Is Hyperbaric Oxygen Therapy Effective for Traumatic Brain Injury? A Rapid Evidence Assessment of the Literature and Recommendations for the Field

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Objective: This systematic review examines the efficacy of hyperbaric oxygen (HBO2) for traumatic brain injury (TBI) to make evidence-based recommendations for its application and future research. Methods: A comprehensive search was conducted to identify studies through 2014. Methodological quality was assessed and synthesis and interpretation of relevant data was performed. Results: Twelve randomized trials were included. All mild TBI studies demonstrated minimal bias and no statistically significant differences between HBO2 and sham arms. Statistically significant improvement occurred over time within both groups. Moderate-to-severe TBI studies were of mixed quality, with majority of results favoring HBO2 compared with “standard care.” The placebo analysis conducted was limited by lack of details. Conclusions: For mild TBI, results indicate HBO2 is no better than sham treatment. Improvements within both HBO2 and sham groups cannot be ignored. For acute treatment of moderate-to-severe TBI, although methodology appears flawed across some studies, because of the complexity of brain injury, HBO2 may be beneficial as a relatively safe adjunctive therapy if feasible. Further research should be considered to resolve the controversy surrounding this field, but only if methodological flaws are avoided and bias minimized. Key words: hyperbaric oxygen therapy, placebo, sham, systematic review, traumatic brain injury
HYPERBARIC OXYGEN (HBO₂) has been proposed as an adjunctive therapy for traumatic brain injury (TBI).1 Reports from the 1960s claimed improvement for patients with TBI undergoing HBO₂; however, much of the clinical trial work then and now shows mixed results and the field remains controversial.1–3 Potential adverse events of HBO₂, such as barotrauma and oxygen poisoning, and the clinical significance of these risks versus benefits are not yet fully understood.2,3 HBO₂ treatment for TBI has not been cleared or approved by the US Food and Drug Administration.4,5 There remains a lack of solid clinical evidence supporting the standard use of HBO₂ for TBI.

Appropriate and effective treatments for patients suffering from the consequences of TBI remain unknown to date. With the increasing rate of reported TBI cases, and our soldiers returning home from war, effective treatments are imminently needed. The purpose of this systematic review is to provide an independent, objective, and transparent analysis of the research conducted to date exploring the evidence base for HBO₂ as a therapeutic option for TBI. Specifically, this review (1) examines the quantity and quality of the research on HBO₂ for TBI; (2) provides a concise understanding as to the safety, dosing variations, and efficacy of this intervention, as it is currently reported in the literature; (3) compares the differences noted across studies and gaps that currently exist in the research; (4) explores the use of sham and placebo in these randomized controlled trials (RCTs) to the extent that data are available for analysis; and (5) draws conclusions upon which recommendations can be made concerning next steps for the field of study.

METHODS

Concepts and definitions were clearly defined and agreed upon by a diverse steering committee (SC) and subject matter experts (SMEs) for both TBI and HBO₂ therapy.6–8 The authors considered all literature looking across the spectrum of TBI sequelae from acute to chronic effects and from mild-to-severe TBI, both in military and civilian populations. Although the definition for TBI seems intuitively correct for moderate-to-severe TBI, it is significantly nuanced for mild TBI (mTBI). Because of the entanglement of symptoms and comorbidity between and within TBI, patients with posttraumatic stress disorder (PTSD) were not excluded as a population from the review.

Study eligibility criteria

Studies were included if they met the following criteria: (1) peer-reviewed study designs presented in the English language; (2) involving subjects suffering from the consequences of TBI in both military and civilian populations; and (3) HBO₂ is being used as the intervention without preexisting conditions. All clinical outcomes, as well as types of control/comparators, were considered for inclusion to ensure capturing all studies relevant to the research question.

Data sources and search strategy

PubMed, CINAHL, PsycINFO, and Cochrane, as well as the Database of Randomized Controlled Trials in Hyperbaric Medicine, were searched from their database inception through December 2014 using the following search terms in PubMed and adapted as necessary in other databases: brain injuries or head injuries or cranioencephal trauma AND hyperbaric oxygenation or “hyperbaric oxygen therapy” or “hyperbaric oxygen” or “hyperbaric therapy” or HBO or HBO₂ or HBOT. In December 2014, an updated search was conducted and the additional search terms were included to ensure any entanglement with PTSD was captured in the review: “stress-disorders, post-traumatic” and “PTSD.”

All searches were restricted to the peer-reviewed, English language studies involving human subjects.

Study selection

Two investigators independently screened titles and abstracts of the citations yielded from the literature search using the predefined study eligibility criteria. Interrater reliability as measured by Cohen’s κ9 was more than 0.90 throughout the screening phase.

Quality assessment and data extraction for RCTs

Methodological quality was independently assessed by 3 reviewers in duplicate using the Scottish Intercollegiate Guidelines Network Checklist for RCTs (SIGN 50).10 External and model validity were assessed using Samueali Institute’s External Validity Assessment Tool (EVAT, Alexandria, Virginia).11 Descriptive data were also extracted and study designs other than RCTs were qualitatively described.

Placebo analysis

When available, information about the sham or control arm for each study was abstracted for placebo analysis as predefined attributes by the SC and SMEs. Parameters included expectation, conditioning, and social learning attributes.

Outcome categorization

Outcome assessment tools for included RCT studies were obtained. These tools were grouped into higher order categorization relevant to TBI and examined using the Common Data Element Project’s classification of outcomes.12
Data synthesis and analysis

Meta-analysis was not performed because of lack of data available. The authors instead synthesized the evidence at the outcome level to (1) determine the confidence in the estimate of the effect; (2) assign a safety grade; and (3) provide an overall recommendation concerning the benefit/risk for the intervention based solely on the evidence being presented through an expert roundtable convening with the diverse SC and SMEs. 

RESULTS

Study selection

The initial database search yielded 393 distinct citations. An additional 2 eligible clinical trials, published after August 2014, were also included. Twelve unique RCTs, published in 16 reports, met the inclusion criteria (see Figure 1). Of these studies, 4 RCTs involved mTBI populations suffering persistent symptoms over many months; 7 RCTs involved moderate-to-severe TBI populations treated acutely; and 1 article did not clearly define the severity of the TBI. (See Supplemental Digital Content A, available at: http://links.lww.com/JHTR/A182.) An additional 27 non-RCT studies met the eligibility criteria and are described in Supplemental Digital Content B (available at: http://links.lww.com/JHTR/A183). 

Quality assessment and risk of bias for included RCT studies

Eight of the included RCTs were rated as acceptable (+) and 4 were rated as low (0) methodological quality according to SIGN 50 criteria. (See Supplemental Digital Content C, available at: http://links.lww.com/JHTR/A184.) According to EVAT criteria, recruitment was predominately adequately covered, indicating that the majority of studies identified the source population and described recruitment methods. Participation was largely well covered, indicating study participants seemed representative of the source population. Model validity was mainly poorly addressed, indicating the manner by which the patients were treated in the studies was not described well enough to understand whether they are representative of the care model (staff, processes, facility, equipment, other resources) as they are in practice. (See Supplemental Digital Content D, available at: http://links.lww.com/JHTR/A185.)

When considering model validity, there was a wide dispersion in the staff conducting HBO2 treatments: an attending physician in 1 study; chamber technicians, study coordinators, and supervising physicians in a second study; and a research crew (crew chief, chamber operator, inside attendant, and medical monitor) in a third study. The specific type of researcher was not noted in the remaining 9 studies. For clinical setting, 7 studies were carried out at a hospital, of which was a multisite study conducted at 4 military hospitals. Additional sites included a naval medicine operational training center, the US Air Force School of Aerospace Medicine, and a hyperbaric institute and research unit of a medical center. Two studies did not describe the study location. Ten of the 12 studies included details of equipment used to some extent; this mostly entailed describing the type of HBO2 chamber used. Five studies used multiplace, and 3 used monoplace chambers. An additional study placed some subjects in a multiplace chamber and others in a monoplace chamber. No studies compared outcome changes between a monoplace and a multiplace chamber or described in detail the environment within the chambers.

Characteristics of mTBI studies

Four studies, involving 250 participants, investigated the use of HBO2 for those suffering the consequences of mTBI. All 4 studies received a SIGN 50 score of acceptable (+). Three of these studies were supported by the Department of Defense, involved military populations and compared HBO2 to a type of sham. The final study was a nonmilitary crossover study with a 2-month washout period. These studies involved various dosages in terms of exposure time to HBO2, with 30 to 40 sessions ranging from 60 to 117 minutes, over 8 to 10 weeks, and pressure ranging from 1.5 to 2.4 ATA (atmospheres absolute).

The types of sham arms differed in terms of pressure and oxygen levels. The first study administered 10.5% oxygen at 2.0 ATA as the sham. The second study administered 21% oxygen (the same concentration as room air) at 1.3 ATA with a slow drift to 1.2 ATA. The third study administered 21% oxygen at 1.2 ATA. Two of these studies were 3-armed trials. One study compared HBO2 plus “TBI-care” to sham plus “TBI-care” and “TBI-care” only. The other study exposed all groups to 2.0 ATA, but inspired oxygen was varied (100%, 75%, and 10.5%) to represent the tissue partial pressure of oxygen equivalent to the inspiration of 100% oxygen at each of 2.0, 1.5, and 1.0 ATA, respectively.

All mTBI studies assessed cognitive performance; 3 assessed psychological outcomes and 2 assessed postconcussive symptom severity. Overall these studies showed no statistically significant results between groups for any of these outcomes. However, there were some statistically significant within-group
improvements within both HBO2 and sham groups across all relevant outcomes17–22 and both groups in the crossover study for cognitive performance.23 In one study, both HBO2 and sham groups improved for cognition and postconcussion symptom severity whereas the “TBI-care” only group worsened or stayed the same respectively.21 (See Supplemental Digital Content A, available at: http://links.lww.com/JHTR/A182.)

**mTBI reported adverse events**

Adverse events were categorized into 4 categories: ear problems, pulmonary complications, other minor, and other serious adverse events. Three of the 4 mTBI studies describe adverse events19–21,23 whereas the remaining study did not mention adverse events.17,18,22 All 3 studies describe minor ear problems including barotrauma and ear pain. Two studies reported minor adverse events: nausea, sinus squeeze and sinus pain,
claustraphobia, headache, musculoskeletal chest pain, latex allergy, tooth pain, and transient worsening of myopia. Pulmonary complications and serious adverse events were not reported. (See Supplemental Digital Content A, available at: http://links.lww.com/JHTR/A182.)

Characteristics of moderate-to-severe TBI studies

Seven studies, involving 486 participants, investigated the use of HBO2 for those who had suffered moderate-to-severe TBI and were being treated acutely. Four of these studies received a SIGN 50 score of acceptable (+)25–27,29,31 and 3 were rated low (0) quality.24,28,32 The HBO2 dosages for these studies involved a large range with 3 to 84 sessions, ranging from 40 to 120 minutes and over a period of 3 to 52 days. All studies used 100% inspired oxygen and varied pressure with a range of 1.5 to 2.5 ATA.

One study compared HBO2 with no HBO2 treatment.26 Six other studies compared HBO2 with various forms of "standard care,"24,25,27–29,31,32 one of which was a 3-armed study comparing HBO2 with "standard care" to normobaric hyperoxia.28 These studies were conducted over nearly a 40-year time span, and what was referred to as standard of care was inconsistent across the studies. None of the studies fully described the standard neurotrauma care in enough detail to compare the control arms across the studies.

Four studies involved outcome measures of consciousness24,26,27,29,31; 5 utilized the Glasgow Outcomes Scale (GOS) and/or measures of mortality24–27,31; and 2 involved intracranial pressure (ICP).25,28 In all 3 acceptable (+) quality studies reporting on consciousness, the HBO2 groups experienced statistically significant better recovery compared with "standard care" (n = 2) and no HBO2 treatment (n = 1). The low (0) quality study found no statistically significant difference between groups overall. Results for the 3 acceptable (+) quality studies reporting on the GOS and/or mortality conflicted: in 2 studies, the HBO2 groups showed statistically significant better scores versus "standard care,"25,27 whereas the third study found no statistically significant differences between the HBO2 and no HBO2 groups overall.26 Both low (0) quality studies reporting on mortality rate found no statistically significant differences between HBO2 and "standard care" groups.24,30 Two studies measured ICP: the acceptable (+) quality study reported no statistically significant difference between the HBO2 and "standard care" groups.25 whereas the low (0) quality study found ICP measurements significantly lower after each treatment in the HBO2 group compared with the "standard care" group.28 (See Supplemental Digital Content A, available at: http://links.lww.com/JHTR/A182.)

Moderate-to-severe TBI reported adverse events

Three of the 7 included studies involving moderate-to-severe TBI populations describe adverse events,24–26 whereas the remaining 4 do not.27–31 These reports describe various ear problems including severe ear pain (resolved by tympanostomy) and hemotympanum; 2 describe pulmonary adverse events; and 2 report the occurrence of seizures. (See Supplemental Digital Content A, available at: http://links.lww.com/JHTR/A182.)

Study in which TBI severity not described

One study, in which severity was not described, was a low (0) quality study comparing HBO2 with a “standard medication” for 320 patients with postbrain neural injury.25 The HBO2 group was exposed to 0.1 megapascal (MPa) pressure, nearly equivalent to 1.0 ATA, while inspiring 96% oxygen. The authors did not report on or mention adverse events nor did they report on any outcomes relevant to this review.

Placebo analysis

Placebo might explain the paucity of between-group and the preponderance of within-group differences (improvements) between HBO2 and sham arms in the mild TBI review group studies. Placebo effects are context sensitive, dependent on expectancy, conditioning, and social learning.61 The placebo analysis was limited by the lack of details describing the sham or control arms of the studies (see Table 1).

The majority of studies report that informed consent was obtained from the patient or family member; however, only one study details the informed consent process.23 The description of the therapeutic intervention and sham during informed consent process can influence expectancy and meaning for the patient and affect the results of the study.62 Although the setting/patient experience and study location was reported in almost all studies, the full description was missing preventing context analysis. Studies reported whether a multipurpose chamber or monochamber was used but did not describe the environment either inside or outside the chambers or whether patients studied in multipurpose chambers were alone or with others (attendants or patients). The social experience or encounter is not discussed in these studies. The characteristics of the researcher/practitioner and their credentialing (believability), relationship between the research team and patient, and instructions given to the patient are not reported on in the majority of the studies. The human interaction element was not described in any of these studies. Only 2 studies described instructions given to subjects.17,18,22,23 None of the studies stated that expectation of the patient, attendant, or researcher www.headtraumarehab.com
was predetermined in the study’s method or measured during the encounter. Eight of the 12 studies conducted follow-up on specific outcome measures; 6 studies report whether blinding took place, however only 2 describe this procedure and state only that blinding was successfully carried out. One of these studies followed up with participants by asking whether they thought they participated in the HBO2 or sham arm.21 Compliance measures were reported in 6 of the studies, the majority of which simply reported subject dropout percentages. Patients dropped out because of scheduling conflicts, HBO2-related adverse events, medical issues, or incurring an additional mTBI injury. Fewer than half of the studies described the reason for selecting specific attributes in the control/comparator arm. Two studies failed to describe the comparator arms in any detail.24,30 Three of the 12 studies concluded their results are likely due to the placebo effect.17–22 Two studies reference that a placebo and/or Hawthorne effect can be expected in randomized sham-controlled trials of interventions in populations with depression, anxiety, and PTSD because of the “intense nature of the intervention.”17–22 Researchers in one study opined that participants, who were Marines, benefited from the study because they traveled to a resort town, and took leave from work.17,18,22 Another study detailed that it was not designed to evaluate components of a placebo response, including response expectancy, verbal suggestion, and stimulus conditioning.21 This placebo analysis demonstrates that although some of these components bearing on placebo were reported in the studies, no study described in the detail necessary to understand the influence on outcomes of any placebo response that may or may not have occurred.

**Discussion**

What are the clinical and policy implications of the current evidence for HBO2 therapy in regard to TBI? To better understand these implications, a synthesis of the evidence was conducted by the SC and SMEs for each outcome category (see Table 2). Recommendations balance safety with efficacy of the procedure for a particular outcome. These recommendations are based solely on the presented systematic review results and are not to be construed as clinical practice guidelines in any way. Systematic review is the first step needed to make evidence-based informed decisions, but further key ingredients are necessary for a clinical practice guideline to be established.

**Synthesis of the evidence—mTBI populations**

Of the 3 SIGN 50 acceptable (+) studies involving a total 183 patients with mTBI suffering persistent symptoms, comparing HBO2 with sham, synthesis and analysis was conducted on cognitive, postconcussive symptom severity, and psychological outcomes. Two of the 3 studies report on adverse events. There seem to be infrequent adverse events and HBO2 appears to be safe for mTBI populations. However, given there are only 3 studies, safety is not well understood.

Although no statistically significant differences were noted across studies when comparing HBO2 with sham for the outcomes assessed, they did show a trend that patients improved over time within both the HBO2 and sham treatment groups. Effect sizes were not reported for any outcomes within the individual studies, and although validated tools were used, the sensitivity of these tools specific to TBI populations is questionable. Consensus was reached that further research is very unlikely

### Table 1: Placebo analysis of essential components for reporting

| Component                                      | Reported n (%) | Not reported n (%) |
|------------------------------------------------|----------------|--------------------|
| Informed consent                              | 8 (66.7)       | 4 (33.3)           |
| Setting/participant experience                 | 11 (91.7)      | 1 (8.3)            |
| Multichamber                                   | 5              |                    |
| Monochamber                                    | 3              |                    |
| Both multi- and monochamber                    | 1              |                    |
| Unclear                                        | 2              |                    |
| Study location                                 | 9 (75.0)       | 3 (25.0)           |
| Placebo effect                                 | 3 (25.0)       | 9 (75.0)           |
| Characteristics of researcher/practitioner     | 3 (25.0)       | 9 (75.0)           |
| Relationship between researcher and subject    | 2 (16.7)       | 10 (83.3)          |
| Follow-up                                      | 9 (75.0)       | 3 (25.0)           |
| Reason for control/comparator                  | 5 (41.7)       | 7 (58.3)           |
| Blinding                                       | 6 (50.0)       | 6 (50.0)           |
| Compliance                                     | 6 (50.0)       | 6 (50.0)           |
| Expectancy                                     | 0 (0.0)        | 12 (100.0)         |
| Instructions to patients during time within chamber | 2 (16.7)   | 10 (83.3)          |
### TABLE 2  Quality assessment of the overall literature pool by outcome category

| Treatment | Number of participants completed (number of studies) | Confidence in estimate of effect | Reported studies safety grade (number of studies)/overall safety | Strength of the recommendation |
|-----------|--------------------------------------------------------|---------------------------------|---------------------------------------------------------------|-------------------------------|
| **Mild TBI** Cognitive | | | | |
| HBO₂ vs sham | 183 (3) | A | +2(2)/0 overall | Weak recommendation against |
| HBO₂ vs “standard care” | 72 (1) | B | +2(1) | No recommendation |
| **Postconcussive symptom severity** | | | | |
| HBO₂ vs sham | 133 (2) | A | +2(1)/0 overall | Weak recommendation against |
| HBO₂ vs “standard care” | 72 (1) | B | +2(1) | No recommendation |
| **Psychological** | | | | |
| HBO₂ vs sham | 183 (3) | A | +2(2)/0 overall | Weak recommendation against |
| HBO₂ vs “standard care” | 72 (1) | B | +2(1) | No recommendation |
| **Moderate-to-severe TBI** Alterations in the level of consciousness | | | | |
| HBO₂ vs “standard care” | 219 (4) | C | 0 | Weak recommendation in favor |
| Intracranial pressure | | | | |
| HBO₂ vs “standard care” | 237 (2) | B | 0 | No recommendation |
| Glasgow Outcome Scale and mortality | | | | |
| HBO₂ vs “standard care” | 357 (5) | C | 0 | Weak recommendation in favor |

Abbreviations: HBO₂, hyperbaric oxygen; TBI, traumatic brain injury.

*This analysis usually also incorporates overall effect size according to outcomes; because of the heterogeneity among the studies and poor reporting, this was not done.

Further research is very unlikely to change our confidence in the estimate of effect (A); further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate (B); further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate (C); any estimate of effect is very uncertain (D).

Safety grade ranges from +2 appears safe with infrequent adverse events and interactions to −2 appears to have serious safety concerns that include frequent and serious adverse events and/or interactions.

Strength of recommendation ranges from “strong recommendation in favor”—SME is very certain that benefits do outweigh risks and burdens—to “strong recommendation against”—SME is very certain that benefits do not outweigh the risks and burdens.

to change the team’s confidence in the estimate of the effect. Given the evidence for HBO₂ for patients with mTBI when compared with a sham intervention for cognition, postconcussive symptom severity, and psychological impairment, a *weak recommendation against* the use of HBO₂ was reached at the expert roundtable. There does not appear to be added value of this therapy over and above the sham treatment, if in fact these are truly “sham” treatments.

Only one study compared HBO₂ with “standard care” within the mTBI population. Highly powered, strong methodological studies with multiple comparator arms are needed to begin to draw conclusions to make recommendations in regard to the risk/benefit for HBO₂ versus “standard care.”

**Synthesis of the evidence—moderate-to-severe populations**

Of the 4 SIGN 50 acceptable (+) and 3 low (0) quality studies involving a total of 486 patients at the acute stage of moderate-to-severe TBI, synthesis and analysis was performed across alterations in the level of consciousness, GOS and mortality, and ICP where applicable, in comparison to no HBO₂ treatment or some type of “standard care.” The low (0) quality studies suffer from methodological flaws reported on by this systematic review, which leaves us to question the validity of the results being reported. Given the nature of these patients’ condition, research becomes quite challenging. Many of these studies are also quite old dating...
to the early 1970s. Clinical approaches to neurotrauma have evolved greatly over the last 45 years. Only 3 of the 7 studies reported on adverse events. It was challenging to assess whether these adverse events occurred because of the nature of the population’s condition or were caused by the intervention. The SC experts agreed that safety is not well understood for HBO2 for moderate-to-severe TBI populations.

Alterations in the level of consciousness were assessed in 4 studies, and the GOS and mortality were assessed in 5 studies. Given the mixed quality of these studies, the fact that many of the studies are quite old, and the sample sizes are relatively small, the SC experts agreed that for consciousness and mortality outcomes further research is very likely to have an important impact on the confidence in the estimate of the effect. Given the nature and severity of these patients’ brain injuries, however, and noting that HBO2 does appear to improve patients’ outcomes versus “standard care,” the SC experts gave a weak recommendation in favor of HBO2 as compared with “standard care” for both consciousness and GOS/mortality. As discussed earlier, there was no one “standard of care” in these studies; because of this and the desperate need for interventions at the acute stage, the SC experts agreed that, if feasible and acceptable to the family and patient, it could be an option for consideration. Feasibility and acceptability will need to play a large role in the moderate-to-severe TBI populations once the methodological flaws are addressed in future studies.

ICP was assessed across 2 studies. The quality of the research and the results were mixed. The SC experts debated whether this outcome is clinically relevant to the purposes of this systematic review. No conclusion and recommendation was made.

Addressing the placebo issue

The scientific research community continues to test and debate specific and nonspecific effects of therapies in medicine on outcomes generally, and the placebo debate is no less important in the field of HBO2 for TBI. There exists an important debate of what constitutes a true sham for HBO2 therapy. A “true” sham may involve normal atmospheric pressure; all sham arms in the studies reviewed had some pressure over atmospheric for the sake of blinding. The shams utilized in HBO2 research require validation. Is it the oxygen, the pressure, or both making a difference in the actual intervention? Or are the effects specific only to the ritual being performed? If the ritual is the critical treatment element, can we enhance it safely at an affordable cost?

The placebo response is complex and is not unique to HBO2 studies in TBI. Acupuncture has been fiercely debated with mixed results from studies comparing real acupuncture to different shams. Is it the context and meaning of the intervention that is producing the effect? Is it worth investing in further research to uncover which parts of the therapeutic experience and what ritualistic components contribute to the efficacy/effectiveness of HBO2 for those suffering the consequences of TBI?

The placebo analysis confirms that clinical ritualistic components of the therapy, actual and sham, are not being reported fully in the reviewed literature. The SC experts were not able to fully understand the meaning and contextual effects imparted by the encounter itself and concluded that a set of specific criteria for the creation of uniform reporting guidelines is necessary. The acupuncture research community developed Standards of Reporting Intervention of Clinical Trials in Acupuncture (STRICTA). Similarly, guidelines for conducting placebo-controlled clinical trials (STRICT-P) could enhance our understanding of how placebo components play a role in the interventions being investigated and what constitutes the most appropriate sham controls to use in future trials. This is a clear gap that needs further investigation in medicine at large, not only related to HBO2 and mTBI. A collaboration of placebo experts is required to push this forward. By reframing the concept of “placebo effects” into these components, we can begin to measure the meaning and contextual effects in clinical trials. The placebo components used in the placebo analysis reported here can be used as a building block for such standard reporting criteria development.

Aside from the challenge with placebo effects, the definition and criteria for diagnosing mTBI and moderate-to-severe TBI is associated with a variety of symptoms and seems to be multidimensional. A holistic, patient-centered care approach should be considered to enhance patient response and improve well-being.

The following suggestions emerged:

1. Support placebo research to provide further insight on the specific and nonspecific effects of HBO2 on mTBI.
2. Investigate the effects of pressure and pressure changes to differentiate the specific pressure effects from the contextual nonspecific effects of pressure during sham treatments.
3. Explore the most validated and sensitive clinical outcome tools for mild and moderate-to-severe TBI therapy research to ensure consistent reporting and the availability of data for future study and consideration of meta-analysis.
4. Compare other types of interventions to HBO2 treatments for TBI. Even if placebo is the influential factor in mTBI therapeutic studies, direct head-to-head comparative effectiveness research and cost-benefit analyses comparing other modalities are necessary.
5. Commission a network meta-analysis to evaluate all treatments for TBI, allowing for the assessment of the relative effectiveness of treatments when they have not been compared directly but have been compared with other treatments.

CONCLUSIONS

This systematic review revealed that for patients suffering from persistent symptoms after mTBI, HBO2 therapy is no better than sham treatment, if, in fact, the “sham” treatment is a true sham, and hence not recommended as a therapy. In comparison to “standard care,” there is not enough evidence to draw conclusions at this time. However, the improvements in outcomes shown within groups for both HBO2 and sham treatment cannot be ignored. The ritual of the therapeutic encounter involved with HBO2/sham HBO2 may be of therapeutic value to TBI. It is also possible another confounding specific effect has yet to be identified. We will not be able to answer questions about specific HBO2 attributes (inspired oxygen levels, pressure, dosing duration, frequency, and length), nonspecific placebo effects, and how to maximize therapeutic benefit for patients with mTBI at the lowest possible risk and cost without further research.

For moderate-to-severe TBI at the acute phase of treatment, although the methodology appears flawed across the studies, and specific outcomes may be unstable, because of the complexity and severity of brain injury in this population, HBO2 may be of value and could benefit these patients as a relatively safe adjunctive therapy if feasible and acceptable to the individual and family. Further research outlined in this report should be considered to resolve the controversy surrounding this field, but only if methodological flaws are avoided and bias minimized.

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