Hypobaria-Induced Oxidative Stress Facilitates Homocysteine Transsulfuration and Promotes Glutathione Oxidation in Rats with Mild Traumatic Brain Injury

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ABSTRACT

BACKGROUND: United States service members injured in combat theatre are often aeromedically evacuated within a few days to regional military hospitals. Animal and epidemiological research indicates that early exposure to flight hypobaria may worsen brain and other injuries. The mechanisms by which secondary exposure to hypobaria worsen trauma outcomes are not well elucidated. This study tested the hypothesis that hypobaria-induced oxidative stress and associated changes in homocysteine levels play a role in traumatic brain injury (TBI) pathological progression caused by hypobaria.

METHODS: Male Sprague Dawley rats were exposed to a 6h hypobaria 24h after mild TBI by the controlled cortical impact. Plasma and brain tissues were assessed for homocysteine levels, oxidative stress markers or glutathione metabolism, and behavioral deficits post-injury in the absence and presence of hypobaria exposure.

RESULTS: We found that hypobaria after TBI increased oxidative stress markers, altered homocysteine metabolism, and promoted glutathione oxidation. Increased glutathione metabolism was driven by differential upregulation of glutathione metabolizing genes. These changes correlated with increased anxiety-like behavior.

CONCLUSION: These data provide evidence that hypobaria exposure after TBI increases oxidative stress and alters homocysteine elimination likely through enhanced glutathione metabolism. This pathway may represent a compensatory mechanism to attenuate free radical formation. Thus, hypobaria-induced enhancement of glutathione metabolism represents a potential therapeutic target for TBI management.

KEYWORDS: Hypobaria, homocysteine, oxidative stress, glutathione, traumatic brain injury

Introduction

Traumatic brain injury (TBI) is a public health problem with more than 175 000 victims among United States service members during the Iraq and Afghanistan wars and this number continues to grow as some of them remain in combat zones.1 Wounded service members, including approximately 10% who suffered from psychiatric disorders such as TBI, were aeromedically evacuated to hospitals in select countries for advanced care critical for survival between 2001 and 2013.2 During these flights, the cabin pressure is usually maintained at the equivalent to 8000 ft altitude. The mild hypoxia that accompanies such hypobaric conditions is thought to be one of several stressors that could cause secondary injury to TBI patients during these flights.3,4

Over the past decade, there has been significant interest in understanding the impact of hypobaria on TBI pathological progression using various injury models and hypobaria conditions. Mice who underwent 5h hypobaria (8000 ft) after a closed head injury 3h earlier presented worse outcomes in cerebral inflammation and neuronal death than those exposed to hypobaria a day later.5 In a recent report, rats subjected to blast overpressure and exposed 48h later to hypobaria for 12h presented with exacerbation of multi-organ injuries, including cerebral lesions.6 Work from our laboratory showed that 6h exposure to 8000ft hypobaria worsened axonal injury in rats subjected to under-vehicle blast.7 Furthermore, hypobaria-exposure aggravated inflammation, neuronal death, and cognitive deficits in rats subjected to TBI by lateral fluid percussion and exposed to hypobaria up to 7 days post-injury.8 Despite current advances in understanding the impact of hypobaria-exposure on TBI pathophysiology, hypobaria-associated biochemical changes and their effects on injury pathological outcomes remain unclear.

Elevated levels of the non-proteinogenic amino acid homocysteine (HCY), known as hyperhomocysteinemia (HHCY), is a stress biomarker and stress inducer with substantial
neurotoxic properties. Homocysteine metabolism is tightly regulated to ensure it is continuously remethylated to form methionine, its precursor, or transsulfurated to contribute to glutathione formation, an essential endogenous antioxidant. There is very limited data on homocysteine metabolism and pathological impact in hypobaria. A previous study by Sharma and colleagues reported that prolonged stay at high altitude (7500 ft) for 3 months, showing an increase in homocysteine levels. We recently reported in our model of methionine-induced moderate HHcy worsens TBI-induced lesion volume, BBB disruption, inflammation, anxiety-like behavior, and working memory in a rat model of mild TBI by controlled cortical impact. In a prior retrospective study involving 150 participants, Rahmani et al found a significant correlation between plasma HCY levels, TBI severity, and associated mortality. In the current study, we tested the hypothesis that hypobaria-induced oxidative stress and associated changes in homocysteine levels play a role in traumatic brain injury (TBI) pathological progression caused by hypobaria.

Mild traumatic brain injury (mTBI) model used in this study is unlike moderate and severe TBI, defined in rodents by no breaking of the dura, subtle pathological changes around the impact region and fast recovery from behavioral deficits.

Materials and Methods

Animals and housing

Male Sprague Dawley rats (300-350 g; Envigo, CA) were used. They were housed in pairs in a temperature-controlled environment, maintained at 23 ± 2°C, with a 12 h light/dark cycle, with continuous access to food and water ad libitum. All animal procedures were approved by The University of Maryland, Baltimore, Animal Use and Care Committee (IACUC # 1118014) and, the USAF Animal Use Programs Office of Research Oversight & Compliance, and the USAMRMC Animal Care and Use Review Office (ACURO). Animals were randomly assigned to experimental groups composed of 4 to 6 rats/group for biochemical assays and histological analyses, and 8 rats/group for behavioral studies.

Induction of controlled cortical impact injury

The Controlled Cortical Impact (CCI) method was used to induce mTBI. Animals were maintained under general anesthesia with 2.5% isoflurane throughout surgery. Rat body temperature was maintained at 37°C using a heating pad coupled with a rectal probe. Following craniotomy, CCI was performed over the left parietal cortex. Using a stereotaxically positioned 4 mm diameter piston tip, impact-injury was performed with an electromagnetically controlled impactor (Leica Biosystems, IL) at a piston velocity of 5 m/sec and depth penetration of 1.5 mm. The bone flap was replaced, the skin closed, and the anesthesia terminated. Sham-operated rats were also anesthetized, followed by craniotomy and replacement of the bone flap in the absence of cortical impact.

Rat exposure to hypobaria

At twenty-four hours post-surgery, rats were housed in cages with ad libitum food and water access and placed in a hypobaria chamber. A vacuum pump connected to the chamber was gradually adjusted to depressurize the chamber to 568 mmHg, which is the ambient pressure at 8000 ft, altitude. As previously reported, the chamber was kept normoxic (21% O2) with supplemental O2, were at a near-normal inspired oxygen level of 30%, the SpO2 was 97.0% at a hypobaria level 8000 ft. Rats were maintained under these conditions for 6 h, followed by depressurization every 15 min to sea level ambient pressure (757 mm Hg). As control, other rats were placed in the chamber under normobaric conditions, equivalent to 80 ft, altitude, 21% O2 for the same duration. Some sham or CCI rats (6/group) received intraperitoneal injections of 2-dihydroethidium (2-DHE) in 50% DMSO (6 mg/kg) to track superoxide formation during hypobaria or normobaria-exposure.

Behavioral testing

All behavioral tests were performed at time points described in the experimental time (Figure 1A).

Beam walk test. This test was performed to assess fine motor movements, as previously described. Briefly, animals were trained to walk on a 1.5 mm wooden beam for 3 consecutive days, and baseline data determining the number of foot faults over a total of fifty steps were collected on day 4. Test to assess the effect of hypobaria-exposure on fine motor movement in sham and CCI rats overtime was performed on days 3, 9, 16, and 30 post-surgery.

Elevated Plus maze test. The Elevated Plus maze was used to assess rats for anxiety-like behavior on days 2, 8, 15, and 29 post-surgery using a Plus-sign shape device as previously described. Briefly, rats were individually placed in the central area of the device and allowed to explore the maze for 10 min. Animal movements were analyzed by an overhead camera and data collected and analyzed by Any-Maze software (SD instrument, CA) to provide several behavioral parameters including the time spent in open arms. Anxiety-like behavior was interpreted as inversely proportional to the time an animal spent in the open arms.

Tissue collection and processing

Brain and blood were collected at 2 h, 24 h, or 30-days post-surgery. Rats were anesthetized and blood collected and processed to obtain plasma for biochemical assays as previously described. Rats were thereafter euthanized and brain
collected and processed for slicing in 40 µm thick coronal sections stored at −20°C for histology as described. 17,24

**Plasma homocysteine measurement**

Homocysteine levels were measured in rat’s plasma collected 2 h or 24 h post-hypobaria or normobaria exposure (n = 4/group). The assay was performed using an enzymatic assay kit (BioVision Inc., CA) following the manufacturer’s instructions.

**Determination of oxidative stress markers in plasma and brain**

**Malondialdehyde measurements.** Lipid peroxidation levels were determined by measuring the amount of thiobarbituric acid reactive substance (TBARS) malondialdehyde (MDA) in rat plasma samples (n = 6/group). The assay was performed using a kit (Cayman Chemical, MI) following the manufacturer’s instructions.

**Superoxide production.** The formation of superoxide anion was monitored in rat brains during hypobaria using 2-dihydroethidium (Molecular Probes, OR), a fluorescent probe that incorporates between nucleic acid strands of superoxide producing cells. 20 DHE was dissolved in 50% DMSO and administered intraperitoneally at 6 mg/kg immediately before hypobaria or normobaria exposure (n = 4/group). Rats were euthanized at two-hours post-exposure and brain removed, processed, sectioned, and preserved as described. 17,24 DHE incorporated cells were microscopically visualized with a 20× objective at 594 nm wavelength.

**Measurements of reduced and oxidized plasma glutathione levels**

Plasma samples were collected during euthanasia from sham or brain-injured rats at 2 h, 24 h, or 30 days post-hypobaria or normobaria exposure. Total and oxidized glutathione concentration was measured in rat plasma samples (n = 4/group) using a colorimetric assay kit (Life Technologies Inc, MD). In brief, the assay used a colorimetric substrate that reacted with the free thiol group on glutathione to generate a highly colored product. Reduced glutathione concentration was the difference between total and oxidized plasma glutathione concentration measured separately, following the manufacturer’s instructions.

**Transcriptional analysis of glutathione metabolizing genes**

Total RNA was extracted from snap-frozen samples of the ipsilateral prefrontal cortex, parietal cortex, and hippocampus from sham and CCI-rats euthanized 2 h post-hypobaria or normobaria exposure (n = 4/group).
normobaria exposure (n = 4/group). RNA extraction was done using the PureLink RNA Mini kit (Ambion/Life Technologies Inc., CA) following the manufacturer’s instructions. Extracted RNA concentration was determined using a Spectrophotometer Nanodrop 2000C (Thermo Scientific, CA), and RNA tubes were stored immediately at −80°C until used.

First, an Access RT-PCR system kit (Promega, WI) was used to amplify 2 µg RNA and determine the optimal annealing temperature of primer sets designed for the quantitative amplification of glutathione metabolizing genes GS, GPx-1, and GPx-4, as well as the housekeeping control gene glyceraldehyde 3-phosphate dehydrogenase (GAPDH) (Table 1).

Quantitative reverse transcriptase-polymerase chain reaction (Q-RT-PCR) was performed using the Power SYBR Green RNA-to-CT 1-Step kit (Applied Biosystem, MA). In brief, a complete RT-PCR mix containing 150 nM of primer (IDT, CA) that yielded the best annealing temperature for the respective GS, GPx-1, or GPx-4 gene of interest and GAPDH control was mixed with approximately 15 µg of individual RNA template and placed in a 96 wells plate. Q-RT-PCR test of samples in the 96 wells plate was run in a QuantStudio-3 machine (Applied Biosystems, CA) for a total of 35 cycles after reverse transcriptase reaction at 48°C for 30 min and enzyme activation at 95°C for 10 min (n = 4/group). Each of the 35 Q-PCR cycles consisted of 15-s denaturation at 95°C, and 1 min annealing/extension at the temperature noted in Table 1. Each sample was run in triplicate for the target genes GS, Gpx-1, or GPx-4 and duplicate for the reference gene (GAPDH). At the end of the run, the amplification of a single gene product was verified by the presence of a single peak in the melting curve analysis. Normalized cycle threshold (ΔCt) values were obtained by subtracting the Ct of the reference gene from that of each of the target genes. The double delta Ct (ΔΔCt) for each target gene was determined by deducting the mean ΔCt of the target gene in the sham normobaric group from the ΔCt of the respective target gene. The fold-change of each target gene compared to the normobaric sham group was calculated as 2−ΔΔCt.

Lesion volume assessment

Hematoxylin and eosin (Sigma, MO), stained brain sections were dehydrated, cleared, and mounted for lesion volume estimation. Injury lesion volume was quantified from 8 sections corresponding to every 24 serial expanding from bregma 1.05 to −6.30 based to cover the complete lesion area (n = 5/group). Quantification was done using the Cavalieri estimator method of unbiased stereology using the stereo-investigator software (MBF Biosciences, VT). The contusion region (impact-induced damaged region resulting in tissue loss) and penumbra region (the pathologically affected region around impact-induced lesion that could be salvaged with an effective therapeutic intervention) were outlined in the ipsilateral hemisphere of all sections with a grid spacing of 100 µm, and the injury lesion volume was determined and expressed in mm3.25,26

Quantitation and statistical analysis

Image J software (NIH) was used to determine the percent area covered by DHE incorporated cells. To achieve that, 30 microscopic fields of each serial section from DHE or saline-injected rats were captured with SRS fractionator of the stereo-investigator software (MBF Biosciences, VT). Each image was optically segmented with a threshold greyscale value of 0.3 μm, and the percent DHE-positive area was automatically measured. The mean of all DHE positive microscopic fields was determined.

Statistical analyses were performed using GraphPad InStat-3 software (GraphPad Software, Inc., CA). Data from biochemical and histological tests were analyzed using One-Way ANOVA and repeated ANOVA for behavioral data. Both tests were coupled to the Tukey-Kramer post-test to compare differences among the various groups. Results were expressed as mean ± standard error of the mean (SEM). Statistical significance was defined as P<.05. The individuals performing histologic, biochemical, and behavioral assays were blinded to animal group identifications, using codes revealed after data collection.

Table 1. Presentation of forward (F) and reverse (R) primer sequences used for transcriptional analysis of glutathione metabolizing genes GS, GPX-1, and GPX-4. The table also shows the annealing temperature of these primers, their respective target amplicon size, and the mRNA sequence’s access code from which each primer set was derived.

| MRNA  | RAT PRIMER SEQUENCE (5′-3′) | ANNEALING TEMP (°F) | AMPLICON (BP) | ACCESS CODE |
|-------|-----------------------------|---------------------|---------------|-------------|
| GS-F  | ACA CAT CTT CAT TGC CCT CTC | 57                  | 101           | NM_012962.1 |
| GS-R  | TCT ATG GCC ATC TAC CCT CTT |                     |               |             |
| GPx-1-F | GTT CCA GTG CGC AGA TAC A | 57                  | 111           | NM_030826.4 |
| GPx-1-R | CCA GAT ACC AGG AAT GCC TTA G |                     |               |             |
| GPx-4-F | CCC ACT GTG GAA ATG GAT GA | 58.5                | 105           | NM_017165.4 |
| GPx-4-R | ACG CAG CCG TTC TTA TCA A |                     |               |             |
| GAPDH-F | CCC TTC ATT GAC CTC AAC TAC | 55                  | 102           | NM_017008.4 |
| GAPDH-R | GAT GAC CAG CTT CCC ATT CT |                     |               |             |
Results

Hypobaria exposure reduces circulating homocysteine levels

To determine the effect of acute hypobaria-exposure on HCY metabolism, we measured plasma HCY levels at 2 h and 24 h after 6 h of 8000 ft hypobaria-exposure using normobaria-exposure as control. We found that mTBI alone significantly reduced plasma HCY levels in normobaria-rats after 2 h after exposure, 1.41 ± 0.723 µM, compared to normobaria-shams, 4.32 ± 0.431 µM (Figure 1; $Q = 4.31, P < .05$). At the same 2 h time point, HCY levels were further reduced or undetectable in the plasma of TBI rats exposed to hypobaria to 0.27 ± 0.28 µM and were significantly lower than levels in normobaric (3.12 ± 0.63 µM) and hypobaric (4.32 ± 0.431 µM) shams (Figure 1; $Q = 0$, $P < .05$; $Q = 5.88$, $P < .01$ respectively).

HCY levels in CCI-hypobaria animals were still significantly reduced 24 h after hypobaric 1.85 ± 0.63 µM compared to normobaric shams (Figure 1; $Q = 4.23$, $P < .05$). Hypobaria-exposure also caused a non-significant decrease in HCY levels among non-TBI sham rats with levels of 3.12 ± 0.63 µM at 2h post-exposure and 2.07 ± 0.52 µM at 24h post-exposure compared to normobaria-shams 4.32 ± 0.431 µM (Figure 1; $P > .05$).

Hypobaria-induced increases oxidized glutathione

We measured reduced and oxidized glutathione levels in that plasma. This showed that hypobaria-exposure caused a modest increase in oxidized glutathione in sham rats at 2 h post-exposure (1.25 ± 0.23 µM) compared to normobaria-shams (0.76 ± 0.19 µM; $P > .05$; Figure 2A). TBI alone had no effect on glutathione oxidation among normobaria-rats at 2h post-exposure (0.723 ± 0.158 µM) compared to normobaria-shams (0.76 ± 0.19 µM; $Q = 0.56$, $P > .05$; Figure 2A). However, the combination of TBI and hypobaria significantly increased plasma oxidized glutathione to 1.998 ± 0.296 µM compared to CCI-normobaria and CCI-hypobaria 24-h and 30 days post-surgery (Figure 2A; $Q = 4.41$, $P < .05$ and $Q = 9.23$, $P < .01$ respectively).

The ratio of oxidized to reduced glutathione is an important metric of the metabolic impact of TBI and hypobaria on glutathione metabolism. When we assessed the ratio of oxidized to reduced glutathione, we found that hypobaria-exposure substantially enhanced oxidized glutathione over reduced glutathione in both sham and CCI rats resulting in 1.57 ± 0.202 and 2.38 ± 0.323-ratios respectively, compared to 0.741 ± 0.164 µM and 0.99 ± 0.22 µM for normobaria-sham and CCI rats (Figure 2B, $P < .05$). This ratio decreased significantly overtime at 24 h and 30-days (Figure 2B; $Q = 4.61$, $P < .05$ and $Q = 6.79$, $P < .01$) post-exposure in a time-dependent manner. The elevation of oxidized glutathione implies increased free radical scavenging through glutathione oxidation,27 which could activate the endogenous glutathione oxidizing metabolic pathways.

Hypobaria induces upregulation of glutathione metabolizing genes

We examined the transcriptional profile GS, GPx-1, and GPx-4 in 3 regions of the ipsilateral hemisphere, the prefrontal cortex, parietal cortex, and hippocampus, at 2h post-exposure, to determine how their expression responds to hypobaria-exposure. All results are presented in Table 2. These

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**Figure 2.** Hypobaria exposure enhances glutathione oxidation: (A) quantitative graph of oxidized glutathione levels showing that glutathione oxidation shows a non-significant increase by hypobaria at 2h post-exposure in sham rats ($n=4$). It is significantly intensified by the combination of mTBI and hypobaria exposure ($n=4$, **$P < .05$, *$P < .01$) and (B) histogram of the ratio of oxidized to reduced glutathione showing a 1.8-fold increase among hypobaric shams relative to normobaric shams ($n=4$; $P < .05$) at 2h post-exposure. In contrast, there was more than a two-fold difference between hypobaric-CCI rats compared to normobaric-CCI rats ($n=4$; *$P < .05$). Glutathione oxidation reduced significantly overtime after hypobaria (*$P < .05$, **$P < .01$, ***$P < .001$).
results show a heterogeneous expression pattern in different brain regions. In the prefrontal cortex, the expression of all 3 genes was enhanced by the cortical impact-induced brain injury (Table 2; Figure 3A–C; $Q=4.51$, $P<.05$; $Q=6.90$, $P<.01$; $Q=9.83$, $P<.001$) with no significant effect of hypobaric-shams.

In contrast, the combination of cortical brain injury and hypobaric-exposure upregulated the expression of GS and GPX-1 in the parietal cortex compared to normobaric brain-injured rats as well as hypobaric and normobaric shams (Table 2; Figure 3D, E; $Q=6.26$, $P<.01$). There was no significant change in GPX-4 expression in the parietal cortex of rats in any group (Table 2; Figure 3F; $P>.05$). In the hippocampus, GS expression was significantly upregulated by hypobaria alone in shams compared to other groups (Table 2; $Q=15.32$, $P<.001$). Similar to the parietal cortex, hypobaria or CCI only increased the expression of GPX-1 in the hippocampus and the combination of both resulted in more upregulation, although all are non-significant (Table 2; Figure 3H, $P>.05$). The expression of GPX-4 in the hippocampus was increased by injury or hypobaria alone, and no additional effect of the 2 combined (Table 2, $Q=7.1$, $P<.01$; $Q=9.96$, $P<.001$). In summary, the expression of glutathione metabolizing genes was differentially affected by brain injury, hypobaria, or by the combination of both factors, dependent on the targeted brain region.

**Hypobaric conditions increase oxidative stress**

The biochemical markers above indicate that there may be considerable oxidative stress causing a compensatory reduction in HCY and elevation in the processing of glutathione antioxidant systems. Consistent with this conjecture, we found that TBI caused a non-significant $12.3\pm5.23\%$ increase in plasma malondialdehyde levels in normobaria-CCI rats at 2 h post-exposure compared to normobaria-shams (Figure 4; $P>.05$). In addition to the modest increase in TBI, when combined with hypobaria-exposure, there was a significant increase in MDA formation of $55.38\pm10.5\%$ over baseline (Figure 4; $Q=6.83$, $P<.01$). Although there is generally a good correlation between blood and brain biomarkers in TBI studies, these measures of oxidative stress do not give information on where in the body they originate. To verify which brain regions were impacted by hypobaria induced free radical formation during hypobaria exposure, we used 2-DHE for in vivo tracking of superoxide production in brain cells (Figure 5A) and compared conditions by measuring the percentage of pixels in defined brain regions that are 2-DHE positive. There was significant increase in 2-DHE incorporated cells in normobaria-CCI rats ($2.42\pm0.18\%$ area covered) compared to normobaria-shams ($1.02\pm0.16\%$; Figure 5B; $Q=0.46$, $P<.05$). Hypobaria-exposure alone did not cause a significant change in superoxide production in

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**Table 2.** Presentation of transcriptional analysis results of glutathione metabolizing genes GS, GPX-1, and GPX-4 in the prefrontal cortex, the parietal cortex of shams, and CCI rats exposed to normobaric or hypobaric conditions. These results represent the fold expression change of each of these genes in different experimental groups compared to normobaric-shams used as control.

| STUDY GROUPS | PREFRONTAL CORTEX | PARIETAL CORTEX | HIPPOCAMPUS |
|--------------|--------------------|-----------------|-------------|
|              | GS                 | GPX-1           | GPX-4       |
| Sham, NB     | $1\pm0.086$        | $1\pm0.035$     | $1\pm0.06$  |
| Sham, HB     | $1.33\pm0.24$      | $1.28\pm0.15$   | $1.22\pm0.14$ |
| CCI, NB      | $7.23\pm0.53$      | $4.5\pm0.32$    | $3.69\pm0.95$ |
| CCI, HB      | $8.43\pm1.51$      | $14.5\pm0.58$   | $1.46\pm0.15$ |

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$Q$ values are given in parentheses.
sham rats resulting in 1.59 ± 0.18% versus normobaric-shams (1.02 ± 0.16%; Figure 5B; P > .05). However, in the presence of hypobaria with CCI, there was a marked increase in 2-DHE positive cells present with 5 times the area compared to sham (TBI/CCI 4.58 ± 0.62% of the brain region; Figure 5B; Q = 0.68, P < .05) and normobaria-CCI rats (Q = 0.46, P < .05). Thus, hypobaria-exposure exacerbates TBI associated oxidative stress.

**TBI lesion volume was not significantly affected by hypobaria exposure**

To assess the effect of hypobaria-exposure on brain injury lesion, we used the Cavalieri method of unbiased stereology to measure injury-induced lesion volume at 2h and 24h post-exposure. At 2h post-exposure, impact-induced injury lesion volume was approximately the same between normobaric, 6.56 ± 2.25 mm³, and hypobaric-CCI rats, 7.09 ± 1.42 mm³ (Figure 6A, B; P > .05). But at 24h post-exposure, corresponding to 48h postsurgery, brain injury lesion volume presented a non-significant increase following hypobaria exposure, 13.68 ± 3.05 mm³ compared to 6.78 ± 2.48 mm³ among normobaric-rats (Figure 6A, B; P > .05). This suggests that hypobaria exposure has no acute negative impact on TBI brain tissue loss.
Next, we wanted to determine whether hypobaria exposure following mTBI in animals exacerbates behavioral deficits. We assessed the effect of hypobaria-exposure on anxiety-like behavior usually associated with TBI-related psychiatric impairment such as posttraumatic stress disorder. Anxiety was assessed with a standard elevated plus-maze in which reduced time spent in the open arms of the maze is interpreted as elevated anxiety-like behavior. As a control, fine motor behaviors classically associated with cerebellar function, a brain region removed from the injury site, were not significantly affected as assessed by the Beam Walk test. On day 2 post-exposure, which is an early time point when changes due to hypobaria are mostly likely detectable, normobaric sham and CCI rats respectively made $1.28 \pm 0.98$ and $1.87 \pm 0.62$ foot faults over 50 steps. In comparison, hypobaric sham and CCI rats made $1.39 \pm 0.75$ and $1.77 \pm 1.02$ foot faults ($P > .05$, all compared to normobaric shams). However, brain-injured rats exposed to hypobaria spent significantly less time in the open arms of Plus maze on days 2 and 8 post-injury, indicative of increased anxiety compared to normobaria-shams. These animals spent $48.21 \pm 10.74$ s and $94.08 \pm 21.49$ s in the open arms of the Plus maze on days 2 and 8 respectively versus $133.04 \pm 25.28$ s and $241.82 \pm 29.12$ s for normobaria-shams (Figure 7; $Q = 5.21$ and 5.58 respectively; $P < .05$). To a lesser
extent, hypobaria-sham and normobaria-CCI rats also exhibited some deficits in anxiety-like behavior on days 2 and 8 post-surgery (Figure 7; Q < 4.96 thresholds for significance; P > .05), compared to normobaria-shams. After 8 days, rats in all groups recovered from this deficit and exhibited Plus maze performance close to normobaria-shams.

**Discussions**

Reports from several investigators suggest that flying at high altitude is associated with increased physiological stress, thus representing a risk for TBI patients who require aeromedical evacuation. Results from the present study indicate that exposure of rats with mTBI, which represents a non-significant in anxiety-like behavior. The combination of mTBI and hypobaria significantly increased anxiety-like behavior on days 2 and 8 post-surgery compared to normobaria-shams (n = 8/group; P < .05). Two and 4 weeks post-HB, this behavioral deficit was resolved, and rats in all groups spent closely the same time in the plus-maze open arms (P > .05).

**Figures**

**Figure 7.** Traumatic brain injury-induced anxiety-like behavior is aggravated by hypobaria exposure. Graph depicting the time spent by rats in the open arms of the elevated plus maze. Both brain injury and hypobaria alone caused a non-significant increase in anxiety-like behavior. The combination of mTBI and hypobaria significantly increased anxiety-like behavior on days 2 and 8 post-surgery compared to normobaria-shams (n = 8/group; P < .05). Two and 4 weeks post-HB, this behavioral deficit was resolved, and rats in all groups spent closely the same time in the plus-maze open arms (P > .05).

with substantial cytotoxic potential when accumulated. Therefore, its metabolism is tightly regulated to ensure it is continuously eliminated. This occurs via its remethylation to methionine, its precursor, or by its transsulfuration to cystathionine, contributing to the downstream formation of glutathione, a very potent antioxidant.

Therefore, we postulated that the drastic reduction of homocysteine levels under oxidative stress at 2-h post-hypobaria was indicative of its flush through the transsulfuration pathway. This hypothesis was potentiated by a significant increase of glutathione oxidation resulting in a two-fold increase of oxidized glutathione versus reduced glutathione at 2h post-hypobaria exposure (Figure 2A, B). The return of oxidized glutathione to baseline levels at day 30 post-exposure suggests an attenuation of oxidative stress. This observation correlates with the significant improvement in anxiety-like behavior to that of normobaria-sham rats. The therapeutic potential of homocysteine transsulfuration to promote glutathione formation has been underscored in several neurological disorders. For instance, the supplementation of N-acetylcysteine to promote glutathione biosynthesis through the transsulfuration pathway was neuroprotective in models of Huntington’s disease, Alzheimer’s disease, and TBI. Moreover, this therapeutic approach is in a trial for pediatric TBI management.

Glutathione has free radical scavenging and anti-inflammatory properties. Its metabolism is regulated by genes, including GS, which controls its biosynthesis through the transsulfuration pathway and GPx, which catalyzes glutathione oxidation to attenuate free radical formation. GPx has 8 isozymes, among which isozymes 1 and 4 are abundant in the brain. We analyzed the transcriptional profile of these genes in the prefrontal cortex and the hippocampus that are very vulnerable to oxidative stress and the parietal cortex around the cortical injury penumbra region. The upregulation of GS and GPx in the prefrontal cortex (Figure 3A–C) attests to its vulnerability to oxidative stress, relative to its distant location to the injury region in the parietal cortex. In the parietal cortex, the combination of CCI and hypobaria resulted in an increase in GS and GPx that was not present with CCI or hypobaria alone (Figure 3D–F). This suggests that the CCI penumbra region may be uniquely stressed by a synergy between the injury and hypobaria. To our knowledge, this study is the first to describe changes in glutathione and associated gene expression in a model of TBI and Hypobaria exposure.

Although the mechanism for the distinctive expression patterns in different brain regions is not fully elucidated, increased GPx expression in the cerebral cortex of aged rats was associated with higher astrocyte density. Furthermore, GPx activity was derived primarily from astrocytes relative to neurons in chicks’ forebrain. Also, increased GPx expression or increased glutathione oxidation was reported in several oxidative stress-associated neurological disorders.
In summary, this study provides evidence that homobaria exacerbates oxidative stress in rats with mTBI. This was associated with an enhanced flush of homocysteine through the transulfuration pathway to promote glutathione biosynthesis, which is necessary to attenuate free radical formation. Increased glutathione metabolism was putatively driven by the upregulation of glutathione metabolizing genes, suggesting that enhanced glutathione metabolism represents a therapeutic target for TBI management. Further studies would be needed to understand how this antioxidant redox system would respond to interventions during and after homobaria-exposure following moderate to severe TBI.

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Author Contributions

Experimental design: FT, GF, CM; Laboratory experiments and data collection: FT, MG; Data analysis: FT, AP, GF, MG; Manuscript writing: FT, GF; Manuscript editing: AP, MG, FT.

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