FIRST POSTER SESSION STROKE/VASCULOPATHY

POSTER **ABSTRACTS**

CLINICAL-TRANSLATIONAL RESEARCH SYMPOSIUM

Elucidating Subtypes and Risk Factors of Brain Arteriolosclerosis

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lular demands and stressors, and play an essential role in cell tial sensitivities to Ca2+ and respond differently to TBI over signaling, differentiation, and survival. There is clear evidence of time. Second, the levels of miR-150 and miR-146a have been compromised mitochondrial function following traumatic brain shown to decrease in mitochondria isolated from the hippocaminjury (TBI), however, the pathological consequences are not pal formation following TBI. In a separate set of experiments, well known. MiRNA are small non-coding RNA molecules that rat primary cortical astrocytes were treated with various conregulate post-transcriptional gene expression, and serve as im- centrations of glutamate followed by mitochondrial isolation portant mediators of neuronal development, synaptic plasticity, and single tube RT-qPCR measurement of select miRNA. These and neurodegeneration. Our recent studies suggest that mito- studies revealed that mitochondria levels of miR-146a and miRchondria may serve as regulators of cellular miRNA expression 150 significantly decreased while levels of these same miRNAs following TBI. Here, we extend our initial observations by re- were unchanged in cytoplasmic fractions. Taken together, we porting the effect of TBI related secondary injury events on mi- hypothesize that TBI-related secondary injury events, such as tochondria associated miRNA expression. Synaptic and non- high Ca2+ and exposure to excitotoxic levels of glutamate, lead synaptic mitochondria were isolated from naïve adult rat cortex to altered expression levels of certain mitochondria-associated and exposed to concentrations of Ca2+ that inhibit State III mi- miRNAs in a manner similar to that observed following TBI. tochondrial respiration by 15-30%. RT-qPCR analysis demonstrated that Ca2+ treatment significantly decreased miR-150 and miR-146a levels in synaptic, but not non-synaptic mitochondria. This observation is particularly intriguing for two reasons.

Mitochondria serve as the powerhouse of cells, respond to cel- First, synaptic and non- synaptic mitochondria exhibit differen-

Supported by the Kentucky Spinal Cord and Head Injury Research Trust 15-12A (JES).