Effect of oculomotor rehabilitation on vergence responsivity in mild traumatic brain injury

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Abstract—A range of dynamic and static vergence responses were evaluated in 12 individuals with mild traumatic brain injury (age: 29 +/- 3 yr) having near vision symptoms. All measures were performed in a crossover design before and after oculomotor training (OMT) and placebo (P) training. Following OMT, peak velocity for both convergence and divergence increased significantly. Increased peak velocity was significantly correlated with increased clinically based vergence prism flipper rate. Steady-state response variability for convergence reduced significantly following OMT. The maximum amplitude of convergence, relative fusional amplitudes, and near stereoacuity improved significantly. In addition, symptoms reduced significantly, and visual attention improved markedly. None of the measures were found to change significantly following P training. The significant improvement in most aspects of vergence eye movements following OMT demonstrates considerable residual brain plasticity via oculomotor learning. The improved vergence affected positively on nearwork-related symptoms and visual attention.

Key words: acquired brain injury, mild traumatic brain injury, nearwork symptoms, neuroplasticity, oculomotor dysfunction, oculomotor learning, oculomotor rehabilitation, traumatic brain injury, vergence, vergence dysfunction, visual attention.

INTRODUCTION

Traumatic brain injury (TBI) is defined as any structural damage caused by an external force to the brain and its associated structures (e.g., cranium) resulting in physiological disruption of brain function [1]. The Centers for Disease Control and Prevention estimated that approximately 1.7 million people have experienced a TBI in the United States, with it being the leading cause of death and disability [2]. TBI is a major medical, optometric, social, economic, and public health issue in the United States [3]. Motor vehicle accidents, falls, assaults, sports-related concussion, gunshot wounds, work-related injuries, etc., are some of the most common causes of TBI [1], with 70 to 80 percent of all TBI being classified as mild TBI (mTBI) [1,4].

Based on the severity and location of the injury, TBI results in a spectrum of dysfunctions involving sensory, motor, perceptual, physical, behavioral, cognitive, linguistic, and emotional aspects [5]. Being a primary modality of sensation, vision and its deficits following TBI will likely have an adverse effect on many activities of daily living (ADLs). Due to the pervasive nature of TBI (e.g.,

Abbreviations: Δ = prism diopter, ADL = activity of daily living, BI = base-in, BO = base-out, CI = convergence insufficiency, CISS = Convergence Insufficiency Symptom Survey, LED = light-emitting diode, mTBI = mild traumatic brain injury, NFV = negative fusional vergence, NPC = near point of convergence, OMT = oculomotor training, OR = oculomotor rehabilitation, P = placebo, PFV = positive fusional vergence, PRII = Power Refractor II, SEM = standard error of mean, SS = steady-state, SUNY = State University of New York, TBI = traumatic brain injury, VSAT = Visual Search and Attention Test.

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of coup-contrecoup), numerous vision-related areas can be adversely affected [3,6]. For example, approximately 90 percent of individuals with mTBI having vision-related symptoms examined in an optometric clinic setting were diagnosed with one or more oculomotor dysfunctions following their acute care phase and natural recovery period [7]. Of the sample population, 70 percent manifested nonstrabismic types of oculomotor deficiencies involving version, vergence, and accommodation. Such deficits could adversely affect reading and other nearwork ADLs [8]. Identifying these abnormalities and rehabilitating them are essential in improving reading ability and overall quality of life [8]. In this article, only the oculomotor system subcomponent of vergence is considered.

Vergence refers to the disjunctive movement of the eyes used to track objects varying in depth over the range of one’s binocular visual field [9]. The goal is to rapidly obtain and maintain fusion, or singleness, of the object of interest by placing the foveally bifixated object on corresponding retinal points within Panum’s fusional area [10]. Furthermore, the vergence system acts in synchrony and precision with the versional system to track objects laterally in one’s visual space accurately and singly, with the accommodative system continuously activated to maintain target clarity [10].

There are several separate subsystems believed to be involved in the neural control of vergence [11]. While the midbrain comprises the majority of neurons [12], evidence for the existence of neurons that also discharge during vergence have been located in the pons [13–14], cerebellum [15], and some areas of the cerebral cortex, such as the frontal eye field [16], parietal lobe [17–18], middle temporal [19] and medial superior temporal visual areas [20], and primary visual cortex [21]. Since the vergence neural pathway is extensive, any injury to the multitude of related brain and contiguous structures may adversely affect the vergence system.

REVIEW OF VERGENCE DYSFUNCTIONS IN TRAUMATIC BRAIN INJURY

Clinical Studies

Retrospective Studies

Five retrospective studies have assessed the prevalence of oculomotor abnormalities in patients with mTBI, both in a civilian clinic population and in Department of Veterans Affairs and military populations [7,22–25]. The results are remarkably similar across these populations, in which the etiology of the TBI included both blast and non-blast injuries. Vergence dysfunctions ranged from 24 to 48 percent. While convergence insufficiency (CI) was the main clinical vergence dysfunction (42.5%), other vergence deficits that were also found with a relatively high frequency included binocular instability (10.0%), convergence excess (2.5%), basic exophoria (2.5%), and divergence insufficiency (<2.0%) [7]. These general findings have been confirmed in four recent related investigations [26–29].

Clinical Studies Involving Nonstrabismic Vergence Dysfunctions

One of the earliest formal studies on the presence of binocular vision abnormalities following TBI was by Cross in 1945 [30]. Observations were made from several hundred cases examined at a military hospital. Convergence dysfunction was found to be one of the most common oculomotor anomalies. General body fatigue following TBI was presumed to be the cause of reported “ocular muscle fatigue,” thus resulting in “defective convergence” in these individuals [30–31].

A number of more recent studies conducted in clinic populations have evaluated vergence function following TBI [32–39]. These studies have also found a range of vergence dysfunctions, including CI, reduced fusional ranges, and increased near exophoria, with percentages ranging from 25 to 75 percent.

Laboratory Investigations

An early study by Ron et al. objectively recorded vergence eye movements to a constant-velocity ramp stimulus, in which 28 patients with unspecified categories of TBI binocularly tracked a small target at near that moved continuously in depth from 30 to 5 cm along their midline [40]. Abnormal dynamic vergence responses were found in 71 percent of patients.

More recently, a wide range of static and dynamic vergence parameters were tested in 21 visually symptomatic adult patients with mTBI (mean ± 1 standard error of mean [SEM] age: 45.7 ± 3.1 yr), as related to nearwork, by the State University of New York (SUNY) acquired TBI research group [41]. Five static parameters were found to be significantly different and abnormal between the mTBI and the visually nondisabled groups: near point of convergence (NPC) break and recovery values were
receded, positive fusional vergence (PFV) break and recovery values were reduced, and the stereocuity threshold was elevated (presumably related to inaccurate vergence) in the group with mTBI [10]. While the transient response amplitudes for convergence and divergence did not differ significantly between the nondisabled subjects and those with mTBI, because they were already normal at baseline, all of the dynamic parameters (i.e., peak velocity, time constant, and latency) were significantly different ($p < 0.05$) between the two groups for both convergence and divergence. Responses were all slowed, delayed, and more variable in the group with mTBI than the nondisabled group.

Lastly, in a recent pilot study, objective recordings of vergence were obtained in two young adults with self-reported mTBI and nearwork symptoms [42]. Vergence dynamics were markedly slowed (i.e., reduced peak velocity) for convergence but not for divergence, as has been found earlier in larger populations [42].

OVERVIEW OF OCULOMOTOR REHABILITATION IN TRAUMATIC BRAIN INJURY

Several clinical case studies and a few population studies have evaluated the effect of oculomotor rehabilitation (OR) in mTBI. One of the earliest studies involved with the treatment of vergence and accommodative disorders was conducted by Chandler in a hospital-based setting in a series of World War II-related TBI cases ($n = 33$) [43]. OR (unspecified, but presumably “orthoptic” fusional training) commenced anywhere from 3 wk to 5 yr postinjury. While 73 percent of the patients treated were either fully remediated or markedly improved, 12 percent failed to improve and only 6 percent exhibited spontaneous recovery (from 3 d to 2 wk postinjury). These results are consistent with later studies [44–48]. Evidence to support the fact that carefully programmed OR remedies binocular vision anomalies in those with mTBI also comes from several clinical population studies [38–39,49]. In each study, reading difficulty was one of the most common symptoms.

From these studies, there is abundant evidence from the literature in both laboratory-based and clinically based studies supporting the notion that targeted, specific, programmed OR procedures, which all incorporate the principle of motor learning [50–51], can remediate patients with a range of binocular vision disorders as a consequence of mTBI. Symptoms were ameliorated concurrent with improvement or normalization of clinical signs. This is important information because improved oculomotor abilities and related visual-perceptual skills can hasten progress in the patient’s other rehabilitative programs [52–53]. This includes cognitive therapy, for example, which requires complex visual saccadic scanning and fine detail discrimination.

The purpose of the current investigation was to evaluate comprehensively clinically and laboratory-based vergence parameters in individuals with mTBI reporting nearwork-related symptoms of an oculomotor nature before and after oculomotor training (OMT) performed in the clinic, without a home-based component. The OMT involved all three main oculomotor subsystems: vergence, accommodation, and version. The measures were also compared after placebo (P) training. For the purpose of the present article, only the oculomotor subsystem of vergence is considered.

METHODS

Subjects

Twelve subjects (8 females and 4 males) between the ages of 23 and 33 yr (mean ± standard deviation: 29 ± 3 yr) participated in the study (see Appendix 1 [available online only] for demographics). The training effects for the study were hypothesized to be moderate to large based on our earlier related laboratory studies [8,54–55], as well as the extensive clinical experience of the second author (K.J.C.). Sample size was calculated using a power analysis program (G*Power 3; Institut für Experimentelle Psychologie, Heinrich Heine Universität Düsseldorf, Düsseldorf, Germany) at an alpha level of 0.05, with a power set at 0.80 using two key parameters of vergence (i.e., NPC and peak velocity). All subjects had documented mTBI, with injury onset of greater than 1 yr (1–10 yr postinjury) to avoid possible contamination from the natural recovery process [56]. All manifested several nearwork-related symptoms (e.g., intermittent diplopia) and at least one clinical sign of vergence dysfunction (e.g., receded NPC) of an oculomotor nature. All had stable general health. None had a significant cognitive dysfunction. Subjects were identified by their university-based healthcare provider and were recruited from the Raymond J. Greenwald Vision Rehabilitation
Center at the SUNY College of Optometry University Optometric Center of New York. Subjects received a comprehensive vision examination prior to participating in the experiment, which included a detailed refractive, oculomotor, and ocular health assessment.

**Study Design**

We conducted a crossover, interventional experimental design of a single-blind nature (for the subject) (Figure 1). In this design, each subject acted as his or her own control, thus negating undesirable intersubject variability. Each subject received both OMT (treatment A) and P training (treatment B). During phase 1, every odd-numbered subject first received treatment A, every even-numbered subject first received treatment B, and vice versa during phase 2. This interventional study had a duration of 15 wk. It consisted of 12 wk of the two treatment phases, 6 wk each phase separated by 1 wk, for a total of 9 h of OMT and 9 h of P training. In addition, there was a 3 wk measurement period: 1 wk before phase 1 treatment, 1 wk after phase 1 treatment, and 1 wk after phase 2 treatment. During these training and measurement periods, subjects did not perform any other OR to avoid contamination of test results.

The study consisted of the following phases:

1. **Week 1**: Initial baseline measures—all evaluative procedures (described later) were recorded over two separate test sessions (each session lasting for up to 1.5 h, including rest periods to prevent fatigue), each separated by at least 2 d.
2. **Weeks 2–7**: Phase 1 treatment—6 wk of either OMT or P training. Subjects received two training sessions per week. Each session was 60 min in duration, involving 45 min of actual training with the remainder of time consisting of short and interspersed rest periods for the subject. Total training time was 9 h over the 6 wk.
3. **Week 8**: Repeat baseline measures—same as step 1.
4. **Weeks 9–14**: Phase 2 treatment—same as step 2.
5. **Week 15**: Repeat baseline measures—same as step 1.

**Evaluative Procedures**

Several general areas of testing were performed; these included clinical and laboratory vergence measures, as well as visual attention and near-vision symptoms. All clinical measures were assessed using standardized clinical techniques [57]. All laboratory-based objective measures were performed using commercially available instrumentation with well-established test protocols developed in our laboratory for version [54], accommodation [58], and vergence [41]. All measures were noninvasive and were recorded with the subject’s habitual distance correction in place. Order of testing was randomized over the 2 d of measurements.

1. **Clinical measures**: Selected binocular vision-related parameters were tested with randomization under standard clinical room illumination (80 Lux). They included NPC break and recovery, horizontal near phoria using the von Graefe prism dissociation method, horizontal near PFV and negative fusional vergence (NFV) ranges, vergence prism facility (with 12 prism diopter [Δ] base-out [BO] and 3Δ base-in [BI] prism flippers), and stereoacuity using the Titmus stereo test.
2. **Laboratory-based objective measures**: Vergence dynamics to symmetric step vergence stimuli was recorded using the Power Refractor II (PRII) (Plusoptix Inc; Atlanta, Georgia) based on the principle of infrared videography and dynamic retinoscopy, with a sampling rate of 12.5 Hz (resolution: <0.9°) for binocular recording, as described elsewhere [41]. This sampling rate exceeds the Nyquist criterion [59]. Targets comprised the contiguous red and green fixation light-emitting diodes (LEDs) (angular size: 0.28°) located on the measuring head of the PRII at 1 m and a white LED (angular size: 0.86°) placed at 0.3 m, both aligned along the midline. Mean response amplitude, peak velocity, time constant, and steady-state (SS) response variability were calculated separately for both convergence and divergence.
3. **Subjective visual attention test**: A subjective correlate of global visual attention was assessed using the Visual
Search and Attention Test (VSAT). It involves a search (for a letter or a symbol) and cancellation (cross-out) task that was developed by Trencerry et al. [60]. The VSAT was performed binocularly at the subject’s habitual nearwork distance. Percentile scores were calculated from the age-matched normative table for the two test sheets.

4. Symptom scale: Individual symptoms related to nearwork were rated by the subjects using the Convergence Insufficiency Symptom Survey (CISS) [61]. It is composed of a 15-item questionnaire probing reading-related symptoms (e.g., intermittent diplopia). The severity of symptoms is scaled from 0 to 4, i.e., from least symptomatic to most symptomatic. The total score was compared before and after each of the two training phases. A reduction in overall score of $\geq 10$ was considered to reflect a significant reduction of symptoms. A score of $\leq 16$ was considered to represent being relatively asymptomatic.

**Treatment Protocol**

*Treatment A: Oculomotor Training*

OMT was performed along the midline at 0.4 m, two sessions per week, for a total of 6 wk. OMT was performed with constant verbal and visual feedback, motivation, and repetition and involved active participation of the subject to maximize attention. At a session, each oculomotor component (version, vergence, and accommodation) was trained for 15 min, interspersed with 5 min rest periods. Each session lasted for 1 h, with 45 min of training and 15 min of rest periods, for a total of 9 h of training over the 6 wk OMT phase, 3 h for each oculomotor subsystem. For the purpose of the present article, however, only the vergence training and related results are discussed (Table 1).

For step vergence amplitude training, BO and BI prisms were used. The basic principle behind the training was to maintain the accommodative demand constant at 0.4 m (2.5 D), while increasing the vergence demand [57,62]. The fusional targets were composed of pictures, symbols, numbers, letters, tumbling E, and colors displayed on a computer screen at 0.4 m. As treatment progressed and the subject demonstrated improvement, task difficulty was increased by reducing target size (from 10° to 2°) and manually increasing the vergence demand prismatically. The total amount of prism depended on the subject’s task performance level. After introducing each BO prism, subjects were instructed to fuse the target as rapidly as possible. This trained the fast vergence mechanism [63].

<table>
<thead>
<tr>
<th>Stimulus Parameter</th>
<th>Duration (min)</th>
<th>Total Duration (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disparity Step Amplitude</td>
<td>7</td>
<td>15</td>
</tr>
<tr>
<td>Step Facility (BO/BI)</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>Ramp</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>No Disparity Placebo-Step</td>
<td>10</td>
<td>15</td>
</tr>
<tr>
<td>Placebo-Ramp</td>
<td>5</td>
<td></td>
</tr>
</tbody>
</table>

*BI = base-in, BO = base-out.*

The fused percept was then maintained for 15 to 20 s. This sustained viewing trained the slow vergence mechanism that maintained the vergence response [63]. Such response maintenance reflects the vergence adaptation mechanism [63–64]. BO training was terminated at the point at which subjects could no longer fuse (and/or focus) with their maximum effort. This was repeated for BI prisms, which stimulated relative divergence. The order of BO and BI training at each session was randomized.

For step vergence facility training [62], combinations of BO and BI prism flippers ($\Delta BO/\Delta BI$, $6\Delta BO/2\Delta BI$, $9\Delta BO/3\Delta BI$, and $12\Delta BO/3\Delta BI$) were used while maintaining accommodation constant at 0.4 m (2.5 D). The fusional targets were similar to those used in the previously mentioned amplitude training. Based on the subject’s initial ability to fuse, the magnitude of prism flipper was chosen. Subjects bifixated targets on a computer screen, and they were instructed to fuse and focus as rapidly as possible and to achieve the maximum number of cycles possible. As the treatment progressed and the subject demonstrated improvement, task difficulty was increased by increasing the prism flipper magnitude and by reducing target size (from 10° to 2°).

For ramp vergence training, subjects binocularly tracked an isolated, high-contrast (>90%), Snellen 20/30 letter controlled by an XY plotter and function generator moving continuously over a range of 0.5 to 0.2 m at the rate of 0.1 to 1.0 Hz. Task difficulty was increased by tracking at closer distances, with the combination of increased speed. Subjects were instructed to maintain the target clear and single.

*Treatment B: Placebo Training*

P training was performed as described previously for OMT. P training did not involve any disparity stimulation,
because this is the primary drive for the vergence system [10]. For the P-step, binocular and monocular plano-pow-
ered loose prism and prism flippers and/or monocular ver-
tical prism (0.5 or 1 ΔD) flippers were used as the P
training analog to OMT. Training was performed both
monocularly (5 min) and binocularly (5 min). For the P-
ramp, subjects tracked a difference of Gaussian (0.2
cycles/°) target through a 0.5 mm pinhole monocularly for
5 min (2.5 min each eye) in an otherwise darkened room,
which did not provide any disparity (or blur or accommo-
dative vergence) drive [10].

Data Acquisition and Analyses for Objective Recordings

The recorded video files were saved to the PRII hard
drive and converted into .txt files. They were then trans-
ferred into Excel (Microsoft; Redmond, Washington) for
detailed analysis. Three artifact-free (free of blinks and/or
saccades) convergence and three divergence responses
were selected for analysis from the right eye position
traces for each subject from a sample of seven to eight
responses in each direction. The middle three responses
were used for analysis, and the initial and final responses
were omitted to avoid possible learning and fatigue
effects, respectively [41]. An exponential decay function
was fit to the traces, and the response amplitudes and time
constants were obtained using GraphPad Prism (Graph-
Pad Software Inc; La Jolla, California). The peak veloc-
ities were derived from first-order differentiation of the
exponential equation. The SS response variability was cal-
culated from the standard deviation of the measured time
window (~5 s) of response after SS was attained. The
goodness of fit was assessed from the $r^2$ values of each fitted
response. The mean $r^2$ value for both increasing and
decreasing steps was always >0.8 for each subject, thus
demonstrating a good fit. The mean amplitude, time con-
stant, and peak velocity of the responses at baseline, post-
OMT, and post-P training were compared statistically
using GraphPad Prism at $p \leq 0.05$.

STATISTICAL ANALYSES

Combined Group

The key objective of the study was to evaluate the
effect of OMT in individuals with mTBI and oculomotor-
based near-vision symptoms. Therefore, the main analy-
ses included a comparison of baseline measures before
and after OMT using paired two-tailed $t$-tests. Data from
all 12 subjects were analyzed and presented as the “com-
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abnormal. All four abnormal parameters (100%) improved significantly following OMT. There was a significant decrease (i.e., improvement) in both the NPC break ($t(11) = 4.07; p = 0.01$) and NPC recovery ($t(11) = 3.64; p = 0.01$) after OMT, but they did not normalize (Figure 4(a)). In addition, this increase in maximum vergence amplitude (NPC break) was significantly correlated ($p < 0.05$) with reduction in symptoms ($r = 0.57$), as well as with improved visual attention ($r = 0.40$). Both the PFV break ($t(11) = 2.80; p = 0.01$) and PFV recovery ($t(11) = 4.71; p = 0.01$) values significantly increased with OMT. Prism vergence facility ($t(11) = 4.22; p = 0.01$) (Figure 4(b)) and stereoacuity ($t(11) = 2.34; p = 0.03$) also improved significantly following OMT. The NFV break increased significantly ($t(11) = 3.40; p = 0.01$) and normalized, while the NFV recovery exhibited a predicted trend ($t(11) = 2.04; p = 0.06$). There was no significant change in the horizontal near phoria value ($t(11) = 0.49; p = 0.62$), which ranged from 14 exophoria to 1 esophoria in the group. See Table 3 for the group mean ($±1$ SEM) values at baseline and post-OMT.

There was no statistically significant effect ($p > 0.05$) of the P training on any of the vergence parameters tested. See Appendix 2 (available online only) for details.

Other Subjective Tests
The CISS total score significantly reduced ($t(11) = 3.69; p = 0.01$) from a mean value of $37 ± 4$ to $28 ± 3$ following OMT. This quantitatively indicated a reduction in nearvision-related symptoms following OMT.
With respect to visual attention at baseline, the group mean VSAT percentiles increased significantly ($t (11) = 4.43; \ p = 0.01$) from the 32nd ($\pm 9$) to the 50th ($\pm 10$) percentile following OMT, with increases in 10 of the 12 (80%) subjects. This indicated quantitatively increased visual attentional aspects concurrent with OMT.

**DISCUSSION**

The primary aim of the present investigation was to evaluate a range of objective laboratory and subjective clinical measures of vergence before and after vergence-based OMT in individuals who reported nearwork-related

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**Table 2.**

Mean ±1 standard error of mean laboratory-based objective parameters of symmetric vergence before (baseline) and after oculomotor training (post-OMT).

<table>
<thead>
<tr>
<th>Dynamic Parameter</th>
<th>Baseline</th>
<th>Post-OMT</th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>C: Peak Velocity (°/s)</td>
<td>13.0 ± 1.9</td>
<td>18.0 ± 0.9</td>
<td>0.01*</td>
</tr>
<tr>
<td>D: Peak Velocity (°/s)</td>
<td>11.6 ± 1.1</td>
<td>13.5 ± 0.8</td>
<td>&lt;0.01*</td>
</tr>
<tr>
<td>C: Time Constant (ms)</td>
<td>399 ± 68</td>
<td>228 ± 14</td>
<td>0.01*</td>
</tr>
<tr>
<td>D: Time Constant (ms)</td>
<td>378 ± 35</td>
<td>312 ± 22</td>
<td>&lt;0.01*</td>
</tr>
<tr>
<td>C: SS Variability (°)</td>
<td>0.90 ± 0.07</td>
<td>0.75 ± 0.04</td>
<td>0.04*</td>
</tr>
<tr>
<td>D: SS Variability (°)</td>
<td>0.81 ± 0.05</td>
<td>0.78 ± 0.02</td>
<td>0.54</td>
</tr>
<tr>
<td>C: Response Amplitude (°)</td>
<td>3.93 ± 0.07†</td>
<td>3.96 ± 0.08</td>
<td>0.43</td>
</tr>
<tr>
<td>D: Response Amplitude (°)</td>
<td>3.93 ± 0.06†</td>
<td>3.93 ± 0.08</td>
<td>1.00</td>
</tr>
</tbody>
</table>

*Statistically significant.
†Already normal at baseline.

C = convergence, D = divergence, SS = steady-state.
symptoms of an oculomotor nature following their mTBI. With only 3 h of total vergence training distributed over 6 wk, significant \((p < 0.05)\) improvements were found in the vast majority (>80%) of the key laboratory and clinical aspects of vergence that were abnormal at baseline. The results were also compared with an equal dosage and distribution of P training. None of the vergence measures were found to have a significant group effect from the P training \((p > 0.05)\).

Although most of the initially abnormal parameters significantly improved with OMT, many did not normalize. This may suggest that the OMT should be increased, perhaps twofold or greater, to obtain a yet more robust result, assuming that the underlying neurology is sufficiently intact to yield a normalization. This critical area needs to be explored in the future. Lastly, as discussed for dynamic OMT aspects, the question remains whether additional hours of training would yield normalization of all static vergence parameters. Future studies in this critical area are needed.

**Training Effect on Vergence Dynamics**

At baseline, both convergence and divergence eye movements consistently demonstrated slowed dynamic trajectories in all subjects. This was evident from the reduced peak velocity along with the correlated increased time constant values. The group mean peak velocity in the population with mTBI was reduced by ~45 percent for convergence and ~25 percent for divergence [65–66]. The slowed but accurate responses suggest the presence of normal visual feedback with respect to disparity detection and processing. This is consistent with both laboratory and modeling findings, suggesting the dual-mode control of vergence [67]. That is, the initial response component (i.e., the first 200 ms) is preprogrammed (i.e., open-loop response) for the estimated amplitude of the step disparity input, followed by completion of the movement over the next several hundred milliseconds via visual feedback control (i.e., closed-loop response), with the overall response being completed in approximately 800 to 1,000 ms. These findings suggest that the slowed but accurate responses were primarily the result of improvement in the pulse subcomponent of the neural signal and not caused by its step subcomponent.

Following OMT, there was a significant increase in peak velocity by ~40 percent for convergence and ~15 percent for divergence from their mean baseline value. Concomitantly, the time constant for both convergence and divergence exhibited correlated and proportional decreases, as expected due to their inverse interrelation.

The prism flipper facility rate is the clinical analog for the overall laboratory-based vergence response incorporating and combining all dynamic parameters (i.e., peak velocity, time constant, and latency) into a global, validated metric [62]. Thus, peak velocity and prism flipper rate were found to correlate significantly with each other both before and after the OMT. At baseline, the mean vergence facility rate was ~65 percent less than the mean clinic norm [62]. With OMT, subjects could now fuse both the BO and BI prisms rapidly, with a large and significant twofold increase in facility rate, but it did not normalize.
In the present study, the SS response was assessed for ~5 s. Within this measured window of time, the SS variability for convergence decreased significantly following OMT as assessed at 30 cm. This suggests improved convergence sustainability involving the slow vergence mechanism [63]. Our previous study in this area [41] found abnormal, reduced vergence adaptation in those with mTBI, which is also typically found in those without mTBI but with vergence-related dysfunction and correlated symptoms [68]. In contrast, the SS variability did not change markedly for divergence at the 1-m test distance. This finding may not be surprising given the fact that the OMT was performed at the conventional near reading distance of 40 cm. This suggests lack of generalization of the rehabilitation effects (i.e., oculomotor learning) to the overall vergence system. In the future, vergence training should also be conducted at different distances and gaze directions to attain a more generalizable improvement in vergence responsivity.

**Training Effect on Static Measures of Vergence**

The NPC is the main static diagnostic measure used in the clinic for assessment of vergence dysfunctions [62]. At baseline, the subjects with mTBI demonstrated markedly receded NPC components (break and recovery), thus suggesting poor maximal convergence amplitude fusional ability. Following OMT, the NPC amplitude and recovery improved significantly but did not normalize.

The relative vergence amplitude increased in both the convergent and divergent directions following OMT. This was evident from the increased PFV and NFV break values. While the PFV recovery value significantly improved following OMT, it did not for the NFV recovery value. The relative vergence system has several response nonlinearities (e.g., amplitude, dynamics) between PFV and NFV [62], and this may reflect one such difference [66]. This is consistent with the fact that training relative convergence (PFV) is easier than relative divergence (NFV) [69]. It is also consistent with neurophysiological evidence demonstrating more convergence-related cells present than divergence-related cells [15].

The overall improvement in convergence ability was also reflected in the improved near stereoacuity with OMT. Presumably, this is caused by improvement in vergence response accuracy and stability, which would reduce the mean fixation disparity vergence error at near, hence improving stereoacuity [10]. This is consistent with the recent finding that increased fixation disparity was significantly correlated with reduced stereoacuity at near in individuals with mTBI [41].

However, OMT did not seem to have an effect on the near horizontal phoria. In the nonadapted state of vergence, this value reflects the horizontal position of eyes in the absence of fusional vergence [70]. This value would be expected to change only if the cross-link ratio (response accommodative convergence/accommodation) changed [71]. However, past studies have reported constancy of this cross-link following vergence training in both visually nondisabled subjects [64,72] and in symptomatic individuals manifesting binocular vision dysfunction [73].

From the present findings, as well as from the previous studies that assessed objective and clinical measures of...
vergence, it appears that the laboratory-based peak velocity and clinically based prism flipper facility, along with NPC, are key diagnostic measures in the population with mTBI. This is consistent with recent suggestions in the literature based on the SUNY research group findings [41,74].

Neurophysiological Implications

Although several areas of the brain have been identified in the control of vergence, the midbrain houses the majority of vergence-related neurons [11]. The motoneuronal controller of vergence has been found to be somewhat similar to saccades, because the final neural signal consists of a small and broad pulse combined with a step [75–76]. The pulse signal, which is produced by the midbrain vergence “burst cells” that fire in relation to vergence velocity, is responsible for rapidly displacing the eyes in a time-optimal manner to a new binocularly fixated target position. In contrast, the step signal, which is produced by the vergence tonic cells that fire in relation to vergence angle, maintains the SS eye position (i.e., vergence angle) on the binocularly fixated target accurately [77]. A neural integrator (i.e., nucleus reticularis tegmenti pontis) [78–79] has been proposed to process the velocity signal to a step signal. Then, the combined signal is sent via the oculomotor neurons to innervate the extraocular muscles to make an appropriate vergence eye movement [11].

Based on the results of the present study at baseline, and earlier studies [41–42], the primary neural deficit in the patient with mTBI is believed to be the pulse. This is reflected in the consistently slowed dynamics (e.g., reduced peak velocity and increased time constant) for both convergence and divergence in the present study prior to OMT, which can be accounted for by a reduction in pulse height. Thus, the overall time course of the vergence dynamic trajectory was slowed. Since the appropriate vergence amplitude was eventually attained both before and after OMT, this suggests that the step component had the appropriate mean height. However, the subsequent vergence SS level exhibited increased variability, which suggests that the presence of increased neural noise could produce a more variable step signal. Tonic vergence cells constantly fire to maintain this SS level. The increased SS variability could reflect a higher degree of variability in the neural firing of such cells. Following OMT, the increased peak velocity can be attributed to an increase in pulse height (presumably because of the increased firing rate), thus resulting in faster motor responsiveness to attain the final SS position. Following OMT, the reduced SS variability during convergence could be attributed to reduced step gain variability as a result of normalization of tonic cells firing.

Mechanisms of Neuroplasticity and Oculomotor Learning

Under normal circumstances, repeated synaptic stimulation, along with its coincident activation, results in an increased synaptic strength and memory storage [80–82]. This experience-dependent neuroplasticity is composed of biochemical-, cellular-, physiological-, and structural-level changes [83]. Recovery following an insult to the brain has been categorized as “spontaneous reorganization” (or natural recovery) and “training-induced recovery” [56]. The former occurs immediately following injury. It is believed to involve restoration of neurotransmission in the adjacent spared area and regions distant from the injury location. This natural recovery period following TBI occurs over the first 6 to 9 mo [84]. However, training-induced recovery appears to be relatively independent of the amount of time elapsed after the injury. Significant oculomotor improvements can occur even 5 to 10 yr after the first injury [8]. This involves functional recovery via a “relearning” process. Remapping and reconfiguration of neural circuits both within and across relevant regions play a significant role in the recovery process [83].

Following TBI, the decreased vergence response peak velocity may be attributed to diffuse axonal injury. The compromised white matter integrity causes slowed conduction of nerve impulses [85–86], thus resulting in an overall slowed response (e.g., slowed vergence). In addition, the decreased number of synapses, reduced firing rate, reduced neural synchrony, and lack of correlation within and across the specific brain regions cause loss of automaticity and an overall reduction in the system’s maximum amplitude (e.g., NPC) [42]. Figure 5 shows the schematic representation of the proposed neurological mechanisms involved based on the aforementioned laboratory findings.

OMT acts as a relearning process, in which the system being trained or conditioned regains its automaticity through repetition, which then becomes preprogrammed with much practice. In the present study, an overall improvement in oculomotor behavior was observed in all individuals with mTBI. It is believed to be a consequence of “oculomotor learning” involving the relearning processes described earlier [83]. Figure 6 shows the schematic representation of the proposed mechanisms of OR based on
Figure 5.
Proposed neural mechanisms of traumatic brain injury causing vergence dysfunction. WM = white matter.

Figure 6.
Proposed underlying mechanisms of vergence-based oculomotor rehabilitation (OR). BI = base-in, BO = base-out.
the aforementioned laboratory findings. A combination of repeated stimulation with various amounts and types of disparity (crossed and uncrossed), increasing task-level difficulty, active participation of the subjects, increased attention, presence of visual and verbal feedback, and high motivation of the subjects to perform the task over the 6 wk training period resulted in a significant OMT effect. These ideas are further supported by a recent study [42] that evaluated the neurological changes using the functional magnetic resonance imaging technique in two individuals with mTBI before and after intensive vergence-based OMT. Their results showed increased amount of voxels and correlation within specific regions of interest (brain stem, cerebellum, frontal eye fields, and supplementary eye fields) following a total of 18 h of clinically based and laboratory-based vergence OR, similar in nature to that conducted in the present study. Their results also correlated with increased vergence peak velocity, as found in the present study. The increased convergence peak velocity was found to correlate with an increase in amount of active voxels and correlation within the brain stem, cerebellum, and frontal lobe regions. While the NPC was correlated with the brain stem activity, the PFV amplitude was correlated with frontal, parietal, and cerebellar regions. Increased cortical activity was suggested to be due to “neural recruitment” in the previously specified regions, and the correlation was attributed to “improved synchronization” of the involved subsystems vergence neurons [42]. These findings provide further direct neurological support for the proposed neurological process involved in our OMT (Figures 5–6).

CONCLUSIONS

Vergence-based OR was effective in individuals with mTBI who reported nearwork-related symptoms. Overall improvement in nearly all of the critical, abnormal measures of vergence was observed both objectively and clinically. Improved vergence motor control was attributed to residual neural visual system plasticity and oculomotor learning effects in these individuals. Concurrently, nearwork-related symptoms reduced, and visual attention improved.

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