Decreased neuroinflammation and increased brain energy homeostasis following environmental enrichment after mild traumatic brain injury is associated with improvement in cognitive function

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Abstract

Background: Persistent neuroinflammation and disruptions in brain energy metabolism is commonly seen in traumatic brain injury (TBI). Because of the lack of success of most TBI interventions and the documented benefits of environmental enrichment (EE) in enhancing brain plasticity, here we focused our study on use of EE in regulating injury-induced neuroinflammation and disruptions in energy metabolism in the prefrontal cortex and hippocampus. Adult male Wistar rats were used in the study and randomly assigned to receive either: mild TBI (mTBI) using the controlled cortical injury model or sham surgery. Following surgery, rats from each group were further randomized to either: EE housing or standard laboratory housing (CON). After 4 weeks of recovery, cognitive testing was performed using the non-matching-to-sample and delayed non-matching-to-sample tasks. After completion of behavioral testing, levels of the pro-inflammatory cytokines IL-1β and TNF-α and the anti-inflammatory cytokine IL-10 were measured. In addition, levels of AMPK (adenosine monophosphate-activated protein kinase), phosphorylated AMPK and uMtCK (ubiquitous mitochondrial creatine kinase) were assessed as measures of brain energy homeostasis.

Results: Our results showed that EE: (1) decreased the pro-inflammatory cytokines IL-1β and TNF-α and enhanced levels of the anti-inflammatory cytokine IL-10 after mTBI; (2) mitigated mTBI-induced cognitive impairment; and (3) attenuated mTBI-induced downregulation in pAMPK/AMPK ratio and uMtCK levels.

Conclusions: Our data demonstrated the potential of EE to modulate the persistent: (1) neuroinflammatory response seen following mTBI, and (2) persistent disturbance in brain energy homeostasis. It is possible that through the mechanism of modulating neuroinflammation, EE housing was able to restore the disruption in energy metabolism and enhanced functional recovery after mTBI.

Keywords: IL-1β, TNF-α, IL10, AMPK, Non-matching-to-sample task

Background

Traumatic brain injury (TBI) is of major public health significance, affecting almost 1.7 million people in the United States each year [1]. Globally, the incidence of TBI is also increasing, particularly in developing countries where road traffic accidents are on the increase as a result of widespread motor vehicle use. Mild traumatic brain injury (mTBI) comprises about 80% of all TBI cases and most individuals exhibit disabilities associated with cognitive problems [2]. Common cognitive impairments seen in mTBI include difficulties in attention, episodic memory, executive functions (such as higher-order planning, initiating and directing, monitoring, problem solving, and inhibitory control), working memory, information-processing speed, language functions, and visual-spatial processing that can last for months or even years [3]. Large
transient increases in excitatory neurotransmitter efflux [4] resulting in excitotoxicity, ionic imbalance, ATP depletion, proteolysis, and oxidative stress [5] happens in the acute phase of TBI. The complex array of responses to injury can result in energy crisis that compromises the capacity of the brain to cope with challenges, thus regulation of energy homeostasis after TBI is a critical step in maintaining brain function.

While the acute effects of mTBI can be devastating, increasing evidence suggest that mTBI also initiates long-term effects in some segment of survivors [6]. The acute effects of mTBI stemming from the primary insult can cause secondary injuries that can evolve over minutes to days and even months after the initial traumatic event. Secondary injury events such as neuroinflammation can also cause neurodegeneration and persistent cognitive impairment [7]. The neuroinflammatory process involves the movement of activated microglial cells to the site of injury in response to extracellular adenosine triphosphate (ATP) released by the injured tissue [8-10]. The microglial processes then fused to form an area of containment between healthy and injured tissues, suggesting that microglia may represent the first line of defense following TBI [11]. However, when microglia become over-activated or reactive they can induce detrimental neurotoxic effects by releasing multiple cytotoxic substances, such as pro-inflammatory cytokines (e.g. interleukin-1 beta and tumor necrosis factor-alpha) and arachidonic acid metabolites [12-14]. These reports suggest that clear beneficial effects can be achieved if neuroinflammation is controlled in a regulated manner and for a defined period of time. However, there is increasing number of evidence that chronic microglial activation is present in most cases of TBI leading to neuroinflammation that can persist for months or years after injury [14,15].

To date, most interventions that target a specific pathological consequence of TBI have failed clinically. Thus, it is possible that using an intervention with known general effects on enhancing brain plasticity and resiliency may be beneficial in mTBI and one such strategy is environmental enrichment (EE). In general EE refers to conditions that provide increased social, cognitive, and physical stimulation [16,17]. Reports show that EE can exert positive effects on behavioral recovery after cerebral ischemia [18-20], mild TBI [22-25], and neurodegenerative disorders [26,27]. Even though the effects of EE in modulating neuroinflammation is just beginning to be explored [28], the impact of EE in modulating brain energy homeostasis after mTBI has not been studied. Here we focused our study on EE effects in regulating injury-induced neuroinflammation and disruptions in energy metabolism in the prefrontal cortex and hippocampus, as these are the brain regions commonly affected in mTBI. The main goal of the study was to determine the effects of EE in attenuating the long-term consequences of mTBI relating to neuroinflammation, alterations in brain energy metabolism, and cognitive impairment. The rationale for examining long-term effects in the present was based on the dearth of information available in the persistent consequences of mTBI.

Methods

Animal model
Adult male Wistar rats were obtained from Harlan Laboratories (Madison, Wisconsin) weighing approximately 375 to 400 grams. Animals were housed in pairs in a pathogen-free vivarium under controlled condition (temperature 22 ± 1°C and humidity 70 ± 5%) and a 14:10 hour light: dark cycle was maintained. All animals were housed in the same room so that temperature, humidity, and lighting conditions are similar for all groups. Animals had free access to food (regular rat chow) and water delivered through an automated and filtered system. Animals were also handled daily throughout the study so that they could get acclimated to the research personnel thereby decreasing stress. Experiments started one week after arrival of the animals from the breeder and all experimental protocols in this study were approved by the Institutional Animal Care and Use Committee (University of Illinois at Chicago) and in accordance with the National Institutes of Health guidelines. All efforts were made to minimize animal distress and to reduce the number of animals used.

Mild traumatic brain injury
Following induction anesthesia with isofluorane inhalation (1%), animals were placed in a Cunningham stereotaxic frame (Stoelting, Wood Dale, IN) in a prone position and stabilized using the ear bars and incisor bar. The head was held in a horizontal plane with respect to the interaural line. Throughout the procedure, continuous isofluorane anesthesia was maintained at 2.5%, aseptic technique was used and the rats’ body temperature was maintained at 37°C ± 1°C using the feedback-regulated water heating pad. Once the animals were fully anesthetized and the head stabilized, a midline incision was made and the soft tissues were retracted. Then two 10 mm diameter craniotomies were made adjacent to the central suture, midway between lambda and bregma. Injury was induced in one of the craniotomies and the second one allowed for lateral movement of the brain during injury. The dura matter was kept intact over the cortex and injury was induced by striking the left or right (ipsilateral) cortex with a pneumatic piston with a 6 mm diameter tip at a rate of 4 m/sec, with 1.5 mm of compression. Velocity was measured using a linear velocity displacement transducer. Any bleeding from the skull during the craniotomy was controlled with bone wax. After the injury, the scalp was closed with 6–0 silk suture, anesthesia was discontinued, and the rat’s temperature was maintained at 37°C ± 1°C until recovery of
locomotion. Sham animals were subjected to the same anesthesia and craniotomy, but no cortical injury.

Animals were assessed every hour × 8 hours then daily × one week, for postoperative complications such as excessive weight loss (>20% of preoperative body weight), bleeding, seizures, and infection. Animals were euthanized immediately and excluded from the study when postoperative complications occurred (n = 0). Liquid Tylenol was given orally with drinking water at 200 mg/Kg after surgery for 48 hours as postoperative analgesia. Sluggishness, extreme aversion to being touched, and weight loss were also assessed as indication of persistent pain (n = 0).

Animal housing
Immediately upon recovery from anesthesia, rats were randomly placed in either: enriched environment (EE) housing or paired housing (social controls). The rats remained in their assigned housing condition throughout the duration of the study. Animals in the EE group (n = 9 mTBI and n = 8 shams) were housed together in a sensory-rich living condition (wire cage measuring 2 m × 1 m × 1.65 m) consisting of a variety of objects as described previously [19,29]. In addition, these rats were placed each day in an open field (1.2 × 1.2 × 1.2 m) during the evening hours with a novel arrangement of toys and objects and allowed to explore for 30 minutes while the objects in the home cage were being changed. Objects in both EE housing and open field were changed daily to maintain novelty.

Animals assigned to the social control (C-S) group (n = 9 mTBI and n = 8 shams) were housed in pairs in standard laboratory cages (16.5 × 22.5 × 13.5 cm). Although rats in this group were able to observe ongoing activity of the room, they did not receive any stimulation and contact was limited to daily handling and routine cage changing. Paired housing was used to control for the social interaction effect of the enriched environment.

Non-matching-to-sample and delayed non-matching-to-sample tasks
Behavioral testing was conducted 4 weeks after EE or control housing and approximately 2 hours prior to the onset of the dark cycle (this is close to the rats' active period for a period of 9 days (including habituation) and performance was recorded using the Ethovision XT® v.8.2 video tracking program (Leesburg, VA). The non-matching-to-sample task consisted of a series of paired sample and test trials. At the beginning of each sample trial, either black or white cylinder was suspended directly above the submerged platform in the water maze. In subsequent test trials, both cylinders were present but the cylinder not present during the preceding sample trial was suspended over the platform and served as a cue for the location of the goal. Thus, if on a given sample trial, the black cylinder cued the platform, and then on the succeeding test trial, the white cylinder was used to cue the platform. The black and white cylinders were selected as sample stimuli for each pair of trials according to a semi-random schedule that ensured each cylinder served as the sample stimulus on 50% of the trials over the phase of the experiment. For each test trial, the platform was moved to another quadrant with the non-sample cylinder located directly above it. The sample stimulus was moved to a different quadrant for each trial. Based on a random schedule, the position of the submerged platform was changed after each sample and test trial to eliminate the use of spatial cues. All quadrants were used equally for locating cues in the sample and test trials and the platform was positioned randomly. The day before testing, rats were allowed a habituation swim for 10 seconds without the submerged platform. The constant water temperature in the pool and habituation swim helped decrease animal stress associated with the task.

At the beginning of each sample trial, the rat was placed in the pool at the same location (in the center of the southeast quadrant), facing the wall of the pool, and allowed to swim to the submerged platform under the sample cylinder. The rat was allowed to remain on the platform for 10 seconds. If the rat failed to find the platform within 60 seconds, it was picked up and placed on the platform for 10 second then removed and placed under a heat lamp while the platform is moved and the cylinders put in position for the test trial. The heat lamp allowed for the rats to get dry before the test trial. Getting the cylinders and platform ready for the test trial took ≈10 seconds. Rats received five daily sessions and each session consisted of four pairs of sample and test trials. The day after completion of the non-matching-to-sample task, rats were tested in the delayed non-matching-to-sample task for three additional daily sessions. Each session consisted of three paired trials, with intervals of 60, 120, or 180 seconds between the sample and test trials (the intervals do not include the 10 seconds required for repositioning the cylinders and platform). The order of the delays varied each day according to a random schedule.

Tissue preparation
Rats were euthanized using CO2 asphyxiation the day after behavioral testing, the brains removed, and the prefrontal cortex and hippocampus were manually dissected then immediately placed in liquid nitrogen and kept frozen until processed. The ipsilateral regions were analyzed for cytokine levels (interleukin-1β and tumor necrosis factor-α) and measures of brain energy homeostasis such as AMPK (adenosine monophosphate-activated protein kinase) and uMtCK (ubiquitous mitochondrial creatine kinase).
Cytokine protein quantification

The concentration of IL-1β, TNF-α, and IL-10 protein levels was determined using commercially available ELISA assays. Briefly, 0.5 g of frozen tissues were homogenized with a glass homogenizer in 1 ml buffer containing 1 mol/liter phenylmethylsulfonyl fluoride, 1 mg/liter peptatin A, 1 mg/liter aprotinin, and 1 mg/liter leupeptin in PBS (pH 7.2) and centrifuged at 12,000 × g for 20 minutes at 4°C. The supernatant was collected and total protein was determined by bicinchoninic acid (BCA) protein assay reagent kit (PIERCE, Milwaukee, WI). Samples were used for ELISA to determine IL-1β, TNF-α, and IL-10 protein levels. The procedure was performed according to manufacturer’s specification using the Quantikine rat-specific ELISA kits (R&D Systems, Minneapolis, MN) and the color reaction was detected using the chromogen tetramethylbenzidine. Color reaction was stopped by an equal volume of stop solution (provided by the manufacturer) and read in a microplate reader (Bio-Tek, Winooski, VT) at a wavelength of 450 nm (650-nm reference wavelength). The color change was proportional to the concentration of the cytokines measured and all samples measured were within the range of the standard curve. This ELISA system detects both natural and recombinant rat IL-1β, TNF-α, and IL-10. Assays were sensitive to 5 pg/ml for IL-1β and TNF-α, and 10 pg/ml for IL-10; the intra-assay and inter-assay coefficients of variation were < 5% and < 10% for TNF-α and IL-10, and <6% and <9% for IL-1β. Assays were performed in triplicates and measurements were averaged and used as one individual data point for statistical analysis.

Western blot

To detect measures of brain energy homeostasis, 0.5 g of frozen tissues was used in the Western blot procedure. Tissues were homogenized and centrifuged at 25,000 × g for 20 minutes as previously described [18,30]. Aliquots from the supernatant were removed for protein determination. Protein concentration in samples was determined using the BCA-Protein assay (Pierce, Rockford, IL). Equal amounts of protein (40 μg) from each rat were loaded and separated by SDS-PAGE and electrophoresis in 8% - 16% acrylamide gradients. Protein bands were electrophoretically transferred to nitrocellulose membranes (Amersham, Piscataway, NJ) stained with 0.5% Ponceau Red to visualize total proteins, then destained. Non-specific binding sites were blocked then nitrocellulose membranes were incubated overnight at 4°C with gentle agitation in the following primary antibodies: (1) monoclonal mouse anti-AMPK (1:1000, Santa Cruz Biotechnology, Santa Cruz, CA); (2) polyclonal rabbit anti-phosphorylated AMPK (1:1000, Cell Signaling, Billerica, MA); (3) monoclonal mouse anti-uMtCK (1:1000, Santa Cruz Biotechnology); (4) polyclonal rabbit anti-β-actin (1:2000, Santa Cruz Biotechnology). The secondary antibodies used were horseradish peroxidase-conjugated immunoglobulin (Sigma, St. Louis, MO) and the Super Signal chemiluminescence substrate kit (Pierce, Rockford, IL) was used to visualize immunoreactive bands. After visualization, the membranes were then stained with Amido-Black to qualitatively verify protein loading. Band visualization was obtained by exposure of membranes to autoradiographic film (Kodak Biomax”). Samples were analyzed in quadruplicates and measurements were averaged and used as one individual data point for statistical analysis.

Quantification of differences in protein bands between samples was done using densitometric analysis (Image software v1.47). The internal control β-actin was used to standardize experimental values in densitometric analysis. Densitometric values were calculated as: density of sample/density of background.

Statistical analysis

The SAS general linear model (SAS Institute, North Carolina) procedures for two-way analysis of variance (ANOVA) were used to examine effects of experimental conditions (pre, post, sham) and housing condition (enriched environment vs. control) on neuroinflammatory state (IL-1β, TNF-α, and IL-10) and brain energy homeostasis (AMPK, p-AMPK, and uMtCK). The SAS CONTRAST statement was used for planned comparisons when appropriate. Repeated measures ANOVA was used to analyze behavioral data. All error bars represent ± standard error of the mean (SEM) of the sample size used in the study.

Results

EE effects on inflammatory state

We examined the expression of the pro-inflammatory cytokines interleukin-1β (IL-1β) and tumor necrosis factor-α (TNF-α) as well as expression of the anti-inflammatory cytokine interleukin-10 (IL-10). Our results showed significant main effects on IL-1β (F(3,30) = 9.21, p < 0.05) and TNF-α (F(3,30) = 9.67, p < 0.05) in the prefrontal cortex where mTBI led to the upregulation of these inflammatory cytokines even after more than a month of recovery from injury (Figure 1A and B). However, housing rats in EE immediately after brain injury significantly decreased levels of IL-1β and TNF-α in the prefrontal cortex to 36% and 32%, respectively, when compared to the mTBI animals housed in regular laboratory cages. No significant changes in IL-1β and TNF-α levels were seen in the sham groups. Similar pattern of significant effects of IL-1β (F(3,30) = 10.07, p < 0.05) and TNF-α (F(3,30) = 9.71, p < 0.05) were seen in the ipsilateral hippocampus (Figure 1D and E).

In addition, we also examined levels of IL-10 and found significant main effects of injury (F(3,30) = 10.99, p < 0.05) in IL-10 levels where a persistent downregulation of this anti-inflammatory cytokine was seen in the prefrontal cortex weeks after mTBI. But housing rats in EE immediately after
mTBI significantly modulated the reduction of IL-10 levels (F(3,30) = 10.02, p < 0.05). IL-10 level in the mTBI rats housed in EE was 41% higher when compared to the mTBI animals housed in regular laboratory cages. No significant differences were seen in IL-10 levels in the sham rats (Figure 1C). Again, a similar pattern of significant effects of injury (F(3,30) = 11.05, p < 0.05) and EE housing (F(3,30) = 12.03, p < 0.05) were seen in the hippocampus (Figure 1F). Together, these results suggest that neuroinflammation can endure weeks after injury but housing rats in EE during recovery can modulate this persistent consequence of mTBI.

EE effects on pAMPK/AMPK ratio
We measured total AMPK and pAMPK to obtain the pAMPK/AMPK ratio as an indication of cellular energy status [31,32]. We found significant main effects of injury (F(3,30) = 10.93, p < 0.05) and EE housing (F(3,30) = 10.48, p < 0.05) on pAMPK/AMPK ratio in the prefrontal cortex where mTBI resulted in significantly decrease pAMPK/AMPK ratio in comparison to the sham groups (Figure 2A). However, housing rats in EE after injury significantly lessened the effects of mTBI on pAMPK/AMPK ratio. Posthoc comparison showed that the pAMPK/AMPK ratio of mTBI rats housed in EE was not significantly different from the sham control group. Comparison of the sham groups also showed significant effects of EE housing on pAMPK/AMPK ratio where sham rats housed in EE had 33% higher ratio compared to the sham control animals. Examination of pAMPK/AMPK ratio in the hippocampus also showed significant main effects of
mTBI \( (F(3,30) = 11.33, \ p < 0.05) \) and housing condition \( (F(3,30) = 12.11, \ p < 0.05) \) with the pattern of changes similar to that seen in the prefrontal cortex (Figure 2B).

**EE effects on uMtCK**

Measurement of uMtCK was also done in the present study because of the involvement of this kinase in energy transduction [33]. Significantly decreased uMtCK expression was seen in the prefrontal cortex after mTBI \( (F(3,30) = 9.61, \ p < 0.05) \) compared to the sham groups; but housing rats in EE after injury mitigated the reduction in uMtCK levels \( (F(3,30) = 10.31, \ p < 0.05) \) induced by mTBI (Figure 3A). Posthoc comparison showed that the uMtCK levels of mTBI rats housed in EE was not significantly different from the sham CON group. Comparison of the sham groups also showed significant effects of EE housing on uMtCK expression where sham rats housed in EE had 28% higher levels compared to the sham control animals. When uMtCK levels were examined in the hippocampus, significant main effects of mTBI \( (F(3,30) = 12.17, \ p < 0.05) \) and housing condition \( (F(3,30) = 11.22, \ p < 0.05) \) were also seen with expression levels similar to that seen in the prefrontal cortex (Figure 3B).

**EE effects on mTBI-induced cognitive impairment**

Rats were allowed to recover for 4 weeks in their assigned housing condition after injury before undergoing behavioral testing. All rats remained in their assigned

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**Figure 2 Ratio of pAMPK/AMPK.** Representative Western blots of total AMPK and phosphorylated AMPK (pAMPK) in the hippocampus (upper panel). Bar graphs show that the ratio of pAMPK/AMPK is significantly decreased in the prefrontal cortex (A) and hippocampus (B) of mTBI rats assigned to control housing. Housing rats in EE after mTBI restored ratio of pAMPK/AMPK to normal levels. Notably, an overall increase in pAMPK/AMPK is evident in the sham animals housed in EE. *p < 0.05, **p < 0.01. Legend: 1 – mTBI control housing, 2 – mTBI EE housing, 3 – sham control housing, and 4 – sham EE housing.

**Figure 3 uMtCK Expression.** Upper panel show representative Western blots of uMtCK expression in the prefrontal cortex (A) and hippocampus (B). Bar graphs show significantly decreased uMtCK expression in both prefrontal cortex and hippocampus of mTBI rats assigned to control housing. Housing rats in EE after mTBI restored uMtCK to normal levels. Remarkably, an overall increase in uMtCK is evident in the sham animals housed in EE. *p < 0.05, **p < 0.01. Legend: 1 – mTBI control housing, 2 – mTBI EE housing, 3 – sham control housing, and 4 – sham EE housing.
environment throughout the testing days until the end of the study. Our results showed that overall mTBI control rats demonstrated significantly increase swim latency ($F(3,30) = 11.22, p < 0.05$) and made more errors ($F(3,30) = 10.23, p < 0.05$) in the non-matching-to-sample task even after 4 weeks of recovery from injury (Figure 4). We also found that the number of errors made in the delayed non-matching-to-sample task increased with increasing time delay between the sample and test trials ($F(3,30) = 10.14, p < 0.05$). Furthermore, performance in the delayed non-matching-to-sample tasks showed impairment in the mTBI control rats overall. Housing rats in EE after injury significantly mitigated the mTBI-induced cognitive impairment where their performance did not differ from those of the sham control group in both the non-matching-to-sample and delayed non-matching-to-sample tasks. Also, the sham EE groups demonstrated better performance overall in both the non-matching-to-sample and delayed non-matching-to-sample tasks.

Discussion

In the present study we show that mTBI-induced neuroinflammation, disruptions in brain energy metabolism, and behavioral impairment may be modifiable by enriched environment housing after injury. These findings are supported by our data that EE: (1) modulated mTBI-induced upregulation of the pro-inflammatory cytokines IL-1β and TNF-α and enhanced levels of the anti-inflammatory cytokine IL-10; (2) mitigated mTBI-induced cognitive impairment; and (3) attenuated mTBI-induced downregulation in pAMPK/AMPK ratio and uMtCK levels. Ultimately, our findings extended the work of others [22-25,34-37] suggesting that EE housing after either mild or severe mTBI may be an effective therapeutic intervention that can optimize behavioral recovery after injury.

Here we show that the neuroinflammation induced by mTBI persisted for 4 weeks after the injury and that housing rats in EE immediately after injury significantly reduced the inflammatory response. The persistent neuroinflammation seen in the mTBI control group is similar to reports from clinical and experimental studies that neuroinflammation resulting from mTBI can last from months to years after injury [14,38,39]. Neuroinflammation in the acute phase of mTBI triggers the release of pro-inflammatory cytokines essential for mounting a defense mechanism associated with neutralization of the insult and restoration of normal structure and function following injury [40]. However, if neuroinflammation is not regulated, it can result in a self-propagating and deleterious process [14]. This is evident in our data showing that the continuing neuroinflammation seen in mTBI rats assigned in control housing is associated with persistent cognitive impairment, downregulation of pAMPK/AMPK ratio, and uMtCK levels in the prefrontal cortex and hippocampus. The modulation of IL-1β and TNF-α expression in the mTBI rats housed in EE may be partially mediated by the increased release of an anti-inflammatory cytokine IL-10.

One of the contributing factors to cognitive impairment following mTBI is neuroinflammation. For example, the pro-inflammatory cytokine IL-1β in particular is reported to affect hippocampal-dependent memory tasks (reviewed in [41]). As well, increased pro-inflammatory cytokine TNF-α is reported to reduce synaptic plasticity and contribute to neurodegeneration [42-44]. Thus, it is logical to think that the modulation of IL-1β and TNF-α expression in the hippocampus and prefrontal cortex of mTBI rats housed in EE seen in this study led to the sparing of disruptions in cognitive function. This line of thinking is supported by our findings that the mTBI rats housed in EE performed as well as the sham control animals in both the non-matching-to-sample and delayed non-matching-to-sample tasks. However, our behavioral findings are at odds with a recent study where only partial mitigation of cognitive impairment in mice housed in EE after Influenza-induced inflammation [28] is reported. Some possible reasons in the conflicting findings are: (1) different triggering

Figure 4 Behavioral performance. Overall, mTBI rats assigned to control housing showed longer mean swim latency (A) and made more errors (B) in the non-matching-to-sample. Increasing time delay between the sample and test trials also resulted in more errors made (C). Remarkably, providing enrichment after injury enhanced functional recovery as seen in the similar performance of the mTBI rats housed in EE and the sham control animals. EE housing in sham animals also led in enhanced behavioral performance. *$p < 0.05$, **$p < 0.01$. 

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events for inflammation, wherein we used mTBI and the other study used Influenza virus. It is possible that Influenza also induced fatigue as part of the 'sickness behavior' commonly seen in peripheral inflammation [45], which affected the animals’ performance in the water maze; (2) time of behavioral testing wherein we started the non-matching-to-sample test 4 weeks after mTBI while in the other study, mice were tested in the water maze 48 hours after inoculation with the Influenza virus. The possibility exists that our data may have been influenced by some degree of spontaneous recovery [46,47] reported to occur after brain injury. That is, in our study the rats were housed in EE for 4 weeks and throughout the behavioral testing period, thus spontaneous recovery may have been enhanced.

In this study we also show that mTBI resulted in persistent downregulation of pAMPK/AMPK ratio, which is tempered by EE housing suggesting that providing enrichment may initiate mechanisms that help conserve ATP levels that are depleted following injury. AMPK is an evolutionarily conserved signaling molecule that is considered one of the most important energy sensors in the body [48]. AMPK is activated via phosphorylation when ATP is depleted, which makes it a good marker for cellular energy [49]. The response of AMPK then to changes in neuronal energy status and metabolic stress makes the pAMPK/AMPK ratio a reasonable measure of neuronal responsiveness to changes in energy status. AMPK controls cellular energy through phosphorylation of metabolic enzymes that leads to the restoration of energy balance by activation of catabolic pathways that produce ATP and inhibition of anabolic pathways that use energy [50]. This role of AMPK in restoring energy balance is apparent in our data showing that the ratio of pAMPK/AMPK in the mTBI rats housed in EE is similar to the sham control group. Furthermore, it is not surprising that in the cases where pAMPK/AMPK ratio significantly increased in sham rats housed in EE in relation to the sham control group given the role of AMPK in maintaining energy homeostasis in concert with reports that EE housing can enhance synaptic plasticity, an energy consuming process.

In addition to its role in controlling cellular energy, AMPK also regulates a variety of transcription factors involved in learning and memory [51]. Accordingly, it is possible that EE housing may engage AMPK to prevent cognitive impairment resulting from mild energy depletion state such as that seen in mTBI. Indeed, this line of reasoning is supported by our data showing that the performance of mTBI rats housed of EE is similar to the sham control group. Furthermore, it is possible that the overall enhanced behavioral performance and upregulation of pAMPK/AMPK ratio seen in the sham EE housed rats in this study may be explained by increased activation of AMPK linking intracellular energy levels and synthesis of proteins involved in learning and memory.

We also measured uMtCK levels in the present study since this molecule is highly expressed in hippocampal granule and pyramidal cells. UMtCK is functionally coupled to oxidative phosphorylation since mitochondrial-derived ATP is preferentially used by this molecule to transfer high-energy phosphates to creatine. The resultant phosphocreatine acts as the cytosolic transport and storage form of energy and used to regenerate ATP from cytosolic ADP, which is essential in maintaining the ATP supply during energy-consuming brain activity. Our data show that mTBI resulted in decreased uMtCK levels but housing rats in EE after injury regulated this injury-induced downregulation leading to levels similar to the sham control group. We also show that sham rats housed in EE have the highest levels of uMtCK overall. These expected findings suggest that EE restored the decreased levels of uMtCK following mTBI and intensified uMtCK expression in EE, a condition that requires high energy because of increased synaptic activity, thus providing a mechanism of energy transduction in which ATP synthesized in the mitochondria is transferred to sites of ATP utilization.

Conclusions

Our data show the potential of EE to modulate the persistent: (1) neuroinflammatory response seen following mTBI, and (2) persistent disturbance in brain energy homeostasis. It is possible that through the mechanism of modulating neuroinflammation, EE housing is able to restore the disruption in energy metabolism induced by mTBI. Furthermore, the modulation of energy metabolism seen in the mTBI rats housed in EE seemed to correlate with functional recovery. Collectively, these results suggest that EE housing can be effective in providing neuroprotection after mTBI consistent with previous reports [22-25,34-37]. Our findings point to the importance of using therapeutic strategies directed at modulating the normal compensatory mechanism seen after CNS insult so that its harmful effects can be minimized.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
TLB conceived of the project, analyzed the data, and wrote the manuscript. JW performed the behavioral testing and ELISA procedures. MR performed the surgeries and Western blots. All authors read and approved the final manuscript.

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