Understanding the Molecular Links Between Exercise and Brain Health Using Pyrosequencing Technology

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# Understanding the molecular links between exercise and brain health using pyrosequencing technology

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## Abstract

Alzheimer’s disease (AD) is a neurodegenerative disease and the 6th leading cause of death in the U.S. Although no cure exists for AD, exercise has been shown to have neuroprotective effects, possibly through protein regulation. Epigenetics – the study of heritable modifications that influence gene expression without changing the underlying DNA sequence – can impact protein regulation and may link exercise with AD protection. Our study focuses on an epigenetic modification called DNA methylation.

We plan to utilize a novel rat model of AD to study how exercise training affects behavioral and physiological outcomes. In the geriatric stage, rat brain tissue (n=36) was collected for DNA methylation analysis. We selected AD-related genes for analysis including brain derived neurotrophic factor (BDNF), insulin-like growth factor (IGF-1), and vascular endothelial growth factor (VEGF). Here, we will demonstrate how a cutting edge technology called pyrosequencing will be used to measure differences in DNA methylation between exercise and control groups. We expect to find hypomethylation in BDNF, IGF-1, and VEGF in response to exercise, and this can influence protein production. Findings from this study could improve our understanding of the neuroprotective effects of exercise for AD patients.

## Project Overview

### Methods

1. **Exercise and Behavior Protocol**
   - AD rat model: TgF344-AD replicates human AD pathology and behavioral impairment
   - Four treatment groups: Wild type sedentary (WS), wild type treadmill training (WTT), Alzheimer’s disease sedentary (ADS), Alzheimer’s disease treadmill training (ADTT)
   - At 12 months, WTT and ADTT groups begin treadmill training 5 days per week
   - At 3, 6, 9, 12, 15, and 18 months, functional tests were given to measure memory (Morris Water Maze), motor coordination (Rotarod), and strength (Grip Strength)

2. **Tissue collection/tissue disruption**
   - At 18 months, brain, blood and skeletal muscle were collected
   - Hippocampal cells were dissected from brain tissue
   - Tissues were stored at -80°C

3. **DNA/RNA extraction**
   - DNA and RNA were extracted from tissues using QiaGen AllPrep DNA/RNA Kit

4. **DNA bisulfite treatment**
   - Used the QiaGen EpicTect Fast Bisulfite Kit
   - Unmethylated cytosines are converted to uracil
   - Methylated cytosines are not converted
   - Allows us to determine which cytosines are methylated

5. **Primer design**
   - The genes we chose to analyze for methylation were BDNF, IGF-1, and VEGF
   - Within these genes, regions of variable methylation were selected based on literature findings
   - Primers were designed using PyroMark Assay Design software to amplify region of interest

6. **Polymerase Chain Reaction (PCR)**
   - Amplification of target region using PCR
   - Designed primers, Taq polymerase, and nucleotides are used
   - Steps of PCR:
     1. Denaturation: double stranded DNA is separated into two single strands of DNA
     2. Annealing: primers bind to complementary DNA strands
     3. Extension: Taq polymerase adds nucleotides to synthesized strands

7. **Gel Electrophoresis**
   - Agarose gel electrophoresis was used to verify amplified DNA

8. **Pyrosequencing run design**
   - Used PyroMark Assay Design software to generate a run file including:
     1. Sequence to analyze
     2. Dispensation order of nucleotides

9. **Pyrosequencing**
   - Used PyroMark OA48 Autoprep software to run pyrosequencing
   - One nucleotide is added at a time based on the sequence to analyze and the dispensation order
   - At possible methylated CpG sites, a cytosine is dispensed, followed by a thymine
   - Methylated cytosines will add a cytosine to the sequence
   - Unmethylated cytosines will add a thymine to the sequence because the bisulfite treatment converted cytosine to uracil
   - DNA methylation patterns determined by percent methylation at each CpG site within the region of interest for each gene
   - DNA methylation patterns will be analyzed for differences between treatment groups

## Candidate Genes

### Brain-derived neurotropic factor (BDNF)

- BDNF is involved in neurogenesis, neuronal survival and differentiation, and helps regulate synaptic plasticity, learning, and memory
- Aerobic exercise increases levels of BDNF in the hippocampus, which is associated with better memory and larger hippocampal volumes
- In the next column, a study shows a significant difference between exercised and sedentary rats in DNA methylation in the BDNF exon IV promoter at the -148 CpG site.

### Vascular endothelial growth factor (VEGF)

- VEGF is involved in angiogenesis, vasculogenesis, and hippocampal neurogenesis
- Exercise has been shown to increase mRNA expression of VEGF
- Below a study shows significant methylation difference between exercised and sedentary rats in exon 1 of VEGF

## References