and progress rostrally towards the frontal lobes, caudal-laterally towards the temporal lobes and dorsally into the sensory motor cortex (4). Within these affected regions there is massive neuronal loss leading to neocortical atrophy in these regions. There is no known cure for AD and diagnosis is based upon clinical and familial
observation of the resulting dementia and impending inability to lead a normal life. Positive diagnosis of the disease with absolute certainty is currently only available with histological and pathological analysis of post mortem tissue samples during autopsy.

1.1.1 – Genetic mutations involved in Alzheimer’s

With the mapping of the human genetic sequence and the ability to quickly and accurately complete genetic screening of affected individuals and their family members numerous genetic mutations associated with the disease have been discovered in the last two decades. Approximately 10 percent of AD cases are believed to be of the early onset familial AD (FAD) variety, which exhibit autosomal dominant inheritance, with the remaining 90 percent being sporadic AD with a later clinical onset (5). The role of each mutation is still being elucidated within the scientific community with each playing a role is what is currently being understood as the disease process. The etiology of AD is multifaceted and amazingly complex with numerous genetic factors that each playing an individual role in the whole disease process.

1.1.1.1 – Amyloid Precursor Protein – APP

Amyloid precursor protein (APP) is an integral type I membrane protein expressed in numerous tissue types and predominantly is found concentrated near the synaptic cleft of neurons. The function of is not currently known but it has been implicated in both cholesterol metabolism (6) and synaptic function and
as such with neural plasticity having been demonstrated that hippocampal
euron cells in cultures that lack APP have enhanced excitatory postsynaptic
currents and increased synaptic vesicle pools (7). In addition, APP knockout
mice have been produced that are viable and have the phenotypic effect of
impaired long-term potentiation illustrating the importance of APP in memory
function (8). APP is sequentially processed by at least three proteinases termed
α, β and γ-secretases along two pathways, one amyloidogenic (producing beta-
amyloid) and the other non-amyloidogenic. Cleavage by α-secretase or β-
secretase occurs within the luminal/extracellular domain and results in the
shedding of nearly all the ectodomain to yield two different peptides termed
APPsα or APPsβ. The sequence is then cut by γ-secretases within the
membrane to produce either p3 (α-secretase cleavage, non-amyloidogenic) or
Aβ (β-secretase cleavage, amyloidogenic) (9) (Figure 1.1.1.1). The genetic
sequence for APP is found on chromosome 21 and contains an estimated 18
exons in 240 kilobases of DNA. Individuals with Down syndrome invariably form
similar plaques and classical AD pathology by the age of 50 years due to the fact
that they have a trisomy of chromosome 21 that leads to the overproduction of
APP, pointing to the importance of APP in plaque formation (10). Dysregulation
of APP processing leads to increased levels of Aβ which induces cell death
processes and disturbed axonal transport, leading to defects in cognition,
synaptic plasticity and development of tau pathology (9). There are several
isoforms that have been found in humans ranging from 365 to 770 amino acids
long due to several alternative splicing pathways, with specific isoforms being
expressed in certain cell lines presumably due to alternative functionality (11). The amino acid sequence for APP has been functionally conserved across species and denotes the underlying importance of the yet to be determined conserved physiological function of APP (9). There are many known genetic mutations in APP, with the most widely recognized being a double mutation in position 670/671 known as the “Swedish mutation” due to its prevalence in the Swedish population resulting in a 695 amino acid long peptide (12). Many individuals who harbor APP mutations develop familial AD rather than the more common sporadic AD, which is not linked to any known APP mutation. Research has shown that virtually any mutation in the APP gene sequence leads to an overproduction of beta-amyloid and enhanced plaque formation.
Figure 1.1.1.1

Schematic diagram of APP sequential processing (not drawn in scale). EC: extracellular; TM: transmembrane; IC: intracellular. Aβ domain is highlighted in red. For simplicity, only one cleavage site is shown for each enzyme. The APP peptide is cleaved extracellularly by α-secretase or β-secretase and is then cut by γ-secretases within the membrane to produce either p3 (α-secretase cleavage, non-amyloidogenic) or Aβ (β-secretase cleavage, amyloidogenic).

(Figure from Zheng et al., 2006 (9))
1.1.1.2 – Presenilin 1 and 2

Presenilin 1 and 2 (PS1 and PS2) are multi-pass transmembrane proteins that assemble with other proteins (nicastrin, PEN-2, and APH-1) to form into an active gamma-secretase complex (13). PS1 is a 467 amino acid long polypeptide that is believed to contain between 7 to 9 trans-membrane spanning domains and includes a large hydrophilic loop region located within the cytosol (5). Vertebrate systems possess two presenilin genes, PSEN1 on chromosome 14 in humans that encodes for PS1 and PSEN2 on chromosome 1 which encodes for PS2 (10). Homologues of the presenilins have been found in other mammals and evolutionarily lower life forms including fruit flies (Drosophila), frogs (Xenopus), nematodes (Caenorhabditis elegans), fungi, and plants (Arabidopsis thaliana) (14), and indeed PS1 and PS2 have a high degree of homology between themselves (10,15). The connection between the presenilin proteins and AD was exposed when it was established that mutations in PS1 and PS2 are causative in approximately 50% of patients with familial AD (5), and are the primary cause of AD with onset before the age of 55 years (10). Within the last 25 years more than 50 different mutations in PS1 have been described in families with FAD, relaying the importance of these proteins in the pathogenesis of AD (5). All mutations in the presenilins are missense mutations, with the majority in hot spots encoded by exon 8 from residue 260 – 290, except for the mutation resulting in the deletion of exon 9 (PS1ΔE9) (10). The different severe structural changes due to multiple mutation sites results in a paradoxical phenomena where some cause a partial loss of γ-secretase function and others
cause an increase in function (10,16). It appears that the difference occurs based on where the mutation resides. Within the cell the presenilins are located in the membrane of the endoplasmic reticulum and the cis-Golgi apparatus (10). The generation of beta-amyloid from APP is known to be a normal processing event in cells, with roughly 90 percent of that normally formed being Aβ40 and 10 percent terminating at amino acid 42 (Aβ42). Pathogenic mutations in presenilins modify APP processing leading to the selective amplification of Aβ42 production (10,17). Currently the mechanism by which mutant presenilins exert their effect on APP processing is not known. Indeed, the normal mechanism by which the presenilins function in the γ-secretase complex is not known either. One model proposes that presenilins are involved in the intracellular trafficking of APP guiding it to specific catalytic binding domains in the γ-secretase complex (10). It is a matter of debate as to the theory of how APP is processed by γ-secretase and the presenilins, but it is known that the last step in processing the APP peptide to produce either Aβ40 or Aβ42 is carried out by this complex. This leads to the strong suggestion that mutations in presenilins resulting in the alteration of γ-secretase function is one of the main culprits in AD pathogenesis (18). The role of the presenilins is not limited to the cleavage of the APP peptide sequence, as it has been linked the cleavage of numerous other proteins (Notch-1, caspases, β-catenin, to name a few) involved in neuronal development, skeletal formation, and apoptosis. The importance of presenilins is highlighted by the fact that homozygous knockout mice for PS1 do not survive, whereas heterozygous knockout mice are viable without any sign of a phenotypic abnormality (10).
Figure 1.1.1.2.1

Schematic diagram of PS1 membrane topology based on the eight-transmembrane-domain model. Amino acid residues that are mutated in the various FAD PS1 mutations are depicted as closed circles. (Figure from Thinakaran, 1999 (5))
1.1.1.3 – Apolipoprotein E4

Apolipoprotein E4 (ApoE4) is one of the three major ApoE 299 amino acid isoforms whose polymorphic genetic sequence has been mapped to chromosome 19 in humans. The other forms consist of ApoE2 and ApoE3 and are found throughout the human population in differing densities. ApoE3 is considered the neutral genotype, is found in approximately 60 percent of the population, and is not associated with any genetic disorders, unlike ApoE2 and ApoE4. Individuals whom are homozygous for ApoE4 have an estimated 20 times risk of developing AD at some point in their lives (19). Approximately 40 – 65 percent of individuals with AD have at least one copy of the ApoE4 isoform (19). The normal function of the apolipoproteins is to bind lipids and other triglyceride-rich lipoprotein constituents in the liver to form lipoprotein complexes for transport throughout the bloodstream for storage. It has been found that the apolipoproteins are specifically expressed by astrocytes and microglia in neural tissue and are involved in the enhanced direct protein-to-protein proteolytic break-down of the Aβ peptide and disposal through the blood brain barrier (Figure 1.1.1.3.1), supporting a protective role in AD (20-23). The exact mechanism by which the apolipoproteins interact with beta-amyloid is unknown but it has been proposed that they act as an intracellular and extracellular chaperone molecule that influence Aβ deposition, fibril formation and clearance (22,24). The three isoforms have different abilities to perform this role with the ApoE4 variant being the less effective of the three in breaking down the beta-amyloid peptides (20). Apolipoproteins have been found co-localized with beta-
amyloid plaques in human AD tissue samples, both in newly forming and established plaques (25). This first lead scientists to believe that apolipoproteins were involved in the generation of beta-amyloid plaques, but while it has been also been surmised that they are involved in plaque clearance due to clearance studies using plaque generating mouse models (24). Interestingly, ApoE positive plaques are seen more frequently in homozygous ApoE4/4 AD brains that in non ApoE4 carrying and heterozygous ApoE4 AD brains (25). This supplicates the question as to whether the ApoE4 protein specifically helps induce the formation of beta-amyloid plaques by potentially concentrating Aβ in brain regions inundated with too much beta-amyloid. If this is the case then the apolipoproteins have a duality in mechanism based on the variant(s) the individual has in their genome (23). ApoE2 and more so for ApoE3 are viewed as proteins that aid in the clearance of extracellular beta-amyloid while the ApoE4 genotype, via a yet to be determined mechanism, aids in the chaperoning of beta-amyloid to facilitate plaque formation. It is a matter of speculation into the mechanism by which ApoE4 aids plaque formation, but the affinity of the different ApoE variants for beta-amyloid seems to be at the core of the issue. ApoE association with Aβ compromises the fundamental lipoprotein binding function of apoE such that it selectively binds beta-amyloid instead of lipids (26). It could be that ApoE4 has such a high affinity for Aβ that it is not able to release the peptide for disposal through the blood brain barrier. This would concentrate the ApoE4-Aβ complex throughout affected brain regions such that it ultimately would find its
way to newly formed plaques where the extremely high affinity for self assembly of the Aβ peptides would strip the peptide from the apolipoprotein (21).
Figure 1.1.1.3.1

ApoE and Aβ plaque formation. ApoE has been implicated in the clearance and deposition of Aβ with pleiotropic effects such that genes encoding ApoE2,3 and 4 influence multiple phenotypic traits. Complexed with Aβ, ApoE facilitates the cellular uptake of Aβ, and ultimately disposal of the peptide through the blood brain barrier. ApoE is also required for the extracellular deposition of Aβ as amyloid via an unknown mechanism. (Figure from Hirsch-Reinshagen et al., 2008 (23)).
1.1.2 – Beta-Amyloid Plaque Formation

Several human proteins have been identified that form amyloid fibrils in vivo (27). More than twenty degenerative conditions affecting the central nervous system feature the deposition of proteins in the form of amyloid fibrils and plaques (28) (Table 1.1.2.1 for a list of amyloid associated diseases). Through the study of these diseases, it has become apparent that the formation of amyloid like structures in not an unusual feature of these proteins but rather a general misfolding property of many polypeptide chains (29). The formation of beta-amyloid plaques is a traditional hallmark associated with the AD brain. However, despite extensive study the exact mechanism by which plaques form is currently unknown. The constituent peptide players, environmental subtleties, and genetic mutations have been associated with plaque formation but their interaction between each other in vivo has not been clearly elucidated.

The formation of beta-amyloid from the amyloid precursor protein has been relatively well worked out, but still remains elusive as far as the selective production of Aβ<sub>40</sub> verses Aβ<sub>42</sub>. As discussed earlier in the APP and presenilin sections, beta-amyloid is the product of the cleavage of APP to form either the amyloidogenic beta-amyloid or p3 (Figure 1.1.1.1). APP is first synthesized in the lumen of the endoplasmic reticulum and then undergoes extensive post-translational modification (30). The peptide is then selectively spliced in the extracellular domain as it is excreted by proteases in the secretase family, α-secretase and β-secretase. When sliced by the alpha variant at amino acid 612 the end result is known as the p3 peptide and is the non-amyloidogenic pathway.
Splicing by beta at the 596 amino acid site results in the amyloidogenic formation of beta-amyloid, generally either 40 or 42 amino acids long but can also be truncated either on the N-terminal or C-terminal end to form a variety of potential peptides (31). Following alpha and beta cleavage the peptide is then cleaved further down the C-terminus direction by γ-secretase in either the 637 or 639 amino acid locations (Figure 1.1.2.1 for splicing locations along the APP peptide). The cleaved resulting peptide is then released into the cytosol where it is either maintained intracellularly or excreted by a yet unknown mechanism into the extracellular domain (30). Figure 1.1.2.2 illustrates the cleavage sequence. Figure 1.1.2.3 illustrates a potential secretory pathway for Aβ to the extracellular domain.

Once in the extracellular domain beta-amyloid undergoes specific protein misfolding and self polymerization to form amyloid fibrils. The 42 amino acid variant of the peptide has been shown to be much more willing to misfold and self aggregate to form fibril bundles. This is hypothesized to be due to the additional two hydrophobic amino acids found in the Aβ42 peptide. When hydrophilic amino acid substitutions are made for hydrophobic ones during bench synthesis they reduce the ability of the peptides to aggregate upon themselves (32). It has been shown that 10-14 AA residues are sufficient for amyloidal fibril formation (33). Substitution of these hydrophobic residues with other random hydrophobic residues still results in fibril formation with differing levels of affinity for one another. These views point toward the notion that the hydrophobic regions are responsible for the fibril formation (34) because generic
hydrophobicity rather than specific side chains promote aggregation of Aβ_{42}. As would be expected with protein tertiary structure, the same is also true when substitutions are made for charged amino acids with non-charges ones, the charge-charge interactions function in concert with the non-ionic hydrophobic interactions (35). When the peptides aggregate together they form both parallel and anti-parallel β-sheet structures with strong hydrogen bonding between the peptides (Figure 1.1.2.4). In reality the beta-amyloid peptides form a twisted β-sheet due to the side chains on the amino acids and because the twisted β-sheet represents a lower energy state than a flat β-sheet (36). The separation between stacked β-sheets is approximately 4.7 Å which is seen to dominate during X-ray diffraction studies giving rise to β-strands when multiple peptides form β-sheets next to one another (37). The β-sheets are then arranged stacked upon one another in such a way that the β-sheets are arranged perpendicular to the axis of the fibrils to form protofilaments (Figure 1.1.2.5) (38). The diameter of the protofilaments is approximately 30 Å, which then arrange upon themselves parallel to each other to form a beta-amyloid fibril of approximately 100 Å in diameter (Figure 1.1.2.6). The beta-amyloid fibril core structure is helical with multiple protofilaments twisting around one another giving a helical repeat (one full twist) of 115.5 Å composed of 24 β-strands (36) (Figure 1.1.2.7). These mature fibrils then aggregate upon themselves via hydrophobic and charged attraction to form beta-amyloid plaque seeds which then overtime accumulate more Aβ fibrils to form beta-amyloid plaques.
Table 1.1.2.1

<table>
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<tr>
<th>Disease</th>
<th>Protein Featured</th>
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<td>Alzheimer's disease</td>
<td>Beta amyloid</td>
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<td>Type 2 diabetes mellitus</td>
<td>IAPP</td>
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<td>Alpha-synuclein</td>
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<td>Transmissible spongiform encephalopathy aka &quot;Mad Cow Disease&quot;</td>
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<td>Cardiac arrhythmias</td>
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<td>Serum amyloid A</td>
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<td>Dialysis related amyloidosis</td>
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Figure 1.1.2.1
Cleavage locations of α-, β-, and γ-secretase. Beta cleaves APP after Met596 (β) and Tyr606 (β'), whereas alpha processes APP within the amyloid β-peptide sequence (α), thereby generating the p3 peptide. γ-Secretase cleavage in the transmembrane region generates amyloid β peptides of mainly 40 and 42 amino acid residues long (γ40 and γ42). The asterisks indicate the locations of mutations causing familial AD. (Figure from De Strooper and Annaert, 2000 (30)).
Figure 1.1.2.2

Illustration of beta-amyloid splicing and excretion. Beta secretase cleaves the APP peptide in the extracellular domain followed by gamma secretase in the transmembrane region to form the beta-amyloid peptide. (Figure from the National Institute on Aging, National Institutes of Health)
Figure 1.1.2.3

Proposed mechanism for APP and Aβ transportation throughout the cell and into the extracellular space. Putative localization of APP processing in neurons. Following translation in the RER, APP is transported through the Golgi and into the plasma membrane. Full-length APP and C-terminal fragments are re-internalized via the endosomal system and either directed back to the trans-Golgi or incorporated into secondary lysosomes for degradation (solid arrows). C-terminal fragments are cleaved to release the intracellular domain of APP which may modulate gene transcription in the nucleus (dashed arrows). β-Secretase activity is found at the cell surface as well as in secretory vesicles and the trans-Golgi. α-Secretase activity is found in secretory vesicles and endosomes/lysosomes; its activity at the cell surface is uncertain. (Figure from Turner et al., 2003 (31))
Figure 1.1.2.4

Schematic of β-sheet structure. Representation of hydrogen-bonding patterns in β-sheet structure, showing antiparallel (a) and parallel (b) arrangements of hydrogen bonding. (Figure from Serpell et al., 1997 (36))
**Figure 1.1.2.5**

Close up view of a proposed amyloid fibril composed of two proto-filaments. The β-sheets are arranged stacked upon one another in such a way that the β-sheets are arranged perpendicular to the axis of the fibrils and are coming out of the paper. (Figure from Nelson *et al.*, 2005 (38))
Figure 1.1.2.6

Illustration of amyloid fibril size and structure. Schematic diagram showing the probable hierarchical assembly of structures making up a mature amyloid fibril from the continuous hydrogen-bonded β-sheet structure within a protofilament to the organization of the protofilaments. (Figure from Stromer and Serpell, 2005 (37))
Figure 1.1.2.7

Molecular model of the common core protofilament structure of amyloid fibrils.
Numerous β-sheets (four illustrated here) make up the protofilament structure. These sheets run parallel to the axis of the protofilament, with their component β-strands perpendicular to the fibril axis. With normal twisting of the β-strands, the β-sheets twist around a common helical axis which coincides with the axis of the protofilament, giving a helical repeat of 115.5 Å containing 24 β-strands.
(Figure from Serpell et al., 1997 (36))
1.1.2.1 – Classification and Morphology of Human Alzheimer’s Plaques

When mature Aβ fibrils aggregate upon one another to eventually form beta-amyloid plaques it does not produce consistent plaque morphology and uniformity throughout the AD brain and thus different plaque types arise. The reason for this has been hypothesized to be due to a number of potential factors. Two main hypotheses have been developed to account for the observation that there are plaques with varying morphology in the AD brain. The first states that one type of plaque gradually is converted into another, that plaque stages represent the morphological history of a single plaque. The second hypothesis is that each plaque type evolves independently of the others and that unique in vivo environmental factors are responsible (39). There have been numerous classifications of plaque type based on morphology improvised by different investigators (40,41) but there is a general agreement that there are three main plaque classifications throughout human AD neural tissue. Plaques can be classified and assigned into three distinct types, either dense-cored, fibrillar or diffuse (39) (Figure 1.1.2.1.1). Dense-cored plaques have also been described as ‘burned out’ or ‘compact’ plaques by other investigators and generally have a solid core composed of a dense arrangement of highly fibrillar Aβ. The core of the plaques is surrounded by a fibrillar halo or coronal region with a relative void of space between the coronal region and the core of the plaque. Both the core and the halo region stain positive for thioflavin-S indicating that they are specifically composed of fibrillar beta-amyloid due to the knowledge that thioflavin-S only is able to bind via intercalation of its aromatic side chains into
the stacked β-sheets of mature fibrils that are long enough (42). Fibrillar plaques stain positive for thioflavin-S and exhibit dense Aβ accumulation within the plaque structure without forming a dense spherical core. Dense core and fibrillar plaques generally are from 30 – 75 µm in diameter, and are found to be as large as 100 µm. Diffuse plaques are composed mainly of less-fibrillar Aβ fibrils that are not as long as those found in the other two classifications and stain only minimally with thioflavin-S and must be viewed microscopically with Aβ antibody staining. This results in a patchy morphology that many have described as a ‘cotton-wool’ plaque and are generally much larger than the fibrillar plaques at upwards of 200 µm in diameter (40,41).

There is support for both hypotheses on the formation of plaques in the literature. All plaque types are found in both preclinical and end-stage Alzheimer’s disease (39). The proportion in which they are found differs though to the stage (43,44) of the AD neural tissue. In the preclinical cases there are a greater proportion of diffuse plaques while in the end-stages of AD there are more fibrillar plaques (dense-core and fibrillar) and fewer diffuse plaque deposits. It is possible that both hypotheses are correct in that plaques form independently of one another but are also able to ‘age’ into other classes over time. The genesis of new plaques is assumed to occur even in the late stages of AD such that early diffuse plaque formation is possible and is seen in tissue samples. Additionally, in preclinical stages dense-cored plaques are found, along with diffuse plaques, but in small quantities (some would say insufficient amounts to cause clinical symptoms, but it remains controversial as to whether plaques are
the cause of or result of the disease). If one were to adhere to the ‘aging plaque’
dogma, then the progression of plaques would first start out as diffuse, then gain
fibrillar Aβ\(_{42}\) over time to form fibrillar plaques and eventually recruit more Aβ\(_{42}\)
and the less fibrillar Aβ\(_{40}\) to form dense cored plaques (40). While this variant
self aggregates less readily upon itself, the recruitment of the less fibrillar Aβ\(_{40}\) occurs because it attaches to the framework of the Aβ\(_{42}\) seeds. It must be
maintained that there are numerous factors in the genesis of plaques, mostly
environment factors involved with the specific brain milieu that they are forming
in. In brain regions such as the basal ganglia more diffuse plaques are formed
with fibrillar plaques being rarely found in late state AD (41). While it is unknown
as to why this occurs, there are two possibilities. The first follows that age
hypothesis and would be that this brain region forms plaques at a later time than
the cortex, and as such only has time to develop diffuse plaques. The second
would follow the environmental hypothesis and state that the diffuse plaques are
only forming due to environmental limitations. As a whole it seems that the
variability in plaques found within the spectrum of AD staging is potentially due to
both factors, aging and environment.
Figure 1.1.2.1.1

Classification of beta-amyloid plaques in human Alzheimer’s disease. (A, i-iii)
Series of three optical sections taken through a dense-cored plaque stained with thioflavin-S in a preclinical AD case (83 years). Images show a distinct central core (arrow) of β-amyloid surrounded by a void or clearing and then an outer spherical rim of β-amyloid (arrowhead). (B, i-iii) A similar set of images sectioned through a fibrillar plaque stained with thioflavin-S in a preclinical AD case (81 years). The fibrillar plaques showed dense β-amyloid accumulations throughout the plaque structure. In the central regions these accumulations often appeared as spoke-like accumulations (arrow), emanating from a denser central accumulation. Distinct pores within the plaque were obvious in deeper sections. (C, i-iii) A set of sequential confocal images sectioned through a diffuse plaque labeled with an antibody to β-amyloid in an end-stage AD case (74 years).

(Figure from Dickson, 2001 (39))
1.1.2.2 – Beta-amyloid Plaques and AD

It was once commonly believed that the mature amyloid fibrils in the form of mature plaques were the sole responsible party involved in the neurodegenerative processes associated with the disease. This hypothesis was brought into a gray area when studies viewed the burden of amyloid plaques in the tissue and compared the number of mature plaques with cognitive decline and neuronal loss in the tissue. The data from several separate studies illuminated that there are inconsistencies between the correlation of amyloid burden in tissue and cognitive decline or neuronal loss (39). There are alternative views that the more soluble form of Aβ may be of greater pathological importance than the insoluble beta-amyloid that plaques are comprised of (45,46). The distinction between intracellular and extracellular beta-amyloid upon AD pathogenesis has also been drawn into the light in recent years with studies and their individual neurotoxic effects.

In cell cultures where soluble extracellular Aβ_{40} and Aβ_{42} is introduced into the culture medium at concentrations believed to be found within the in vivo AD brain the extracellular Aβ did not induce any neurotoxicity (46). This indicates that soluble extracellular peptides are not directly responsible for neurodegeneration, in an in vitro cell culture system at least. In order for extracellular beta-amyloid to cause neurodegeneration the peptides must aggregate to form insoluble fibrils which invokes oxidative injuries followed by neuronal death (47) (more on this in the oxidative stress section). When intracellular concentrations of Aβ_{42} are increased neuronal apoptosis results
albeit with a yet unknown mechanism of action, but it has been speculated to be due to overloading of the endoplasmic reticulum (48) or alterations in axonal structure and transport (49) leading to apoptotic signaling. Apoptotic features have been observed in AD neural tissue but the involvement of apoptosis in AD neurodegeneration has been difficult to observe when looking at lesions produced several years in the past (50).

The relationship between intracellular and extracellular beta-amyloid is slowly being provided, but there are many questions as to the dynamics between their relationship. In mouse models, the appearance of extracellular plaques is proceeded by the accumulation of intracellular beta-amyloid (51). This provides some evidence that the two pools of Aβ are somehow related and the accumulation of extracellular beta-amyloid may be in part dependent upon the intracellular accumulation of Aβ (49,51). The method by which soluble extracellular beta-amyloid accumulates outside the cell and eventually forms insoluble fibrillar plaques is purely speculative. There is some evidence to support that beta-amyloid is directly released during final gamma-secretase cleavage into the extracellular space (51). There is also evidence that it is transported and released into the extracellular space at the synaptic terminals where it is known to accumulate and potentially disrupt neuron to neuron transmission (52). There also remains the possibility that intracellular accumulation of beta-amyloid is caused via receptor or membrane mediated endocytosis of extracellular Aβ where it is brought into the cell in an attempt to degrade extracellular beta-amyloid but overloads the system and results in
intracellular accumulation (53). There is yet another possibility in that when excess intracellular beta-amyloid accumulates and eventually leads to cellular death (autolysis) this causes the release of intracellular beta-amyloid into the extracellular matrix contributing to extracellular stores (54). Clearly the relationship between intracellular and extracellular Aβ is complex and as such is under intense research to elucidate their exact dynamics.

1.2 – Iron and the Body

Iron is fundamental and absolutely required for the normal functioning of most life on Earth. It is an essential element necessary for numerous biological functions within the body due to its biochemical reduction and oxidation properties. Its ability to change valence and donate and accept electrons with surrounding ions gives it a vital role in energetic processes that rely on electron transport. While this property is necessary for numerous biological functions, free iron the body can have extremely toxic repercussions when not properly maintained.

1.2.1 – Normal Metabolism of Iron in the Body

Iron is the most abundant metal in the human body, existing biologically in both ferrous (Fe^{+2}) and ferric (Fe^{+3}) forms (55). Iron plays a vital role in oxygen transportation within hemoglobin, is involved in DNA synthesis with ribonucleotide reductase, electron transport in cytochromes, the citric acid cycle, myelination, and the synthesis of neurotransmitters (56-58). Most individuals
who are maintaining proper iron balance and have adequate nourishment have approximately 4 grams of iron within their bodies. The majority of this is found in hemoglobin within red blood cells to carry oxygen and carbon dioxide during respiration and accounts for 3.5 grams of iron. Within the soft tissue of the body there is approximately 500mg and the skeletal system accounts for 150 mg. Iron circulating through the body is found in blood plasma bound to transferrin and accounts for 4 mg of the total. Iron intake and excretion are approximately 1 mg per day each, although these can vary according to gender due to female menstruation which can increases excretion to 2mg per day (59,60) (Figure 1.2.1.1). There is no pathway in the body for the excretion of large amounts of iron per se, other than the process of bleeding. Iron normally is expelled from the body among shed skin cells, hair fibers and cells along the lower intestinal tract lost during defecation (61). This places emphasis on the body to properly maintain iron intake and storage because excess iron cannot leave the body in rapid manner.

Iron intake is obtained through dietary means by absorption through the intestinal lumen. Duodenal cells along the absorptive villus of the intestinal lumen act as sensors of dietary iron intake and overall total body iron (via unknown mechanisms) and regulate intake or iron accordingly (61). Dietary iron may be taken up as either free iron (ferrous or ferric) or as iron found bound to heme. Free iron is principally present in the diet as ferric iron which is insoluble and must be converted to the soluble ferrous form by a reductase in the luminal surface of the duodenal enterocyte (59). After reduction (if necessary), iron is
transported across the apical membrane by the divalent metal transporter where it is stored in the cell for a short period bound within ferritin or is directly transferred into circulation by the iron transporter ferroportin-1 where it is then bound by transferrin, brought primarily to the liver and where it is stored within ferritin (59) (Figure 1.2.1.2). Transferrin is a blood plasma glycoprotein which binds iron reversibly for iron ion delivery via receptor mediated endocytosis (61). Ferritin is a ~450 kDa globular protein comprised of 24 subunits to create a spherical shell of approximately 8 nm inside diameter that stores up to 4500 iron atoms within its core in the form of crystalline ferrihydrite (62,63). There are two forms of ferritin found within the body depending on the subunits they consist of, heavy (H) or light (L). L-ferritin is primarily found within the liver and is involved in long term iron storage for regulatory purposes. H-ferritin is found throughout the body and is involved in the rapid uptake and release of iron to sustain metabolic processes within cells and to promptly help protect against excessive iron detoxification (64,65).

In the brain, iron is found bound to heme within hemoglobin and in iron containing proteins as non-heme iron. The amount of iron stored in the brain as non-heme iron in ferritin accounts for approximately one-third of the total iron in the brain (66-68). There is a progressive increase of brain iron throughout life. At birth a very low iron concentration is present, then follows an extremely rapid increase occurring during the first year of life due to rapid myelination (58). Iron concentration then slowly increases during the first two decades and eventually plateaus to a maximum level at approximately thirty years of age (67). The
distribution of iron in the brain is not homogenous and is largely dependent upon location and function within brain regions. The highest levels of brain iron is found in the extrapyramidal system within the globus pallidus, substantia nigra, red nucleus and myelinated fibers of the putamen. Moderate levels are found in the majority of the forebrain, midbrain and cerebellar structures and the lowest levels are found in the brain-stem and spinal cord (66,68). The regions in the brain with the highest concentration of iron contain more iron than any other organ in the body with the exception of the liver (67). This renders iron homeostasis and regulation within neural tissue to be extremely important. Iron entering the brain must cross the blood brain barrier via receptor mediated transportation. Although not completely understood, iron bound to transferrin binds to transferrin receptors found on the apical surface of endothelial cells lining the neural blood vessels and is then internalized and ported across the basil membrane via the divalent metal transporter protein into cerebral spinal fluid or via a glial route to awaiting neuronal cells (69).
**Figure 1.2.1.1**

Iron stores in the body. (Figure from J.F. de Wolff, 2004, http://commons.wikimedia.org/wiki/File:Iron_metabolism.svg)
Red blood cells (3500 mg) → Reticulo-endothelial system
Spleen, liver

* Ineffective red blood cell production

Bone marrow (150 mg)  →  Blood plasma (4 mg)

Blood plasma (4 mg)  →  Oral intake (1 mg/day)

Tissues (500 mg)  →  Excretion (1 mg/day)
Figure 1.2.1.2

Iron absorption in the body. Ferric iron is reduced to ferrous iron in the intestinal lumen by a membrane reductase where it is transported across the plasma membrane by the divalent metal transporter (DMT-1). Within cells, iron is stored as ferritin or exported across the basolateral surface by the iron transporter ferroportin. Exported iron is converted to ferric iron by the membrane oxidase and bound to transferrin for distribution to tissues. (Figure from Ezquer, 2006 (59))
1.2.2 – Iron and Alzheimer’s disease

Research in the past two decades has shown that misregulation of iron in the body is associated with a number of human neurological disorders. Post-mortem and *in vivo* magnetic resonance imaging studies have shown that there are significantly increased iron levels in the substantia nigra of patients with Parkinson’s disease compared to age matched control brains (70-72). Iron content has been found to be increased in gray and white matter of patients with amyotrophic lateral sclerosis (73). Reduction of iron due to an unknown cause is seen in patients with restless legs syndrome, especially within the substantia nigra. The symptoms of these individuals are somewhat alleviated with intravenous iron therapy and dopaminergic therapy highlighting the importance of iron in neurotransmitter synthesis (74). Data also strongly implicates a disruption of iron homeostasis in Alzheimer’s disease as demonstrated by neuronal alterations in normal iron distribution and iron regulatory proteins (75).

1.2.2.1 – Iron Imbalance

Iron homeostasis in the Alzheimer’s disease brain has been shown to be regionally disrupted based on cellular and protein observations. Quantitative studies on iron in AD brain have shown that iron is 67 percent higher in gray matter and 27 percent higher in white matter compared to age matched controls (76). In the motor cortex gray matter, iron and ferritin levels are both elevated in AD brains but have a 64 percent decrease in transferrin levels (55). These data suggests that iron mobilization is diminished in this brain region (77). Within the
gray matter of the occipital cortex iron levels do not seem to significantly increase with age while ferritin levels do show an increase in concentration. This is interesting considering that the occipital cortex is one of the least affected regions and shows a high ferritin to iron ratio, alluding to the benefit of ferritins ability to store and compartmentalize free iron. Within white matter, transferrin levels are consistently reduced in AD throughout the entire brain by approximately 40 percent (77). Ferritin in white matter is decreased by upwards of 40 percent with the same relative amount of iron present. Without the ability to regulate iron transport and storage while still containing the same amount of iron presents a danger to neural tissue. This creates an environment where unbound free iron is able to detrimentally interact and cause oxidative damage to tissues it comes in contact with. There is a general tendency in the AD brain to have reduced amounts of transferrin and ferritin with a concomitant increase of free iron (56). It is of considerable interest that brain regions which are less susceptible to AD related atrophy show the highest ferritin/iron ratio while those regions that are most susceptible have the lowest ferritin/iron ratio. It seems apparent that the ability to store and properly sequester free iron in regions with the highest ferritin/iron ratio is beneficial in reducing neurodegeneration (55).

Iron acquisition, sequestration, and management is usually under the control of the oligodendrocytes in normal neural tissue (78). Histological studies on iron, transferrin and ferritin unearth an abnormal pattern of cellular staining in AD brains. It is observed that the normal cellular distribution of transferrin in oligodendrocytes, which is responsible for transportation of iron, is altered in the
AD brain such that the majority of transferrin is found within astrocytic bodies (56,75,77). The absence of detectable transferrin in oligodendrocyte cells suggests a specific disruption of iron mobilization to this population of cells. The loss of ferritin in these cells suggests that myelination could be altered as iron is believed to be requisite during this process (58,75). The unusual detection of transferrin in astrocytes seems to have ties to the transfer of iron through the blood brain barrier. Excessive amounts of ferritin and free iron are found around blood vessels in AD tissue suggesting that iron uptake into the brain is compromised and no longer under tight regulation (77). Overall the collective data suggest that iron homeostasis is disrupted in AD neural tissue and changes in levels of iron and iron regulatory proteins could be highly related to the pathogenesis of Alzheimer’s disease.

1.2.2.2 – Iron and Beta-Amyloid

It is known that metal ions can play a role in Aβ fibril formation, but the association in vivo to how they aid in Aβ fibril formation and plaque genesis is not completely understood. The component fibrils of the Aβ plaques are known metalloproteins and possess metal chelation regions in their amino-acid sequence (79-81). The conversion of Aβ’s secondary protein structure has been shown to occur in the presence of divalent metal ion’s like Fe^{2+} (82). It is well documented that iron is found tightly associated with many beta-amyloid plaques. Microglia cells associated with plaques consistently stain positive for intracellular
iron and these cells are found within the plaque boundary and have processes that extend into plaques (75,83-85). However, it is clear from iron stains of beta-amyloid plaques that there is a considerable amount of free iron not associated with any cellular structure that is bound within the amyloid fibrils of the plaques. The relationship of iron to plaque generation is not currently well understood within the literature. It is largely unknown whether the generation of the plaque causes the aggregation of iron to form a core, or if intracellular iron causes the formation of the beta-amyloid protein around it. It is known that iron can aid in the creation of fibrillar beta-amyloid seeds when iron is introduced in a solution of soluble Aβ\textsubscript{42} (86,87). This highly suggests that free iron found throughout the AD brain aids in the formation of beta-amyloid plaques. However, there is an alternative explanation for the relationship between iron and the ability of the beta-amyloid peptide as a metalloprotein. It has been suggested that beta-amyloid might have a neuroprotective function in the AD brain as it might act as a iron metal chelator (88,89). Therapies involving the chelation of metals in AD have been suggested to reduce the beta-amyloid load and neurocognitive effects of the disease (90). Specifically, the usage of the chelator clioquinol has been shown to reduce beta-amyloid aggregation in solutions and has been used in animal models with modest results at reduction of plaque burden (91). Clinical trials are underway with the usage of this chelator as a therapy for AD and show promise based on the hypothesis that decreasing metal ions are pivotal in the formation and degradation of beta-amyloid plaques (92-94).
1.2.2.3 – Oxidative Stress

The brain is the most aerobically active organ in our bodies consuming 20 percent of the total oxygen we take in due to its high metabolic requirements. Maintenance of the oxidative balance is imperative in the brain and is normally under tight regulation by antioxidants that are present in higher quantities in the brain than any other organ (79). This balance is altered in the AD brain due to the oxidative potential of iron if it is not properly maintained in tissue. In addition to the vital physiological roles that iron has a role in, iron is highly toxic when not regulated properly within the body. Free iron catalyses the production of oxygen free radicals via the Fenton reaction.

\[
(1) \text{Fe}^{+2} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{+3} + \text{OH}^- + \text{OH}^- \\
(2) \text{Fe}^{+3} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{+2} + \text{OOH}^- + \text{H}^+
\]

These reactive oxygen species (ROS) are devastating to neuronal cells and cause a wide variety of cellular dysfunction leading ultimately to cellular apoptosis and death. Oxidative stress can result in a number of deleterious cellular consequences including lipid peroxidation, glycoxidation, protein oxidation, protein crosslinking and DNA fragmentation (95). The half-lives of the ROS varies dependent upon the chemical structure of the radical. Hydroxyl radicals have very fast half-lives and as such their reactions are limited to sites near their generation. There are however other radicals that are generated that have much longer half-lives of a few seconds and thus can diffuse away from their sites of generation and transport their oxidant function to other target sites (96).
There is a clear relationship between beta-amyloid plaques and oxidative stress. Plaques have been shown to be sinks for oxidative ROS created by metal ions where the region surrounding the plaques exhibit oxidative markers (87,97). Ferrous iron bound to beta-amyloid has been shown to facilitate the reduction of $O_2$ to peroxide. The hydrophobic environment in the beta-amyloid plaques acts as a reservoir for the oxygen substrate and creates a microenvironment where bound iron facilities the electron transfer from the reduced metal ion to the dioxygen molecule (97). This creates a redox cycle where iron ions cycle between oxidation states that then repeatedly transfers electrons to successive oxygen molecules (79,87,97). The cycle in hypothesized to follow this pathway:

1. $A\beta + Fe^{3+} \rightarrow A\beta + Fe^{2+}$ (Reduction of the metal ion)
2. $Fe^{2+} + O_2 \rightarrow Fe^{3+} + \cdot O_2$ 
3. $\cdot O_2 + \cdot O_2 + 2H \rightarrow H_2O_2 + O_2$ (production of $H_2O_2$)
4. $Fe^{2+} + H_2O_2 \rightarrow Fe^{3+} + OH^\cdot + OH^\cdot$ (Fenton Chemistry)
5. $\cdot O_2 + H_2O_2 \rightarrow OH^\cdot + OH^\cdot + O_2$ (Haber -Weiss reaction)

The first reaction involves the reduction of the metal ion by beta-amyloid which is then oxidized by oxygen, in turn reducing the oxygen. The reduced oxygen can then combine with hydrogen to produce hydrogen peroxide which is then reduced by ferrous iron to form hydroxyl radicals in the Fenton reaction.
Alternatively, reduced oxygen can bind with hydrogen peroxide to directly produce hydroxyl radicals in a Haber – Weiss reaction (79).

In normal brain tissue the low concentration of ferrous iron and competitive substrates for hydrogen peroxide normally ensure that the production of ROS is not favored in vivo (87). In AD tissue, with imbalances in iron homeostasis and increased free labile iron, the system becomes overwhelmed and unable to maintain an oxidative balance. Measurements of protein oxidation (protein carbonyls) show elevated levels of oxidative stress in the frontal lobes, hippocampus, and superior temporal gyrus with normal results in the cerebellum in AD patients that correlates well with AD histopathology (98-104). Other makers of oxidative stress are indicated in measurements of antioxidant enzyme concentrations. The antioxidant glutathione transferase is altered in several regions of the AD brain including the hippocampus (105) while the antioxidant superoxide dismutase (SOD) displays elevated expression in AD neural tissue (106). The results from numerous investigation teams points to an increase in the production of oxidative species and an alteration in the antioxidant ability of the AD brain. The increased production of reactive oxygen species overwhelms the normal antioxidant capacity of the brain resulting in widespread oxidative stress and neuronal cell death throughout AD neural tissue.

1.3 – Animal Mouse Models of Alzheimer’s Disease

Several mouse models of Alzheimer’s Disease have been developed that attempt to mimic individual components of the disease etiology. The various
models aim at replicating one or a combination of the following aspects of Alzheimer’s; lesions associated with the disease (beta-amyloid plaques and neurofibrillary tangles), symptomatology, or biological mechanisms responsible for the disease. The three levels of the animal models (symptoms, lesions and cause) are independent of one another and different models exhibit distinctive characteristics. Individual types of animal models of Alzheimer’s are shortly reviewed herein for APP, PS1 and APP/PS1 models, specifically, in view of the fact that a APP/PS1 line was utilized in this work.

1.3.1 – Injection of Beta-Amyloid

The reproduction of the lesions associated with AD has been attempted with via the direct injection of β-amyloid (1-42) into the neuronal parenchyma. These injections show some sign of pathological processes in the form of enlarged ventricles and neuronal damage to hippocampal structures. Spatial memory deficits are seen on water maze tests hypothetically accountable to the neuronal damage associated with neuronal loss in the hippocampal structures (107). The injections do not directly reproduce amyloidal plaque or neurofibrillary tangles lesions of the disease.

1.3.2 – Transgenic Expression Models

Through the power of transgenic technology, it is possible to reproduce specific lesions found in AD within animal models. Elevating the levels of transgene product has helped model autosomal dominant human
neurodegenerative diseases mostly in transgenic mouse models. The mammalian prion protein has been utilized as a vehicle to express foreign proteins in the central nervous system. Specifically, the hamster prion protein allows the insertion of a sequencing region up to 45 kb in an open reading frame within a single exon (108). Standard pronuclear injection is used where a linearized DNA cosmid construct is injected into one of the pronuclei during the single cell stage of a fertilized egg. This egg is then transferred into the oviducts of pseudopregnant foster mice and upon which the cosmid replicated along with nuclear division throughout all subsequent dividing cells. The cosmid’s reading frame is transcribed to a functional transgene product utilizing the cells intrinsic nuclear encoding allowing a “gain-of-function”. Upon birth, approximately 20% of the pups have integrated the injected DNA into their genomes. Various DNA sequences, up to 45 kb pairs in length, can be inserted into each cosmid allowing for a variety of end products.

1.3.2.1 – APP Transgenic Models

Seventeen amino acids differ in the mouse and human APP gene sequence, three of which are located in the Aβ sequence (109). However, increasing the level of mouse APP does not produce Aβ deposition, presumably due to the differences in the amino acids between human and mouse APP and Aβ. The first attempts to transflect the beta-amyloid peptide sequence itself resulted in intracellular deposition of beta-amyloid only, resulting in cellular death because the beta-amyloid was neurotoxic (110). It was deemed necessary to
transfect the entire human APP genetic sequence in hopes that it would properly
be synthesized and that the mouse molecular machinery would cleave the APP
to produce beta-amyloid peptide. Three isoforms of human amyloid precursor
protein (hAPP) have been used as transgenes, the 695, 751 and 770 amino acid
variants. Transfecting the APP gene induces overproduction of the protein, from
which the truncated beta-amyloid_{(1-40/42)} peptide is taken. Data from the different
mouse lines that have been derived indicate that the onset and severity of
amyloid deposits in directly linked to levels of Aβ_{42} peptide. APP transgenic mice
generate extremely high amounts of Aβ_{42} and generate beta-amyloid plaques.
However, none of the APP models reflect a complete picture of AD
neuropathology because of the lack of neurofibrillary tangles and prominent
neurodegeneration (111).

1.3.2.2 – PS1 Transgenic Models

Mouse models revolving around PS1 genetics were first created by
knocking out the gene for endogenous mouse PS1. This resulted in a non-viable
mouse line because a null for PS1 results in death late in gestation due to
alternative responsibilities of the γ-secretase complex to cleave other important
proteins necessary for during gestation (111). In an effort to recreate a mouse
model that utilized the point mutations on the PS1 gene a mouse model was
created were transgenes expressing familial AD associated mutations in the PS1
gene sequence, variant A246E, were introduced. Brain measurements of these
mice had highly significant increases in the level of Aβ_{42}. However, since the
altered transgene product was still splicing the endogenous mouse APP peptide
the results were varied as far as plaque deposition. The amino acid difference
between hydrophobic residues on the human and mouse APP amino acide
sequence, especially those on the Aβ sequence, were deemed responsible.

1.3.2.3 – APP/PS1 Transgenic Models

The next logical step in the production of a mouse model that recreated
beta-amyloid plaques and perhaps other AD like neuropathologies was to
combine the FAD mutations in both the APP and PS1 gene into one mouse
model. Animals that co-express both mutations develop numerous amyloid
deposits much earlier than mice that express human APP or human PS1 alone,
at approximately six through nine months of age (112-114). The mice have
elevated Aβ42/Aβ40 ratios in brain homogenates that is not seen in single
transfected APP or PS1 mice (111). AD like pathology was substantially
enhanced as far as plaque production and composition while the mice revealed
deteriorations in brain function as tested with water mazes. The mice still do not
develop numerous other AD like pathologies, namely neurofibrillary tangles and
widespread cerebral atrophy.

1.4 – MRI Quantification of Iron

The basic principles of magnetic resonance imaging was first described
and measured by Isidor Rabi in 1938 when he discovered the magnetic
resonance phenomenon (115), and then was refined independently by Felix
Bloch (116) and Edward Purcell (117) in 1946. Rabi was awarded the Nobel Prize in Physics in 1944 for this work and Bloch and Purcell were co-awarded the prize in 1952. It was noticed that magnetic nuclei, like $^1$H and $^{31}$P, absorbed RF energy at a specific energy when placed in a magnetic field. The phenomenon when absorption occurs was described as having the nuclei in resonance with the magnetic field. It was discovered that different atoms resonate at different frequencies at a given field strength, and that the resonance of atoms in a molecule would describe molecular structure information. When atoms such as hydrogen, carbon$^{13}$, fluorine, sodium and phosphorus are in a magnetic field they align themselves along the dominant magnetic field as then act like small bar magnets with dipoles (118). The breakthrough for the motivation behind medical applications of MR technology came when Raymond Damadian in 1971 showed that the nuclear magnetic relaxation times of tissues and tumors differed.

1.4.1 – MRI 101 – Basics of Magnetic Resonance Imaging

In the majority of clinical MR imaging and the MR experiments in this work, the vast majority of nuclei being imaged are hydrogen in water (protons). Hydrogen is also found in other macromolecules (proteins, lipids, etc.) but the signal acquired from them decays much more rapidly and in general is not detected. A fundamental property of atomic nuclei is that they possess spin. Different atoms possess different spin states, with hydrogen having a spin of $\frac{1}{2}$. This spin may be positive or negative. A nucleus with a spin produces a magnetic field around it, termed the magnetic moment. The strength of the
magnetic moment depends upon the mass and rate of spin of the nucleus. Just as a the magnetic needle of a compass aligns with the earth’s magnetic field, the magnetic moment of the nucleus will also align with an external magnetic field. Depending on the spin state (+ or -), a proton can align either parallel or anti-parallel to external field \( (B_0) \). When a proton with a spin state of + \( \frac{1}{2} \) aligns to the external magnetic field the net result is the generation of a small magnetic moment aligned parallel or in the direction of the \( B_0 \) external magnetic field. When a proton with a spin state of - \( \frac{1}{2} \) aligns to the external magnetic field this results in the creation of a magnetic moment that is opposite or anti-parallel to the external magnetic field (Figure 1.4.1.2). The magnetic moment does not align precisely along the external magnetic field, but is tilted at an angle. The interaction of an external magnetic field on the protons and this angle causes the nuclei to precess about the \( B_0 \) field axis. The speed of precession of the proton about the axis of the static \( B_0 \) magnetic field direction is determined by the Larmor equation, which factors in the strength of the external field and the gyromagnetic ratio of the precessing nuclei. At a field strength of 1 Tesla, the set frequency of procession that is equal to 42.58 MHz/T (termed the Larmor frequency) (Figure 1.4.1.1). The parallel spin state results in a lower energy state because the created magnetic field is in alignment with the external field. The opposite is true for the anti-parallel spin state, as it is in a higher energy state with its created magnetic field in opposition to the external field (Figure 1.4.1.3). Due to the lower energy state, the parallel spin state is found slightly more in protons throughout a sample in an external magnetic field. The
population of spin states at the lower energy level is slightly larger than at the higher level, creating a very small excess of spins in one direction verses another (approximately 1 out of 100,000 at 1.5 Tesla). However, the very small difference in population between the two energy states results in very small signal. Considering the amount of protons in a sample following Avogadro’s number, the net spin excess is great enough to be measurable from an outside source (119).
Figure 1.4.1.1

Proton precession around an external magnetic field. Spinning protons precess about the axis of the external $B_0$ field. The frequency of the precession (Larmor frequency) is directly proportional to the strength of the magnetic field. (Figure from Stark, 1999 (120))
Figure 1.4.1.2

Parallel and anti-parallel quantum spin state. On the left, a proton in a parallel spin state resulting in a net magnetization aligned with the external magnetic field. The proton on the right is in an anti-parallel spin state with the net magnetic moment in the opposite direction of the external magnetic field. (Figure from Stark, 1999 (120))
Figure 1.4.1.3

Spin states of protons. The parallel spin state is of lower energy than the anti-parallel spin state because the parallel state results in a magnetic moment that is parallel or in the same direction as the external magnetic field. (Figure from Hoffman, 2008; http://chem.ch.huji.ac.il/nmr/whatisnmr/whatisnmr.html)
The case of the spin-$\frac{1}{2}$ nucleus

The Nuclei are electrically charged and many have spin that causes them to behave like a magnet.

A nucleus with spin at higher energy generates a magnetic field in the opposite direction to the external magnetic field.

No field

Energy gap corresponds to radio frequency

A nucleus with spin at higher energy generates a magnetic field in the direction of the external magnetic field.

$B_0$

External magnetic field
Tissue magnetization represents a source of potential signal for the generation of an MR image. To accomplish this, the magnetization of the protons must be rotated away from their alignment along the $B_0$ magnetic field. Energy in the form of an RF pulse can be applied from an RF transmit coil that alters the sample’s magnetization. The applied RF pulse must be adjusted to the same Larmor frequency at which the proton being imaged processes in the $B_0$ field. This converts some of the magnetization along the $B_0$ axis into magnetization along the perpendicular axes. Magnetization along the $B_0$ field is termed longitudinal magnetization and that along the perpendicular axis transverse magnetization (118). The RF pulse also exchanges energy with the protons and some of them are lifted to a higher energy spin state. After an RF pulse is applied by an RF transmit coil, the perpendicular magnetization is lost quickly while the longitudinal magnetization recovers back to its original precessional state. Induced transverse magnetization recovers at a different rate back to its pre-RF pulse value of 0. The change of transverse magnetization induces an electrical current causing a net magnetic flux and oscillating current through an RF receive coil. This current is perceived as the MR signal. Recoveries of longitudinal and transverse relaxation are termed relaxation. Longitudinal relaxation is referred to as spin-lattice relaxation and follows an exponential decay with a time constant termed $T_1$. Longitudinal relaxation reflects the process that the energy transferred to the protons by the RF pulse is transferred back to the protons surroundings or lattice (118). $T_1$ for a given sample is defined as the amount of time it takes to recover 63 percent ($1-1/e$) of
its original longitudinal magnetization after the RF pulse. Transverse relaxation, also termed spin-spin relaxation, is referred to as $T_2$ and is defined as the amount of time it takes for 63 percent of the transverse magnetization to dissipate (118-120).

The data collected by the receive coil contains information only on the net amount of change in transverse or longitudinal relaxation but does not relay any information on the spatial location of this change within the sample. To relay this information, the application of other external magnetic fields is required. It was realized that applying a spatially changing magnetic field, or field gradient, across the sample produced a proton magnetization signal with varying frequency components depending on spatial location in the magnetic gradient. The magnitude of the differing gradient of magnetization across the sample must be known and back calculated from the resulting spin frequency to find the spatial location of the spinning proton. This magnetic field gradient is applied by a linear gradient coil that changes the magnetic field linearly in a set direction. The gradient directions are set in three dimensions allowing three dimensional imaging when slices are selected dependent upon pulses chosen at a specific frequency and bandwidth. In multi-dimensional MR imaging, the protons in the imaging slice are selectively excited by a combination of a frequency selective RF pulse and a field gradient perpendicular to the imaging plane. The bandwidth of the RF pulse and the amplitude of the slice-selection gradient determine slice thickness and location. The phase of rotation of the protons along one axis (read axis) are aligned with one another, while the phase of protons precession along
the other axis (phase encoding) is changed such that it adds another dimension to the data collection. This results in protons with different frequencies and phases throughout the selected slice of the sample dependent upon spatial location. The computer of the MR scanner takes this mixture of different signals and sorts through the different frequencies and phases for protons with the same frequency by means of the Fourier transform. This results in an output of how much signal (amplitude) of a specific frequency and phase is present at a specific location and then assigns spatial information based on proton frequency and phase in grayscale contrast seen in MR images (118-120).

Tissue relaxation rate is dependent upon properties of the tissue that cause it to relax along the longitudinal or transverse axis at different rates. This is caused by a number of factors, including the amount of protons present in that tissue and factors that cause protons to relax along either axis at increased rates thus shortening their time of relaxation. These perturbations that cause relaxation to occur more rapidly can be inherent tissue properties, i.e. more protons in gray matter verses white matter and faster relaxation of protons in fat verses water or they can be due to properties of the tissue that is due to an alteration in the normal tissue properties, i.e. more protons in cancerous tissue verses healthy tissue or proximity to paramagnetic particles. Transverse $T_2$ relaxation can occur due to a variety of reasons; magnetic dipole-dipole coupling, electric quadrupole coupling, chemical shift anisotropy, scalar coupling and spin-rotation interaction (120,121). Of all these interactions, the most important upon proton relaxation in MRI is magnetic dipole-dipole coupling. This is because protons do not possess
a quadrupole moment as they do not have more than 2 different spin states. Coupling of the proton nucleus to magnetization variations due to atomic level electron currents (chemical shift, scalar and spin-rotation coupling) are small compared to dipole-dipole interactions due to the small field variations caused by electrons (unless there is an unpaired electron as with a paramagnetic material) (120-122).

1.4.2 – Quantitative Iron Measurement with MRI

In an ideal system within a homogenous magnetic field, all nuclei would transversally relax with the same time constant. In real systems, there are differences in the chemical/molecular environment which can lead to a distribution relaxation time constants. This distribution can lead to a diffusion of the magnetic spin vectors causing a loss of signal. Magnetic field inhomogeneities caused my particles altering the magnetic flux of the static field cause a loss of nuclei frequency and phase coherence. The degree of which is dependent on the location of the nuclei in relation to the magnetic field perturbation. The corresponding transverse relaxation time constant is termed $T_2^*$, and is much shorter than $T_2$. There is a relationship between $T_2$ and $T_2^*$ written as:

$$\frac{1}{T_2^*} = \frac{1}{T_2} + \frac{1}{T'_2}$$

Or

$$R_2^* = R_2 + R'_2$$
Where $T'_2$ is the time constant for relaxation due to magnetic inhomogeneities or $\gamma \Delta B_0$, where $\gamma$ represents the gyromagnetic ratio, and $\Delta B_0$ the difference in strength of the locally varying field.

This important factor causes modifications to tissue relaxation rates. Particles induce magnetic field inhomogeneities in the static $B_0$ that, when in close proximity to protons, cause the protons to loss the energy gained from the RF pulses at an increased rate. There are a variety of particles that cause protons to alter their relaxation properties, some of which are used clinically to differentiate tissue with similar relaxation times i.e. contrast enhancers to visualize tumors that are relatively homogenous with their surroundings. Iron particles cause perturbations in the magnetic field homogeneity throughout the tissue and cause the rapid loss of signal due to the large magnetic moment associated with the iron atom (approximately 1000 times greater than hydrogen protons) (123). A single particle can affect the signal in a dephasing region far away from the particle affecting a volume of proton signals that is much larger than the original particle volume. Iron is ferromagnetic and has permanent domains of electron spin magnetic moments that produce very strong macroscopic magnetic fields. Iron has a high permeability and high susceptibility that causes strong spin-spin coupling with nearby protons producing static dephasing of protons within the vicinity of iron particles. Particles of iron act to concentrate the magnetic field lines (flux), creating field changes in regions. This produces field in-homogenous in the static magnetic field compared to the surrounding tissue (119).
As such, ferric iron (Fe$^{+3}$) causes the rapid reduction in T$_2$ and T$_2^*$ rates in protons in the direct vicinity of the iron particle. The effect of ferrous (Fe$^{+2}$) iron on T$_2^*$ and T$_2$ relaxation is much less prominent than that caused by the ferric form (124). This is believed to be due to the short electronic relaxation time of Fe$^{+2}$, while Fe$^{+3}$ effectively increases T$_2$ relaxivity through the paramagnetic dipole-dipole mechanism (125). It has been well documented that iron will increase proton transverse relaxation rates and that the relaxation rate difference is proportional to the concentration of iron ions (123,126-128). It has generally been considered that ferritin and hemosiderin (degraded ferritin) are the distinct forms of non-heme iron in the normal non-diseased brain present in sufficient quantities to affect MR contrast (123,129).

Ferritin exhibits extremely interesting magnetic characteristics and proton interactions that are not easily described by classical theory. Current theoretical models describe the relaxation of water protons in the presence of magnetized nano-particles. The outer sphere model predicts that relaxation of protons is caused by the diffusion of water protons through an inhomogeneous magnetic field created by the magnetic particles. The model predicts that there is a quadratic dependence of relaxation (1/T$_2$) upon the magnetic moment of a spherical particle and magnetic field strength (63,130-135). It has been found that the contributions of the outer sphere theory for solutions of iron oxide nano-particles underestimates the measured relaxation rates and does not fully describe the proton relaxation. In studies using ferritin solutions at differing magnetic field strengths it was found that relaxation 1/T$_2$ to field strength exhibits
a linear dependence (63,125,133,136). This also correlates well to MR data describing a linear relationship between $1/T_2$ relaxation and the iron found in the globus pallidus as the majority of iron (at least 80 percent) found in this structure is in the form of loaded ferritin (63,137). The inner sphere model predicts the occurrence of proton exchange with hydrated spherical magnetic particles (130). The theory aims to describe the relaxation of bound protons to hydrated iron oxides. Hydrated iron oxides are known to interact with hydroxyl groups (protons) on their surface in an amphoteric way (able to act as an acid or a base). Amphotericism can occur with metalloids that have amphoteric oxides and also with amino acids and proteins, which have amine and carboxylic acid groups. There is a problem with this model describing the relaxation mechanism of ferritin in that the theory predicts a quadratic dependence of relaxation rate with magnetic field strength (130). A model that does describe a linear relationship between field strength and relaxation is the static dephasing regime (130,132,138). This model describes the transverse relaxation of homogeneously distributed static protons in the presence of a magnetic field generated by randomly distributed and oriented dipoles (123,130,138). Studies have shown the this model does not satisfy the experimental results seen in ferritin solutions (130,132). Iron load within the ferritin core has an effect upon relaxation as well. It was assumed that iron load would be proportional and directly affect the relaxation dependent upon the amount of iron present in the core with a quadratic dependence explained by simple paramagnetism. Interestingly, this is true for low iron loading factors within the core but when the
core reaches a threshold amount of iron within it and begins nucleation to form
the ferrihydrite crystalline structure it produces a linear relationship between field
strength and $1/T_2$ relaxation (125) (Figure 1.4.2.1) (Figure 1.4.2.2). This is
because the ferrihydrite core of ferritin is antiferromagnetic; however, because of
the size of small ferromagnetic clusters of the loaded ferrihydrite core, ferritin is
also superparamagnetic (123,139). Clearly, classical theories do not accurately
describe the proton relaxation properties of ferritin. Gossiun et al. has proposed
a new theory to explain the relaxation mechanism of ferritin named the proton
Figure 1.4.2.1

Ferritin and ferrihydrite core. Representation of the ferritin spherical protein shell and the ferrihydrite iron oxyhydroxide mineral core. (Figure from Krispin, 2008; http://www.physik.uni-augsburg.de/exp2/nano/nano.en.shtml)
Protein - Dimers

8 nm

12.5 nm

Ferricydrite
Figure 1.4.2.2

Molecular structure of ferrihydrite iron oxyhydroxide mineral (Fe₅HO₈•4H₂O) found within the ferritin core. Figure from Michel, 2007; http://www.aps.anl.gov/Science/Highlights/2007/20070801.htm)
exchange dephasing model (PEDM) (130). The model accounts for the fast-exchange mechanism between both absorbed protons and diffusing protons each with their own relaxation rates that together describe the relaxation rate of the whole system.

Measuring iron in the body with magnetic resonance imaging has proved to be a difficult endeavor. There is a pressing need for quantitative means of measuring iron in the body that are non-invasive, safe, and accurate for clinical diagnosis and management of patients with iron overload. Diseases that could benefit from quantitative measurement of iron include thalassemia major, sickle cell disease, aplastic anemia, myelodysplasia, restless leg syndrome, Parkinson’s disease, Alzheimer’s disease, and other disorders (140). The majority of MR studies investigating the quantification of iron in the body study hepatic (liver) (127,128,141-143), cardiac (144) and brain (70,72,136,145-148) iron concentrations. Currently the best method of measuring iron status of the body is through liver biopsy but the discomfort and risk of the procedure limit its usage (140). MR procedures to quantify iron load have described a number of methodologies from dual to multi-echo $T_2$ and $T_2^*$ imaging (127,140,141,149), phase imaging (123), susceptibility imaging (127,150-152) to using multiple magnetic field strengths in serial (153). Each method has its own advocates and promising results within the iron quantification field.
1.5 – Basics of RF Coils

The primary function of the RF system is to generate and collect MR signals. When tissue or a sample is placed in the magnetic field, the magnetic dipoles of the protons are forced to align with main field ($B_0$). The sample acquires a slight net magnetization (longitudinal) due to the forced alignment of protons to precess about the $B_0$ field. An RF coil is used to induce a current which creates an external magnetic field ($B_1$) generated at a right angle to the static field at a frequency that is tuned and matched to the Larmor frequency of the proton. This results in the exertion of torque on the precessing protons due to the perpendicular $B_1$ field to the static $B_0$ field. This tilts the protons away from alignment with the $B_0$ field towards the $B_1$ field in the transverse plan. The RF pulse is then stopped and the proton then gradually precesses in a complex spiral back towards the $B_0$ field. While the proton is coming back to alignment with the static $B_0$ field there is a small amount of magnetization produced which induces a small current in the RF receiver coil. This current is the resulting MR signal that contains the frequency and phase data from the sample. The voltage of the signal received is on the order of one tenth of a microvolt while the transmitted signal to induce the $B_1$ field is on the order of hundreds of volts resulting in a few milliTesla (or micro) of magnetization (120).
1.5.1 – Radio-Frequency Coil Design

An RF coil can have three possible functions, transmission of the RF signal to align protons along $B_1$, reception of the transverse signal, or both transmit and receive. To transmit or receive signal, an RF coil must be tuned and matched to the resonate frequency of the proton (or other atom) at the Larmor frequency. The coil must be able to maintain this resonance and efficiently store energy at this frequency. All imaging coils are composed of an inductor and a set of capacitive elements to achieve this goal. The inductance ($L$) and capacitance ($C$) of the inductor capacitor circuit determines the resonant frequency of an RF coil such that:

$$\omega = \frac{1}{\sqrt{LC}}$$

A basic diagram of an RF coil can be seen in figure 1.5.1.1. The inductance coil is represented by ($L$) can be made of various metals, usually copper, which have a natural resistance ($r$). Capacitance is generated by variable capacitors, $C_1$ and $C_2$, which function to tune and match, respectively, the frequency of the coil. With sample loading the frequency of the coil goes down due to coupling with the sample, thus the variable capacitors allow the tuning and matching to the set frequency when a sample is inserted (120).
Figure 1.5.1.1

Simple RF coil circuit design. L is the inductance coil with a natural resistance (r) due to the inherent properties of the metal they are made of. $C_1$ and $C_2$ are variable capacitors to tune and match the coil when a sample is inserted.
A basic principle of RF coil design is that the $B_1$ transmitting coil must generate a rotating magnetic field that is perpendicular to the $B_0$ field to create proton torque that causes the proton to align with along the transverse plane ($B_1$). If the generated $B_1$ field lies parallel to the $B_0$ field then there will not be an induced change in the precession of the proton along the transverse plane. The physical properties of a coil such as coil size around the sample are very important considerations. The magnetic coupling between an RF coil and a sample is maximized by making the coil as physically small as possible to accommodate the sample (154). The inherent noise of a coil is increased as the diameter of the coil increases due to the thermal motion of electrons in the conductor, termed Johnson–Nyquist noise. This affects the overall performance of the coil and reducing resistance increases coil sensitivity which increases the coils signal to noise ratio. There is always inherent noise due to the thermal motion in the conductor, but there is also inherent noise in the sample as due to the high electrolyte content of cells and body fluids. These ions result in a relatively high conductivity for the tissue and eddy currents are created with the application of an RF pulse that dissipate power, which enlarges with an increase in magnetic field frequency (also hence the reason for the variable capacitors). For coils that are a large diameter the majority of the noise is due to the large sample they can accommodate. For coils that are less than 3mm, the major source of noise arises from the coil itself (120,154).

There are a vast amount of coils sizes and geometries that have been produced for various applications. In general, coils are developed for very
specific applications, such as head, body trunk or limb imaging. The premise behind much of this work in this thesis was instigated by the desire to image a specific sample, very thin slices of neural tissue. This presented and challenge with numerous hurdles which had to be overcome to produce a coil that accommodated the sample with enough signal to noise to yield high quality MR images.
Chapter 2

Histological Coil Design and Implementation

2.1 – Direct Magnetic Resonance Imaging of Histological Tissue Samples at 3.0 T

2.1.1 – Abstract

Direct imaging of a histological slice is challenging. The vast difference in dimension between planar size and the thickness of histology slices would require an RF coil to produce a uniform radiofrequency magnetic (B1) field in a two-dimensional plane with minimal thickness. A novel RF coil designed specifically for imaging a histology slice was developed and tested. The experimental data demonstrated that the coil was highly sensitive and capable of producing a uniform B1 field distribution in a planar region of histological slides, allowing for acquisition of high resolution T2 images and T2 maps from a 60 µm thick histology sample. The image intensity and T2 distributions were directly compared with histological staining of relative iron concentration of the same slice. This work demonstrates the feasibility of using a micro-imaging histology coil to image thin slices of pathologically diseased tissue to obtain a precise one-to-one comparison between stained tissue and MR images.

2.1.2 – Introduction

Magnetic resonance imaging (MRI) has been applied to a variety of in vivo applications because of its capability in providing high and versatile soft tissue contrasts. However, interpretations of the changes in observed image contrast
associated with the experimental interventions or pathological conditions need to be validated histologically. When comparing MR images to histological methods, a tissue sample of a given organ in whole or in part is traditionally imaged first and then cut, stained and compared to MR images of selected slices (155). This presents a challenge when correlating histological results with MR images because precise co-registration of tissue samples with MR images is notoriously difficult. This method requires that the histology samples be cut in the same planar orientation as MR images. Alternatively, previous investigations utilized three-dimensional MR scans to create rotatable three-dimensional MR models of tissue blocks, which allowed for the rotation of the image slice into a specific orientation with a visual inspection in order to compare to traditional two-dimensional or confocal three-dimensional histological images (156,157). This kind of co-registering process is time consuming and prone to human errors and subjective variability. In addition, the slice thickness of an MR image is generally much larger than the histological tissue slice resulting in a mismatch between the MRI and histology results. In this case, a number of histological slices need to be cut sequentially and assessed as if stacked upon each other to match the thickness of one MR slice selection. With this approach, it is difficult to perform quantitative comparison and analysis of MR images with histology.

Ideally, precise correlation between tissue histology and MRI studies could be obtained using the same tissue sample for both analyses. The implementation of such an approach would require the design and construction
of a specific RF coil able to directly image a thin slice of histological sample. In optimization of an RF coil for this purpose, it is particularly important to obtain an adequate signal-to-noise ratio (SNR) because of small voxel size and the minute volume of tissue being imaged (on the order of $4.5 \times 10^{-3} \text{ cm}^3$) (154,158). To increase the SNR of an image, it is necessary to maximize the filling factor of the coil, which, in turn, requires an RF coil design conforming to the geometry of the histology slice. The vast aspect ratio between planar size and the thickness of the histology slice require that the coil design criteria include the production of a uniform B1 field in a two-dimensional plane with minimal thickness to maximize the filling factor of the coil (159). In this report, we present a novel RF coil design that is capable of imaging thin histological tissue sections, allowing direct comparison of histology and MR images of the same tissue sample.

2.1.3 – Materials and Methods

The histological coil was designed and optimized with the aid of computer modeling using XFDTD 6.3 (Remcom, State College, PA). The initial coil design was first based on a flat slotted tube resonator by Bobroff et al. (160), consisting of two rectangular parallel plates connected by one 664 $\mu$F capacitor at each of the four corners. The optimized RF coil design consisted of a 24 mm wide U-shape conductor strip wrapped around a dielectric material with relative permittivity of $\varepsilon_r = 5$, simulating two stacked #1 thickness glass coverslips (24 x 30 x 0.150 mm) used for holding histology slices. The open end of the U-shaped coil was connected to a voltage driving port. A square slab (24 x 24 x 0.3 mm) of
dielectric material simulating teflon with a relative permittivity of $\varepsilon_r = 2$ was placed between the perfect conductor strips next to the glass cover slips and driving port, providing distributed capacitance for the coil. This design results in a single loop of 24 mm wide copper strip tightly fitted around the histological slides. The B1 field distribution and electromagnetic properties of the flat parallel plate resonator and the histological coil were calculated and evaluated after tuning the coil models numerically to 125.44 MHz with a sinusoidal wave from a 50 ohm voltage source placed along the driving port. The total mesh size of both models consisted of 2,162,560 Yee cells with 0.3 mm in each dimension. The schematics of the initial flat slotted tube resonator, the optimized histological coil design and their corresponding calculated B1 field distributions are shown in Figs. 2.1.5.1 and 2.1.5.2, respectively.

An RF coil was constructed following the optimized histological coil design. The coil was tuned and matched to 125.44 MHz and evaluated on a 3.0 T MRI system (Bruker BioSpin GmbH, Ettlingen, Germany). The Q factor of the coil was measured with a Hewlett-Packard network analyzer and was approximately 290 when unloaded and remained unchanged when loaded with a uniform phantom. The homogeneity of the coil was evaluated with two phantoms produced by placing either 12.5 µl or 43.2 µl of 0.9 % Type A agar (Sigma, St. Louis, MO) between two cover slips. The agar results in a nearly uniform distribution of water throughout the sample and is an ideal test for image homogeneity. The 12.5 µl phantom was nearly circular and approximates the
dimensions of histological brain samples within the imaging region. The 43.2 µl phantom was used to image the entire 24mm x 24mm coil region as this amount of agar was calculated to fill the space of two 24mm x 30mm coverslips at a 60 µm thickness. To evaluate the B1 field homogeneity of the coil with an experimental B1 field map, two gradient-echo images were acquired from the phantom with 90° and 45° flip angles with 100 ms TR, 5.2 ms TE, 192 NEX with a field of view of 40 mm x 40 mm and a matrix size of 128 x 128 for an imaging time of 41 minutes each. From these two images the B1 field map within the phantom sample was calculated following the method by Stollberger and Wach (161). Subsequently, histological brain tissue sections taken from a donated Alzheimer’s disease brain specimen were used for evaluating the coil. Tissue samples were fixed with 4% paraformaldehyde, sectioned at 60 µm using a Leica cryostat, rinsed in phosphate buffered saline (PBS) to leach out the paraformaldehyde, and then placed between two #1 thickness glass coverslips. To maintain tissue hydration, 25 µl of PBS was placed onto the tissue. A hydrophobic barrier pen was used at the edge of the coverslips to keep the interior region between the coverslips from desiccating during imaging. The field of view of all images was 23 mm x 23 mm with a matrix size of 128 x 128, resulting in an in-plane voxel resolution of 179 µm². The 60 µm through plane resolution, defined by the thickness of the sliced tissue, resulted in an overall voxel size of 179 µm x 179 µm x 60 µm. A fast spin-echo T2 imaging was obtained using the same geometric parameters with 2500 ms TR, 12 ms TE, 128 NEX and 16 segmented acquisitions for a total image time of 92 minutes. Multi
spin-echo images were acquired with 1200 ms TR, 10 ms TE, 40 kHz bandwidth, and 64 NEX with 12 echoes for a total imaging time of approximately three hours. T_2-weighted images were generated by summation of the amplitude images from all the echoes to improve image SNR. T_2 maps were calculated from the multi spin-echo data with a linear regression method (Chips2005 software, Children’s Hospital Image Processing System, Cincinnati, OH) in IDL6.1 (Research Systems, Inc., Boulder, CO). The first echo image was removed for the T_2 calculation to reduce the T_1 effect. After imaging studies, the brain tissue samples were stained for iron load with a modified Perl’s prussian blue stain with 1,3-Diaminobenzene back-staining (Sigma, St. Louis, MO).

2.1.4 – Results

Figure 2.1.5.1 illustrates the design of flat slotted tube resonator and the corresponding calculated B1 field map. With such a small coil size, the B1 field of this design becomes very inhomogeneous because of the connections to the capacitor at each of the corners. Due to the small dimensions of the coil, current distribution near the feeding points and capacitive elements becomes a determining factor for B1 field homogeneity over the entire image plane. This problem was overcome by our optimized coil design depicted in Fig. 2.1.5.2 along with the corresponding calculated B1 field map. The two plates are formed with a continuous copper strip folded above and below the sample. This reduced the field inhomogeneity by eliminating four capacitive element connections. In addition, the plates were extended over a region filled with dielectric material to
form a distributed capacitor. The thickness of the 24 x 24 mm Teflon piece was 0.370 mm creating a total capacitance of 26.5 pf. As demonstrated in Fig. 2.1.5.2b, with such a design, the coil produced a nearly perfect homogeneous B1 distribution over the imaging region outlined by the dashed white box. Comparison of the B1 field profiles along the crosshair in Fig. 2.1.5.1b and 2.1.5.2b indicates that the histological coil’s B1 field has a greater magnitude and is more uniform within the imaging region than the parallel plate resonator in both dimensions. Construction was completed with Delrin around the copper strip as seen in both the computer model and the photograph of the coil in Fig. 2.1.5.3. Additional simulations performed with the Delrin enclosure showed that Delrin does not have a noticeable effect upon the B1 field between the copper plates within the imaging region.

To experimentally demonstrate the sensitivity and B1 field homogeneity of the optimized histological coil, a T2-weighted image and B1 field map were obtained from the two uniform agar phantoms shown in Fig. 2.1.5.4a and 2.1.5.4b, respectively. As indicated by the images and the signal intensity profiles along the two center orthogonal lines, the image intensity and the B1 field of our coil are uniform over the entire imaging plane. The average signal to noise ratio (SNR) of the phantom image in Fig. 2.1.5.4a is approximately 51. Figure 2.1.5.5 shows a T2-weighted image obtained with the coil along with a histology image of the same brain slice. The MR image in Fig 2.1.5.5a yields an average SNR of 65. In the histology image with a modified Perl’s stain in Fig. 2.1.5.5b,
the higher iron concentration regions such as those in the white matter appears as darker red-brown. For finer detail, the magnified regions of interest at the bottom of Fig. 2.1.5.5 were digitally post-processed to 50x magnification for the MRI images (5a) and photographed at 50x optical magnification for the bright field histological image (5b). The figure demonstrates that regions with higher focal iron concentration correspond to those having darker intensity in the MR image. High iron concentration is known to cause faster $T_2$ relaxation leading to hypo-intensities in $T_2$-weighted images. With our approach, the iron concentration shown in the histological analysis correlates $T_2$ contrast within MR images in a straightforward fashion.

With the high B1 field homogeneity produced by the histological coil, an accurate measurement of $T_2$ relaxation distribution in a histological tissue sample is possible. Figure 2.1.5.6 shows a representative $T_2$ parameter map (a) and average relaxation curves (b) from the two ROIs selected within white and gray matter from six tissue samples. The $T_2$ map exhibits a uniform distribution within the same tissue type over the slice, indicating an absence of “bias field” interference due to RF field inhomogeneity. Conversely, the $T_2$ values for different tissue types can be clearly delineated and reliably reproduced over multiple scans. These observations are equivalently evident in the $T_2$ relaxation curves, which demonstrate clear differences in $T_2$ decay between ROIs within the two tissue types. Each echo point along the curve has small standard deviations in signal intensity between the scans indicating that data over the six scans are
uniform and reproducible. The $T_2$ measurements were repeated with the histology coil from six different tissue samples of the same brain and yielded an average $T_2$ of $85.5 \pm 2.4$ ms for gray matter and $66.1 \pm 1.5$ ms for white matter.

With the implementation of a high-field 7.0 T magnet at the Center for NMR Research it seemed a logical path to manufacture another histological to utilizing the increased magnetic field. The same design was used for the 7.0 T histology coil with modifications to both the imaging region and capacitance resulting from the Teflon between the continuous copper plate and the tune and matching circuits such that the coil would tune to the 300.44 MHz Larmor frequency. The coil utilizes a 22mm x 30mm x 0.4mm imaging space that allows two 22mm x 30mm glass coverslips with a histology sample between them to be positioned inside. The coil length was decreased from 48mm to 33 mm which decreased the size of the Teflon between the copper plates to 10mm to decrease the distributed capacitance across plates for tuning and matching. To decrease coupling between the coil and other components and aid in grounding the coil, a balanced and unbalanced circuit (Balun) was implemented into the tuning and matching circuit (162) (Figure 2.1.5.7). The additional coil and tuned capacitor in the balun acts to dissipate any residual voltage source that is present in the ground of the coaxial cable.

The resulting numerical field calculations showed that the 7.0 T coil had similar field uniformity compared to the 3.0 T coil but with an increased signal-to-noise ratio of ~204 for the 7.0 T coil verses ~65 for the 3.0 T coil. The resulting images are able to be taken at higher matrix dimensions allowing for higher
imaging resolution across the sample during similar time of scan durations. Images taken with the 3.0 T coil with a matrix size of 128 x 128 over a 23mm x 23mm imaging space yielded 179µm in-plane voxel dimensions, while images taken with the 7.0 T during a similar six hour time period with a matrix of 512 x 512 over the same 23mm x 23mm imaging region yields an in-plane resolution of 45µm. This represents a sixteen times (four in both X and Y dimensions) increase in imaging resolution over a similar time period as the previous coil design (Figure 2.1.5.8).
2.1.5 – Figures

Figure 2.1.5.1

(a) Schematic of the flat slotted tube resonator. Two separate copper strips are connected by a capacitor at each of the four corners. (b) Calculated transverse $B_1$ field component perpendicular to $B_0$ in the center plane of the coil. The imaging region where the glass coverslips are placed is outlined by the white dashed box. The profiles of the $B_1$ field along the cross hairs through the imaging region are shown above and to the right of the $B_1$ map. The direction of the $B_0$ field is noted to aid in the orientation of the coil within the magnet.
Figure 2.1.5.2

(a) Schematic of the optimized histological coil. A continuous strip of copper is wrapped around the glass coverslips and a piece of Teflon (green). Two glass coverslips with the histological sample encased between them are inserted into the opening of the coil for tissue loading. (b) Calculated transverse B1 field perpendicular to Bo in the center plane of the histological coil. The imaging region where the glass coverslips are placed is enclosed in the white dashed line. The profiles along the crosshairs demonstrate the homogeneity of the coil. The red outlined box represents the region where the Teflon is placed. The color scale in both Fig. 2.1.5.1b and 2.1.5.2b are set the same to facilitate comparison between the two designs. The profiles in Fig. 2.1.5.1b and 2.1.5.2b indicate that the homogeneity of the B1 field is improved with the optimized histological coil design. The direction of the Bo field is noted to aid in the orientation of the coil within the magnetic field.
Figure 2.1.5.3

(a) Computer model of the histological coil. The Delrin acetal resin where the coil is positioned is represented by the semitransparent object. (b) Photograph of the completed coil with the 370 µm opening on the side where the tissue sample and coverslips are inserted along with tuning and matching variable capacitors and BNC connector at the end.
Figure 2.1.5.4

(a) $T_2$ weighted image obtained with the histological coil of the 12.5 µl phantom approximating the size of a tissue sample. Profiles through the horizontal and vertical center of the phantom are presented on the top and to the side of the image. (b) B1 field map of the 43.2 µl phantom with profiles demonstrating the B1 field uniformity of the coil within the imaging region. There are signal dropouts at the bottom two corners due to the drying of the agar sample and two small artifacts due to air bubbles within the sample.
Figure 2.1.5.5

(a) T₂ weighted image of a 60 µm thick slice of human brain tissue. (b) Image of the same tissue section stained after MRI by modified Perl’s Prussian blue for iron content. Dark regions are indicative of higher iron concentration within the tissue. The correlation of iron content with T₂ contrast is seen in the images where the white matter shows up as darker on the T₂ image and high in iron with the histology stain. Conversely, gray matter regions show up as brighter on the T₂ image and lower in stained iron concentration. The 50x digital enlargement of the MR image and 50x optical bright field magnification at the bottom demonstrate the close relationship between iron deposition and T₂ content. The regions of highly focal cellular iron concentration in the magnified histology image are clearly seen as darker spots in the MR image. The relative size of the iron deposition affects the magnitude of the hypo-intensities in the MR image. A typical arrowhead-type imaging artifact is visible around the large hypo-intensity in the digitally enlarged MR image. Staining artifacts at the edges of the tissue sample due to increased edge surface area are not indicative of higher iron content. Scale bars of 2000 µm for the whole tissue images and 250 µm for the 50x magnified images are included for reference. The direction of the Bo field is noted to aid in the orientation of the slice within the histological coil.
Figure 2.1.5.6

(a) \( T_2 \) parameter map generated from the same histology slice data as in Fig. 2.1.5.5. Regions of interest were selected in gray and white matter from six individual maps and are represented by the solid and dashed boxes, respectively. (b) Average \( T_2 \) relaxation curves (N=6) with standard deviation bars of each echo image from the ROIs. The average \( T_2 \) in the ROI is 85.5 ± 2.4 ms for gray matter and 66.1 ± 1.5 ms for white matter. The direction of the Bo field is noted to aid in the orientation of the slice within the histological coil.
Figure 2.1.5.7

Side photograph of the 7.0 T histology coil. The fundamental 7.0 T design is similar to the 3.0 T design except for the shortening of the copper plates to reduce the distributed capacitance and the addition of a balun to reduce coupling with outside interference.
**Figure 2.1.5.8**

Comparison between images obtained with the 3.0 T and 7.0 T histocoils.

(a) 3.0 T histology coil MSME image made up of the addition of 8 echoes with a matrix size of 128 x 128 for an acquisition time of 5h59m. (b) 7.0 T histology coil MSME image made up of the addition of 8 echoes with a matrix size of 512 x 512 for an acquisition time of 6h15m. Both scans are of tissue from an Alzheimer’s disease brain. White matter and gray matter and gray matter can clearly be made out in both images, however the increased resolution in the 7.0 T data allows the clear visualization of focal iron deposition that is not easily discernable in the 3.0 T data.
2.1.6 – Discussion

Numerous NMR micro-imaging studies have been carried out on various samples with attainable resolution on the order of several micrometers. Many of these studies continue to use small traditional birdcage volume coils (163,164), while others have investigated using Helmholtz pair coil designs (150,165). Small multi-turn solenoid volume and surface coils (166,167) remain to be the most prevalent design for high resolution images at ultra-high field strengths. Our coil, however, was designed specifically to directly image a histological sample between two standard histological #1 glass cover slips. For such an application, the coil design must meet two stringent criteria: 1) given the minute amount of tissue in a histology slide, the coil sensitivity must be maximized. This requires a compact coil design to optimize the filling factor (4) such that coil volume is minimized and the tissue is as close as possible to the conductive coil elements over the entire sample. 2) As a result of the vast difference in aspect ratio between imaging planar size and the thickness of the slice, the coil must produce a homogenous field distribution within a planar area of 24 mm x 24 mm in the histological slice. Since the two criteria appear to be mutually exclusive, the difficulty in designing such a coil arises in meeting the two criteria simultaneously. To meet criterion 1, the coil must fit the sample tightly. However, as seen in Fig. 2.1.5.1, the B1 distribution under such a condition can be dominated by the current around the terminals of the conductive elements because of the size of the coil and the close vicinity of the conductor and sample. These effects are generally insignificant for the overall field homogeneity in the
imaging region for human studies because of the relatively large size of the coil and sample. This problem can be alleviated by increasing the planar dimension as seen in the flat slotted tube resonator in Fig. 2.1.5.1. With this approach, however, the filling factor of the coil is significantly reduced since the effective coil size is considerably enlarged. In this case, the coil sensitivity is reduced as a trade off for B1 field homogeneity. In our coil design, the number of termination points is minimized to a single input pair by using a continuous copper foil and a distributed capacitor. The latter condition is created by efficiently utilizing the space that is necessary for RF field homogeneity between the input port and imaging region. The calculated B1 field and experimental image profiles in Fig. 2.1.5.2 and 2.1.5.4b are in close agreement with one another and highlight the nearly perfect homogenous field within the histological coil.

It is evident in Fig. 2.1.5.2 that our design utilizes the far end of the coil region where the electric field is weakest for imaging and the region next to the input port where the electric field is strongest for creating distributed capacitance. A decrease of the electric field in the sample results in a reduction of resistive losses within the sample (6) and an increased Q factor (168-171), which is desirable for SNR enhancement in micro-imaging. Coil design placing the tissue sample as close as possible to the coil’s conductor surfaces (within 150μm, coverslip thickness) optimizes the filling factor and further enhances SNR (4).
One other major concern in designing the histological coil for optimizing RF field homogeneity is the so-called “fringe effect” in a conductor with high frequency alternating current. The current distribution, in this case, concentrates to the edge as the frequency of the alternating field increases. This fringe effect is particularly strong if the conductor is a long flat strip (172). Eddy currents in the conductor that tend to compensate the magnetic flux changes in space following Faraday’s law are responsible for the creation of the fringe effect. As seen in the calculated B1 magnitude map in Fig. 2.1.5.2 and demonstrated experimentally in the phantom image in Fig. 2.1.5.4b, concentration of current along the edges is minimal in our coil design because the copper plates below and above the sample are extremely close and the currents in them are nearly equal in magnitude and opposite in direction. With such a coil configuration, the magnetic flux in the regions outside the coil above and below the plate is nearly zero and thereby minimizing the fringe effect in the conductors. This results in a nearly uniform current distribution within the copper plate and a highly concentrated and uniform magnetic flux density inside the coil. The reduction of eddy currents with our coil design equivalently increases the surface area of the conductor and therefore lowers the resistance to the total current, which increases the coil’s Q factor (168,173).

This work enables the micro imaging and histological staining of the same pathologically diseased tissue sections. Post-mortem high resolution MRI studies of regions of interest and comparisons to histology of disease states are
important for biomedical research. The accurate comparison between MR images and histology of the same tissue slices has not previously been possible. With our coil design, we acquired MR images from a histology slide of Alzheimer’s disease brain tissue sections with high resolution and SNR. The results in Fig. 2.1.5.5 demonstrate that the MR image contrast can associate to a specific histology staining technique. Iron staining patterns of the tissue samples are in excellent agreement with the $T_2$ contrast distribution. Closer histological analysis of the enlarged regions in Figure 2.1.5.5 suggests that the focal iron deposits possibly are abnormal accumulations of iron oxide hemosiderin resulting from microbleeds (174,175). Despite these improvements with our coil, however, discrepancies between the images acquired with these two different imaging modalities could occur due to sample preparation and various image artifacts, particularly in the MR images. For example, there are two visible dark spots in the left corner of the $T_2$ images that are not shown in the histological images. These two dark spots appear to be sandwiched by the two bright pixels above and below and are most likely the magnetic susceptibility artifacts caused by the air bubbles trapped between the two coverslips. Thus, great care must be made during sample preparation to remove the air pockets between the sample and one of the two glass coverslips. Furthermore, such magnetic susceptibility artifacts could also arise around the focal region of high iron concentration. As seen the amplified MR image in Fig. 2.1.5.5a, there are three bright spots forming a triangular shape around the dark circular region. While the dark region is clearly seen in the corresponding histology images indicating a focal high iron
concentration, these bright spots are absent in the histology image. The bright spots are typical arrowhead-type imaging artifacts caused by the Bo field gradient in the proximity of the focal iron deposit (151,167,176). Hence, when comparing MR and histological images, caution must be exercised to identify the artifacts in MR images and potential morphological alterations of the sample during histology staining.

The ability to create $T_2$ parametric maps (Fig. 2.1.5.6a) is of importance for comparison of histology with tissue relaxation times. This allows for quantitative studies between regions of interest within the same histology slide and comparison between normal and diseased tissues. These results were acquired from a field strength used in clinical MR imaging systems which simplified our analysis for establishment of a more direct relationship between tissue relaxation times and various pathologies. In our experiment, the histological coil assessment of $T_2$ in formalin fixed ex vivo human entorhinal cortex (N=6) results in 85.5 ± 2.4 ms for gray matter and 66.1 ±1.5 ms for white matter. In vivo $T_2$ measurements using the same field strength of 3.0 T yields 109 ms and 79 for gray and white matter, respectively (177). The observed reduction in $T_2$ relaxation from in vivo conditions to formalin fixed ex vivo measurements follows previously reported results (178,179). Since the physical and physiological conditions such as temperature, diffusion, perfusion, and oxygenation between in vivo tissue and a fixed histology slide are vastly different, the $T_2$ values obtained with the histological coil are generally not expected to be the identical to the in
vivo measurements (147,178-182). The exact understanding of $T_2$ relaxation drop due to the fixation process is not completely understood on a molecular level. In the literature, a number of groups have studied how fixation with formaldehyde affects the $T_2$ relaxation of tissue samples. It has been observed that the $T_2$ value of freshly excised brain tissue decays exponentially during the first week of formalin fixation and plateaus following this time period (179). It is known that fixation causes a number of tissue microstructure changes that affects water motility (147). The main reaction of fixation is believed to be due to the creation of methylene bridges between amino acids in proteins (180). This cross-linking of proteins in turn affects the motility of water in the tissue sample.

Following fixation, the alterations in the microstructure of the tissue cause additional barriers to water molecules that make it more difficult for them to cross resulting in decreases in water motility and overall water diffusion (181,182).

Imaging of unfixed fresh tissue samples was attempted with the histological coil. However, it was found that thin slices of fresh tissue could not withstand the additional forces exerted upon it by the water surface tension between the coverslips. With all of this in mind caution should be exhibited when comparing quantitative histological coil measurements with in vivo results.

Implementation of the coil design at 7 Tesla (Fig. 2.1.5.7) greatly improved both the signal-to-noise ratio and more importantly the resolution of the MR images. The previous 3 T design had adequate SNR and resolution of 179 µm$^2$ but was greatly improved to over sixteen times this at 45 µm$^2$. This is extremely
beneficial when imaging tissue samples in an attempt to compare the MR images to histological images. Images taken under a microscope are at ultrahigh resolution compared to MR datasets (on the order of well under 1 µm² per pixel). Thus it is favorable to have as high resolution MR data to compare to these microscopic images. Of importance in the scope of this research, the size of beta-amyloid plaques in human neural tissue are approximately 30 – 100 µm in diameter. With the original histology coil design and resolution, the resolution of the voxels in the MR images meant that it was extremely difficult to view these plaques. The increased resolution of the 7 Tesla coil meant that individual plaques could be resolved in the MR images, allowing for greater measurement of individual plaque MR characteristics (Fig. 2.1.5.8).

In summary, novel RF coils at 3.0 and 7.0 Tesla specifically were designed, developed and tested for direct imaging of histology slices. The numerical simulation and experimental data demonstrated that the developed coils were highly sensitive and capable of producing a uniform B1 field distribution in a planar region of a typical histological slice, allowing for acquisition of high resolution T₂ images and T₂ maps from a histology slice. The image intensity and T₂ relaxation map were directly compared with histological staining of relative iron concentration of the same slice. The experimental results demonstrated the feasibility of using our histological coil to image thin slices of tissue and to improve co-registration with histological results. Analysis and comparison between histology of a tissue sample and MRI in a one-to-one
fashion is of importance for validating new imaging methods and contrasts, especially in molecular imaging research. Understanding the relationship between a specific pathology and the resulting MRI contrast or quantitative parameters will aid in interpretation of contrast differentials seen in clinical studies.
Chapter 3

Magnetic Resonance Imaging and Histological Analysis of Beta-Amyloid Plaques in Both Human Alzheimer’s Disease and APP/PS1 Transgenic Mice

3.1 – Abstract

Imaging of beta amyloid (Aβ) plaques in human Alzheimer’s disease (AD) and the APP/PS1 mouse model has been of great interest for AD research. However, the histo-pathological basis of the image contrast and relaxation mechanism associated with Aβ-plaques has not been well-understood. With the aid of the previously developed histological coil, $T_2^*$-weighted images and $R_2^*$ parametric maps were directly compared to histology stains acquired from the same set of Alzheimer’s and APP/PS1 tissue slices. The electronic microscopy and histology images revealed substantial differences in plaque morphology and associated iron concentration between AD and transgenic APP/PS1 mice tissue samples. For AD tissues, $T_2^*$ contrast of Aβ-plaques was directly associated with the gradation of iron concentration. Plaques with considerably less iron load in the APP/PS1 animal tissues are equally conspicuous as the human plaques in the MR images. These data suggest a duality in the relaxation mechanism where both high focal iron concentration and highly compact fibrillar beta-amyloid masses cause rapid proton transverse magnetization decay. For human tissues, the former mechanism is likely the dominant source of $R_2^*$ relaxation; for APP/PS1 animals, the latter is likely the major cause of increased transverse proton relaxation rate in Aβ-plaques. The data presented are essential for
understanding the histo-pathological underpinning of MRI measurement associated with Aβ plaques in humans and animals.

3.2 – Introduction

The formation of amyloid-beta (Aβ) plaques is a neuropathological hallmark and a cardinal feature of Alzheimer’s disease (AD). APP/PS1 transgenic mouse models that mimic the formation of human Aβ plaques in the mouse brain are widely used as an animal model for AD investigations. The development of an imaging technology capable of visualizing and quantifying Aβ plaques in animal models and in the AD brain is critically important for translational, preclinical and clinical research. The ability to delineate Aβ plaques with MRI has been demonstrated ex vivo with human brain samples (183) and in vivo with the mouse model (184-186). Understanding the histological basis of MRI contrast associated with Aβ plaques is essential in this endeavor. To achieve this goal, it is necessary to correlate MRI results with histological stains, which has been technologically challenging because of the limitations in co-registration of planar histology tissue samples with MR images. A prior study involving the innovation of a histological coil has addressed this long-standing difficulty (187). The ability to directly image histological samples is possible when employing the developed histological radio-frequency (RF) coil design. Consequently, MR images and histology data from the same tissue sample can be directly overlaid and compared without uncertainties of co-registration between the two imaging modalities. The goal of the current study is to use this novel technology to a)
further optimize the method for routine imaging-histology studies, b) establish the relationship between MR image contrast associated with Aβ plaques and their histology, and c) compare such relationships in human and transgenic APP/PS1 mouse tissues. Furthermore, to ultimately apply the therapies developed with the animal models to human studies, the Aβ plaque MR image-pathology relationship must be validated in humans and compared to transgenic animal data. It is hypothesized within the literature that iron found in and around the amyloid plaques is the dominant cause of the hypo-intensities seen in the MR images. Examination of the relationship between MR contrast due to Aβ plaque and iron deposition both in human AD and the APP/PS1 model is described. These data suggest that iron load alone does not account for the hypo-intensities that are observed in the T2*-weighted images of the animal pathology. The relationship of plaque morphology and overall globular size with their manifestation in the MR image are described. The method developed in this report can be particularly useful for further validation of the histological basis of MR contrast and the development of animal models for other neurological diseases.

3.3 – Materials and Methods

Human Alzheimer’s and Control Brain Samples:

Entorhinal cortex brain tissue samples from clinically and histologically determined AD subjects (n=5) and age-matched controls (n=4) were donated with consent following institutional guidelines and obtained from both internal and external sources (Harvard Brain Tissue Resource Center, McLean Hospital,
Belmont, MA). Analysis of the tissue obtained from the brain bank indicates that there was not a significant difference between the age of the subjects upon bereavement. The post mortem index between the tissue harvesting at time of death and utilization in this study was significantly different between the tissue samples, with a slightly longer time period for controls. Tissue from the entorhinal cortex was fully fixed in 4% paraformaldehyde in pH 7.3 phosphate buffered saline (PBS) after which blocks of tissue were cryogenically protected in graded sucrose solutions then sectioned at 60 μm on a cryostat. The sections were prepared for magnetic resonance imaging according to previous methods explained below (187).

**Transgenic APP/PS1 and Control Mice:**

Transgenic mice (n=5) inserted with a chimeric amyloid precursor protein (APP) (APPswe_695) and a mutant human presenilin 1 (A246E variant), developed by Borchelt et al. (108,112), were obtained commercially from The Jackson Laboratory (strain type B6C3-Tg(APP695)3DbTg(PSEN1)5Db/J). Animals were kept in the animal facility under veterinary care with normal feeding, environmental, and handling conditions. Age-matched cage mate C57BL/6 mice (n=3) were used as a control group. Upon aging naturally until 24 months, animals were euthanized via an intra-parietal injection of sodium pentobarbital (Nembutal, 200 mg/kg) and were transcardially perfused with cold Lactated Ringer's solution (pH 7.4) followed by buffered 4% paraformaldehyde in PBS. Whole brain tissue was harvested fixed in pH 7.3 buffered 4% paraformaldehyde,
placed in graded sucrose solutions for cryo-protection, then sections were cut at 60µm and prepared for MR micro-imaging using the protocol below.

**Histological slice MRI Scanning Protocols:**

Histological MRI was accomplished on a 7.0 T magnet (Bruker BioSpin MRI GmbH, Ettlingen, Germany) with the histological coil (187). 60 µm tissue samples were floated in PBS for 30 minutes to allow any residual formalin and sucrose in the tissue to leach out and then encased between the two coverslips. The sample was then placed into the histological RF coil and tuned/matched to 300.48 MHz for micro-MR imaging. For T$_2^*$ imaging, multi-gradient-echo (MGE) images were acquired in 6h 32m with a TR=718, TE=6.48, FA=30°, 8 echoes with 9ms echo spacing, 64 averages, FOV=23mm x 23mm, and a matrix of 512 x 512 resulting in a final in-plane resolution of 45 µm x 45 µm.

**Iron and Amyloid Staining:**

After histological MR imaging, the 60 µm tissue sections were free floated and rinsed in deionized water (dH$_2$O) for 15 minutes. Tissue sections were co-stained for both iron and beta-amyloid with a Perl’s Prussian Blue stain followed by an aqueous Thioflavin-S stain. The sections were placed in equal volumes of freshly prepared 4% potassium ferrocyanide (#P236, Fisher Scientific, Waltham, MA) and 4% hydrochloric acid (final combined concentrations 2% for each) for 30 minutes under gentle rocking followed by two rinses in dH$_2$O. Intensification of the iron stain was performed with 5 minutes of 3,3'-diaminobenzidine
tetrahydrochloride (DAB) staining (#D5637, Sigma, St. Louis, MO) (10 mg dissolved in 15ml of PBS with 16 µl of 30% H₂O₂) followed by two five-minute rinses in dΗ₂O. Tissue samples were then free floated in 20 ml of 1% Thioflavin-S (#T1892, Sigma, St. Louis, MO) aqueous solution for 5 minutes, followed by differentiation in 70% ethanol for 5 minutes and two five-minute washes in dΗ₂O. Sections were mounted on histological slides with FluorSave Aqueous mounting media (#345789, Calbiochem, San Diego, CA). To test for the possibility of confounding interactions between the two staining methods during co-staining, each was tested on separate tissue samples to determine their individual efficacy and compared to the co-stained sections. A modified Perl’s stain further tested the transgenic animal tissue for minute amounts of iron within the amyloid masses. The stain was adapted and modified from LeVine (188,189) and employs the usage of proteinase K and detergents to break down the periphery of the plaques forming openings in the highly hydrophobic amyloid plaques. This allows the aqueous Perl’s stain to infiltrate into the amyloid mass and is sensitive enough to detect trace amounts of iron in the Aβ plaques.

Microscopy:

High resolution microscopy of the tissue sections was performed using a Nikon OptiPhot microscope and Nikon Digital Sight camera using NIS-Elements software.—Bright field microscopy under the visible light spectra was used to view iron stains. A FITC fluorescence cube at 430 nm excitation and 550 nm emission was used to visualize Thioflavin-S positive Aβ deposits. Whole tissue image
mosaics were created with the 40x magnification bright field of fluorescent images using the Photomerge option in Adobe Photoshop CS (Adobe Systems Incorporated, San Jose, CA, USA). For transmission electron microscopy (TEM), samples were prepared by cutting tissue sections at 60 µm and staining them with Thioflavin-S to elucidate Aβ plaque location. Known regions of Aβ plaque distribution approximately 1 mm x 1 mm were micro-dissected from the sections and processed for electron microscopy following traditional methods.

**MRI and Histology Post Processing:**

Post processing was performed using in-house software developed with the interactive data language (IDL) 6.1 framework (Research Systems Inc., ITT Industries, Boulder, CO, USA). The program allows for the creation of $T_2^*$ parameter maps from MGE data with a linear regression method. Plaque regions of interest (ROI's) were selected by averaging the relaxation of three voxels of interest within each measured plaque. Averaging of multiple voxels helps eliminate partial volume effects where it is possible and likely that plaques span multiple voxels on the MR data sets. This allowed for the evaluation of relaxation and magnitude measurements from individual plaques in both human AD and APP/PS1 tissue. ROI's for non-plaque regions were selected in regions known to be devoid of Aβ plaques according to the Thioflavin-S staining. For co-registration to the staining methods, $T_2^*$ weighted images were generated by the summation of the amplitude images from all of the echoes. Histology and MRI images were imported into Image J (National Institutes of Health, Bethesda, MD,
USA) to measure Aβ plaque diameter. Freehand boundaries were drawn around the plaques and the average pixel length of the major and minor axes from the resulting ellipsoid were used to estimate overall diameter. A micrometer slide was viewed at different magnifications to obtain precise measurements of the pixel to micrometer dimensions for plaque diameter calculation.

3.4 – Results

Figure 3.5.1 shows a T$_2^*$ weighted MR image and histological images co-stained with thioflavin-S for Aβ plaques and a traditional Perl’s stain for ferric iron (Fe$^{3+}$) within the same tissue section from the entorhinal cortex of an AD subject. The image scales of the MR scan and stain mosaic images for Aβ and iron are the same for ease of comparison. For detailed evaluation, 40x magnification images of four selected regions outlined with red boxes are shown on both sides of the corresponding main images. The T$_2^*$-weighted images of human Alzheimer’s samples show a loss of signal in focal regions that co-registers remarkably well to the histological iron stain. Many dark hypo-intense spots can clearly be observed in the gray matter in the MR image in Fig. 3.5.1a. As seen in the magnification of the thioflavin-S stain images (figure 3.5.1b), those hypo-intensities observed in the MR images correspond to either punctate green fluorescent Thioflavin-S positive Aβ-plaques (red arrows) or dark brown patches indicating regions of high focal iron. Focal regions of high iron that are not associated with Aβ plaques can be attributed to locations high in iron such as capillary vessels, hemosiderin (degraded ferritin) or regions of iron oxide.
(magnetite, hematite or wüstite) deposition (blue arrows) (175,190). Co-staining of Aβ with Thioflavin-S and iron with a Perl's stain allows the direct comparison of ferric iron concentration in and around the beta-amyloid plaques (Fig. 3.5.1c). The Perl's stain of the tissue section demonstrate focal high iron concentration associated with Aβ plaques in exceptional detail. Many of the Aβ plaques seen in the magnified images appear to have a dark core in their centers that are loaded with a higher concentration of iron, as indicated in the Perl's stain. This set of images reveals an unprecedented close relationship with MRI contrast between Aβ plaques and brain tissue iron deposition. The minimal AD plaque size diameter that can be resolved with the $T_2^*$ weighted sequence is 38 µm, demonstrable with the 250 µm scale bars in the figure 3.5.1 magnifications. Smaller human AD plaques that do not visibly contain large amounts of iron are not as discernable on the MR images. The high iron distribution within the white matter tracks of the human tissue is consistent with iron found in oligodendrocyte cells making up the myelin sheaths (191). With the histological coil, the relationship between $T_2^*$ image contrast associated with Aβ plaques and their association with iron deposition are demonstrated unambiguously. The presence of higher iron deposition in the Aβ plaques is a contributing factor to $T_2^*$ image contrast generation. Regions of gray matter that exhibit high iron content are clearly visible on the MR image with varying degrees of signal dropout differing in size based on the amount of iron present at that location. MR image and histology from a normal brain tissue are shown in Fig. 3.5.2 and demonstrate a complete absence of Aβ plaques. The Perl's stained images of the human
control tissue sections demonstrate the known patterns of normal iron staining in regions associated with high iron concentration. Iron stains of the human control tissue (figure 3.5.2c) show the expected high diffuse iron in the white matter tracks due to the high iron concentration within the oligodendrocyte cells (58). This known phenomena can be seen with all histological iron stains and in MR images as a signal dropout. Comparison of white matter iron staining between human AD and control samples shows similar iron deposition illustrating negligible iron loss due to formalin leaching. Fluorescent microscopy shows a lack of Thioflavin-S positive Aβ staining, while nonspecific background fluorescence is present. Having neither Aβ plaques nor high focal iron regions, MR images of human control tissue (Fig. 3.5.2a) exhibit a lack of hypo-intensities in gray matter except those caused by the highlighted focal iron regions (blue arrows in Fig. 3.5.2) and coil noise. Small hypo-intensities that are seen in figure 3.5.2 are related to inhomogeneities within the copper foil used in the RF coil, small micro-pockets of air causing artifacts, or small regions of focal iron.

The corresponding images from APP/PS1 and age-matched C57BL/6 control mice tissue sections are shown in figures 3.5.3 and 3.5.4, respectively. The image scales of the animal MR images and histology mosaics for iron and Aβ plaques are the same as in the human data for ease of comparison. The widespread dark spots (red arrows) seen in the human tissue are also shown conspicuously in the $T_2^*$ image of the transgenic animal tissue within the piriform cortex (Fig. 3.5.3a). Thioflavin-S staining (Fig. 3.5.3b) confirms that these dark
spots are Aβ plaques within the APP/PS1 animal brain. MR imaging of mouse control tissue (Fig. 3.5.4a) shows no evidence of focal hypo-intense regions in the neo-cortex. The iron stain of the control mouse tissue in figure 3.5.4c shows that there are no visible focal iron regions in the neo-cortex or other gray matter regions. Thioflavin-S staining does not indicate any positive amyloid staining for plaques in mouse control sections apart from the expected non-specific background staining. Regions that are known to be high in iron, such as the substantia nigra and caudate/putamen, stain dark as expected (Fig. 3.5.3c) (192). The plaque distribution is consistent with previous studies using the APP/PS1 transgenic mouse line (186,193-195), while plaques are not present in the C57BL/6 control animal tissue. Measurements of the plaques within the APP/PS1 model indicate that MR microscopy observable plaques have a minimum diameter of approximately 40 µm, which is very similar to the human AD samples. Smaller plaques are also visible with Thioflavin-S staining throughout the neo-cortex (barrel fields, somatosensory, motor, and granular cortices), hippocampus, thalamic region, caudate and putamen of the animal brain but are not visible in the MR images. The most prominent difference between the human AD (Fig 3.5.1b and c) and APP/PS1 mouse tissues (Fig. 3.5.3b and c) involves the focal iron concentration found in the vicinity of the Aβ plaques. Large amounts of focal iron are found within and around the Thioflavin-S positive amyloid plaques in the human AD tissue while very little is seen in the APP/PS1 tissue. In the magnified images, the plaques indicated by the red arrows in figure 3.5.3b coincide with the white patches in figure 3.5.3c,
suggesting lower iron content in these areas. This observation seems to be in opposition to what has been demonstrated in the human AD tissues. No dark iron cored plaques were observed in the transgenic animal histological images compared to the human AD sections. This is also at seen at higher magnification, as illustrated at 100x in figure 3.5.5. Aβ plaques in human AD tissue appear to be small cored plaques surrounded by large diffuse coronas while the plaques in the APP/PS1 mouse tissue exhibit larger dense cores made of compact fibrillar Aβ protein deposition with a smaller diffuse corona region. The Alzheimer’s tissue samples demonstrate the localization of iron (Fig. 3.5.5b) within the amyloid plaque mass (Fig. 3.5.5a). Transgenic animal tissue shows a high concentration of large amyloid deposits while co-staining with the traditional Perl’s ferric iron stain does not show regions of high iron within the these plaques. Staining capillaries high in iron serves as a positive control for the effectiveness of the traditional Perl’s stain for detecting iron, indicated by arrows in figures 3.5.5c and 3.5.5d. Minor digestion of the Aβ plaque periphery in the APP/PS1 tissue and staining with the modified Perl’s technique are shown in figures 3.5.5e and 3.5.5f. There is visible evidence of trace amounts of iron in the transgenic animal plaques that was not stainable with the traditional method. The iron load within the APP/PS1 plaques is very minute compared to the human AD tissue. Staining the human AD tissue (not shown) with the modified Perl’s technique rendered the samples completely brown, which is indicative of iron throughout the tissue samples. There is a possibility that the protein digestion also exposed the ferrihydrite and magnetite core of the ferritin complexes in
these samples allowing the Prussian blue reaction to occur throughout the sample (196). This result rendered the microscopic detection of focal iron around Thioflavin-S plaques impractical for human AD sections stained in this manner. The iron stain of both APP/PS1 and control animal tissue does show appropriate tissue iron distribution as evident in the known high iron brain structures such as the substantia nigra, striatum and white matter tracks (197). High magnification 6000x and 46,000x transmission electron microscopy images of amyloid plaques from both APP/PS1 and human AD can be seen in figure 3.5.6a and b, respectively. Each plaque in figure 3.5.6a and b is similar in size, approximately 10 µm in diameter. The 6000x images include the whole plaque within the viewing plane and illustrate that Aβ plaques in the APP/PS1 model have a denser overall structure then the plaques found in human AD. Higher 46,000x magnification of the outlined regions shows fibrillar Aβ strands in both AD and APP/PS1 tissue and further demonstrates the difference in density between AD and APP/PS1 plaques.

For quantitative MRI studies, Fig. 3.5.7a graphically shows average relaxation values for regions of interest that included and excluded Aβ plaques within the AD sample and regions within the gray matter of the control sample. Average $R_2^*$ measurements for the AD tissue samples were $30.33 \pm 3.86$ (1/s) in regions with plaques and $16.10 \pm 2.92$ (1/s) in regions without plaques. Control tissue $R_2^*$ relaxation rate was $14.54 \pm 2.01$ (1/s). $R_2^*$ relaxation rates for the regions with plaques were significantly higher ($p<0.0001$) than both the controls.
and regions without plaques. Relaxation measurements in regions without plaques were not significantly different from control tissue. Average relaxation values for gray matter regions in the APP/PS1 mice that included or excluded Aβ plaques and control animals are seen in figure 3.5.7b. Average $R_2^*$ measurements for APP/PS1 were $24.84 \pm 3.67$ (1/s) in regions with plaques and $18.40 \pm 1.47$ (1/s) in regions without plaques. C57BL/6 controls had an average $R_2^*$ relaxation of $18.75 \pm 1.03$ (1/s). Regions with plaques in the APP/PS1 animals had a significantly higher $R_2^*$ relaxation compared to both control tissue and regions without plaques ($p<0.0001$). Measurements of $R_2^*$ relaxation were not significantly different between regions of interest in the transgenic animal devoid of plaques and control mouse tissue.
3.5 – Figures

Figure 3.5.1

$T_2^*$ weighted MR image (a) and histological images of Thioflavin-S stain for beta-amyloid plaques (b) and (c) Perl's iron stain of the same tissue section from the entorhinal cortex of an Alzheimer's disease subject. For detailed comparisons, images with 40x-magnification of four selected regions outlined with red boxes are shown on both sides of the corresponding main images. A large amount of black spots can be seen clearly in the gray matter in the MR image in (a). As seen in Fig 3.5.1b in the Thioflavin-S stain images, these black spots are shown either as bright green indicating Aβ-plaques (red arrows) or dark brown indicating small blood vessels (blue arrows). The Perl's stain of the brain tissue in (c) demonstrates focal high iron concentration in both Aβ-plaques and blood vessels. For the former, a higher iron deposition in Aβ has been previous demonstrated. For the latter, the elevated iron is likely associated with residual blood, ferritin/hemosiderin or magnetite. As seen in Fig. 3.5.1c, select hypo-intensities in the MR that are due to beta-amyloid plaques are seen with red arrows while signal dropout due to other focal iron regions are highlighted with blue arrows in the four concurrent image magnifications. The figure illustrates that hypo-intensities seen in the $T_2^*$ weighted image correlate to plaque location and focal iron concentrations. Iron deposition is present in the beta-amyloid plaques that are viewable in the MR image sets. Scale bars for the magnifications are 250µm and 1mm for the whole images.
Figure 3.5.2

Magnetic resonance image, thioflavin-S and iron stains in human control subjects. (a) An MGE $T_2^*$ weighted image, (b) thioflavin-S and (c) Perl’s iron stain of the same 60µm thick tissue section from the entorhinal cortex of an Alzheimer’s disease subject. The $T_2^*$ image shows hypo-intensities that are due to micro pockets of air or coil inhomogeneities. The thioflavin-S stain illustrates a lack of beta-amyloid plaques in the human control subjects. Perl’s staining shows high iron in white matter tracks due to oligodendrocytes and one focal iron region in gray matter that is highlighted with the blue arrow in the three images and can be seen as a signal drop in the MR image. Scale bars in all images are set at 1mm.
Figure 3.5.3

Co-registration of (a) magnetic resonance images, (b) beta-amyloid and (c) iron stains in the APP/PS1 mouse model. (a) An MGE $T_2^*$ weighted image of a 60µm slice from a APP/PS1 mouse brain approximately -2.92mm Bregma. Final in-plane image resolution was 45µm x 45 µm. Image magnifications (40x) of the selected regions of interest within the left and right piriform cortex are seen below. Hypo-intensities are noted by red arrows. (b) Thioflavin-S fluorescent mosaic image of the same tissue section in 1a. Beta-amyloid staining is evident and can be seen as the bright green positions. The same regions as in 1a have been magnified with the arrows pointing toward the plaques responsible for the $T_2^*$ weighted hypo-intensities. (c) Perl's iron stain with the same magnifications and regions of interest arrows as in 3.5.1a and 3.5.2a. The figure illustrates that the hypo-intensities seen in the $T_2^*$ weighted image are in the same region as large beta-amyloid plaques approximately 50 – 60 µm in diameter. Iron deposition is not present at the plaque locations, as they seem to be regions of low iron concentration compared to the surrounding gray matter tissue. Scales bars for the magnifications are 250 µm and 1mm for the whole image.
Figure 3.5.4

Magnetic resonance image, thioflavin-S and iron stains from a control C57BL/6 mouse. (a) An MGE T₂* weighted image, (b) thioflavin-S, and (c) Perl's iron stain of the same 60 µm thick section of tissue from a C57BL/6 control mouse at approximately -2.80mm Bregma. The T₂* weighted image shows hypo-intensities and iron staining at regions of known high iron concentration such as the substantia nigra, white matter tracks and the caudate/putamen. Thioflavin-S staining reveals only non-specific background staining with no beta-amyloid plaques in the control animals. There are no MR hypo-intensities that are associated with positive thioflavin-S staining. Scale bars in all images are 1mm.
Figure 3.5.5

High magnification (100x) images of beta-amyloid plaques in both human Alzheimer’s entorhinal cortex (a,b) and APP/PS1 animals piriform cortex (c,d,e,f). The thioflavin-S stain (a) and traditional Perl’s stain (b) illustrate a close relationship between beta-amyloid plaques and focal iron deposition in Alzheimer’s disease. This relationship between plaques (c) and iron (d) is not clearly seen in the APP/PS1 animals with traditional Perl’s staining. The arrow in (c) and (d) illustrate iron deposition in a capillary of the APP/PS1 animal demonstrating the positive stain for iron. Staining with a modified Perl’s technique (e,f) via degradation of the plaques’ periphery proteins allows the aqueous stain to penetrate the plaques more readily. There is an indication of minute amounts of iron in the transgenic animal plaques that is not perceivable with the traditional Perl’s stain. Differences in plaque morphology between the AD and APP/PS1 animals is evident in images (a) and (c,e). The human AD plaques have a dense core of fibrillar amyloid protein with a halo of amyloid protein around them that is less susceptible to thioflavin-S staining. APP/PS1 Aβ plaques show a larger and denser thioflavin-S positive core with a smaller halo region around them. The Perl’s stains indicate that high concentrations of iron found throughout the human AD plaques that associated with the amyloid protein or within cells such as microglia inside the space the plaque has occupied. Compared to the human AD plaques, the APP/PS1 images show a reduction in focal iron within the plaques that is diffusely found throughout the plaque. Scale bars for all images are 100µm.
Figure 3.5.6

Transmission electron microscope images of plaques in (a) APP/PS1 and (b) human AD tissue. The 6000x magnification images of the plaques include the whole plaque within the imaging plane. The plaque diameter for both APP/PS1 and AD tissue is approximately the same, at 10 µm. The lower magnification image illustrates the denser overall structure of the beta-amyloid plaques in the APP/PS1 compared to the human AD tissue. The human AD plaques have numerous gaps present between the fibril bundles throughout them that are rarely found in the transgenic mouse plaques. 46,000x magnification of the outlined regions reveal differences in the fibrillar orientation of the beta-amyloid strands in the APP/PS1 and AD plaques. Magnification of the outlined regions also illustrates the denser nature of the APP/PS1 plaques while the human AD tissue has gaps between amyloid strand clusters.
**Figure 3.5.7**

Bar graphs of average $R_2^*$ rates from ROI’s with plaques, without plaques and control tissue within human (a) and mouse tissue (b). The $R_2^*$ relaxation rate of plaque ROI’s in the AD tissue is significantly higher than both regions without plaques and control tissue sections. A similar trend is found in mouse data. There is also a higher $R_2^*$ relaxation rate for the plaque ROI’s in AD than in the APP/PS1 mouse, hypothesized to be due to higher iron in the AD plaques as shown with the Perl’s stain in Fig. 3.5.5 (b,d and e).
3.6 – Discussion

Understanding the relationship between beta-amyloid plaque pathology in AD and the associated MRI contrast is fundamentally important for MRI study of AD pathology. This work demonstrates the ability to image Aβ plaques in histology sections of both human AD and amyloid-generating APP/PS1 transgenic mice brains. The histological MR images taken at a field strength of 7.0 T clearly show high resolution details of Aβ plaques in human and animal samples. Previous MRI studies of Aβ plaques have been carried out with blocks of human AD brain tissue samples (183). With this approach, accurately matching histological sections of the whole tissue block with the MRI image slice is difficult. Using the histological RF coil, MRI and histological analyses were performed on the same tissue sample so that tissue pathology and associated MRI contrast changes could be precisely compared. This capability allowed us to establish a one-to-one relationship between MRI parameters and histology data acquired from human AD and APP/PS1 transgenic mouse tissue samples. Such a relationship is important for the interpretation of MRI findings in the AD brain and its animal models. Additionally, the information acquired with this technology is valuable in the development of potential therapies for AD research and investigations of other pathologies.

The MR imaging of amyloid plaques in transgenic APP/PS1 mice has been pioneered by several groups using both in vivo and ex vivo imaging methodologies (186,193,194). The hypo-intensities in the MR images of
APP/PS1 animal tissue that coincide with the plaques are similar in signal dropout and size compared to MR images of human AD plaques (Figs. 3.5.1 and 3.5.3). The cause for the hypo-intensities seen in T2* and T2 weighted images of APP/PS1 mice thus has been believed to be caused by high iron accumulation in the plaques (186,193,194). However, when viewing the iron and Aβ stained images in Figs. 3.5.3 and 3.5.5, the iron concentration is considerably reduced in the APP/PS1 Aβ plaques compared to the human AD tissue in Fig. 3.5.1c, while Aβ-plaques in APP/PS1 and human AD samples are perceptibly the same when viewed with MR images. To confirm that the staining methods did not cause aberrant interactions with one another, independent iron and amyloid stains were performed yielding the same appearance as co-staining procedures. To understand this unexpected result, notice that APP/PS1 plaques in Figs. 3.5.3b and 5 show an extremely dense nature, which suggested that the current traditional Perl's staining method might not be able to stain the iron inside these plaques. The hydrophobic nature of the plaques coupled with the acidic aqueous based Perl's stain could prevent the Prussian blue reaction from occurring if the stain cannot infiltrate into the plaque. A modified Perl's stain (188) was tested that utilizes protein degradation and detergents to break down the outer surface of the plaques to allow better peripheral penetration of the aqueous stain. With the modified stain, the images in Fig. 3.5.5f indeed exposed minute focal iron distributions within the APP/PS1 transgenic animal plaques, but much less than that in the human AD plaques. As a control condition, application of the modified stain to human AD tissues resulted in dark brown staining throughout the entire
tissue sample (not shown). This could be due to staining of not only the free labile iron pool but additionally iron bound inside ferritin cores that can be stained only when the proteins are digested open with the modified staining procedure. The above observations suggested that iron in the animal plaques is appreciably less than that in the human plaques and the plaques in the animals are much denser than those in human samples. Since MR images of the plaques in both human and animals appeared essentially the same, the differing characteristics of the animal plaques raised a fundamental question as to the cause of the MR image contrast associated with the animal plaques.

It is well known that $T_2$ and $T_2^*$ relaxation and contrast can be altered in a variety of manners. Iron bound within the ferrihydrite-like core of ferritin, hemosiderin and an assorted number of iron oxide mineral deposits are found in sufficient quantities in the brain to alter MRI contrast (132,133,136,198,199). As a ferromagnetic ion, ferric iron ($Fe^{+3}$) is known to cause a shortening of $T_2$ relaxation due to the creation of microscopic magnetic field inhomogeneities that de-phase the signal of water protons in the vicinity (123,137,197). Thus, ferric iron in brain tissue acts as a natural contrast agent causing faster proton $T_2$ and $T_2^*$ relaxation. There is some evidence to support that small amounts of ferrous iron ($Fe^{+2}$) is also stained with the Perl’s stain reaction (200). The effect of ferrous iron on $T_2^*$ and $T_2$ relaxation is much less pronounced than that caused by the ferric form (124) (our laboratory’s unpublished data). Thus, the signal hypo-intensities seen in the MR images associated with focal iron regions is likely
dominated by the ferric form. One other important factor associated with the MRI appearance of plaques is their size under given experimental conditions (pixel resolution, TE/TR times, etc.). The minimal plaque size viewable in our MR images of the APP/PS1 mice was similar to the AD tissue images, about 40 µm and 38 µm, respectively. Thus, due to the similarity in the plaque size for both species, plaque diameter alone does not account for the hypo-intensities seen in the APP/PS1 data. This is also supported by the fact that human AD plaques of a similar diameter with low observable iron are marginally visible on the MR images, while APP/PS1 plaques with considerably less iron association are clearly discernable on the MR images. Magnification of the thioflavin-S positive plaques (100x seen in figure 3.5.5 and 6000x & 46000x in figure 3.5.6) shows a distinct difference in plaque morphology between AD and APP/PS1 mice. Differences between human AD and the APP/PS1 Aβ plaques are most noticeable in the TEM magnifications. The human AD plaques consist of fragmented patches with random fibrillar orientation in the amyloid core while the structure of transgenic mouse plaques appears to be a highly packed aggregation of long and oriented fibrils. The decreased density of the amyloid core creates gaps that are prevalent in the human AD plaques while rarely found in the transgenic mouse plaques. The APP/PS1 mice Thioflavin-S data illustrate that, while the plaques are a similar size to those in AD tissue, the plaques have a larger center core of dense compact amyloid. In previous studies, comparison of the structure of the amyloid fibril in the transgenic APP/PS1 model and human AD is subtly different due to post-translational modifications leading to alterations.
in the Aβ molecule (109,111,201). Plaques found in the APP/PS1 mouse model have a variable distribution of both human and mouse amyloid in the plaques. Diffuse plaques have an intermingled distribution of both human and mouse amyloid while dense plaques are composed of human amyloid cores surrounded by mouse amyloid (202). Thioflavin-S is known to bind to fibrillar but not to diffuse Aβ deposits (203,204). The difference in Thioflavin-S staining between human AD and APP/PS1 mice could be due to the different primary fibrillar or diffuse composition of the plaques. Many of the amino-acid residue positions within the Aβ protein are hydrophobic and are sufficient for amyloidal fibril formation (33,205). With the additional two hydrophobic amino acids found in the Aβ42, it aggregates more readily than Aβ40 (206). These views suggest that the hydrophobic regions are at least partially responsible for the fibril formation (34). The morphology of the mouse plaques with their large globular nature and the dense center core seem to be responsible for the T2* relaxation associated with these plaques. Unlike smaller proteins, the amyloid mass found in the animal plaques is rigid and fixed in the tissue resulting in a situation where the protein mass would behave similar to a polymer-like solid. This creates a number of consequences that may affect relaxation in the plaques. First, it sets up an environment where a large amount of available surface area is present inside the plaque mass along the component Aβ fibrils. Water molecules then become bound to the hydrophilic regions of the fibril axis while repelled by the hydrophobic regions. Water in the immediate vicinity of the hydrophilic regions is bound via hydrogen bonding such that it is rotationally or irrotationally bound.
Layers of water are affected as they diffuse near the bound water. Cross-relaxation between the protons of bound water and the protein molecules by proton-proton magnetization exchange could lead to a more rapid relaxation of water proton. Protein wobbling due to random tumbling and thermal vibrations also must be considered which would cause altered proton-proton magnetization exchange to occur. As such, the relaxation of bound water on the Aβ fibril chains would act as a relaxation sink for water moving in the surrounding area (207,208). As a result, $T_2$ relaxation shortening occurs. Secondly, the magnetic susceptibility differences between the highly compact Aβ protein mass and surrounding issues could induce static dephasing in addition to that caused by deposited iron. Thirdly, tissue hydration could also be an important contributing factor for plaque contrast. The dense formation of the animal plaques as indicated in the TEM image limits free water from accessing their core. As shown by histology results, staining iron in the plaques in the APP/PS1 mouse was difficult following standard methodology without protein degradation treatment. These experimental data suggest that access of free water molecules to the animal plaque is very limited. For human samples, the gaps seen between the Aβ patches (Fig. 3.5.6b) would accommodate more free water pools. The transverse relaxation of these water molecules is shortened by the magnetic inhomogeneities caused by ferromagnetic iron in the plaques. Thus, with these observations we would suggest a dual relaxation mechanism for the generation Aβ plaques $T_2^*$ contrast associated with both the compact fibrillar protein mass and iron deposition. The focal iron concentration could play a dominant role for
rapid $T_2^*$ relaxation in human AD plaques while the increased fibrillar density and compacted morphology would be more likely a major factor for $T_2^*$ shortening in the APP/PS1 transgenic plaques. Although the potential relaxation mechanisms in Aβ plaques discussed above are preliminary and speculative, it is important to realize that alternative relaxation mechanisms must be considered. To thoroughly develop and validate the model of relaxation in Aβ plaques, more experimental studies are required in future investigations. In addition, caution should be used when comparing *ex vivo* $R_2^*$ contrast obtained in the fixed tissue samples and extrapolating this to *in vivo* conditions (187).

The quantitative comparison of the $R_2^*$ maps from different tissue types supports our hypothesis in regard to the relaxation mechanism associated with Aβ plaques. The mean $R_2^*$ for the selected amyloid plaques in the AD tissues (Fig. 3.5.7a) is on average 90% greater than that from the ROI’s without plaques and control tissue. Regions in AD gray matter with no Aβ plaques and in control tissue samples (Fig. 3.5.7a) have no statistical difference in relaxation time. However, there is a trend of faster $R_2^*$ relaxation rates in regions without plaques throughout the Alzheimer’s tissue compared to control tissue samples. This is indicative of a higher overall iron concentration in the AD neural tissue compared to the control tissue, possibly due to a systemic inability to regulate iron properly within the brain. Iron staining of the tissue samples supports this, which shows an overall increase in high focal iron in the AD tissue samples that is not seen in the control tissue. In the mouse tissue samples (Fig. 3.5.7b), the mean $R_2^*$ for
the selected amyloid plaques is on average 56% higher than that in the ROI's without plaques and in the control tissue. The mean $R_2^*$ of regions without plaques and of control tissue are nearly identical, indicative of similar tissue fixation across tissue sub-types and consistent preparation of tissue samples. This information alleviates the confounding possibility that separate tissue handling procedures are the source of the observed MR contrast differences.

Comparison of $R_2^*$ in ROI's with plaques in both human and APP/PS1 animals tissues reveal that the human plaques have a significantly higher $R_2^*$ value. It is conceivable that human plaques have a duality of relaxation where both high focal iron concentration and fibrillar Aβ masses cause rapid proton transverse magnetization decay. APP/PS1 plaques exhibiting slower decay due to the reduced concentration of iron is a reasonable explanation for this result. The $R_2^*$ for normal human tissues also appeared to be higher than that of animal tissue. This is consistent with the overall higher iron content in human neural tissue compared to the APP/PS1 animal samples. Iron staining for regions without plaques in the gray matter of APP/PS1 tissue and mouse control tissue samples exhibit similar patterns of iron staining.

These findings generate several important questions regarding the difference between Aβ plaques found naturally occurring in the human Alzheimer’s brain and transgenically introduced into the APP/PS1 animal model. Previous studies have revealed that Aβ-plaques in APP/PS1 animal show numerous differences from human AD plaques, including plaque morphology and
composition (109,209). The difference in morphology and the reduction of iron staining in the plaques raises the question for translating the APP/PS1 model’s data to human AD as it is hypothesized that there is a link between Aβ plaques and elevated iron in the brain tissue as the cause of oxidative stress leading to cellular neurotoxicity (210,211). The relationship of iron to plaque generation is not currently well understood within the literature and is a subject of much investigation. An imbalance in the iron regulatory system and a dysfunctional difference in mineralization of iron within the ferritin core in AD subjects has been previously demonstrated (78,175). It is largely unknown whether the initial generation of the plaque causes the aggregation of iron or if aberrant regulation of extra- or intra-cellular iron stores cause the formation of the Aβ protein around it. There is general agreement that the location of Aβ plaques in human AD brain tissue coincides with focal iron deposits. Iron associated with plaques can accumulate from a number of potential sources. As a metalloprotein, free iron will bind to Aβ fibrils and collect within the plaques. Iron from ferritin, and its degradation to hemosiderin, can originate from nearby neurons and microglial cells that have migrated to the Aβ plaques and can become part of the plaque mass (77,212,213). Biogenic magnetite found proportionally higher in AD ferritin cores than normal aged human neural tissue is known to accumulate within Aβ plaques (175,190). While it is unknown precisely how Aβ plaque aggregation occurs, it is known that metal ions do play a role in Aβ fibril formation. The conversion of Aβ’s secondary protein structure to a β-sheet has been shown to occur in the presence of divalent metal ions, including ferrous iron (82). The
component fibrils of the Aβ plaques are known metalloproteins and possess metal chelation regions in their amino-acid sequence (79). The MR and histological images in Fig. 3.5.1 demonstrate that Aβ plaques are consistently associated with elevated iron load in and around their vicinity in human AD brain tissue. Research has shown that the co-localization of iron with the human Aβ plaques is accompanied by endoplasmic reticulum stress induced apoptosis, DNA oxidation and cellular damage in cells adjacent to plaques. Conversely, in regions where Aβ plaques accumulate alone without iron there is no indication of oxidative stress or apoptosis pathway activation (214). It has been demonstrated that the toxicity of Aβ is amplified upon the direct interaction of iron ions with the Aβ peptide, while unbound iron itself has no effect upon toxicity (215). This strongly suggests that Aβ plaques by themselves are not toxic to cells adjacent to plaques, whereas iron accumulation in and around plaques is essential for neuronal damage. The morphology and associated iron dissimilarity between the human AD and transgenic APP/PS1 plaques raises a question regarding differing amyloidal genesis processes for each species. In the APP/PSI mouse model used here, plaques are produced in the brain as early as six months after birth and are believed to continually grow throughout the life span of the animal (216). Current understanding of human amyloidal genesis suggests that plaque generation occurs over years, perhaps decades, and places the plaques into discrete classes based upon the morphology of the plaque and surrounding tissue. It is hypothesized that plaques start out as a seed formations of Aβ peptide and eventually become much more compact and dense over time (217).
The APP/PS1 model generates plaques at a rapid pace and iron staining indicates a reduced amount of iron associated with these plaques compared to AD tissue. It is plausible that the rapid fibrillarization from Aβ oligomers into compacted fibrillar plaques could lead to less of iron deposition within the plaques in the APP/PS1 model.

In summary, imaging of thin slices of tissue samples with the aid of the histological coil allows the one-to-one comparison of tissue pathology as seen in histological stains and different MRI methodologies. With this technique, the relaxation mechanisms for Aβ plaques in human Alzheimer’s disease and the APP/PS1 mouse model were investigated with respect to their morphology and relationship with iron deposition. The histology stains of AD and APP/PS1 tissue samples were directly compared to T2*-weighted images and R2* parametric maps. The histological iron stains presented here support the hypothesis that iron associated with the Aβ plaques in the human AD samples plays a major role in generation of MRI T2* contrast. Histological data from previous studies using post mortem human AD tissue (67,77,192,218) also support the notion that focal iron load in Aβ plaques is the dominant cause for faster T2 and T2* decay. The electron microscopy and histology data revealed that there are important differences in plaque morphology and associated iron concentration between transgenic APP/PS1 mice and human AD tissue samples. In the Alzheimer’s tissue, beta-amyloid plaques with high iron concentration are clearly visible in the T2*-weighted images while others of similar size with less focal iron are not as
discernable. In the APP/PS1 animal tissue, large plaques are equally observable as human plaques in MR imaging while iron load is considerably less than human plaques. This suggests that there is a degree of difference between the amyloid plaques in the APP/PS1 mouse model and human Alzheimer’s, in respect to their morphology and relationship with \textit{in vivo} iron stores. The increased transverse proton relaxation rate in A\textbeta{} plaques in animals is likely caused mainly by the interactions of water with the highly compacted amyloid fibril mass. The improved resolution of the \textit{ex vivo} data set allows for the detailed comparison between tissue histology and MR contrast that is not possible with current lower resolution clinically based protocols. The extrapolation of microscopic MR images to \textit{in vivo} applications is promising in the near future considering the rapid increase in achievable image resolution with the development of higher magnetic fields and the usage of parallel imaging technology (219). The data presented in this report are essential for understanding the histo-pathological underpinning of MRI measurement of A\textbeta{} plaques in humans and animal models for both current and future MR applications.

3.7 – Appendixes

3.7.1 – Correlating Plaque Size and Iron Load with $R_2^*$ relaxation

It was hypothesized that the hypo-intensities resulting from the beta-amyloid plaques in the Alzheimer’s and APP/PS1 mice would correlate to plaque diameter (in the case of both AD and APP/PS1 tissue) and/or iron loading (in the
case of AD tissue). Detailed evaluation of plaque diameter and the optical density of the Perl’s and thioflavin-S stains was undertaken to evaluate if there was any statistical correlation. Eight-echo MGE images of human AD tissue samples were imported into in-house software developed with the interactive data language (IDL) 6.1 framework (Research Systems Inc., ITT Industries, Boulder, CO) and regions of interest were selected inside the plaque to yield detailed $R_2^*$ ($1/T_2^*$) relaxation rate and magnitude measurements of individual plaques. The same plaques were selected on bright field Perl’s iron stain and thioflavin-S 40x magnification images with Image J (National Institutes of Health, Bethesda, MD) to measure Aβ plaque diameter and optical density. Calibrated optical density is measured in Image J as the average gray scale contrast on a 0 – 255 scale for selected regions of interest. An optical density for plaques was calculated by comparing the plaque optical density to the optical density of the background of the sample. This was done to standardize optical density measurements across tissue samples and stains. The relaxation rate $R_2^*$ or magnitude was then graphed against either iron or thioflavin-s optical density or against plaque diameter to compare their correlation.
Figure 3.7.1.1

$R_2^*$ relaxation verses $A\beta$ plaque diameter.
$y = -0.0043x + 30.68$
$R^2 = 0.0005$
**Figure 3.7.1.2**

$R_2^*$ relaxation versus iron optical density ratio.
$y = 9.4305x + 22.274$  
$R^2 = 0.0109$
Figure 3.7.1.3

$R_2^*$ relaxation verses thioflavin-S optical density ratio.
$y = -0.0332x + 30.377$

$R^2 = 3E-06$
Figure 3.7.1.4

Magnitude verses Aβ plaque diameter.
$y = -55.365x + 29068$

$R^2 = 0.053$
Figure 3.7.1.5

Magnitude verses iron optical density ratio.
$y = 26445x + 1980.6$

$R^2 = 0.0608$
Figure 3.7.1.6

Magnitude verses thioflavin-S optical density ratio.
The equation of the line is:

\[ y = -7542x + 35723 \]

with a coefficient of determination \( R^2 = 0.1188 \).
The data sets and scatter plots indicate that there is no significance for any of the correlation data between $R^2$ and magnitude versus plaque diameter, iron optical density or thioflavin-S optical density (Figures 3.7.1.1 – 3.7.1.6). This is theorized to be due to a number of factors. Measurements of optical density are extremely hard to standardize across tissue stains. It was attempted to standardize the optical density by creating a ratio using the background of the stain. The inherent problem with this is that there is potential for different stains over different periods of staining to have slight differences in the optical density. All stains were standardized according to length of the procedures and concentrations of the chemicals used but there is still intrinsic variability in the stains, especially the Perl's iron stain. The Perl's stain relies upon the usage of 3,3'-Diaminobenzidine (DAB) as a chromogen which is oxidized by hydrogen peroxide in the presence of iron to give a brown precipitate. This process is highly variable even with strict time constraints implemented into the staining procedure. The thioflavin-S stain generally is very consistent even with differing timing of the staining period. There is a stage during staining that can alter the outcome of the thioflavin-S brightness when 70 percent ethanol is used to wash the wash out the residual background thioflavin-S that is not bound to beta-amyloid. However, the thioflavin-S that is bound to the beta-amyloid has an extremely high affinity for intercalation into the peptide and ethanol washing has a minimal effect upon it. There is a possibility that the thioflavin-S stain lost some florescence overtime and this affected the results. However, the microscopic images of the tissue samples were taken as soon as possible to try and minimize
this effect. There is the possibility that because the tissue samples were dual stained for thioflavin-S and iron that the Perl's stain is the main culprit of the lack of correlation because even under fluorescent light the dark precipitate in the Perl's stain is still slightly visible.

It is unfortunate that there were no statistically significant results to report for this evaluation. When viewing the hypo-intensities in the magnetic resonance images and comparing these to the amount of iron present at those points, it is evident that regions that stain for more focal iron have larger diameters of signal dropout in the MR images. The hypo-intensities are essentially regions devoid of contrast, and relaxation in the middle of these plaques was hoped to correlate to plaque or focal iron size statistically. Perhaps measurement of the diameter of the hypo-intensities verses iron content and plaque size would be a better comparison. A limitation in the resolution of the MR data sets then becomes an ever increasing factor when attempting to measure the diameter of the microscopic hypo-intensities.
Chapter 4

Morphological and Histochemical Differences
between Human and Mouse plaques

4.1 – Histological and Cellular Differences Between Beta-Amyloid Plaques
in APP/PS1 Transgenic Mice and Human Alzheimer’s Disease

4.1.1 – Abstract

Transgenic mouse models that mimic neuropathology are important in understanding disease etiology and testing new treatment procedures. Mutations in human genetic sequences for the amyloid precursor protein and presenilin 1 have been introduced into mouse models that develop beta-amyloid (Aβ) plaques in a considerably condensed time period compared to Alzheimer’s pathology. The benefit of producing plaques in less than nine months motivates the utilization of these animals for a variety of potential studies such as beta-amyloid clearance, protein processing pathways, and amyloidogenesis. As with any animal model, there exist drawbacks when comparing them to their model system. Understanding the amyloidogenic and morphological characteristics of the APP/PS1 mouse model are important in considering the outcome from studies involving their usage and eventually translating results for treatment of the Alzheimer’s disease brain. Using various histological stains and microscopy aimed at examining plaque morphology, iron distribution and inflammatory response we show that there are distinct differences between the naturally
occurring human Alzheimer’s plaques and the APP/PS1 mouse model. Aβ antibody and thioflavin staining along with electron microscopy demonstrate distinct morphological differences in plaque composition and ultra-structure alluding to differences in plaque genesis and component fibril formation. Microglia and astrocyte staining shows a differing inflammatory response pattern with more pronounced response in the Alzheimer’s tissue. Iron mismanagement and focal iron deposition is found ubiquitously throughout the Alzheimer’s tissue, with plaques showing a close relationship to iron accumulation within the plaques that is not seen in the transgenic animal model. The data presented here emphasizes distinct differences between naturally occurring Aβ plaques in the AD brain and those instituted to produce in the transgenic mouse model. Utilization of these animal models for AD research should be done with this caveat.

4.1.2 – Introduction

The extracellular accumulation of beta-amyloid (Aβ) protein, leading to the formation of amyloid plaques are major defining neuropathological characteristic of Alzheimer’s disease (AD). Genetic screening of individuals with familial Alzheimer’s disease has identified reoccurring alterations in genes believed to be involved in the disease process. The augmented production of Aβ peptides is believed to be integral to the disease process according to the amyloid cascade hypothesis (220). Central to this hypothesis is the knowledge that genetic mutations associated with familial AD cause an increase in aberrant APP
production leading to increased Aβ production. This has lead to the development of transgenic mouse models that produce Alzheimer’s like beta-amyloid plaques in the brain. In particular, animals that develop amyloid plaques age dependently are particularly important in understanding disease etiology and testing new therapies. Several transgenic mouse lines have been fashioned that over-express known human Alzheimer’s mutations and develop fibrillar beta-amyloid plaques (113,221-224).

Transgenic mice that harbor chimeric mouse/human familial Alzheimer’s genes for amyloid precursor protein (APPswe695, K595N and M596L mutations) and a mutant human presenilin 1 (PS1-A246E) under control of the mouse prion promoter develop fibrillar beta-amyloid plaques at approximately nine months of age and continually do so progressively throughout their lifespan (108,112). This model reproduces some of the neuropathological traits of AD, most notably amyloid plaques that exhibit thioflavin-S fluorescence. The exact role of amyloid deposition in Alzheimer’s disease has not been elucidated but it is clear that they are involved in the disease process as they are found ubiquitously throughout the AD brain. A transgenic mouse model that reproduces and encompasses all characteristics of the Alzheimer’s disease pathology has not been created. Although there are known differences in pathology between the various transgenic models and naturally occurring AD tissue, many of these mouse models have been used to study the creation and clearance of plaques with pharmacological or biochemical interventions. Care must be taken when comparing results from the transgenic model and translating these to AD neural
tissue. The assumption that the successful alleviation of plaque burden in the transgenic brain with various methods can be transferred to the AD brain should not be casually accepted without any doubt or reservation. Histological investigation of the similarities and differences between Aβ plaques in the human AD and APP/PS1 brain is the first and essential step in understanding how best to use these models for future human studies. In this report, we have undertaken various histological staining techniques to compare the transgenic plaques to human AD plaques to examine morphology and histological inflammatory response.

4.1.3 – Materials and Methods

Human Alzheimer’s and Control Brain Samples:

Entorhinal cortex brain tissue samples from clinically and histologically determined AD subjects (n=5) and age-matched controls (n=4) were donated with consent following institutional guidelines and obtained from both internal and external sources (Harvard Brain Tissue Resource Center, McLean Hospital, Belmont, MA). Analysis of the tissue obtained from the brain bank indicated that AD tissue samples were highly positive for beta-amyloid and neural fibrillary tangles staining consistent with a post mortem diagnosis of Braak stage VI (43,44). There was not a significant difference between the age of the subjects upon bereavement while the post mortem index between the tissue harvesting at time of death and utilization in this study was slightly different between the tissue samples (longer time period for controls). The tissue was fully fixed in 4%
paraformaldehyde in pH 7.3 phosphate buffered saline (PBS) after which blocks of tissue approximately 1 cm x 1 cm x 0.5 cm were dissected from the entorhinal cortex. For cryogenic protection the samples were placed in graded 10% and 20% sucrose solutions for 48 hours each, and then placed in 30% sucrose. Tissue sections were cut at 16µm on a cryostat, mounted on poly-lysine and gelatine coated slides and prepared for histological staining according to individual protocols described below.

Transgenic APP/PS1 and Control Mice:

Transgenic mice (n=5) inserted with a chimeric mouse/human amyloid precursor protein (APP) and a mutant human presenilin 1, developed by Borchelt et al. (108,112), were obtained commercially from The Jackson Laboratory (strain type B6C3-Tg(APP695)3Dbo Tg(PSEN1)5Dbo/J). The expression plasmid for these animals is composed of cDNA harboring human familial Alzheimer’s disease (FAD) mutations regulated by the mouse prion promoter (MoPrP.Xho). Transgenic material introduced consists of human mutations found in presenilin 1 (PS1-A246E) and the Swedish variant of FAD (APPSwe695, K595N and M596L mutations). Animals were kept in the animal facility under veterinary care with normal feeding and handling conditions. Age-matched cage mate C57BL/6 mice (n=3) were used as a control group. Upon aging naturally until 24 months, animals were euthanized via an intra-parietal injection of sodium pentobarbital (Nembutal, 200 mg/kg) and were transcardially perfused with cold Lactated Ringer's solution (pH 7.4) followed by buffered 4% paraformaldehyde in
PBS. Whole brain tissue was harvested and placed overnight in pH 7.3 buffered 4% paraformaldehyde to allow full tissue fixation. Graded sucrose solutions of 10%, 20% and 30% were used for cryo-protection. Tissue sections were cut at 16µm on a cryostat, mounted on poly-lysine and gelatin double coated slides and prepared for histological staining according to individual protocols described below.

*Iron and Amyloid Staining:*

Tissue sections were co-stained for both iron and beta-amyloid with a diaminobenzidine enhanced Perl’s Prussian blue stain followed by an aqueous Thioflavin-S stain. Sixteen µm tissue sections mounted on lysine and gelatin coated slides were heated in an oven at 50°C to better adhere the samples to the slides then rinsed in deionized water (dH₂O) for 15 minutes. The sections were placed in equal volumes of freshly prepared 4% potassium ferrocyanide (#P236, Fisher Scientific, Waltham, MA) and 4% hydrochloric acid (final combined concentrations 2% for each) for 30 minutes under gentle rocking followed by two rinses in dH₂O. Intensification of the iron stain was performed with 5 minutes of 3,3′-diaminobenzidine tetrahydrochloride (DAB) staining (#D5637, Sigma, St. Louis, MO) (10 mg dissolved in 15ml of PBS with 16 µl of 30% H₂O₂) followed by two five minute rinses in dH₂O. Tissue samples were then placed in 1% thioflavin-S (T1892, Sigma, St. Louis, MO) aqueous solution for 5 minutes, followed by differentiation in 70% ethanol for 5 minutes and two five-minute washes in dH₂O. To preserve fluorescence, sections were covered with
FluorSave Aqueous mounting media and cover slipped (Catalog #345789, Calbiochem, San Diego, CA). To test for the possibility of confounding interactions between the two staining methods during co-staining, each was tested on separate tissue samples to determine their individual efficacy and compared to the co-stained sections. A modified Perl’s stain further tested the transgenic animal tissue for minute amounts of iron within the amyloid masses. The stain was adapted and modified from LeVine (188,189) and employs the usage of proteinase K and detergents to break down the periphery of the plaques forming openings in the highly hydrophobic amyloid plaques. This allows the aqueous Perl’s stain to infiltrate into the amyloid mass and is sensitive enough to detect trace amounts of iron in the Aβ plaques. The staining method consists of the following procedure. Tissue sections were hydrated in PBS for 15 minutes followed by immersion in sodium borohydride (10 mg/ml PBS, # 213462, Sigma, St. Louis, MO) for 30 minutes. They were then rinsed in PBS twice for five minutes and immersed in proteinase K (30 µg/ml, # P6556, Sigma, St. Louis, MO) and 0.01% Triton X-100 in PBS for 20 minutes. Next, the sections were placed in a solution of 1% hydrochloric acid, 1% potassium ferrocyanide and 1% Triton X-100 in distilled water for 30 minutes. Amplification of the iron staining was accomplished with 0.5 mg/ml 3,3'-diaminobenzidine tetrahydrochloride (DAB) and 2µl/ml of 30% hydrogen peroxide in pH 7.6 0.05 M Tris HCl for 15 minutes. The sections were rinsed twice for 5 minutes in dH₂O, stained with thioflavin-S aqueous solution for 5 minutes, differentiated in 70% ethanol for 5
minutes and then underwent two final five-minute dH₂O rinses before being mounted and cover slipped.

Immunohistochemistry Staining:

Slides with tissue samples on them were air dried at 50°C for 30 minutes then placed in 95°C Citrate Buffer (10mM sodium citrate tribasic dihydrate (#C7254, Sigma, St. Louis, MO), 0.05% Tween 20 in dH₂O, pH 6.0) for 15 minutes. Slides were rinsed three times with PBS for five minutes each then a water repellent circle was drawn around each section with a hydrophobic barrier pap pen to localize staining reagents. Nonspecific protein binding was blocked by 30 minute incubation with 1% Bovine serum albumin (BSA) (#A9430, Sigma, St. Louis, MO) in 1x PBS Tween (PBST)(2ml Tween 20 in 1litre PBS pH 7.3). Sections were incubated with primary antibodies at their respective dilution factors in 1% BSA in 1x PBST overnight in an airtight incubation-humidity chamber. After primary incubation, the antibody was decanted and slides rinsed in PBS three times for five minutes each. Sections were incubated with fluorescent secondary antibodies (Alexa Fluor, Invitrogen, Carlsbad, CA) in 1% BSA in 1x PBST for two hours in a humidity chamber. Following this the secondary was decanted and slides rinsed in PBS three times for five minutes each followed by mounting with FluorSave and cover-slippping.

For beta-amyloid (1-40) staining, a mouse monoclonal beta Amyloid 1-40 antibody [BAM-10] (1/100, AB7501, Abcam Inc., Cambridge, MA) was used followed by an anti-mouse red fluorescent Alexa Fluor 555 IgG (A-21422)
secondary antibody. Following secondary incubation and PBS rinsing tissue sections were co-stained with 1% thioflavin S for 10 minutes, differentiated in 70% ethanol, washed twice in dH₂O, mounted with FluorSave and cover-slipped.

For beta-amyloid (1-42) staining, a rabbit polyclonal beta Amyloid 1-42 antibody (1/200, AB10148, Abcam Inc., Cambridge, MA) was used followed by an anti-rabbit red fluorescent Alexa Fluor 555 IgG (A-21428) secondary antibody. Tissue sections were then co-stained with thioflavin S according to the method outlined above.

To stain for ferritin light polypeptides, a mouse monoclonal ferritin light chain (D-9) antibody (1/125, SC-74513, Santa Cruz Biotechnology, Santa Cruz, CA) was used followed by Alexa Fluor 555 IgG (A-21422) secondary. Tissue sections were co-stained with thioflavin S to stain for fibrillar beta-amyloid deposits. Ferritin heavy polypeptides were stained with a rabbit polyclonal ferritin heavy chain (H-53) antibody (1/125, SC-25617, Santa Cruz Biotechnology, Santa Cruz, CA) was used followed by Alexa Fluor 555 IgG (A-21428) secondary antibody. Thioflavin S was used as a co-stain after antibody staining.

To stain for microglial cells, a rabbit Anti IBA-1 (Ionized calcium binding adaptor molecule 1) antibody (1/500, 019-19741, Wako Chemicals USA, Inc., Richmond, VA) was used following by incubation with Alexa Fluor 555 IgG (A-21428) secondary antibody. Thioflavin S was used as a co-stain for fibrillar beta-amyloid after antibody staining.

To distinguish astrocytes, a polyclonal chicken anti-glial fibrillary acidic protein (GFAP) antibody (1/250, AB5541, Millipore, Billerica, MA) was used
followed by Alexa Fluor 555 IgG (A-21437) secondary incubation. Tissue sections were then stained with thioflavin S for fibrillar beta-amyloid visualization.

**Fluoro Jade C Staining:**

To identify degenerating neuronal cells and their relation to Aβ plaques, a Fluoro-Jade C stain was utilized along with immunohistological antibody staining for Aβ$_{40}$. Sections were first stained with Aβ$_{40}$ specific primary antibodies, followed by Alexa Fluor 555 secondary according to the procedure above and then rinsed twice for five minutes in PBS. Slides were then air dried for 30 minutes at 50°C to help tissue sections adhere to the slides. Slides were then rinsed in dH$_2$O for two minutes followed by immersion in 0.06% Potassium Permanganate for ten minutes followed by a rinsing for two minutes in distilled water. Slides were immersed in the working 0.0001% Fluoro-Jade C (AG325, Millipore, Billerica, MA) solution for ten minutes followed by rinsing them three times for one minute each in dH$_2$O. Slides are then air dehydrate for 30 minutes at 50°C followed by immersion in xylene twice for five minutes each, mounting with cytoseal (or another xylene based mounting media), and cover-slipped.

**Microscopy:**

High resolution microscopy of the tissue sections was performed using a Nikon OptiPhot microscope and Nikon Digital Sight camera using NIS-Elements software. Bright field under the visible light spectra and phase contrast using a phase contrast objective and condenser were used to view iron stains. A FITC
fluorescence cube at 430 nm excitation and 550 nm emission (Nikon B-22) was used to visualize thioflavin-S positive Aβ deposits and fluoro-Jade C stains. A Nikon G-2A filter cube was used to visualize the Alexa Fluor 555 secondary antibody which is excited at 550 nm and emits at 570 nm.

4.1.4 – Results

There exists a clear divergence between human AD and APP/PS1 plaque morphology, when viewed with Aβ antibodies (40 and 42 amino acid) and thioflavin-S reactivity (Fig 4.1.5.1, 100x magnification). Thioflavin-S binding of human plaques shows a dense fibrillar core surrounded by a large diffuse halo or corona region. APP/PS1 thioflavin positive plaques exhibit a larger and denser core region with a smaller diffuse thioflavin-S positive corona region. Beta-amyloid 40 antibody bound to human plaques show both the dense core and corona regions are positive for the 40 amino acid Aβ variant with a clear overlap with thioflavin-S staining. APP/PS1 plaques exhibit much more Aβ40 reactivity that extends beyond the thioflavin positive stain. There is still an overlap between thioflavin and Aβ40 staining but there is a non-fibrillar diffuse region outside of traditional thioflavin staining that is evident with the Aβ40 stain. Human AD and APP/PS1 tissue both show positive intracellular Aβ40 reactivity in numerous cells outside the plaques. Antibody staining with Aβ42 shows that much of the dense fibrillar core of the human plaques is composed of the 42 amino-acid Aβ variant, with minimal staining in the corona region. APP/PS1 tissue shows a lack of Aβ42 staining associated with plaques. Intracellular
staining of Aβ\textsubscript{42} is evident inside numerous human AD cells and is not seen in the transgenic tissue samples.

Astrocyte proliferation is illustrated in figure 4.1.5.2 in tissue samples from AD, human controls, APP/PS1 and mouse controls. Control tissue from both human and mouse sources exhibit a nearly random distribution of GFAP positive astrocytes throughout the microscopic field. In the AD tissue samples, there is a high likelihood of astrocytes to be found in close proximity to Aβ plaques. Close examination shows that GFAP positive astrocytes are found on the periphery of nearly every human AD plaque. The astrocytes in the human AD tissue exhibit classical amebic morphology indicative of a phagocytic astrocyte state. Human control and both transgenic and control mouse sections show astrocytes in a traditional star pattern with radiating processes typical of astrocytes acting as neuronal metabolic helper cells. Conversely, mouse APP/PS1 tissue samples show a similar random staining pattern seen in control mouse sections. Beta-amyloid plaques in the APP/PS1 tissue samples are not surrounded by GFAP positive astrocytes unlike the human AD plaques.

Microglial staining and microglial inflammatory state associated with beta-amyloid plaques are seen in figure 4.1.5.3. IBA1 positive stains for microglia in human AD and age-matched controls show microglia in both ramified and active states throughout the tissue samples. AD sections show a close association with Aβ plaques and activated microglia. Microglia bodies and processes are found surrounding and within the fibrillar corona region of the human plaques. Mouse control and transgenic tissue samples both exhibit positive IBA1 microglial
staining. Transgenic and control mouse sections show less microglial staining than both human AD and controls. Microglial activation states for mouse controls tissue resemble microglia mostly in ramified states with minimal activation. Transgenic tissue exhibits both activated and ramified microglial states, but appreciably less than that of AD tissue samples. There is an association between activated microglia and the periphery of Aβ plaques in transgenic tissue, with activated microglia surrounding plaques. Unlike the human AD plaques, there are no activated microglia found within the plaque core or coronal regions.

Light and heavy ferritin antibody staining in human and mouse tissue samples associated with thioflavin-s positive Aβ plaque staining is illustrated in figure 4.1.5.4. Cells positive for L-ferritin staining were found in both human AD and APP/PS1 tissue samples. Human L-ferritin cells were found throughout the tissue sample and also in close proximity to Aβ plaques. Cells positive for L-ferritin appear to be microglial cells in morphology and are found within the coronal region of the human plaques. There was positive L-ferritin staining within the APP/PS1 tissue samples that does not appear to be associated with the beta-amyloid plaques. Furthermore, L-ferritin staining in the animal model does not appear to be microglial in nature. Cells positive for L-ferritin in the transgenic animals appear to be specific for neurons in the granular cell layer of the cortex. H-ferritin was found in cells throughout the cortex of human AD tissue samples in a random manner with minimal association with Aβ plaques. There was no positive staining for H-ferritin in the APP/PS1 transgenic animals.
Fluoro Jade-C staining for degenerating neurons in association with Aβ_{40} staining is demonstrated in figure 4.1.5.5. Human AD and control samples exhibited positive fluoro Jade-C staining that precisely overlap with cells that also exhibited intracellular Aβ_{40}. Cells stain positive for both fluoro Jade-C inside the AD plaques and also outside their halo regions. There was no association in or around the transgenic animal plaques and fluoro Jade-C staining. No positive fluoro Jade-C staining in either APP/PS1 or mouse control tissue samples was observed.

Iron staining compared to thioflavin-S positive plaques in both human AD and transgenic APP/PS1 animals are seen in figure 4.1.5.6 and 4.1.5.7, respectively. Figure 4.1.5.6 shows (a) thioflavin-S staining of plaques, (b) phase contrast and (c) bright field microscopy of Perl's iron stain in the same human AD tissue sample at 40x magnification. Thioflavin-S staining shows numerous dense core plaques throughout the sample that stain positive for focal iron deposition (red arrows). The majority of the human plaques exhibit a focal iron core that can be seen in both the phase contrast and bright field images. The phase contrast image also shows the outline of the beta amyloid plaques as opaque due to the difference in wave length of light passing through the plaques. The APP/PS1 (Fig. 4.1.5.7) tissue sample exhibits positive (a) thioflavin-S staining for beta-amyloid, but does not exhibit focal iron staining in either the phase contrast or the bright field images. Positive iron staining was found in one region (blue arrow) that does not co-register to Aβ plaques. Similar to the human AD data, the APP/PS1 plaques also exhibit a difference in wave length in the phase
contrast images. The phase contrast for the human data however shows a close approximation of size of the plaques compared to the thioflavin-S staining. The phase contrast images for the transgenic mouse data shows that the plaques are larger in diameter then they stain in the thioflavin-S stain. This is similar to the results in figure 4.1.5.1 where the Aβ40 staining of diffuse amyloid is larger in diameter then the thioflavin-S stain, which only stains fibrillar amyloid. A modified iron stain involving protein digestion to enable access of the aqueous Perl's iron stain of the APP/PS1 tissue is seen in figure 4.1.5.8. A minute amount of iron is stained in association with the transgenic plaques. This stain was attempted on human AD tissue but resulted in staining for iron throughout the tissue sample (data not shown).
4.1.5 – Figures

Figure 4.1.5.1

Antibody stains for \( \text{A}\beta_40 \), \( \text{A}\beta_42 \) (red) and thioflavin-S (green) of Alzheimer’s disease tissue from the entorhinal cortex and APP/PS1 cortical tissue viewed at 40x magnification. \( \text{A}\beta_40 \) is found in plaques in the AD and transgenic tissue samples as well as intracellular staining throughout both samples. Thioflavin-S stains for fibrillar beta-amyloid and shows that the core of the AD plaques is composed of highly fibrillar filaments with the halo or coronal region being less compact. APP/PS1 plaques exhibit a much different morphology with a globular fibrillar structure that radially extends from the center of plaque. Antibody staining with \( \text{A}\beta_40 \) illustrates that the size of the APP/PS1 plaques is underestimated with the thioflavin-S stain alone having stained for both fibrillar and non-fibrillar beta-amyloid while the periphery of the plaques is composed of non-fibrillar filaments. The APP/PS1 tissue exhibits much more positive beta-amyloid 40 staining, indicative of the vast over production of beta-amyloid overwhelming the system. Alzheimer’s samples shows positive \( \text{A}\beta_42 \) staining in the core of the plaques and intracellularly throughout the sample with minimal staining of the coronal region. The stains indicate that the core of the Alzheimer’s plaques is composed of both 40 and 42 amino-acid variants while the coronal region is composed primarily of \( \text{A}\beta_40 \). The APP/PS1 tissue did not stain positive for \( \text{A}\beta_42 \) indicating that the transgenic plaques are composed primarily of the 40 amino-acid constituent due to a lack of or reduced production and incorporation
of this beta-amyloid form in the transgenic model plaques. The scale bar is calibrated to 250 µm.
Glial fibrillary acidic protein (GFAP) (red) for astrocytes and thioflavin- S (green) staining for Aβ plaques in Alzheimer’s disease, matched human controls, APP/PS1 and mouse control tissue at 100x magnification. Astrocytes in the AD tissue are seen in a motile phagocytic state surrounding the coronal region of the beta-amyloid plaques. In human control tissue, the astrocytes are observed in their normal highly branched form indicative of their supportive role for neuronal tissue. APP/PS1 and mouse control tissue similarly stain positive for GFAP throughout the tissue samples and illustrate astrocytes in a highly branched normal supportive role. The strong association seen between astrocytes and beta-amyloid plaque in the AD tissue is not perceivable in the transgenic model. The data indicates a differential astrocytic inflammatory response to beta-amyloid plaques in the natural AD milieu compared to the transgenic model. The scale bar is standardize to 100 μm.
Figure 4.1.5.3

Ionized calcium-binding adaptor molecule 1 (IBA1) antibody staining for microglia (red) and thioflavin-S (green) for beta-amyloid plaques in Alzheimer’s disease, human control, APP/PS1 and control mouse tissue samples at 100x magnification. Microglia the AD sample are highly associated with beta-amyloid plaque location and are seen in an activated amoeboid surrounding plaques with processes infiltrating into the coronal region. In the APP/PS1 tissue, microglial cells are also observed in an activated state associating with plaque location. The transgenic microglia are surrounding plaques but do not have exhibit processes that penetrate into the plaques. Human control tissue samples show positive microglial staining and demonstrate microglial cells in a intermediate ramified resting state with numerous processes surrounding their somas. Mouse control tissue exhibits less microglial staining overall and shows cells in a quiescent resting state. The data demonstrates a subtle difference in microglial inflammatory response for AD and APP/PS1 neural tissue samples. The scale bar is 100 µm in length.
IBA1 and Thio-S

AD

Human Control

APP/PS1

Mouse Control
Light and heavy ferritin antibody (red) and thioflavin-S (green) staining for beta-amyloid plaques in AD and APP/PS1 tissue samples at 100x magnification. L-ferritin staining in the AD tissue samples shows intracellular staining in microglial cells that are similar in morphology to those in figure 4.1.5.3. L-ferritin staining in the APP/PS1 tissue shows intracellular staining of neurons in the granular layer of the cortex. There is no positive staining of microglial cells in the APP/PS1 tissue samples. H-ferritin staining of AD samples demonstrates positive intracellular neuronal staining throughout the tissue samples with minimal association to beta-amyloid plaque location. There was no positive staining for H-ferritin in the APP/PS1 tissue samples. The scale bar is set to 100 μm in length.
Figure 4.1.5.5

Beta-amyloid 40 staining (red) and fluoro Jade-C (green) staining in Alzheimer’s disease, APP/PS1, human control and mouse control tissue sections at 200x magnification. Beta-amyloid plaques staining is similar to that seen in figure 4.1.5.1 for both AD and APP/PS1 tissue samples with positive plaques and intracellular Aβ40 staining. Fluoro Jade-C staining is positive for degenerating neurons in the AD tissue sample and overlaps precisely with intracellular Aβ40 staining. There is no positive staining for degenerating neurons in the APP/PS1 tissue sample. Human control samples also show intracellular Aβ40 staining that overlaps with fluoro Jade-C degenerating neurons. Mouse control tissue stains negative for both Aβ40 and fluoro Jade-C. The scale bar is 50 µm in length.
Figure 4.1.5.6

Human Alzheimer’s disease stained with (a) thioflavin-S and (b,c) a traditional Perl’s stain under (a) fluorescence, (b) phase contrast and (c) brightfield at 40x magnification. Red arrows indicate selected plaques that have iron associated with them, while the blue arrow indicates a focal iron region not associated with a beta-amyloid plaque and is composed of hemosiderin or magnetite. The core of many beta-amyloid plaques in AD exhibit high iron deposition within them, as seen in both the phase contrast and brightfield magnifications. The scale bar is set to 250 µm.
Figure 4.1.5.7
APP/PS1 tissue sample stained with (a) thioflavin-S and (b,c) a traditional Perl's stain under (a) fluorescence, (b)phase contrast and (c) brightfield at 40x magnification. The transgenic mouse tissue does not stain for positive for high focal iron associated with beta-amyloid plaques with a traditional Perl's stain. The blue arrow illustrate a focal iron deposit not associated with a plaque that indicates the Perl's stain is effective in staining iron deposits. Phase contrast microscopy show regions where beta-amyloid plaques are located where light passes through the plaque and alters the wavelength of the light. Similar to the AB_{40} stains and thioflavin-S stain in figure 4.1.5.1, the phase contrast allows the viewing of the non-fibrillar beta-amyloid surrounding the transgenic plaques that does not stain with thioflavin-S. The scale bar is set to 250 µm.
Figure 4.1.5.8

(a) Thioflavin S and (b) modified Perl's stain using protein digestion to allow further penetration of the aqueous stain into the hydrophobic APP/PS1 plaques. The stain indicates that there is a very minute amount of iron found in the APP/PS1 plaques that is not stainable with traditional Perl's staining methodology. The scale bar is set to 250 µm.
4.1.6 – Discussion

Alzheimer’s disease (AD) pathology is largely characterized by the hallmark formation of beta-amyloid (Aβ) plaques within gray matter of the cerebral cortex. The creation of transgenic mouse models that generate beta amyloid plaques help mimic this pathological aspect of Alzheimer’s disease. These animal models offer the ability to examine forthcoming pharmacological treatments and assess imaging technologies that potentially could be translated to human AD research and eventual treatment regimens. The time of formation of plaques in these animal models range from approximately six months to a year, and as such greatly accelerate the time of plaque development for research purposes compared to the in vivo disease process in AD. While these transgenic mouse models recreate in vivo plaque formation, plaque formation and morphology are considerably different compared to the neuronal milieu of the AD brain. The histological comparison between human and transgenic mouse Aβ plaques is critical in elucidating experimental results obtained with transgenic Aβ plaque models.

Since the first relationship between cognitive decline and Aβ plaques by Alois Alzheimer in 1907 (3) to the discovery of the 40-42 amino acid constituents of the plaques 75 years later by Allsop et al. (225), the exact role of Aβ in the disease process of AD is still not currently understood. Indeed, the generation of plaques in the human AD brain from the constituent Aβ peptides has not been worked out. Research has shown that there is a relationship between elevated levels of Aβ plaque burden and cognitive decline (226,227) and as such it never-
the-less remains a central feature in the neurobiological study of the disease. The progression of beta-amyloid plaque formation is vastly different in the transgenic model compared to the human AD brain. The mouse brain variant of the amyloid precursor protein differs from human APP by seventeen amino acids, three of which are located within the region coding the Aβ peptide. Non-transgenic normal mouse neural tissue does not produce an amyloidogenic peptide sequence and increasing the level of mouse APP does not lead to the production of Aβ plaques (109). Transgenic insertion of human familial APP<sub>695</sub> into the mouse leads to an increase in the production of Aβ<sub>42</sub> but is insufficient to cause plaque formation. The same is true for transgenic models utilizing single human presenilin (PS) 1 or 2 mutations of the γ-secretase protein. When mice are co-transfected with both APP<sub>695</sub> and PS1, the formation of plaques occurs at approximately six months and increases the level of Aβ<sub>42</sub> which is directly linked to the onset and concentration of amyloid deposits in these transgenic models.

In order to compare data obtained with the APP/PS1 model to human AD, evaluation and comparison of the beta-amyloid plaques in the transgenic models to human AD plaques must be carried out. There appear to be numerous distinct differences between the plaques in the naturally progressing AD and the transgenic mice. Thioflavin-S staining of the plaques shows that the animal model has plaques which are globular in nature and are composed of filaments that branch out radially from a central location (Fig. 4.1.5.1). Compared to the human AD plaques they lack the typical dense core and large surrounding halo region of the human plaques. Antibody staining of human AD and APP/PS1
plaques with both 40 and 42 amino acid variants shows apparent differences in staining patterns, which allude to fundamental differences in plaque composition, formation and morphology. It is important to note that the beta-amyloid plaques in the transgenic model also are recognized by human antibodies to Aβ, as the transcribed peptides are derived from human mutations (228). There is an underestimation of transgenic plaque size with thioflavin-S staining alone compared to Aβ40 staining. The core of the transgenic plaques exhibit a high binding affinity for the thioflavin-S molecule, which is indicative of a highly fibrillar amyloid core within these plaques. The large coronal region that is stained only with the Aβ40 antibody indicates that this region is composed of more diffuse beta-amyloid, which is less fibrillar in nature. Aβ40 antibody staining also shows positive staining in regions where no or very little thioflavin-S staining is evident. These areas could be indicative of regions that are composed of non-fibrillar Aβ during the formation of new plaques. Human plaques show high amounts of fibrillar Aβ42 mixed with lower amounts of Aβ40 within the core of the plaques, surrounded by a halo region predominantly composed of Aβ40. Transgenic plaques do not stain positive with Aβ42 antibody illustrating that these plaques are not composed of the 42 amino-acid variant of beta-amyloid. These differences in morphology between human AD and transgenic plaques could be due to a number of factors. The decreased time of amyloidogenesis in the transgenic APP/PS1 model is one culpable factor. Although chronologically beneficial for performing studies using the animal models, the decrease in time for amyloid plaque formation creates an uncharacteristic environment outside that seen in
the AD neuronal milieu (111). Normal Aβ plaque progression in the AD brain takes a number of years and perhaps decades. The APP/PS1 transgenic model produces plaques at a greatly accelerated rate within nine months from birth. The accelerated time in amyloid fibril formation and plaque growth are important factors to reflect upon when considering the difference between the AD and transgenic plaque morphology. The growth of the human AD plaques incorporates the surrounding neuropil and cells within the fibrillar coronal region. Transgenic plaques appear to rapidly grow radially from a central seed and force the surrounding neural tissue away from the growing plaque mass. The factors involved in natural AD amyloidogenesis are enormously complex and multifaceted. Ex vivo synthetic beta-amyloid peptide like that found in AD undergoes conformational changes dependent upon environmental factors such as pH and solution conditions (36). Transgenic beta-amyloid peptide is known to undergo different post-translational modifications compared to when it is produced in the normal AD environment (229). This explains why work looking at the solubility of AD and transgenic amyloid fibrils shows that transgenic fibrils are completely soluble in buffers containing sodium dodecyl sulphate whereas human AD fibrils are completely insoluble (229), further illustrating the chemical and morphological dissimilarity between the two fibril types.

Intracellular staining of Aβ deposits shows a clear difference between AD and transgenic tissue samples (Fig 4.1.5.1). Human AD samples show intracellular staining for both Aβ40 and Aβ42, while there is a decrease in Aβ40 intracellular staining and a lack of apparent staining for Aβ42 in the transgenic
model. It is not clear from these stains if the intracellular beta-amyloid stained in the tissue samples is of the soluble type which has been associated with neuronal apoptosis (46). Staining of AD and APP/PS1 tissue with Fluoro Jade-C (Fig. 4.1.5.5) to view neuronal degeneration tissue shows a clear overlap in intracellular neuronal staining with Aβ40 in AD and aged human controls. This is congruent with data suggesting that intracellular beta-amyloid induces neuronal apoptosis. The fact that transgenic tissue, which stains positive for amyloid plaques, and control mouse tissue do not stain with fluoro-Jade C suggests that neuronal degeneration is not occurring in regions surrounding Aβ plaques and that neurons are not undergoing apoptotic signaling associated with intracellular beta-amyloid. This represents a clear and fundamental difference between the transgenic mouse model and AD. The neuronal degeneration associated with beta-amyloid is one of the underlying mechanisms responsible for the disease process. The relationship between intracellular and extracellular beta-amyloid and the mechanism by which extracellular amyloid is produced in the normal AD disease process could be due to a number of factors and seems to be different from the method by which it is produced in the transgenic APP/PS1 model.

Amyloid precursor protein processing is believed to occur in the endoplasmic reticulum and trans-Golgi network (49), producing beta-amyloid through the gamma and beta-secretase cleavage pathway. This would result in intracellular amyloid accumulation when the peptide is released into the cytosol of the cell. While the exact biochemical mechanism by which extracellular amyloid is transported outside the cell is unknown, there does appear to be a dynamic
relationship between intracellular and extracellular beta-amyloid in the AD disease process (49,51). It has been hypothesized that intracellular Aβ is the initiating lesion in the development of Alzheimer’s disease and extracellular Aβ can also increase de novo production of intracellular species (230). The lack of intracellular amyloid staining in the transgenic samples implores the question as to the mechanism by which extracellular amyloid accumulates in the APP/PS1 model and how it differs from the natural AD process. It is plausible that the transgenic APP is processed in the extra-cellular membrane and as such expels amyloid directly into the extracellular environment, or by other means including exocytotic excretion (52).

Microglial and astrocyte cells are important in Alzheimer’s disease because they respond to neuronal environmental changes. These glial cells produce numerous regulatory proteins, inflammatory cytokines, and many proteases and protease inhibitors molecules associated with inflammatory function (231). Neuronal inflammatory response viewed with microglia and astrocyte migration and proliferation is also strikingly different between the APP/PS1 and AD samples. The involvement of microglia in the pathogenesis of Alzheimer’s disease has been well studied and collectively there is support that inflammatory microglia involvement is an important component of the disease. Although generally considered a positive presence during inflammatory response, there is growing support as to the detrimental role that microglia play during the AD disease process. The recruitment and activation of microglial cells in and around neuritic plaques leads to the production of various cytokines and
neurotoxins, which are known to cause neuronal injury and death (83). There is a marked association between activated microglial cells with beta-amyloid plaques in the human AD samples (Figure 4.1.5.3). Microglial cells are seen surrounding and throughout the neuritic AD plaques infiltrating within the outer halo region in close proximity to the dense core. Microglial cells are known to assimilate high amounts of intracellular beta-amyloid within them due to the phagocytosis of stressed and dying neurons. There is evidence to support that the recruitment and resulting close proximity of microglial cells to initial beta-amyloid plaque formation results in the delivery of additional fibrillar peptide encouraging further plaque enlargement (85). While there is also an association with microglial cells and amyloid plaques in the APP/PS1 model, there are distinctive differences to the human AD samples. Microglial cells are fewer in number throughout the APP/PS1 samples compared to AD tissue and are visually similar in number and activation state to the control mouse tissue samples. This perhaps is indicative of less neural inflammation occurring in the APP/PS1 brains overall. Microglial cells are seen surrounding transgenic plaques but are not found within the perimeter of the plaques as in the AD samples. While there is speculation that microglial cells in the transgenic model also contribute to plaque formation (232), it is suspicious that during amyloid plaque growth the microglial cells are not engulfed and incorporated into the perimeter of the amyloid star. With the decrease in time of amyloidogenesis, there is a lack of assimilation of surrounding neural tissue as the transgenic plaques radially progress possibly as a result of the diminished time period.
While microglia are found adjacent to and surrounding APP/PS1 Aβ plaques, it seems that overwhelming of the regulatory/degradation systems due to the overabundance of extracellular beta-amyloid seems to be the source of raw Aβ peptide and the driving force toward transgenic plaque formation.

Comparison between AD and transgenic tissue samples demonstrate that there is a weak astrocytic inflammatory response in the transgenic model. The AD tissue samples exhibit a pronounced astrocytic response with a clear relationship between plaque and astrocyte location (Fig. 4.1.5.2). There are well defined astrocyte somas found along the periphery of the plaques with processes extending into the coronal halo region. This differs from the human control sections that show normal astrocyte distribution and processes for an aged human sample. The APP/PS1 tissue shows a similar distribution of astrocytes as found in the control mouse tissue samples, with little to no association of astrocytes with Aβ plaques. Astrocytes are classically well known to be involved in inflammatory response to regions of neural-cellular distress and neurodegeneration to aid in distribution of metabolites, provide support and aid in the repair of effected regions. In the AD tissue samples, it is expected that neural tissue will respond as such to the accumulation of beta-amyloid due to the foreign nature of the plaque. Astrocytes in the AD tissue typically show an reduced star like state with a large somo and few processes. This is typical of an astrocyte in a state when responding to cellular injury. In the transgenic model, the samples do not demonstrate this anticipated response, and are viewed as resting astrocytes in a metabolic helper state. It is plausible that the formation of
the transgenic material into beta-amyloid plaques in the APP/PS1 occurs in such a way that the neural tissue doesn’t recognize the material as foreign and as such fails to initiate an inflammatory response. It is of considerable interest that the APP/PS1 tissue samples do show a microglial reaction in close proximity with the periphery of the plaques while not presenting an astrocytic response. There remains the possibility that in the human AD tissue the astrocytes are responding not to the beta-amyloid in the region but to cellular distress associated with beta-amyloid. Astrocytes are known to respond to neuronal cellular stress and death in an attempt to aid in neuronal and cellular repair. In the APP tissue the apparent lack of astrocytic response could be due to a lack of cellular distress and death occurring in the vicinity of the plaques.

There is a considerable difference between the amyloidal genesis in the APP/PS1 mouse model and human Alzheimer’s, in respect to its relationship with in vivo iron stores. Localized focal iron is frequently found in AD tissue samples both in the central core of the human AD neuritic plaques and also in an apparently random fashion in the form of large focal iron deposits in the form of hemosiderin or magnetite (190,233). The Perl’s co-stain with thioflavin-S show an exact overlap between the focal iron deposits within the central core of numerous AD plaques (Fig. 4.1.5.6). There is a reduced amount of iron associated with the APP/PS1 transgenic model (Fig. 4.1.5.7). The phase contrast microscopy images allow the halo region to be visualized in the AD tissue samples and the non-fibrillar halo around the transgenic plaques as the wavelength of the light is altered when it passed through this region, while clearly
viewing iron only in the AD tissue samples. Only when stained with a modified Perl’s stain that breaks down the periphery of the APP/PS1 plaques are traces of iron found (Fig. 4.1.5.8). Similar staining of AD tissue samples with the protein lysis results in a complete staining of iron throughout the tissue. This is presumably due to protein degradation of iron storage proteins (ferritin) and staining of their iron oxide cores. Ferritin antibody staining for tissue sections indicates differential control of iron storage and association with beta-amyloid plaques in transgenic APP/PS1 and AD neural tissue. Microglial cells are known to harbor high quantities of the light ferritin isoforms for long term iron storage, assumedly due to phagocytosis of cells undergoing apoptosis caused by oxidative stress due to iron induced free-radical production (64,213,234). Human AD beta-amyloid plaques exhibit a tight association with L-ferritin as the pattern of staining is indicating that the L-ferritin isoform is found within microglial cells that are surrounding and infiltrating the plaques. The pattern of staining in the AD tissue for L-ferritin is similar to the microglial antibody stain in figure 4.1.5.3. In the APP/PS1 model, L-ferritin staining was found throughout the tissue sample in the granular cell layer and appears to not be associated with microglial cells based on the cellular morphology and pattern of staining compared to the microglial cells in figure 4.1.5.3. The APP/PS1 having no light isoform associated with microglial cells is indicative of a system that is not undergoing excessive iron build up like in the AD tissue where microglial cells are responding to oxidative apoptotic cell death and sequestering high amounts of iron during the phagocytic process. Heavy ferritin staining exhibited positive staining in neurons
within the AD tissue and was not observed in the transgenic mouse tissue. H-
ferritin is known to rapidly take up and release iron and is involved in excessive
iron detoxification protecting against iron-induced oxidative damage (64,65). The
ferritin staining data sets are congruent with an over abundance of iron in the AD
tissue and less overall iron in the APP/PS1 neural tissue. Overall, the iron
staining data suggest an overall misregulation of iron homeostasis in the AD
diseased brain that is not seen in the transgenic mouse tissue.

While it is unknown precisely how Aβ plaque aggregation occurs, it is
known that metal ions can play a role in Aβ fibril formation. The conversion of
Aβ’s secondary protein structure has been shown to occur in the presence of
divalent metal ion’s (82). Aberrant regulation of iron leading to its focal
concentration in brain tissue is believed to be associated with the
neurodegenerative process in neurological disorders such as Alzheimer’s
disease and Parkinson’s Disease (55). Abnormal regulation of iron is believed to
be associated with the neurodegenerative processes in Alzheimer’s disease.
Regions of focal iron concentration leading to potential oxidative stress can lead
to cell death and apoptosis (87). The relationship of iron to plaque generation is
not currently well understood within the literature. It is largely unknown whether
the generation of the plaque causes the aggregation of iron to form a core, or if
intracellular iron causes the formation of the beta-amyloid protein around it.
There is a general agreement that the location of some focal iron regions in brain
tissue coincides with some Aβ plaques in certain brain regions. Numerous
studies support the hypothesis that oxidative damage surrounding the Aβ
plaques is involved in the human Alzheimer’s disease process. Research has shown that the co-localization of iron with the Aβ plaques is accompanied by endoplasmic reticulum stress induced apoptosis, DNA oxidation and cellular damage in cells adjacent to plaques (104,235,236). Conversely, in regions where Aβ plaques accumulate alone without iron there is no indication of oxidative stress or apoptosis pathway activation (214). It has been demonstrated that the toxicity of Aβ is amplified upon the direct interaction of iron ions with the peptide, while unbound iron itself has no effect upon toxicity (215). This strongly suggests that iron accumulation in and around plaques is essential for neuronal damage. Indeed, there has been speculation as to the positive neuro-protective roll that Aβ might possess in the disease process. The high affinity for human Aβ to bind iron might be the result of homeostatic pressure to bind and eventually expel excess iron from the human brain (89). Regardless, overall it is a widely held belief that the Aβ peptide is involved the neuro-pathogenesis of AD. Numerous studies support the hypothesis that oxidative damage surrounding the Aβ plaques is involved in the disease process. This is furthered by the knowledge that cytotoxic effects can be ameliorated with the usage of free-radical scavengers and antioxidants (237). The redox-metal dependent generation of free radicals and highly reactive hydroxyl radicals through a Fenton type reaction is believed to trigger the neurotoxic effect in cells that are close proximity to Aβ plaques (238). Neuropathological and biochemical studies have shown increased activity of free radical defense enzymes, elevated levels of
oxidative damage to mitochondria, DNA fragmentation credited to apoptotic signaling and oxidative stress markers in AD patients (104,238).

The data in this report suggest that there are numerous differences between normal AD disease process and the APP/PS1 transgenic model, which aims at mimicking the genesis of beta-amyloid plaques formation. The morphological differences between the plaques in the transgenic and AD tissue sample are of considerable importance to researchers whom are considering using the model for studies involved in amyloid clearance based on specific beta-amyloid peptide sequence targeting. The difference in plaque composition (Aβ 40 or 42 subtype) and gross plaque morphology in the APP/PS1 model could result in misleading results when translating pharmacological or antibody methodologies to plaque degradation in the AD brain. This also is true due to the increased in size of the transgenic plaques and highly fibrillar hydrophobic core surrounded by thioflavin-S negative coronal region in the transgenic plaques, which warrants caution when considering using this model for these studies. Other methodologies for AD therapeutics involve the augmentation of the immune system inflammatory response to alter the production and/or clearance of the beta-amyloid peptide sequence. This again should be done under the caveat that there is a considerable divergence between the transgenic model and the normal AD inflammatory response as far as with microglial and astrocytic proliferation and plaque association. It is largely unknown as to why human AD neuronal tissue produces beta-amyloid resulting in plaque formation. For the most part, the same does not hold true for the transgenic animal tissue for whom
beta-amyloid and plaque formation is governed by the introduction of two known human mutations. Herein is the cause of the vast increase in Aβ production found in the transgenic model. While in the normal AD disease process, the increase in the amyloidosic beta-amyloid peptide pathway leading to the disease process is highly complex and represents a multi-faceted discoordination or miss-regulation of numerous endogenous systems. Of those systems, three are outlined throughout this work upon which each shows considerable differences in morphology, inflammatory response, and iron management between the natural AD process and transgenic model. It should be understood that these systematic biochemical errors represent a small proportion of the potential plethora associated with the normal AD disease process.
Chapter 5

General Conclusions

5.1 – Utilizing Magnetic Resonance Imaging for the Visualization of Beta-Amyloid Plaques

Magnetic resonance imaging is an extremely powerful technology which has pushed forward our ability to diagnose and treat injury and disease into what many would call the modern age of medicine. With the advancement of MR technology, magnets with higher strength fields are becoming commonplace in medical institutions around the world. There is a hope that these magnets will usher in an era with the ability to image human tissue at higher resolutions in less time than ever before. Nevertheless, even at these high field strengths we are pushing the technological envelope with microscopic magnetic resonance imaging.

The small size of beta-amyloid plaques and the current clinical resolutions available to the majority of health care providers is a limitation in our ability to image Aβ plaques. With this said, there are a number of researchers who have investigated imaging beta-amyloid plaques with MRI, in both human AD (183,187,219,239) and transgenic APP/PS1 mice (184-186,193,194,239-243). There has been a great deal of success in imaging neural tissue at high resolution with microscopic MRI techniques (155,163,166,187). The majority of these studies involve the usage of home build RF coils that are very specific and conform to the dimensions of tissue being imaged. Coil specificity aids in creating higher resolution images with increased signal to noise ratios with less
imaging time. There is a downside to this however; transferring these results to other systems utilizing traditional volume, surface, or even multi-array coils. These apparati cannot produce comparable high resolution images in an adequate time frame for in vivo imaging (whole or partial brain). Here we are left with a conundrum; the technology exists to microscopically MR image tissue with sufficient resolution to view individual plaques (186,187) but the inability to currently translate this to clinical systems. Attempts to compare the transverse relaxation value of large regions of interest and then measure plaque burden in these regions have largely proven to be marginally successful in detailed quantification of plaque burden (195,241,244). The resolution used in these studies would be comparable to clinical in vivo studies of AD patients, so the results would hypothetically be directly comparable. The main problem with this line of research is that the plaques in the transgenic model are not only distinctly different in morphology from the human AD plaques but also have differences in iron load. These factors are pivotal in the relaxation associated with the transgenic plaques. It is also of note that within the AD brain there is a sizeable amount of focal iron that is not associated with plaques. This iron would invariably account for a large portion of the transverse relaxation of a large region of interest containing number plaques and iron deposits. As such, the measurement of relaxation in this region would not be a direct measure of plaque burden but instead a mixture of both plaque and iron load. With the knowledge that the mouse model does not have considerable amounts of iron present; the quantification of regions in the transgenic mouse would be a more direct
measure of plaque burden. This would be in large part due to the fact that the plaques themselves in the APP/PS1 model have transverse relaxation associated with them due to proton relaxation in the vicinity of the protein mass. The human AD Aβ plaques also have relaxation associated with them but it seems that iron found in and around the plaques is the dominant cause of transverse relaxation.

The data collected in these studies emphasize the progression of MR technology to microscopically MR image tissue samples. Translating these results to current in vivo MR scanners in a clinical setting is challenging. The main limitation, at the moment, exists due to the current resolution of clinical scanners. Ideally, with enough imaging time and image averaging, a lower field MR system could produce a comparable high resolution image. However, in a clinical setting there are time constraints on how long a patient may remain in the scanner for both safety and feasibility reasons. While the scans taken in this work took approximately 6 hours to image at 7.0 Tesla, the time to produce such images at a clinical 3.0 T system are not allowable (on the order of multiple days). In the future, as higher field strengths are used for clinical imaging there is the possibility of acquiring images with similar resolution. There remains the question; Why would we want to image beta-amyloid plaques in the human brain in vivo? The easy answer is to be able to detect plaques in the brain for earlier diagnosis of AD. This in itself is challenging, as the work here shows that there is a general iron overload in the AD brain as well as beta-amyloid plaques. These both represent sources of potential transverse T₂ relaxation, and as such
are difficult to tease apart. Early in MR imaging it was realized that T2 weighted images were sensitive to brain pathologies, and it was hoped that this would lead to the ability to utilize T2 relaxation to specifically address disease pathology (245). This was not fully realized after extensive research for a number of reasons, most notably because proton transverse relaxation is multi-dimensional and occurs due to a number of reasons but ultimately is shown on a single dimension scale as image contrast. Teasing apart the different relaxation curves that make up the summated transverse relaxation would help to understand the mechanisms associated with proton compartmentalization. Comparing these results to different brain pathologies, specifically beta-amyloid plaques, might aid in understanding and translating these results to a clinical setting. This would first be accomplished on microscopic scale, and then broadened to the larger voxel resolutions in a clinical setting with the understanding that partial volume effects would be present.

5.2 – Using MRI to diagnose early AD and Mild Cognitive Impairment

There have been a number of proposed mechanisms to determine the clinical onset and progression into Alzheimer’s disease utilizing magnetic resonance imaging. They include utilizing various MR methodologies such as functional imaging (246-252), diffusion tensor (253-255), quantitative measurements of regions of interest (252,256,257) and direct imaging of plaques (183,185,187,190). The hope is that there are measureable biomarkers such as olfactory deficits, cortical thinning, white matter track alterations or changes in
relaxation that can predict changes in cognition that eventually lead to clinical onset of Alzheimer’s. Individually these studies have shown that there are structural changes in the AD brain that differ from normal aged matched controls. Going from the realm of research to a comprehensive clinical test has proven quite difficult as individual markers lack the ability to accurately predict progression into Alzheimer’s. If a cross modality approach to the data is taken and it is organized in a regression analysis there is prospect in their ability to accurately predict disease progression. This is difficult, as a battery of MR and cognitive tests would be required at the same institution or cross institutionally. Many have placed emphasis on the utility of imaging and quantification of plaques and associated iron deposition for potential clinical examination. As accentuated in the previous section, there are a number of considerations in this approach that are large hurdles to overcome.

5.3 – The APP/PS1 mouse model of Alzheimer’s disease

The majority of the studies involving MR imaging of plaques have assumed that the ability to view them has been due to the localization of iron within and around these plaques (193,258). Detailed analyses of plaques in the transgenic APP/PS1 model have revealed that there are distinct differences in the relationship between iron and their plaques and also in plaque morphology. This difference has been mentioned in the literature relatively recently (109,193,201,229,259), but these dissimilarities and the cause of transverse relaxation have not been drawn fully into the magnetic resonance discipline. The
reason for the clear difference in plaque morphology is at first glance relatively straight forward; the attempt to generate Aβ plaques in a mouse model that naturally does not do so. In reality the circumstances are complex and multifaceted, with a number of factors that seem to have individual roles that coalesce. The natural APP processing in the murine brain does not result in the production of fibrillar amyloid, and as such does not generate plaques. With the introduction of human genetic mutations into the APP/PS1 mouse model we have created plaques, but not in the same way they are produced in the human AD brain. In the APP/PS1 brain there are two systems are play during APP processing, the endogenous APP and γ-secretase (PS1) and then the introduced human FAD mutations in APP and PS1. Unlike the human AD brain, the APP/PS1 model now has the ability to produce both a natural non-fibrillar product and the fibrillar AD Aβ. When initial seeding of fibrillar Aβ occurs in the transgenic model it provides scaffolding that allows the natural product to potentially bind. Indeed, it has been shown that APP/PS1 animal plaques can be composed of both human and mouse amyloid (202,260). There are also different post translational modifications to the beta-amyloid that is produced in the mouse model (209). The environment within the mouse brain is very different from the human brain in a number of aspects. The levels of various metals, especially iron, are a great deal reduced in murine neural tissue compared to the human brain (192,261).

A very important consideration when using the APP/PS1 transgenic model is the ability to transfer acquired datasets to the natural AD process. There has
been great emphasis placed on using various methods to reduce plaque burden, chelate metals coupled to plaques or institute pharmacological methods in an effort to decrease neuronal damage associated with Aβ plaque deposition. While the model does show some similarity to AD pathogenesis, there is an inherent caveat that must be disclosed. The mouse model generates beta-amyloid plaques in a different manner than those produced in the AD brain. The APP/PS1 model generates plaques in response to the drastic overproduction of fibrillar beta-amyloid. The over production of APP in the murine model is evident when looking at both microscopic images and quantification of beta-amyloid peptides with protein quantification. The murine model possesses an endogenous APP variant that does not get processed into amyloidogenic beta-amyloid fragments to product plaques. The increase in amyloidogenic peptides is directly related to the introduction of human familial Alzheimer’s genetics. A noteworthy question is why does the murine model over produce the introduced human variants to such a degree? It is possible that the mouse model is unable to break down the amyloidogenic human peptides as well as the endogenous murine forms, leading to accumulation of human forms. This seems unlikely however with recent data suggesting that wild-type mouse apolipoprotein is able to efficiently degrade both soluble and insoluble beta-amyloid and eliminate thioflavine-S positive amyloid deposits (262). It does seem that overproduction or accumulation of the human peptides is not specific for certain neural structures in the mouse brain. In the human AD tissue, plaques form predominately in the middle cortex layers and plaque production starts near the entorhinal cortex and
eventually spreads throughout the brain. In the APP mouse model, plaques for throughout the brain. They are non-specifically produced not only in cortical tissue but also in deep brain matter and white matter tracks. It seems that there is a vast overproduction of Aβ in all APP neural cell types that is clearly not seen in the naturally occurring human AD brain. The direct cause behind why human AD tissue produces plaques is not known, but there are theories that support the current available data. Beta-amyloid is a natural product of APP processing, and is found in normal cognitively intact individuals and also in the murine brain but not in the fibrillar form. The formation of beta-amyloid plaques is not limited to the AD brain, as the normal aging process produces plaques in aged individuals (217,263). The divergence point between a cognitively intact individual and a person with AD is hard to place a finger on. Many would place emphasis on the genetic mutations found in familial AD cases. There is a wealth of data to support this, but the majority of individuals diagnosed with AD do not have mutations that are currently associated with the disease process. The foundation of the APP/PS1 model focuses on the genetic mutations of AD, and generation of plaques is a direct result of this. However, there are clearly other factors that have not been accounted for in the natural AD brain. A threshold of cognitive decline has been suggested in that individuals may appear cognitively intact because they have yet to reach a specific point of atrophy in the brain where it can no longer compensate for lost tissue. This would suggest that it is not the generation of plaques, per se, but the loss of neural tissue that is responsible for cognitive decline. The literature currently agrees with this, as
there have been clear results showing cortical loss associated with cognitive
decline in AD (264,265). The reason for the cortical loss is the key element in
the disease process. Staging of AD has clearly shown that cognitive decline is
associated with an increase in Aβ plaque and NFT burden (4,43,44). This has
led the majority of the field to conclude that these hallmarks are responsible, in
some way, for the loss of neurons in the AD brain.

5.4 – Plaques and Iron, what is their relationship and which comes first?

The involvement of iron in the pathogenesis of Alzheimer’s has been well
established. The incorrect regulation of iron management proteins and irregular
storage of iron within AD neural tissue results in the accumulation of excess
amounts of free iron. It appears that these conditions are not separate from the
AD process, as they are found in the majority of AD cases. Increased stored iron
is associated with a number of medical conditions, such as type 2 diabetes and
vascular disease, which increase the risk for progression into AD (266). Body
iron stores increase with age in both men and women with markers for iron
stores increasing sharply between the teenage years until iron levels plateau in
middle age. Alzheimer’s is generally considered a disease of aging with
increased risk of developing AD after 60 years of age. The age related increase
in iron precedes the disease associated cognitive decline by decades. It seems
highly probably that the natural increase in iron precedes clinical AD symptoms.
Other diseases such as heart disease and atherosclerosis have been shown to
escalate in parallel to the increase in iron storage in the body (266-269). It has
been proposed that the age-associated increase in body iron stores could be responsible for the start of the pathophysiological cascade that promotes Alzheimer’s disease progression. Genetic and environmental factors associated with abnormal iron homeostasis may go unnoticed for decades, as the body is able to maintain excessive iron storage for a period of time. However, there comes a moment when the body can no longer maintain this threshold of iron maintenance and the system overloads. Excessive iron has been shown to mediate and increase the production of the APP peptide, which ultimately then is cleaved to form the Aβ peptide (270). It has been hypothesized that this is due to the metal binding properties of the beta-amyloid peptide. It has been suggested that neural tissue produces the peptide to help aid in the sequestering of excessive iron that the body’s normal storage mechanisms can no longer keep up pace with (88,89). This might help in the short term, but as the Aβ peptide binds more iron and plaque aggregation occurs the accumulation of iron in plaques increases and produces focal regions of iron deposition. In the midst of this, more iron is continually being shuttled into the brain, which cascades to produce more Aβ peptide which amasses into additional beta-amyloid plaques. As this ‘backup’ system of sorts fails to sequester errant iron, accumulation of hemosiderin and magnetite occurs in the AD brain (271). The plaques and other regions of highly focal iron now function as reduction and oxidation centers; reactive oxygen species and free radicals are produced, cells undergo excessive amounts of oxidative stress, they tumble into apoptotic signaling and neuronal atrophy occurs.
5.5 – Potential Follow-Up Studies

It is important to understand that the transgenic mouse models of AD created to date do not exhibit iron imbalances in their neural tissue. There is no focal iron found throughout their brains and Aβ plaques that form have only a minimal amount of iron. This is a strict limitation of the animal model, and ultimately might prove to be a cause for the lack of atrophy on the scale seen in AD. This also might explain why plaques form differently within the APP/PS1 brain with a completely different morphology. It would be very interesting to see how increased iron in neural tissue would affect plaque generation and morphology along with markers of oxidative stress and apoptotic signaling. Increasing iron storage in neural tissue is not a normal process in the murine brain. An alternative means at increasing iron systemically throughout the body is required. This can be done with lipophilic iron compounds such as TMH-ferrocene [(3,5,5-trimethylhexanoyl)ferrocene] (272). A similar study of increasing iron stores in the mouse for quantification and measurement via MRI has been undertaken and is found within the appendix of this thesis. Iron concentration levels do increase in the neural tissue with a concomitant increase in neuronal L-ferritin (272-277). The next step would be to feed APP/PS1 mice food with TMH-ferrocene from birth. A detailed evaluation of iron load and plaque genesis compared to litter-mate transgenic mice feed a normal or low iron diet would be essential. Magnetic resonance imaging sequentially throughout this period would help understand how relaxation associated with plaques is changing. It is hypothesized that iron load would increase throughout specific
regions in the neural tissue as viewable with MRI (view appendix) and that plaque generation would be altered in the mice feed the lipophilic iron compound. Plaques would contain more focal iron and potentially would form more akin to the way they do in the Alzheimer’s brain. Oxidative stress markers would increase in neural tissue surrounding the plaques and neuronal death would occur in close proximity to beta-amyloid plaques. This would help create a more realistic animal model of AD that incorporates not only the generation of plaques but also some of the iron imbalances known to be associated with the disease.

Understanding the differences in inflammatory response between the APP model and human AD would be important in future development and testing of therapies for AD. The data here show that there are differences between neuronal stress markers and cellular death in the APP and AD brains around the plaques. This might be why there are histological differences between astrocytic and microglial response between the two systems. Detailed measurements of neuronal stress proteins are reactive oxygen species in the vicinity of the plaques would help tease apart these differences.

Another line of study would be the further usage of the histological coil in aiding to understand the MR relaxation associated with Aβ plaques. At the time of the studies pursued here we were unable to quantify the amount of iron present at the plaque locations. There are two available methods to measure this. Using laser dissection of individual plaques followed by atomic-absorption spectroscopy (AAS) of the samples or another method that directly uses a laser to vaporize the tissue in a XY plotter like fashion that then measures the gases
produced during vaporization to evaluate metal ion concentration. The first method would require individual plaque dissection, which would be very time consuming to undertake, and involves a certain amount of error when measuring such small pieces of dissected tissue with AAS. The second method would vaporize an entire sample, but would give a detailed three dimensional graph with XY being spatial locations and Z being iron (or another metal) concentration. The latter method would seem to best available option but would require the equipment, which would be best done with collaborative efforts considering the considerable resources involved. There is also the possibility that the second method could measure beta-amyloid concentration. It would be hypothesized that quantitative measurements of transverse relaxation would correlate to the concentration of iron or amyloid present. This would generate a noteworthy finding in understanding how transverse relaxation is affected by both protein and metal ion concentration in both human AD and APP/PS1 tissue samples.

Potential follow-up studies to further evaluate the mechanism by which the beta-amyloid protein in the plaques elicits transverse relaxation would be beneficial. It was hypothesized that the close interaction (i.e. rotational and irrotational binding) of the protein with the protons causes the rapid relaxation of other protons in close proximity. Further testing of this hypothesis is possible by altering the exchange rate of proton-proton interactions via heating or cooling of the sample followed by subsequent measurement of the transverse relaxation rate for plaque regions of interest. Heating the sample would result in increased proton (water) movement and thus increased proton-proton dipole-dipole
interactions resulting in an increased relaxation rate. The opposite holds true for cooling of the sample in that this would decrease the relaxation rate due to decreased proton-proton interactions. It is unknown if heating or cooling at the possible temperature range available would be able to appreciably alter the transverse relaxation of these regions; however, if heating and/or cooling does elicit a difference in relaxation for these regions of interest this would point towards a dipole-dipole interaction of relaxation.

There has been a large push toward molecular and contrast enhanced imaging of beta-amyloid plaques. With the discovery that positron emission topography (PET) imaging of plaque burden is possible with a thioflavin-T molecule with a radioactive C\textsuperscript{11}, otherwise known as Pittsburgh compound B, many have considered an MR alternative. MRI has the unique benefit of spatial resolution over PET imaging, which could allow individual plaque imaging. Testing potential contrast mechanisms in animal models is time consuming and takes considerable resources. Potential contrast agents can come in the form of chelated metal ions (gadolinium or iron oxide particles) bound to proteins or antibodies that bind to specific peptide sequences or other new novel agents (chemical exchange-dependent saturation transfer (CEST)) that also increase proton relaxation. This technology is not limited to study just within the realm of Alzheimer’s research, as a great deal of potential MR molecular probes count be tested for specific antigens found on cancerous tissue, specific cell types, or specific regions in the brain or body. Testing both the efficacy on proton relaxation and binding of the probe is important is understanding the utility of this
technology. The most effective probes would elicit proton relaxation when bound to their respective antigens and would have a potential proton shielding effect when not bound such that they would not elicit relaxation. The probe must be able to bind with the antigen with high enough specificity but also be able to release from the antigen overtime as to not interfere with other potential \textit{in vivo} mechanisms and be excreted over time. The probe must also be able to cross the blood brain barrier successfully without opening the tight junctions (i.e. mannitol).

There also is the investigation of other diseased tissue samples with the histological coil. Alterations in proton relaxation have been associated with a number of neurological diseases and other diseases of the body. Detailed MR evaluation and subsequent histological staining of these tissues with some of the methods outlined in this work could help in the understanding of how tissue contrast in many disease processes is related to tissue histology. With careful changes to MR sequences to allow for a large amount of echoes it is possible to extrapolate the multiple relaxation curves from multiple proton (water) compartments in the average T$_2$ relaxation curve. This would allow for comparison of multiple exponential T$_2$ curves in tissue samples with the histological coil. Comparing these values to histological stains would be of great interest. Specifically, viewing the ratio the myelin water fraction (MWF) (278) to Luxol Fast Blue staining of white matter in Amyotrophic Lateral Sclerosis and control tissue would be interesting to analyze. Also, it would be of considerable interest to see if multiple MR T$_2$ relaxation due to differing water
compartmentalization differs between human AD and APP plaques. This would further aid in the understanding of the mechanism of MR relaxation associated with the plaques.
Appendix A

Quantitative Measurement of Iron with Magnetic Resonance Imaging

6.1 – Purpose of Appendix Material

The data that follows was a study designed to evaluate and test the hypothesis that quantitative iron measurements in a phantom and in vivo model system was possible with magnetic resonance imaging. The data collection was nearly complete, except for difficulties in quantification of protein levels using slot blot quantification. This work represents a nearly complete study that has not been completely written in manuscript format. A short introduction, materials and methods, figures and new discussion of the results are included. It is incorporated as an appendix in support of using MR for iron imaging.

6.1.1. – Introduction

The role of iron for normal cellular function and body homeostasis has been well established. Normal neuronal functioning requires a tightly regulated iron repository and delivery system in order to maintain optimal amounts of free iron. Normal iron concentration in the body is approximately 40 – 50 mg Fe/kg body weight in women and men (140). Improper management of iron can lead to its progressive accumulation and storage in ferritin and hemosiderin. However, the body’s ability to sequester and store iron in these proteins can be overwhelmed eventually leading to toxic accumulation and systemic overload in
organ systems. This mismanagement of cellular iron can lead to decreased cellular metabolic activity and provides an environment of vulnerability for oxidative damage (78). Accumulation of iron can lead to the production of free radicals through a Fenton-type reaction where reactive oxygen species are produced (279,280). Increasing amounts of evidence over the last decade has pointed toward the causational role of oxidative stress and the pathogenesis of Alzheimer's disease (281). Oxidative damage has been observed in Alzheimer's tissue relative to controls with the up-regulation of antioxidant enzymes indicative of the cellular response to oxidative stress pointing toward the relevance of free radical reactions in the disease process (210,279). Data has shown that iron is required in the pathway leading to amyloid-beta (Aβ) toxicity, suggesting that iron catalyzed production of hydroxyl radicals occurs and excess free iron aids in Aβ induced cell damage (279). The concept of the relationship between iron and Aβ protein aggregates has been addressed within the literature. Ferric iron has been shown enhance the aggregation of synthetic soluble Aβ_{42} (282-284).

Quantification of tissue iron load has been proposed with magnetic resonance imaging techniques for some time. The non-invasive in vivo measurement of hepatic iron stores has occurred with some success (128,141,142). The limitation in the technique revolves around the extremely high iron content of the liver where transverse relaxation rates no longer are able to linearly follow the increasing iron concentration (126,130). Even considering this, MRI has proven to yield reliable estimates of in vivo tissue iron content (123,140). The presence of iron in tissue acts as a natural contrast agent. Iron
is a ferromagnetic compound that causes the dephasing of protons in its vicinity, effectively increasing the relaxation rate of these protons (123). This effect is seen far away from the particle of iron and the size of the hypo-intensities associated with focal iron is an over-exaggeration of particle size (120). The majority of iron in the brain is hemoglobin bound iron (approximately two-thirds), the amount of iron stored in the brain as non-heme iron in ferritin accounts for approximately one-third of the total iron in the brain. The aim of the current study was to measure the concentration of iron in an iron loaded mouse model and compare this measurement to transverse relaxation rates of the same brain structures.

### 6.1.2. – Materials and Methods

*Iron Loading Model*

Thirty-six male C57BL/6 mice were divided into six groups; Controls, 0 week, 2 weeks, 4 weeks, 6 weeks and 8 weeks. Animals were fed ad libitum a diet composed of 0.1% 3,5,5-trimethylhexanoyl-Ferrocene (TMH-Ferrocene) (272) mixed into Teklad Global 18% protein rodent diet (Harlan Teklad, Indianapolis, Indiana, USA) for a period of 0, 2, 4, 6, or 8 weeks for iron loading. Control animals were free fed a normal iron 18% protein rodent diet for the eight week period. Experimental animals from the 2, 4, 6, and 8 weeks groups given the TMH-Ferrocene diet were MR imaged after the last day of feeding. Control animals were imaged on time point zero and at week eight for comparative
purposes. Time point zero-week animals were scanned immediately at the beginning of the study.

**MRI Data Acquisition**

Animals were imaged on a 3.0 T Medspec S300 MR imaging-spectrometer (Bruker Biospin, Ettlingen, Germany) using a home-built gradient (9.5 cm aperture and 100 G/cm). Animals were anesthetized with 1.5% isoflurane (induction and maintenance) in compressed air. An in house built slotted-tube RF coil (160) was used during scans that incorporates a heated water jacket to maintain the animals temperature at 37ºC during scanning protocols. $T_2$ weighted fast spin echo RARE images were taken with a $TR = 2500$, $TE = 8$, 128 x 128 matrix, FOV of 23 x 23 mm, RARE factor of 12, with eight averages for an imaging time of 4 minutes. For $T_2$ parametric mapping, a multi-slice multi-echo $T_2$ protocol with a $TR = 2078$ ms, $TE = 10.0$ ms, 128 x 128 matrix, FOV of 23 x 23 mm, 12 echoes with six averages for a imaging time of 28 minutes. The mGESEPI (285) $T_2^*$ parametric mapping protocol was utilized to minimize tissue boundary artifacts. The parameters for the 3D $T_2^*$ protocol were $TR = 250$ ms, $TE = 5.34$ ms, 128 x 128 x 16 matrix, FOV of 23 x 23 x 10mm, 12 echoes with four averages for an imaging time of 34 minutes. The dimensions of the individual voxels in the protocols were 179 x 179 µm in the in-plane directions and 625 µm in the through plane. Total anesthesia time for each animal during scanning was under 100 minutes. Upon completion of scans animals were taken out of the magnet and allowed to fully recover on a 37ºC heating pad. For the
phantom imaging, a 25cm birdcage head volume coil was used with the internal
body gradient system. Temperature of the phantoms was maintained at 37°C
with a pneumatically driven vortex heater to mimic the temperature of an in vivo
system. When up to set-point temperature was reached and maintained, three
imaging protocols were utilized. A \( T_2 \) weighted multi-slice multi-echo scan was
used to provide \( T_2 \) measurements with a 2000ms TR, 15.32ms TE, 256 x 256
matrix, 250mm x 250mm FOV, composed of two slices with one average at a
slice thickness of 4mm for an imaging time of nine minutes. For \( T_2^* \) relaxation
measurements, a 3D \( T_2^* \) mGESEPI protocol with a TR = 322ms, TE = 8.0ms,
256 x 196 x 16 matrix, FOV of 250mm x 250mm x 8.8 mm, over two slices with
two averages for an imaging time of 33 minutes. For \( T_1 \) relaxation
measurements a variable timing rate multi-slice multi-echo scan was used with a
128 x 128 matrix, 250mm x 250mm FOV, 10mm single slice thickness, with
variable echo times of 30, 50 100, 200, 500, 1000, 2000, 3000, 4000, 6000 and
10,000 ms with one average for a total imaging time of 57 minutes.

**Animal Tissue Preparation**

Upon completion of the final set of imaging protocols animals were deeply
anesthetized with a sodium nebutal barbiturate intraperitoneal injection (100
mg/kg), and weighed. When no pinch response was seen in the animals, they
were transcardially perfused with cold Ringer’s solution till the solution ran clear
then followed by cold 4% paraformaldehyde for 15 minutes. The whole brain
tissue and livers were then excised and weights recorded for each. Tissue
samples were placed in 4% paraformaldehyde overnight at 4°C to allow full fixation. After fixation, tissue samples were re-weighed and placed in two rinses of 0.9% phosphate buffered saline (PBS) for 24 hours each to fully leach out the fixative. Following rinsing the tissue was weighed again and the brains placed in a 0.5mm tissue cutting matrix. The whole brain was cut coronally at a thickness of 500µm with a razor blade and slices were then placed into a twelve well plate with PBS. Regions of interest (RIOs) encompassing the motor/sensory cortex, caudate/putamen, lateral globus pallidus/internal capsule, anterior cerebellum and corpus callosum white matter tracks were selected from these sections following a mouse brain atlas and the T₂* parameter maps. These regions were precisely excised under a dissecting microscope being careful to include only the desired RIOs and then placed in pre-weighted 1.5ml conical tubes to obtain a final wet tissue weight.

Iron Quantification with Atomic Absorption Spectrophotometry

Samples of specific regions of interest and livers for each mouse were pre-weighted and placed in 400 µl of 70% fuming nitric acid at 65 °C for 24 hours until the samples were completely digested. Each sample was then diluted in duplicate 1:100 in 1% Nitric acid in ultra-pure water and standard graphite furnace atomic absorption spectroscopy (AAS) with a Perkin Elmer spectrophotometer was used to measure iron concentrations in the corresponding excised brain regions. Two measurements of each samples iron concentration were completed. A standard curve was generated from serial
dilutions of a calibrated iron standard. If the samples concentration was found to be outside the standard curves upper or lower limits the sample was re-diluted to increase or decrease its concentration to within this limit. Standards were rerun throughout each analysis to verify measurement accuracy over the analysis time. The variation for each replicate and standard was taken and if the measurement varied more than 3% for each duplicate over both measurements the sample was re-diluted and measured again.

*Protein Quantification*

Samples from each RIO and livers for individual week mouse groupings were combined and placed into 400 µl of homogenization buffer consisting of Lysis Buffer A with 1/100 (b/v) Protease Inhibitor Cocktail (PIC) (Sigma # P8340, St. Louis, MI, USA) for deep freeze storage until subsequent processing. Tissue samples were then homogenized with draws through 18g, 21g and 23g needles followed by complete homogenization at 4°C with a Branson S-250A ultrasonic sonifier. Following homogenization samples were centrifuged at 3000g for 15 minutes such that the supernatant containing the protein samples was collected. Pellets were rinsed twice and centrifuged again to collect the remaining protein. Total protein quantification was completed using a Bicinchoninic Acid (BCA) Protein Assay Kit (Pierce, Rockford, IL, USA) with serial Albumin standards and read with a Molecular Devices Emax 96 well precision microplate reader at a wave length of 565nm. Samples were diluted to 1µg total protein/100µl in Lysis Buffer A with PIC. Precise quantification of L-Ferritin, H-Ferritin and Transferrin
proteins was accomplished using a 72 well slot-blot array system (Whatman, Middlesex, UK) transferred onto Protran BA85 0.45µm nitrocellulose film. Protein standards were commercially obtained for Apo-Transferrin (Calbiochem # 616419, Darmstadt, Germany), Recombinant L-Ferritin (Calbiochem # 341491, Darmstadt, Germany) and Recombinant H-Ferritin (Calbiochem # 341490, Darmstadt, Germany). Primary antibodies for L-Ferritin Anti-Rabbit IgG (Santa Cruz Biotechnology Inc. #SC-25616 (H-45) 1/2000, Santa Cruz, CA, USA) and Transferrin Anti-Goat IgG (Santa Cruz Biotechnology Inc. #SC-22599 (M-20) 1/1000, Santa Cruz, CA, USA) were commercially obtained. The primary antibody for H-Ferritin was a monoclonal Anti-Mouse IgG that was generously given to us from Dr. Paulo Avosio and used at a dilution of 1/40,000. Anti-Rabbit IgG Horseradish Peroxidase linked antibody (from donkey) (Amersham Biosciences # NA934V 1/2500, GE Healthcare, Pittsburgh, PA, USA) was used as the secondary antibodies for the L-Ferritin primary. Anti-Goat IgG Horseradish Peroxidase linked antibody (from donkey) (Santa Cruz Biotechnology Inc. #SC-2020 1/2500, Santa Cruz, CA, USA) was used as the secondary antibody for the polyclonal Transferrin. Anti-Mouse IgG Horseradish Peroxidase linked antibody (from sheep) (Amersham Biosciences # NA931V 1/2500, GE Healthcare, Pittsburgh, PA, USA) was used as the secondary antibody for the monoclonal H-Ferritin primary. Western Lightning Chemoluminescence Reagent (PerkinElmer Life Sciences, Boston, MA, USA) was used as a light emitting reaction with horse radish peroxidase, images taken
with a Fujifilm LAS-3000 imaging system and analyzed for density measurement with Fujifilm’s MultiGuage software.

*Phantom Iron Model*

To determine the relationship between iron quantity and $R_2$, $R_2^*$ and $R_1$ relaxation measurements a phantom model was designed and implemented. Relaxation measurements for both average gray and average white matter $T_1$ and $T_2$ were determined from both empirical data and that found in the literature (177,286). The phantoms were made to mimic human gray and white matter $T_1$ and $T_2$ relaxation measurements. Average relaxation values for white matter were determined to be 832ms for $T_1$ and 79ms for $T_2$. Similar average relaxation values for gray matter were determined to be 1330ms for $T_1$ and 109ms for $T_2$. The $T_2$ relaxation parameter was lowered by creating the phantom with differing base percentages of boiled agar in distilled water, specifically Type A agar (Sigma # A6549, St. Louis, MI, USA). To lower the $T_1$ relaxation parameter, gadolinium DTPA (Magnevist, gadopentetate dimeglumine, Berlex, Bayer Schering Pharma AG., Berlin, Germany) was employed in different percentages. The phantoms were poured into 15ml conical tubes and floated in a water bath just above the melting temperature of the agar to allow any bubbles that formed to rise to the surface. After solidification of the agar, they were placed inside a specially built insulated holder and brought up to the equivalent *in vivo* body temperature of 37°C. The temperature of the apparatus inside the MR systems bore was maintained during the scanning protocol with a pneumatically driven
vortex heater. Calculation and regressional analysis based on the differing
gadolinium and agar percentages were used to create equivalent $T_1$ and $T_2$
values of white and gray matter for the two separate phantom types. To mimic
differing brain iron concentrations and approximate this within the phantoms, iron
(III) chloride ($\text{FeCl}_3$) (Sigma # F7134, St. Louis, MI, USA) was used. A stock
solution of $\text{FeCl}_3$ was prepared and mixed into the agar to create final iron
concentrations ranging from 0 µg/g to 240 µg/g wet weight. The $\text{FeCl}_3$ solution
was added to the phantoms just above the melting temperature of the agar and
allowed to slowly solidify to prevent $\text{FeCl}_3$ precipitation and to allow bubbles
formed during mixing to come to the surface.

*Relaxation Measurements and Statistical Analysis*

For the phantom study, MR protocol results were loaded into the
Chips2005 software (Children's Hospital Image Processing System, Cincinnati,
OH) in IDL6.1 (Research Systems, Inc., Boulder, CO). $T_2$, $T_2^*$ and $T_1$
parametric maps and relaxation rates were calculated from the corresponding data with a
linear regression method. Regions of interest were selected with the Chips2005
software and statistically compared. For the $T_2$ maps, the first echo image was
removed for the $T_2$ calculation to reduce the $T_1$ effect.
6.1.3. – Figures

Figure 6.1.3.1

T$_2^*$ measurement of the gray matter phantom doped with FeCl$_3$ to mimic physiological iron concentrations. A combination of 1.475% type A agar and 0.02% gadolinium DTPA created an ideal phantom, which precisely mimicked both T$_1$ and T$_2$ of human gray matter, with values of 109.8 ± 0.145ms for T$_2$ and 1333.5 ± 6.178ms.
T2* - Fe Gray Matter

[Fe] ug/g Wet Weight

T2* (ms)
Figure 6.1.3.2

$T_2^*$ measurement of the white matter phantom doped with FeCl$_3$ to mimic physiological iron concentrations. It was determined that a phantom consisting of 2.20% type A agar and 0.041% gadolinium DTPA accurately reproduced both $T_1$ and $T_2$ relaxation values of human white matter, resulting in values of 79.5 ± 0.0975ms for $T_2$ and 833.9 ± 5.53ms for $T_1$. 
Figure 6.1.3.3

$T_2^*$ parameter map of the iron loaded 37 °C Gray Matter phantom. Higher iron concentrations result in lower signal intensities and thus are darker on the map.
**Figure 6.1.3.4**

$T_2^*$ parameter map of a control mouse (A) and one fed the high iron diet for eight weeks (B). High iron regions show as dark hypo-intensities on the maps. $R_2^*$ relaxation rates for the caudate/putamen regions of interest are included. The two parameter maps were created using the same threshold values and are thus comparable for color/contrast.
Figure 6.1.3.5

$R_2^*$ relaxation rates for the caudate/putamen. Iron loading with the TMH-Ferrocene over the eight week course resulted in increases in $R_2^*$ relaxation. The data is the average relaxation rate for the groups indicated with standard error bars.
Figure 6.1.3.6

$R_2^*$ relaxation versus iron concentration. The average $R_2^*$ relaxation and iron concentration for the caudate/putamen are included from animals feed the high iron diet for 2 through 8 weeks. The average values have a high correlational coefficient for the dependence of $R_2^*$ transverse relaxation on iron concentration.
The graph shows the relationship between $R_2^*$ relaxation and iron concentration. The equation of the line is $y = 4.6259x - 58.501$ with an $R^2$ value of 0.9717. The graph plots $\mu g$ Fe/Wet Weight (g) on the y-axis and $R_2^*$ (1/sec) on the x-axis.
6.1.4. – Brief Discussion and Results

The creation of the phantoms in retrospect was a rather challenging task. The co-variation of both $T_1$ and $T_2$ relaxation with agar and gadolinium concentration required extensive testing of different concentrations of both substances. It originally was believed that gadolinium would affect $T_1$ relaxation while leaving $T_2$ relaxation relatively unchanged and conversely for agar concentration which was hoped to affect $T_2$ more profoundly then $T_1$. While this was generally true, it was found that both changed $T_1$ and $T_2$ substantially enough that they were co-variables in the consideration of overall effect on transverse and longitudinal relaxation. After careful experimentation along with trial and error, the concentrations of agar and gadolinium were found that approximated the white matter and gray matter relaxation values very accurately and reproducibly over different experiments. It was then realized that these values were obtained at room temperature, which was not congruent with the desire to create phantoms that approximated near in vivo conditions. To account for this, the phantoms were heated in a specially built chamber and retested at 37 °C. It was found, to no surprise, that the data no longer held as approximations of white and gray matter $T_1$ and $T_2$ relaxation. The phantoms were recalculated at physiological human temperature with new agar and gadolinium concentrations.

Doping of the white and gray matter phantom with iron proved to be very reproducible over multiple experiments. The data suggests that there is a nearly linear relationship between iron and transverse $R_2^*$ relaxation within the
physiological range of about 80 – 200 µg/g wet weight. \( R_2 \) and \( R_1 \) relaxation rates were found to not respond in a linear fashion to iron doping of the phantoms, suggesting that \( T_2^* \) relaxation was the best option to measure and quantify iron with MR imaging technology (Figure 6.1.3.1 – 6.1.3.3). After careful consideration of the data, it was believed that there is an inherent drawback to the phantom data. Iron in the form of ferric chloride was used to dope the phantoms. This would represent iron in a free form within the brain while the majority of iron is found within the core of the ferritin complex. Transverse relaxation responds differently to the ferritin complex with its crystalline ferricydrite core. Attempts were made to create a phantom that used variable amounts of iron loaded horse sleep ferritin along with specific concentrations of gadolinium and agar. The hope was to create a phantom that more accurately represented the iron milieu of the brain. The trek along this path was started but ultimately not concluded due to the considerable difficulty there was in including four variables (agar, gadolinium, ferritin and \( \text{FeCl}_3 \)) in the calculation of each individual phantom. The existing phantom showed that \( T_2^* \) relaxation was more sensitive than either \( T_1 \) or \( T_2 \) in measuring iron concentration with MRI.

Loading of the mice with the lipophilic iron compound TMH-Ferrocene proved to be moderately successful. The \( R_2^* \) relaxation measurements for regions of interest largely proven unsuccessful and erratic; mostly due to difficulties in imaging cortical regions with the imaging protocol due to errant susceptibility artifacts caused by the air/tissue boundary in the ears. This was not the case for deep brain structures such as the caudate/putamen which were
generally free of artifacts and produced clear results of a linear increase in $R_2^*$ with iron loading time. Figure 6.1.3.4 shows $T_2^*$ parameter maps of two mice, one feed a normal iron diet and the other loaded with iron. The relaxation values for the caudate and putamen are seen in figure 6.1.3.5 which shows a nearly linear increase in $R_2^*$ relaxation in this region with differing iron loading times. Iron concentration measured with AAS proved to be difficult but the results were promising as far as reproducibility. There was some variability in the iron concentration data due mostly to the minute amounts of tissue present. Weighing and measuring iron concentrations in such small amounts of tissue can introduce some variability into the dataset, based on accuracy of the detection methods (spectrophotometer and scale). Even with these considerations; there is a high correlational coefficient between $R_2^*$ relaxation dependence upon iron concentration (Figure 6.1.3.6) for the caudate and putamen region of interest. Overall the data here are evidence of the potential for quantitative magnetic resonance imaging to measure iron concentration is tissue. There is a limit to the range of the iron concentration that could be measured, as the relaxation largely plateaus or becomes statistically insignificant when a threshold of iron is imaged. These data was collected at a field strength of 3.0 T, which is currently found in many clinical scanners worldwide. It was hoped that a calibration curve would be produced to accurately quantify iron concentration based on $R_2^*$ relaxation rate. The data show there this has promise but clearly further study would have to be undertaken in a human system. It is inherently
difficult to create an accurate calibration curve with a phantom system or a mouse model to be used within human clinical imaging.
APPENDIX B
Histological Staining Techniques

7.2 – Histological Staining Techniques

The mechanism of action of the various histological staining techniques used throughout this work are described in detail, along with staining procedures for each.

7.2.1 – Thioflavin-S

Thioflavin can refer to either of two benzothiazole salt dyes used for fluorescent histology staining. The two derivatives are known as Thioflavin T (Figure 7.2.1.1) and Thioflavin S (Figure 7.2.1.2). Thioflavin T is obtained by the methylation of dehydrothiotoluidine with methanol in the presence of hydrochloric acid. Thioflavin T is made from the methylation and sulphonation of dehydrothiotoluidine with methanol and sulfonic acid (203). Thioflavin S synthesis results in a mixture of planar asymmetric dyes with charged groups at the end of the molecule. They are believed to interact with the crossed β-sheet structure that is common to amyloid plaques. The component β-strands that make of the β-sheets have alternating side chains that when stacked upon one another form channels that run along sheet perpendicular to the strands (Figure 7.2.1.3). Binding studies suggest that the thioflavins bind to the amyloid fibers in these channels. The molecular axis of the thioflavin lies parallel to the amyloid fiber surrounded by charged side chains (287). The elongated planar molecular
structure and negative charge at the ends of thioflavin S allow it to form multiple bonds with numerous side chains along the channel giving it a particularly high affinity for amyloid fibers that have a β-sheet conformation (42,203,288,289). The dyes will both bind to fibrillar strands of beta-amyloid that are composed of protein strands that are 28, 40 and 42 amino-acids long (Figure 7.2.1.4). The charged side chain region of the amyloid peptide resides in the first 28 amino acids. In addition, a change in pH of the aggregation solution outside physiological boundaries and thus conformational changes in protein structure and amino-acid charges cause a reduction in thioflavin binding. This suggests that both bind due to the multiple charges found on the side chains of the amino-acids and not due to hydrophobic amino-acid interaction (288). Thioflavin S fluorescence is stimulated by excitation at 385 nm and emits at 482 nm, allowing the usage of standard FITC filter cubes for microscopy. The emission spectra for thioflavin S is unchanged from that of the dye in a free solution. Thioflavin T in contrast undergoes an excitation spectra change when in the presence of aggregated Aβ with a peak shift towards 450 nm. This suggests that thioflavin T undergoes a slight conformational change when bound to the parallel axis channels. Thioflavin T thus has less background fluorescence than thioflavin S due to this shift in emission from the native dye and is used for the quantification of fibrillar amyloid in solutions due to this. The additional binding characteristics of thioflavin S make it ideal for usage in histological staining and imaging of beta-amyloid plaques due to its resistance to wash out of the channels during staining, but it cannot be used for quantification of fibrillar amyloid formation in solution.
Of considerable interest to this research project is the fact that thioflavin S and T will not bind to amyloid amino acid monomers while they will bind to longer oligomeric species. This is understandable considering that the dyes bind to the channel regions of the long fibrillar Aβ strands. This holds true for diffuse verses fibrillar Aβ species also. Thioflavin will favorably bind to fibrillar Aβ plaques but binds poorly to diffuse plaque formations.
Figure 7.2.1.1

The molecular structure of thioflavin-T; a benzothiazole salt obtained by the methylation of dehydrothiotoluidine with methanol in the presence of hydrochloric acid.
The molecular structure of thioflavin S; a benzothiazole salt obtained by the methylation of dehydrothiotoluidine with methanol in the presence of sulfonic acid. The structural formula for thioflavin S can be altered from a tri-ring structure, like that of thioflavine T, to contain five or seven rings.
Figure 7.2.1.3

Thioflavin-T and diagram of a β-sheet. (A) Structure of thioflavin-T. The molecule is 4.3 ± 0.1 Å thick. (B) A schematic representation of a β-sheet (290). Indicated are the backbone atoms (N, C, and Ca) and the side chain (R) for one residue. No hydrogen bonds are indicated and this scheme is valid for both parallel and anti-parallel sheets. If part of a fibril, the long axis of the fibrils would be perpendicular to the surface of this page. One of the binding channels is indicated with a double headed arrow. It is thought thioflavin-T binds with its long axis parallel to the long axis of this arrow. In a fibril it is likely only one of the faces of the sheet would be accessible from the solvent. (C) Schematic representation of a protofilament with, arbitrarily, three β-sheets. Individual β-strands are shown by ‘zig-zag’ lines in black, dark and light grey. Sidechains accessible from the solvent are shown as black circles and point out of the plane of the paper. Note that sidechains on the third, light grey, β-sheet are also accessible from the solvent and point below the plane of the paper (not shown for clarity). Dye molecules are represented by double headed arrows. When several protofilaments form a fibril, some of these surfaces will be obscured, resulting in impeded dye binding in those areas. (Figure and figure text from Krebs, 2005 (287))
Figure 7.2.1.4

Amino Acid sequence of Human Beta-Amyloid 1-40 and 1-42 with expected charges on AA’s at pH 6.
DAEFRHDSGYEVHHQKLVFFAEDVGSNKGAIIGLMVGGVIA

β(1-40)

β(1-42)
7.2.1.1 – Thioflavin-S staining procedure

Staining procedure is adapted from traditional thioflavin S staining procedures and is as follows:

1. The tissue sections are brought to up to dH$_2$O
2. The sections were stained with 1% Thioflavin-S aqueous solution for 5 minutes.
3. Differentiated in 70% ethanol for 5 minutes with slight agitation throughout.
4. Two final five minute dH$_2$O rinses.
5. For aqueous mounting: Mount with FluorSave Aqueous mounting medium and coverslip.
   For dehydrated mounting: Dehydrated mounting of Thioflavin S stained tissue sections is not possible. This is because the ethanol and xylene completely leech the Thioflavin S from the section.

The 1% Thioflavin S solution is mixed with dH$_2$O and allowed to mix completely with a vortex. A slight residual amount of insoluble material is left which can be filtered through a syringe attachment.

7.2.2 – Perl’s Stain for Iron

Perl’s stain is a highly sensitive and widely used classic metallic-staining method to demonstrate ferric iron (Fe$^{3+}$) found in tissue samples. Dilute mineral acid hydrolysis with dilute hydrochloric acid releases ferric ions from protein bound tissue deposits. These ions then react with (potassium) ferrocyanide ions
(potassium hexacyanoferrate) to produce a dark blue colored and highly water-insoluble complex known as Prussian blue or chemically as ferric ferrocyanide (ferric hexacyanoferrate) (291).

It has generally been considered that the Prussian blue reaction only reacts with ferric (Fe$^{+3}$) iron, but there is some speculation that it also reacts with ferrous (Fe$^{+2}$) iron as well but to a lesser degree (200,292). The Prussian blue reaction of potassium ferrocyanide with ferric iron (Fe$^{+3}$) to create ferric ferrocyanide (Prussian blue)(ferric hexacyanoferrate) is shown in equation 1 below. Utilizing a 3,3'- diaminobenzidine (DAB) (3,3',4,4'-Tetraamino-diphenyl) (Figure 7.2.2.1) enhanced method of the Perl’s stain helps to visualize trace amounts of iron in tissue samples (293). DAB acts as a chromogen dye as ferric ferrocyanide catalyzes the oxidation of benzidine to a polymer by hydrogen peroxide, which gives the brown colored precipitate. The insoluble precipitate may range in color from a light golden brown to dark brown depending upon the amount of catalyst present. The brown precipitate is collectively known as benzidine brown and is composed of many potential stereo-isomer configurations of an azopolymer like that in figure 7.2.2.2 (294). Care must be taken not to allow tissue samples to reside in non-buffered formalin solutions as an acidic environment will leech iron out of tissue into the liquid. Using ~7.3 pH buffered para-formaldehyde or reducing the fixation time is recommended. The acidic nature of the Perl’s stain itself also can leech iron out of the tissue samples if the concentration of the hydrochloric acid is too great or the time in solution in too
long. DAB is a suspected carcinogenic agent as its basic benzidine core is linked to bladder cancer and pancreatic cancer.

The Prussian blue reaction of potassium ferrocyanide with ferric iron (Fe$^{3+}$) to create ferric ferrocyanide (Prussian blue) in an acidic environment is shown below:

$$4 \text{FeCl}_3 + 3 \text{K}_4\text{Fe(CN)}_6 \rightarrow \text{Fe}_4[\text{Fe(CN)}_6]_3 + 12 \text{KCl}$$

The oxidation of DAB with H$_2$O$_2$ in the presence of ferric ferrocyanide is shown below:

(1) $$2 \text{Fe(CN)}_6^{-4} + \text{H}_2\text{O}_2 \rightarrow 2 \text{Fe(CN)}_6^{-3} + 2 \text{OH}^{-1}$$

(2) $$\text{Fe(CN)}_6^{-3} + \text{OH}^{-1} + \text{DAB} \rightarrow \text{Fe(CN)}_6^{-4} + \text{Oxidized DAB (benzidine brown)}$$
Figure 7.2.2.1

Molecular structure of 3,3'-diaminobenzidine (DAB) (3,3',4,4'-Tetramino-diphenyl)
Figure 7.2.2.2

Representative stick bond molecular structure of potential bonding seen in the polymerization of DAB into the water insoluble precipitate benzidine brown. There are any number of possible polymers formed as each amine group can form a double bond to a neighboring amino group (X).
7.2.2.1 – Traditional Perl’s Stain Procedure

The basic Perl’s stain procedure for sliced tissue (10 – 60 µm) is as follows:

1. Fix tissue in 4% Paraformadehyde for 10 minutes to complete fixation.
2. Tissue must be hydrated in dH₂O to leech out residual formalin in the samples for at least 5 minutes.
3. Place the samples in solution of 2% Hydrochloric Acid and 2% Potassium Ferrocyanide for 30 minutes.
4. Rinse in dH₂O for 5 minutes.
5. Immerse in diaminobenzidine / H₂O₂ solution for 2 – 4 minutes.
6. Rinse twice in dH₂O for 5 minutes each.
7. For aqueous mounting: Mount with FluorSave Aqueous mounting medium and coverslip.
   
   For dehydrated mounting: Dehydrate in ethanol gradient, 95%, 100% x2 for 5 minutes each, followed by 100% xylene for 5 minutes twice. Mount with Cytoseal 60 mounting medium and coverslip.

Necessary chemicals and preparation for stain:

2% Hydrochloric Acid and 2% Potassium Ferrocyanide:

Consists of 100ml of 4% Hydrochloric Acid and 100 ml of 4% Potassium Ferrocyanide made separately until mixed together just prior to staining. It is recommended to use H₂O that is relatively free of iron contamination and using
well washed glassware, possibly washed with nitric acid to eliminate all trace iron.

3,3'-diaminobenzidine (DAB)

Twenty ml final volume consists of 13.33 mg DAB (Sigma-Aldrich, D5637) + 20 ml PBS, when ready to use mix in 16 µl of 30% H$_2$O$_2$. Care must be taken when staining and disposing of product as it is carcinogenic. Dispose of product with sodium hypochlorite (bleach)

7.2.2.2 – Modified Perl’s Stain with Proteinase K

Modified Perl’s Stain Procedure

The stain was adapted and modified from LeVine (188,189) and employs the usage of proteinase K and detergents to break down the periphery of the plaques forming openings in the highly hydrophobic amyloid plaques. This allows the aqueous Perl’s stain to infiltrate into the amyloid mass and is sensitive enough to detect trace amounts of iron in the Aβ plaques.

The staining method consists of the following procedure.

1. Tissue sections were hydrated in PBS for 15 minutes

2. Immersion in sodium borohydride (10 mg/ml PBS, # 213462, Sigma, St. Louis, MO) for 30 minutes.

3. rinsed in PBS twice for five minutes
4. then immersed in proteinase K (30 µg/ml, # P6556, Sigma, St. Louis, MO) and 0.01% Triton X-100 in PBS for 20 minutes.
5. sections were then placed in a solution of 1% hydrochloric acid, 1% potassium ferrocyanide and 1% Triton X-100 in distilled water for 30 minutes.
6. Amplification of the iron staining was accomplished with 0.5 mg/ml 3,3′-diaminobenzidine tetrahydrochloride (DAB) and 2µl/ml of 30% hydrogen peroxide in pH 7.6 0.05 M Tris HCl for 15 minutes.
7. Rinse twice for 5 minutes in dH₂O.
8. For aqueous mounting: Mount with FluorSave Aqueous mounting medium and coverslip.

For dehydrated mounting: Dehydrate in ethanol gradient, 95%, 100% x2 for 5 minutes each, followed by 100% xylene for 5 minutes twice. Mount with Cytoseal 60 mounting medium and coverslip.

7.2.3 – Thioflavin S and Perl’s co-staining.
Traditionally, individual stains for ferric iron and beta-amyloid plaques would be carried out on sequential slices of tissue samples and compared to one another. It was found necessary to develop a co-staining technique that allowed both Perl’s staining and thioflavin S staining on the same tissue piece. This was required because it permitted the imaging of a piece of tissue with the MR system and subsequent co-staining, which has proved to be essential for this study. The procedure follows sequentially through the Perl’s stain then proceeds
to the thioflavin-S stain, both procedures are outlined above. To test for any confounding variables that occurred during the co-staining method, tissue samples were stained with the Perl’s and thioflavin-S methods alone and compared to the co-stained tissue. The procedures used during the thioflavin-S staining do not alter the Perl’s results, as the Prussian blue reactant and DAB precipitate are not soluble in the aqueous or ethanol based solutions used in the thioflavin-S stain.

7.2.4 – Antibody Staining

The basic principle behind immunohistochemistry is the usage of specific antibodies to localize antigens in tissue through antigen-antibody interactions. Its usage is advantageous because it involves very specific primary antibody-antigen binding, and thus allows the staining of specific antigens that are restricted to one type of protein or cell for instance. With the usage of this technique with fluorescent secondary antibodies multiple stains for different antigens is possible. Various immunohistochemistry techniques have become a widely used with the medical research community and have become vital in numerous scientific discoveries.

The selection of primary and secondary antibodies is critical to successful staining of tissue samples. Primary antibodies must be chosen for a target antigen and able to bind with selective specificity. Different antibodies are raised in different host systems, and selection of the host system is important for secondary antibody specificity for the primary antigen. For multiple staining
techniques, primary antibodies should be chosen from two (or more, depending on the total number of co-stains) host systems such that the secondary antibodies will be specific for the host system used. Fluorescent secondary antibodies should not only be chosen for a specific house match to the primary antibody, but also must be chosen to fluoresce under a different wavelength (color) such that they can each individually be viewed under the microscope at their respective wavelengths. Other important considerations are the blocking of tissue samples and antigen retrieval. To allow the specific binding of the primary and secondary antibodies to their targets, the tissue must be washed in a mixture of protein wash. This allows the selective binding of the primary and secondary antibody due to the extremely high binding affinity of these antibodies to their antigen targets, they essentially outcompete other proteins for these target sites. The majority of tissue samples stained with these methods are fixed prior to staining which creates cross-links between and within proteins. In order to have the antibodies bind specific proteins that have been cross-linked, proteins must have these cross-links broken to open up their binding sites. This can be done in a variety of ways, with the two most popular methods being heat induced epitope retrieval and proteolytic induced epitope retrieval. Heat induced retrieval literally heats the tissue sample up to break the cross-links, which the proteolytic approach uses enzyme digestion.

7.2.4.1 – Antibody Staining Techniques

The immunohistochemistry staining technique used in these studies is as follows:
1. Warm slides with tissue sample on them at 50 °C to help bind fatty tissue to the slides, place slides in slide holder.

2. Place slides in 95 °C citrate buffer for 15 minutes (heat induced epitope retrieval).

3. Rinse slides three times in PBS for five minutes each.

4. Use a hydrophobic barrier pen (PAP pen) to create individual wells around each tissue sample.

5. Incubate tissue with 1% bovine serum albumen (BSA) (A9430, Sigma Aldrich, St. Louis, Mo). in PBS-Tween (PBST) for 30 minutes to block unspecific binding of antibodies.

6. Decant the blocking mixture, and incubate tissue with the mixture of properly diluted primary and secondary antibodies in 1% BSA in PBST overnight at room temperature.

7. Decant mixture, and rinse with PBS three times for five minutes each.

8. Incubate tissue with the mixture of fluorescent antibodies in 1% BSA for at least one hour at room temperature. This must be done in the dark.

9. Decant mixture and rinse three times with PBS for five minutes each.

10. Mount tissue with an anti-fade fluorescent mounting media and coverslip.

The solutions used in these stains are as follows:

Phosphate Buffered Saline with Tween:

Mixture of PBS with 0.1% Tween 20x (P1379 Sigma Aldrich, St. Louis, Mo).
Citrate buffer 10mM:
1.92g / l of Sodium citrate tribasic dehydrate (C7254 Sigma Aldrich, St. Louis, Mo) in dH$_2$O. Adjust pH to 6.0.

7.2.5 – Fluoro-Jade C

Fluoro-Jade C (FJC) is a poly-anionic fluorescein derivative whose exact molecular formula and chemical structure currently remains to be elucidated. Being a derivation of the basic fluorescein molecule, Fluoro-Jade C must have at least some similarity to the known structures of fluorescein (Figure 7.2.5.1) and Fluorescein isothiocyanate (FITC) (Figure 7.2.5.2). Recently, the compound has been reported as useful for detecting neuronal specific degeneration in brain slices (295). Based on the fact that Fluoro-Jade C is a known anionic compound, it is speculated that the biochemical molecules that the compound binds to would include molecules with multiple positive charges on them. There is some evidence that molecules with a strong electrostatic attraction to Fluoro-Jade –C include spermidine, putrescine, and cadaverine (296,297). Putrescine and cadaverine are both organic chemical compounds consisting of four (putrescine) or five (cadaverine) carbon atom chains with amine groups at the first and last carbons. Both molecules are produced as a result of the breakdown of amino acids in living cells. Spermidine is a similar polybasic amine with an amine group situated between three and four carbon chains with amine at the terminal ends. Although it has not been resolved what exactly FJC binds to or its mode of
biochemical action, it is known that it binds to (believed to be intracellular) and thus detects dead neurons regardless of the cause of death.
**Figure 7.2.5.1**

Molecular structure of fluorescein, a fluorophore commonly used in microscopy. Fluorescein dissolved in water has an absorption maximum at 494 nm and emission maximum of 521 nm. Fluoro-Jade C has an emission peak at 450 nm and excitation at 530 nm.
Figure 7.2.5.2

Molecular structure of fluorescein isothiocyanate (FITC). The molecular structure of FITC used the original fluorescein molecule as a backbone with a different isothiocyanate functional group (\(-N=\text{C}=\text{S}\)), replacing a hydrogen atom.
7.2.5.1 Fluoro-Jade C Staining Procedure

Basic Tissue Preparation:

Tissue samples were fixed with 4% paraformaldehyde and cryogenically protected with a gradient of sucrose solutions up to 30% sucrose in distilled water. Tissue samples were frozen at -18 °C and cut at a thickness of 16 µm with a cryostat. Cut samples were thawed on slides double coated with gelatin and poly-lysine and stored at -80 °C until usage. The following procedure was adapted from the manufacture’s (Chemicon International) instructions:

1. If co-staining with an antibody stain (fluorescent or otherwise), follow normal antibody procedure until after the secondary antibody ready to be discarded from the slides.
2. Rinse the slides twice for five minutes each in PBS.
3. Slides were dried for 30 minutes at 50°C.
4. Rinse the slides for two minutes in distilled water.
5. Immerse the slides in 0.06% Potassium Permanganate for ten minutes.
6. Rinse the slides for two minutes in distilled water.
7. Immerse the slides in the working 0.0001% Fluoro-Jade C solution for ten minutes.
8. Rinse the slides three times for one minutes each.
9. Air dehydrate the slides for 30 minutes at 50°C.
10. Immerse the slides in xylene twice for five minutes each.
11. Mount with cytoseal, or another xylene based mounting media, and coverslip.

Necessary chemicals and preparation for stain:

Fluoro-Jade C stock solution consists of 0.01% in distilled water. The working Fluoro-Jade C solution is made by the addition of 1ml of stock solution in 99 ml of 0.1% acetic acid vehicle.

Potassium Permanganate solution is made is distilled water to a final concentration of 0.06%.

7.2.6 – Transmission Electron Microscopy Procedure

A. Fixation

Fixative used for EM is 2% paraformaldehyde and 2% gluteraldehyde in 1x PBS

1. Harvest the tissue as quickly as possible and place it in cold fixative. Ideal size of tissue pieces is 1mm cubed.
2. Fix the tissue for 2 hours in cold fixative on ice.
3. Rinse with cold PBS on ice three times for 10 minutes each.
4. Fix the tissue in 1% PBS buffered osmium tetroxide overnight in the refrigerator.
5. Rinse with cold PBS on ice three times for 10 minutes each.

B. Dehydration

1. 50% ethanol for 15 minutes.
2. 70% ethanol for 30 minutes.
3. 90% ethanol for 15 minutes.
4. 95% ethanol for 15 minutes.
5. 95% ethanol for 15 minutes.
6. 100% ethanol for 30 minutes.
7. 100% ethanol for 30 minutes.
8. 100% ethanol for 30 minutes.
9. 100% propylene oxide for 30 minutes.
10. 100% propylene oxide for 30 minutes.

C. Embedding

Embedding resin is composed of 10ml Embed 812, 8ml Dodecenyl Succinic Anhydride (DDSA), 4ml Methyl-5-Norbornene-2,3-Dicarboxylic Anhydride (NMA) and 0.35ml 2,4,6-Tri(dimethylaminomethyl) phenol (DMP-30). Any volume made with these volumetric ratios is sufficient. All chemicals can be purchased as a kit from Electron Microscopy Sciences (#14120 - EMbed-812 Kit, EMS, Hatfield, PA).

1. Propylene oxide : resin (2:1, volume/volume) for 1 hour.
2. Propylene oxide : resin (1:1, volume/volume) for 1 hour.
3. Propylene oxide : resin (1:2, volume/volume) for 1 hour.
4. 100% resin for two hours.
5. 100% resin for two hours.
6. 100% resin overnight.
7. Place specimens in molds, pour fresh vacuumed resin over them and place them in a vacuum chamber for 2 hours.

8. Place molds in 60°C oven overnight to cure resin.

D. TEM Tissue Preparation

1. Cut tissue from hardened resin with a jeweler’s saw and mount onto chuck.

2. Cut tissue at 90nm with a diamond knife on an ultra-microtome.

3. Float slices on dH₂O and place them on 200 mesh copper grids.

4. Stain with 2% aqueous uranyl acetate for 10 minutes.

5. Stain with Reynolds lead citrate for 5 minutes.

6. Rinse with dH₂O and dry sections prior to TEM imaging.


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