Brain-derived neurotrophic factor, epigenetics in stroke skeletal muscle, and exercise training

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Abstract

Objective

(1) To compare paretic (P) vs nonparetic (NP) skeletal muscle brain-derived neurotrophic factor (BDNF) and the effects of resistive training (RT) on systemic and skeletal muscle BDNF mRNA expression in stroke; and (2) to compare the DNA methylation profile for BDNF and BDNFAS (BDNF antisense RNA) between P and NP muscle and the effects of aerobic exercise training (AEX) on DNA methylation in stroke.

Methods

In this longitudinal investigation, participants (50–76 years) with chronic stroke underwent a fasting blood draw, a 12-week (3×/week) RT intervention (n = 16), and repeated bilateral vastus lateralis muscle tissue biopsies (n = 10) with BDNF expression determined by RT-PCR. Five stroke survivors completed 6 months of AEX (3×/week) and had bilateral muscle biopsies. DNA methylation status in gene BDNF and BDNFAS was assessed by Illumina 450k methylation array.

Results

P muscle had ~45% lower BDNF mRNA expression than NP muscle (6.79 ± 1.30 vs 10.52 ± 2.06 arbitrary units [AU], p < 0.05), and P muscle exhibited differential methylation status in the DNA sequences of BDNF (3 CpG [5′-C-phosphate-G-3′] sites, p = 0.016–0.044) and BDNFAS (1 CpG site, p = 0.016) compared to NP. Plasma BDNF and muscle BDNF messenger RNA (mRNA) expression did not significantly change after RT. BDNFAS DNA methylation increased after AEX in P relative to NP muscle (p = 0.017).

Conclusions

This is the first evidence that stroke hemiparesis reduces BDNF skeletal muscle expression, with our findings identifying methylation alterations on the DNA sequence of BDNF and BDNFAS gene. Preliminary results further indicate that AEX increases methylation in BDNFAS gene, which presumably could regulate the expression of BDNF.
Glossary

AEX = aerobic exercise training; BDNF = brain-derived neurotrophic factor; BDNFAS = BDNF Antisense RNA; NP = non-paretic; P = paretic; RT = resistive training.

It is well established that stroke leads to long-term disability and cognitive dysfunction. Physical activity improves motor and cognitive ability perhaps, in part, due to enhanced neurotrophic factor signaling. Brain-derived neurotrophic factor (BDNF) is a protein that promotes neuroplasticity, including the growth and survival of neurons, regulation of axonal and dendritic branching, and synaptic transmission. BDNF may mediate the improvement in cognition after acute exercise. Yet, it is unknown whether BDNF changes with exercise training in neurologic conditions, such as chronic stroke. We hypothesize that BDNF is involved in the skeletal muscle changes poststroke and that exercise (resistive training [RT] or aerobic exercise training [AEX]) exerts beneficial effects through epigenetic regulation of BDNF expression. Therefore, in this 2-part preliminary investigation, we sought to compare paretic (P) vs nonparetic (NP) skeletal muscle BDNF and the effects of RT on systemic and skeletal muscle BDNF mRNA expression in stroke and to compare the DNA methylation profile for BDNF and BDNF antisense RNA (BDNFAS) between P and NP muscle and assess the effects of AEX on DNA methylation in stroke.

Methods

Twenty-one participants who had a prior stroke aged 43–81 years with body mass index between 21 and 39 kg/m² who had mild-to-moderate hemiparetic gait deficits and completed conventional rehabilitation therapy provided written informed consent to participate in this longitudinal investigation at the Baltimore Veterans Affairs Medical Center and the University of Maryland School of Medicine. Evaluations included medical history, physical examination, and fasting blood profile. Participants with stroke were excluded for unstable angina, congestive heart failure (NYHA II), severe peripheral arterial disease, major poststroke depression, dementia, severe receptive aphasia, and orthopedic or chronic pain conditions. All methods and procedures were approved by the institutional review board of the University of Maryland and the VA R&D Committee.

Tests

Height, weight, fat mass, lean tissue mass, and %body fat were determined by dual-energy x-ray absorptiometry (Prodigy LUNAR GE version 7.53.002). Exercise testing with open-circuit spirometry was conducted to measure peak aerobic capacity (VO₂ peak) using a graded treadmill test. One-repetition maximum strength tests assessed leg press and leg extension strength of each leg (Keiser, Fresno, CA).

Resistive training

Participants trained 3×/week for 12 weeks, performing 2 sets of 20 unilateral repetitions on the leg press, leg extension, and leg curl machines (Keiser; pneumatic resistance, Fresno, CA) at every session.

Aerobic exercise training

The progressive treadmill protocol consisted of three 40-minute sessions per week at a target aerobic intensity of 60%–70% heart rate reserve performed over a 6-month training period. Training started at low intensity (40%–50% heart rate reserve) for 10–20 minutes and gradually progressed to target levels.

Plasma and muscle analyses

Plasma BDNF was measured in duplicate using Human BDNF Emax ImmunoAssay Kit. Vastus lateralis biopsies of the P and NP muscle were performed under local anesthesia after a 12-hour fast at baseline and after RT (n = 10) and after AEX (n = 5), 24–36 hours after the last bout of exercise. Muscle was immediately freeze-clamped and stored at −80°C. RNA extraction, reverse transcription, and quantitative real-time PCR for gene expression of BDNF was conducted. The Illumina 450k methylation array was used for DNA methylation according to manufacturer’s protocol. Briefly, DNA purified from muscle was bisulphite-converted using EZ-96 DNA methylation kit (D5004, Zymo Research), amplified, and then enzymatically fragmented followed by hybridization with the 450k array and fluorescent staining.

Statistical analyses

Baseline BDNF levels between P and NP thigh muscle was determined using paired Student t tests. Changes were assessed using repeated measures analysis of variance using SPSS 22.0. The 450k array image file (IDAT files) processed using the minfi package to calculate intensity and perform Subset-quantile within array normalization method was used to reduce the technical variability between type I and type II assay designs. Methylation scores are calculated as β values, ranging from 0 (totally unmethylated) to 1 (totally methylated). M values were derived as logit (β), which have properties suitable for statistical tests. Differential methylation between P and NP thigh muscle and between post- and pre-AEX thigh muscle was evaluated by paired Student t test on the M values. Data are presented as means ± SEM. p values < 0.05 are statistically significant.

Data availability statement

De-identified data and the study protocol may be available. More information regarding the data is available from the corresponding author on reasonable request.

Clinical trials identifiers

NCT02347995, NCT00387712, NCT00891514.
Results

Resistive training

Leg press strength increased by 26% for the P leg (304 ± 40 vs 363 ± 41 lbs derived from pneumatic resistance equipment) and 25% for NP leg (449 ± 36 vs 560 ± 33 lbs) (both p < 0.0001) with RT. Leg extension strength increased by 47% and 25% (P: 57 ± 10 vs 84 ± 11 lbs, NP: 113 ± 10 vs 141 ± 9 lbs respectively, both p < 0.0001). Plasma BDNF protein increased by 25% with RT but this was not significant (0.64 ± 0.08 vs 0.87 ± 0.20 ng/mL). P muscle has 45% lower BDNF mRNA expression (p < 0.05) than NP muscle (figure 1). Skeletal muscle BDNF mRNA did not change with RT (P: 7.21 ± 1.38 vs 7.06 ± 1.85 AU and NP: 11.39 ± 2.12 vs 7.84 ± 1.32 AU).

Aerobic exercise training

VO₂ peak increased by 18% with AEX (22.5 ± 2.5 vs 26.6 ± 3.5, p < 0.05). P and NP muscle have differential methylation status for DNA sequences of BDNF (3 CpG sites, p = 0.016 to 0.044) and BDNFAS (1 CpG site, p = 0.016) (table 1) at baseline (prior to exercise training). Table 1 also provides the β coefficients and the probe set ID. After AEX, there were significant changes in methylation of P muscle at 2 CpG sites (table 2). DNA methylation in BDNFAS increased in P relative to NP muscle (p = 0.017, figure 2). DNA methylation in BDNFAS did not change significantly in NP muscle.

Discussion

Our results provide the first preliminary evidence of reduced BDNF expression in P skeletal muscle as well as differential methylation status for BDNF and BDNFAS between P and NP muscle and suggests that AEX modulates methylation for BDNF in P muscle. We are unaware of any studies examining BDNF expression in skeletal muscle of stroke patients, or changes with exercise training in this population. The difference in BDNF expression between the P and NP limb in both gene expression and methylation provide important evidence for epigenetic modifications after stroke.

There is limited work in human skeletal muscle for BDNF. Acute exercise increases BDNF mRNA and protein expression in young healthy men. Evidence remains inconclusive regarding the effectiveness of exercise training on serum BDNF. Our results indicate that circulating BDNF did not change after RT. Other studies report that 8–12 weeks of RT either increase or do not change serum BDNF in older adults. Postulated mechanisms for improved cognition with exercise includes increased blood flow or oxidative stress leading to increased endothelial function and release of BDNF from the brain microvascular endothelial cells. Indeed, we have shown

Table 1 Baseline P and NP skeletal muscle BDNF methylation

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<tr>
<th>Probeset ID</th>
<th>Gene</th>
<th>β (NP muscle)</th>
<th>β (P muscle)</th>
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<tr>
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Abbreviations: BDNF = brain-derived neurotrophic factor; BDNFAS = BDNF Antisense RNA; P = paretic; NP = nonparetic.
increased cerebral vasomotor reactivity after aerobic training in chronic stroke providing evidence of changes in cerebral blood flow, which could potentially influence BDNF levels.

Epigenetic regulation of numerous genes, specifically methylation at the BDNF gene, was reported in a recent systematic review related to neurodegenerative diseases. Peripheral BDNF methylation is promoted as a surrogate for central BDNF methylation, given the direct relationships observed between BDNF promoter methylations in the hippocampus and quadriceps tissue of patients with bipolar disorder. Patients with mild depressive disorder not only have higher BDNF promoter methylation than healthy controls but also thinning of several cortical regions of the brain. The latter finding is inversely correlated with BDNF promoter methylation. We are unaware of studies in chronic stroke. Our finding that methylation decreased the gene expression of BDNFAS supports an increase in BDNF.

Limitations of the current trial include the relatively small sample size and limited human muscle sample available for analysis. In addition, a broader analysis on genome-wide DNA methylation and transcription profile changes in P vs NP muscles would be informative in future studies. However, the current studies were highly controlled and included supervised exercise training programs and standardization of muscle biopsy timing and conditions and novel with respect to skeletal muscle methylation in chronic stroke. Our earlier work has established that P muscle after chronic stroke has metabolic and structural differences relative to the NP muscle. The clinical relevance of the difference in DNA methylation in P vs NP muscle and following exercise training cannot be determined in this preliminary investigation.

A larger randomized controlled trial is necessary to establish the effects of exercise on BDNF methylation and other epigenetic modifications in stroke skeletal muscle. Future research could also examine changes in epigenetics with exercise training and whether training-induced changes associate with improved cognitive ability after stroke.

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Disclosure
Disclosures available: Neurology.org/NG.

Publication history
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Appendix Authors

<table>
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<tr>
<th>Name</th>
<th>Location</th>
<th>Role</th>
<th>Contribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alice S. Ryan, PhD</td>
<td>University of Maryland, Baltimore</td>
<td>Author</td>
<td>Design and conceptualized study; performed and collected the data; drafted the manuscript for intellectual content</td>
</tr>
<tr>
<td>Huichun Xu, MD, PhD</td>
<td>University of Maryland, Baltimore</td>
<td>Author</td>
<td>Performed statistical analysis and revised the manuscript</td>
</tr>
<tr>
<td>Frederick Ivey, PhD</td>
<td>University of Maryland, Baltimore</td>
<td>Author</td>
<td>Performed and collected the data; revised the manuscript</td>
</tr>
<tr>
<td>Richard Macko, MD</td>
<td>University of Maryland, Baltimore</td>
<td>Author</td>
<td>Revised the manuscript</td>
</tr>
</tbody>
</table>

References
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