First- and second-order contrast sensitivity functions reveal disrupted visual processing following mild traumatic brain injury

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ABSTRACT

Vision is disrupted by traumatic brain injury (TBI), with vision-related complaints being amongst the most common in this population. Based on the neural responses of early visual cortical areas, injury to the visual cortex would be predicted to affect both 1st order and 2nd order contrast sensitivity functions (CSFs)—the height and/or the cut-off of the CSF are expected to be affected by TBI. Previous studies have reported disruptions only in 2nd order contrast sensitivity, but using a narrow range of parameters and divergent methodologies—no study has characterized the effect of TBI on the full CSF for both 1st and 2nd order stimuli. Such information is needed to properly understand the effect of TBI on contrast perception, which underlies all visual processing. Using a unified framework based on the quick contrast sensitivity function, we measured full CSFs for static and dynamic 1st and 2nd order stimuli. Our results provide a unique dataset showing alterations in sensitivity for both 1st and 2nd order visual stimuli. In particular, we show that TBI patients have increased sensitivity for 1st order motion stimuli and decreased sensitivity to orientation-defined and contrast-defined 2nd order stimuli. In addition, our data suggest that TBI patients’ sensitivity for both 1st order stimuli and 2nd order contrast-defined stimuli is shifted towards higher spatial frequencies.

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1. Introduction

Traumatic brain injury (TBI) is one of the most common causes for disability amongst the North American population affecting approximately 3.2–5.3 million people (Corrigan et al., 2010; Merigan, Nealey, & Maunsell, 1993; Schiller, 1993). For example, a lesion to the macaque visual area V2 resulted in a mild 1st order contrast sensitivity loss within the lesioned cortical region whereas perception of orientation-defined 2nd order stimuli was severely impaired (Merigan et al., 1993). Chemical lesions to macaque monkey V4 resulted in deficits in both 1st order contrast sensitivity and 2nd order contour discrimination and these findings were in notable agreement with human data from stroke patients with lesions in corresponding cortical area of vision—patients who are unaware of symptoms may nonetheless suffer from sub-clinical disruptions to visual performance.

While total loss of the primary visual cortex (V1) results in effective blindness (blindsight) (Cowey, 2010; Stoerig & Cowey, 1997), injury to the rest of the visual cortex results in contrast sensitivity loss for both 1st and 2nd order stimuli—stimuli that vary in a dimension other than luminance such as texture, motion and contrast, thought to involve extra-striate cortical regions (El-Shamayleh & Movshon, 2011; Larsson, Heeger, & Landy, 2010; Merigan, 2000). First-order or luminance modulation losses are smaller in magnitude than the 2nd order losses, suggesting that the extra-striate cortex may be specifically involved (Hayes & Merigan, 2006; Merigan, Nealey, & Maunsell, 1993; Schiller, 1993). For example, a lesion to the macaque visual area V2 resulted in a mild 1st order contrast sensitivity loss within the lesioned cortical region whereas perception of orientation-defined 2nd order stimuli was severely impaired (Merigan et al., 1993). Chemical lesions to macaque monkey V4 resulted in deficits in both 1st order contrast sensitivity and 2nd order contour discrimination and these findings were in notable agreement with human data from stroke patients with lesions in corresponding cortical area.
(Hayes & Merigan, 2006). Thus, the processing of 1st and 2nd order stimuli (non-luminance modulation) can be affected in TBI, suggesting that the putative diffuse injury involves both extra-striate as well as striate processing.

Describing a deficit in terms of 1st and 2nd order processing is challenging for two reasons. For example, contrast perception for 1st order stimuli might be affected by whether the stimulus is static or moving. Second-order stimuli can be defined in a number of ways, e.g., being defined solely by contrast variation, texture variation, or dynamic variations over space. Independent of the stimulus type, it is imperative that a range of stimulus parameters be tested so as to not obtain biased estimates of group differences—for instance, TBI and normal subjects may have a difference in performance at only high or only medium spatial frequencies. This information is important to identify the affected mechanisms as well as the potential means of treatment. Critically, 2nd order stimuli all have equi-detectable carriers (i.e. all carriers were set to a contrast factor above threshold). We do this to ensure that any 2nd order loss in sensitivity is not simply a consequence of a less detectable carrier (i.e. a first order loss).

Previous findings with fixed stimulus parameters suggest that sensitivity, particularly for 2nd order contrast modulated stimuli, can be affected by TBI. While sensitivity to a 1st order low spatial frequency luminance grating was not affected, sensitivity to both static and dynamic contrast-defined 2nd order stimuli at the same spatial frequency was lower in children who suffered a mild TBI (Brosseau-Lachaine, Gagnon, Forget, & Faubert, 2008). Another study showed that reaction times on a motion direction discrimination task were longer in mild TBI participants for both 1st and 2nd order stimuli using parameters comparable to a previous study. However, unlike in the control group, the reaction times for 2nd order stimuli were longer compared to 1st order stimuli in the TBI group (Piponnier et al., 2015).

Electrophysiological results appear to corroborate the psychophysical findings. Lachapelle, Ouimet, Bach, Pitto, and McKerrall (2004) recorded visual evoked potentials (VEPs) to 1st and 2nd order visual stimuli and assessed the delays as well as the amplitudes of the low- and high-level VEP components. While the amplitudes did not significantly differ between the two groups in either condition—albeit on average being diminished in the TBI group—the delay was significantly longer for motion- and texture-defined 2nd order stimuli. A later study by the same group showed a prolonged event-related potential latency to motion-defined texture (2nd order) but not simple (1st order) motion or pattern reversal (Lachapelle, Bolduc-Teasdale, Pitto, & McKerrall, 2008).

A particular challenge in interpreting previous findings is that the spatial frequencies tested are often limited, for example some studies used only low spatial frequency (0.5 cpd) for both 1st and 2nd order stimuli (Brosseau-Lachaine et al., 2008; Piponnier et al., 2015). In addition, the carriers contrast of the 2nd order stimuli were fixed at a constant contrast (usually 50% or 100%) and were not scaled by the 1st order sensitivity of each participant (Brosseau-Lachaine et al., 2008; Lachapelle et al., 2008; Piponnier et al., 2015). We have addressed these issues by estimating the full contrast sensitivity function (CSF) for both static and dynamic 1st and 2nd order stimuli. Our approach—utilizing the quick contrast sensitivity method (qCSF; (Lesmes, Lu, Baek, & Albright, 2010; Reynaud, Tang, Zhou, & Hess, 2014))—allowed us to match the 2nd order stimulus presentation parameters to their 1st order detectability across the spatial frequency range, allowing us to accurately measure alterations in 2nd order contrast perception that are independent of any 1st order performance deficit. We also measured the 2nd order sensitivity for three fundamentally different types of stimuli—stimuli defined by contrast, orientation, or motion. Using this unified approach, we observed changes to both 1st order and 2nd order visual perception, with particular differences relating to dynamic vs. static stimuli.

2. Methods

2.1. Participants

A group of 26 mild TBI participants (17 females, 9 males, mean age 34.69 years ± 14.7 SD) was recruited either from the McGill University Health Center Out-Patient TBI Program or via public advertisements. The criteria of mild TBI were as follows: (1) any amnesia of events immediately before or after the accident lasting no longer than 24 h and (2) a Glasgow Coma Score ranging between 13 and 15. If loss of consciousness was present, it had to be shorter than 30 min. Mild TBI could be sub-classified as trivial, simple or complex (presence of a positive acute intracerebral bleeding in CT scan). The time between the TBI and the testing session varied between 35 days and 96 months. All participants had normal or corrected-to-normal visual acuity and wore their habitual refractive correction during the experiment. All procedures were in accordance with the Code of Ethics of the World Medical Association (Declaration of Helsinki) and were approved by the Research Ethics Board of the McGill University Health Centre. Informed consent was obtained from all participants prior to data collection. A short verbal screening for relevant medical history e.g. visual and psychiatric disorders, recurrent migraines, or vertigo was administered prior to participation. The exclusion criteria were: general anesthesia within the past six months, other acquired brain injuries in the past, severe tremors, and/or epilepsy. All participants successfully completed a quick neuropsychological screening of visual attention—the Trail Making Test A (Giovagnoli et al., 1996), the Bells Test (Gauthier, Dehaut, & Joanette, 1989)—and spatial neglect—the Clock-drawing test (Ishiai, Sugishita, Ichikawa, Gono, & Watabiki, 1993) (see Table 1).

2.2. Subjective visual complaints

In order to evaluate how the TBI affected vision of our group of participants we used a modified version of the questionnaire included in the Defense Centers of Excellence guidelines for assessment of visual dysfunction associated with mTBI (Defense Centers of Excellence for Psychological Health & Traumatic Brain Injury, 2013). The questionnaire is included in Table 2. In brief, the questionnaire probes for common complaints after concussion, including blurred vision, reading difficulties, discomfort during use of computer screens, etc. Twenty two participants completed the questionnaire, and were asked to rank their responses on a scale from 1 to 10 where 1 = “not at all” and 10 = “totally”. There were 11 ranked questions therefore the minimum total score was 11 and the maximum total score was 110.

2.3. Stimuli and experimental procedure

The stimulus generation procedures have been previously described in detail (Gao et al., 2014; Reynaud et al., 2014). The 1st order orientation-defined stimuli were created by filtering a white noise with horizontally- or vertically-oriented Gabor filters with a half-response spatial frequency bandwidth of 1.84 octaves, resulting in horizontally- or vertically-oriented patterns (Fig. 1B). The motion-defined stimuli were created by filtering the white noise by both orthogonal filters and were drifted either along the horizontal or vertical directions at a temporal frequency of 2 Hz. The 2nd order stimuli are best described in terms of a carrier (high-frequency texture) and an envelope (lower-frequency constraint on the carrier contrast variations over space). Thus, the
measurable spatial frequency for 2nd order is necessarily lower than the 1st order stimuli—the 2nd order envelope must include 1st order modulations, hence it must be larger. The 1st order stimuli were used as carriers for the 2nd order stimuli that were defined by orientation, motion, and contrast (Fig. 1B). The carrier-to-envelope modulation sensitivity function (Supplementary Fig. 1) was not established for first-order (Lesmes et al., 2010) and second order (Gao et al., 2014; Reynaud et al., 2014) sensitivities.

Participants performed a two-alternative forced choice (2AFCh) task of identifying the orientation of the grating in the 1st order orientation condition, orientation of motion (horizontal vs. vertical) in the 1st order motion task, and the orientation of the envelope in all 2nd order conditions (Fig. 1B). The order of conditions was pseudo-random, following previous schemes—for half of the participants, the order consisted of 1st order orientation, 2nd order orientation, 1st order motion, 2nd order motion-defined, and finally 2nd order contrast modulation, while for the other half of the subjects the order was 2nd order contrast modulation, 1st order motion, 2nd order motion modulation, 1st order orientation, 2nd order orientation (Gao et al., 2014; Reynaud et al., 2014). Each participant completed one repetition per condition, with each qCSF estimate requiring 100 trials preceded by five training trials that were discarded from analysis. All stimuli were created by Psychtoolbox (Brainard, 1997; Pelli, 1997) under Matlab 2012a (MathWorks, Natick, US) installed on a PC (Intel Core i7 processor, 4 GB RAM, 2.67 Hz, ATI Radeon HD 3400 8 bit graphics card) and viewed on a gamma-corrected CRT screen (LG Flatron F900P, 1024 × 768, 85 Hz). The monitor was positioned at 60 cm and viewed monocularly (half of the participants used their right eye) with an opaque patch over one eye.

2.4. Analysis

The qCSF method provides estimates of five parameters (Fig. 1) the maximum gain, the peak frequency, the bandwidth, truncation, and the cut-off frequency of the CSF (Gao et al., 2014; Lesmes et al., 2010; Reynaud et al., 2014). In line with our previous application of qCSF (Gao et al., 2014; Reynaud et al., 2014), the truncation parameter was discarded from our analyses.

Data from this study were compared to an extension of the normative dataset of Reynaud et al. (2014) which expands the age-range of the dataset. The extended normative dataset consisted of qCSF measurements in 102 heathy adults (49 males, 53 females, mean age 46.3 ± years 22 SD). All parameter estimates for each condition were compared between the TBI and the normative dataset using the non-parametric Mann–Whitney U-test. To evaluate the relationship between the 1st order orientation and motion CSFs, we calculated sensitivity difference in decibels between the maximum gain estimates for the 1st order sensitivity functions. This measure was then subjected to the Mann Whitney U-test. Alpha level of 0.05 was adopted for all analyses. In addition, we conducted a non-parametric Bayes factor analysis (Holmes, Caron, Griffin, & Stephens, 2015) (Supplementary material). Spearman’s correlation was used to assess the relationships between time-since-TBI, neuropsychological test measures, and summed score from the visual complaints questionnaire and maximum gain estimates for each condition (Supplementary Figs. 2–6). To account for multiple comparisons (time-since-TBI, all neuropsychological measures, and the questionnaire score were subjected to 20 correlations – 5 conditions × 4 qCSF parameters), we adopted a Bonferroni-corrected alpha 0.0025 for this analysis.

Prior to the group analysis, all individual data were visually inspected. In one participant (T19), the 2nd order motion-modulation sensitivity function (Supplementary Fig. 1) was not log-parabola shaped. For the purpose of the group analysis of 2nd order motion condition, this participant’s data were excluded.

Table 1

<table>
<thead>
<tr>
<th>ID</th>
<th>Age</th>
<th>Gender</th>
<th>Trail making test time</th>
<th>Trail making test errors</th>
<th>Trails test time</th>
<th>Trails test missed</th>
<th>TBI type</th>
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<tr>
<td>T1</td>
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<td>F</td>
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<td>0</td>
<td>106.27</td>
<td>0</td>
<td>Self-reported</td>
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<tr>
<td>T2</td>
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<td>M</td>
<td>56.22</td>
<td>0</td>
<td>157.88</td>
<td>2</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T3</td>
<td>19</td>
<td>F</td>
<td>37.82</td>
<td>0</td>
<td>126.03</td>
<td>1</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T4</td>
<td>22</td>
<td>M</td>
<td>30.02</td>
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<td>100.01</td>
<td>0</td>
<td>Mild simple</td>
</tr>
<tr>
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<td>28</td>
<td>M</td>
<td>22.42</td>
<td>0</td>
<td>114.72</td>
<td>2</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T6</td>
<td>23</td>
<td>F</td>
<td>37.24</td>
<td>0</td>
<td>79.22</td>
<td>1</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T7</td>
<td>48</td>
<td>M</td>
<td>39.77</td>
<td>0</td>
<td>40.08</td>
<td>8</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T8</td>
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<td>F</td>
<td>25.933</td>
<td>0</td>
<td>76.599</td>
<td>4</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T9</td>
<td>55</td>
<td>F</td>
<td>39.204</td>
<td>0</td>
<td>80.945</td>
<td>7</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T10</td>
<td>50</td>
<td>F</td>
<td>23.363</td>
<td>1</td>
<td>68.466</td>
<td>6</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T11</td>
<td>20</td>
<td>F</td>
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<td>0</td>
<td>50.909</td>
<td>13</td>
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</tr>
<tr>
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<td>89.576</td>
<td>10</td>
<td>Self-reported</td>
</tr>
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<td>M</td>
<td>33.646</td>
<td>0</td>
<td>82.928</td>
<td>6</td>
<td>Mild complex</td>
</tr>
<tr>
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<td>F</td>
<td>19.98</td>
<td>0</td>
<td>46.6</td>
<td>6</td>
<td>Mild simple</td>
</tr>
<tr>
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<td>F</td>
<td>19.98</td>
<td>0</td>
<td>46.6</td>
<td>6</td>
<td>Mild complex</td>
</tr>
<tr>
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<td>F</td>
<td>32.72</td>
<td>0</td>
<td>101.05</td>
<td>3</td>
<td>Mild simple</td>
</tr>
<tr>
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<td>F</td>
<td>19.93</td>
<td>1</td>
<td>32.18</td>
<td>5</td>
<td>Mild simple</td>
</tr>
<tr>
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<td>24</td>
<td>F</td>
<td>33</td>
<td>1</td>
<td>60.02</td>
<td>11</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T19</td>
<td>31</td>
<td>F</td>
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<td>0</td>
<td>78.41</td>
<td>0</td>
<td>Self-reported</td>
</tr>
<tr>
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<td>F</td>
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<td>1</td>
<td>62.04</td>
<td>1</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T21</td>
<td>33</td>
<td>F</td>
<td>20.23</td>
<td>0</td>
<td>38.09</td>
<td>10</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T22</td>
<td>28</td>
<td>M</td>
<td>32.35</td>
<td>1</td>
<td>119.65</td>
<td>2</td>
<td>Mild complex</td>
</tr>
<tr>
<td>T23</td>
<td>26</td>
<td>M</td>
<td>27.73</td>
<td>0</td>
<td>125.43</td>
<td>7</td>
<td>Mild complex</td>
</tr>
<tr>
<td>T24</td>
<td>50</td>
<td>M</td>
<td>26.18</td>
<td>1</td>
<td>87.71</td>
<td>3</td>
<td>Self-reported</td>
</tr>
<tr>
<td>T25</td>
<td>23</td>
<td>F</td>
<td>24.49</td>
<td>1</td>
<td>88.21</td>
<td>1</td>
<td>Mild simple</td>
</tr>
<tr>
<td>T26</td>
<td>44</td>
<td>M</td>
<td>22.35</td>
<td>0</td>
<td>88.36</td>
<td>1</td>
<td>Mild simple</td>
</tr>
</tbody>
</table>
While the results reported here reflect the exclusion, the results were unaffected by the inclusion of this subject’s data (see Supplementary material).

3. Results

Average sensitivity functions and measured model estimates for all sensitivity functions based on pseudomedian estimates of qCSF parameters for both groups are depicted in Fig. 2. Values for frequencies smaller than 1 cpd in the 1st order conditions and 0.5 cpd in the 2nd order conditions are plotted as the truncation parameter was discarded from our analysis. Broadly, the sensitivity functions are clustered in two groups corresponding to the 1st and 2nd order vision whereby the 1st order functions show higher maximum gain and sensitivities at higher spatial frequencies.

3.1. First order perception

On average there was a significant trend for decreased sensitivity for the orientation condition and increased sensitivity for the
motion condition in the TBI group across all tested spatial frequencies. This resulted in a separation of the two functions in the TBI group which is in contrast to the normative data where the two functions practically overlap. Quantified as decibel ratio of the maximum gain estimates, the separation of the two functions was significantly larger between the TBI compared to the normative groups ($U = 1619, p = 0.001$). Furthermore, direct comparison of the maximum gain estimates revealed a significantly higher 1st order motion sensitivity in the TBI group ($U = 2017, p = 0.047$); the individual parameters are reported in Fig. 3. There was no significant correlation between the maximum gain estimate and time-since-TBI, visual complaints questionnaire score, or any of the neuropsychological measures.

There was a small but significant shift of the CSF peak towards higher spatial frequencies in the TBI group for both orientation ($U = 1529, p < 0.001$) as well as motion 1st order stimuli ($U = 1906, p = 0.02$). In addition, the variance of the peak spatial frequency values was higher in the TBI group compared to the normative dataset ($W > 7.0, p < 0.001$). The bandwidth estimate for the motion condition ($U = 1363, p < 0.001$) and cut-off spatial frequency ($U = 718, p = 0.001$) were significantly larger compared to the normative dataset (Fig. 3).

### 3.2. Second order perception

Analysis of the CSF parameters for the 2nd order conditions revealed a decreased sensitivity for orientation-defined ($U = 1710, p = 0.003$) and contrast-defined stimuli ($U = 1211, p < 0.001$) but not for motion-defined stimuli ($U = 2532, p = 0.954$) as estimated by the maximum gain parameter (Fig. 3). However, for the motion-defined stimuli, there was a larger variance in the maximum gain parameter estimates in the TBI group $W = 1.71$, $p = 0.04$). Similarly to 1st order conditions, we found no correlation between the maximum gain and the time since TBI, visual complaints questionnaire score, or the neuropsychological measures.

In terms of peak spatial frequency, there was an average shift of the CSFs towards higher spatial frequencies in all 2nd order conditions, but the effect was significant only for the contrast-defined condition ($U = 1736, p = 0.04$). The variance of the peak spatial frequency estimates was higher in the TBI group for all conditions ($W > 2.5, p < 0.006$). Similarly, the cut-off spatial frequency of the CSFs was higher in all conditions on average but reached significance only for the 2nd order contrast-defined ($U = 1911, p = 0.02$) and motion-defined stimuli ($U = 1779, p = 0.014$). Finally, the bandwidth was significantly broader for contrast-defined stimuli ($U = 1566, p = 0.001$); Fig. 3.

### 3.3. Visual complaints questionnaire

Twenty two out of the 26 TBI participants completed our visual complaints questionnaire (Table 2). Five of these participants (23%) reported a change in vision associated with the TBI (score of 5 and above); Table 2. The most common complaint was sensitivity to light and glare (59%) followed by headaches and/or difficulties associated with work with a computer screen (45–54%). Also, nine participants reported that they had to change their visual habits post-concussion. On the other hand, only three and four participants reported blurred vision at near and distance, respectively.

In addition to the Spearman ranked correlation analysis reported above, we carried out a further exploratory analysis of the most vs. the least symptomatic patients based on their questionnaire responses. Based on their ranking on the aggregate score from the questionnaire, we did not observe any significant differences between the five most versus the five least symptomatic patients on the qCSF parameters ($U > 4, p > 0.095$).
Carriers for 2nd order orientation and motion contrasts were set for spatial frequencies. Because of our experimental design in which order stimuli scaled for 1st order performance. The three most order stimuli, with both dynamic and static stimuli, and with 2nd order contrast modulation.

Fig. 3. Pseudomedian estimates of the sensitivity function parameters. Error bars represent ± nonparametric 95% confidence intervals. *p < 0.05 Mann–Whitney U test. 1st Ori – 1st order orientation, 1st Mot – 1st order motion, 2nd Ori – 2nd order orientation modulation, 2nd Mot – 2nd order motion modulation, 2nd Cont – 2nd order contrast modulation.

4. Discussion

In this study, we evaluated several aspects of 1st and 2nd order visual processing following TBI using a unified approach that measured the full contrast sensitivity function for both 1st and 2nd order stimuli, with both dynamic and static stimuli, and with 2nd order stimuli scaled for 1st order performance. The three most notable results are that (1) TBI patients have altered relative sensitivity between dynamic and static 1st order stimuli, (2) TBI patients’ sensitivity to orientation-defined and contrast-defined stimuli is lower, and (3) TBI patients’ sensitivity for both 1st order and 2nd order contrast-defined stimuli is shifted towards higher spatial frequencies. Because of our experimental design in which carriers for 2nd order orientation and motion contrasts were set to be a constant times their 1st order contrast threshold, we can be sure that these deficits for 1st and 2nd order stimuli are independent.

Visual disturbances are prevalent after TBI, and many patients complain of blurred vision, increased sensitivity to visual motion (e.g. watching TV, scrolling on computers and tablets etc.) and/or sensitivity to flicker (e.g. photosensitivity to fluorescent lighting) (Ciuffreda, 2008; Kapoor & Ciuffreda, 2002). Altogether, these observations suggest an altered temporal processing of visual information, this is in agreement with previous psychophysical studies as well as our results. For example, TBI patients have been shown to have elevated thresholds for global motion, as assessed by the random dot kinematogram (Patel, Ciuffreda, Tannen, & Kapoor, 2011), and the increase may be related to visual motion sensitivity and vertigo—increased sensitivity to local motion would amplify the iso-directional dots in a global motion task, thereby decreasing sensitivity. Similarly, it has been shown that TBI patients—at least within the first 30 days post injury—have impaired adaptation to optic flow (Slobounov, Tutwiler, Sebastianelli, & Slobounov, 2006), suggesting heightened sensitivity to this motion signal. The sensitivity to optic flow motion can be so severe within the three days post injury so as to prevent the patients from performing the task as it produced sickness, disorientation and nausea (Slobounov et al., 2006). Interestingly, another index of temporal visual processing, the critical flicker frequency (i.e. the highest temporal frequency allowing a distinction of flickering vs. steady stimulus), does not seem to differ in TBI population (Chang, Ciuffreda, & Kapoor, 2007; Schrupp, Ciuffreda, & Kapoor, 2009). However, the critical flicker frequencies may be related to severity of light sensitivity symptoms in mild TBI patients compared to TBI patients without light sensitivity symptoms or controls (Chang et al., 2007).

Increase of intracortical excitation and/or decrease of GABAergic inhibition—a well-known sequel of TBI (Cantu et al., 2014; Guerriero, Giza, & Rotenberg, 2015; Spiegel, Lague-Beauvais, Sharma, & Farivar, 2015)—may be the cause of this increased motion sensitivity. Support for this proposition can be found in studies investigating neural excitation in the motion visual area MT+/V5 in normal participants. Anodal transcranial direct current stimulation (tDCS)—a noninvasive brain stimulation technique that can increase excitation (Antal, Kincses, Nitsche, & Paulus, 2003) and reduce GABAergic inhibition (Spiegel, Hansen, Byblow, & Thompson, 2012; Stagg et al., 2009)—improved motion direction discrimination task performance for fully coherent motion but lowered performance with decreased coherence (Antal et al., 2004). These findings suggest that changes in the global excitation/inhibition balance affect signal extraction from noise, likely by amplifying the noise.

Migraineurs, who are also known to have increased cortical excitability (Aurora & Wilkinson, 2007), also exhibit superior performance for fully coherent motion condition (Antal et al., 2005) but perform worse than the control group in the non-coherent motion condition (Antal et al., 2005; McKendrick & Badcock, 2004). These findings are notable for two reasons. Firstly, migraineurs are also a very common consequence of TBI (Mayer, Huber, & Peskind, 2013) suggesting that motion sensitivity alteration in both conditions may be driven by similar neuronal mechanisms. Secondly, the findings explain the seemingly contradictory findings between our results (in particular increased sensitivity for the 1st order motion condition) and the previous study on motion sensitivity in TBI participants (Patel et al., 2011). Whereas Patel and colleagues used the random dot kinematogram, i.e. incoherent environment, our 1st order stimuli represent fully coherent motion.

We observed a lower gain for orientation- and contrast-defined 2nd order stimuli in the TBI group. This corroborates and extends...
previous studies. For example, Brosseau-Lachaine et al. (2008) showed reduced sensitivity to 2nd order contrast-defined stimuli of low spatial frequency (0.5 cd/). Lachapelle et al. (2004) reported a significant time delay of the later VEP peak that is believed to reflect higher-order visual processing (Lachapelle et al., 2004, 2008) and the patients showing more pronounced higher-order visual deficits measured by evoked potentials had lower expectations of returning to their normal occupational activities.

It is unlikely that refractive and/or undiagnosed ocular pathologies would explain our observations. While we did not specifically carry out a full optometric evaluation, patients wore their up to date refractive corrections. Furthermore, the carrier contrast was set at ten times its contrast threshold to be fully visible. Therefore, any ocular disorders that might have resulted in an elevation of contrast thresholds would have had negligible impact on the 2nd order sensitivity measurements.

The observed CSFs' shift towards higher spatial frequencies indicated by increased peak and cutoff spatial frequency in both 1st order and 2nd order contrast-defined stimuli in TBI participants is intriguing. This may suggest that in our sample of mild TBI patients, low spatial frequencies may be more affected. This observation is in agreement with some previous findings showing that a proportion of patients with cerebral lesions or injury exhibit mid-to lower-spatial frequencies impairment (Bodis-Wollner & Diamond, 1976; Hess, Zihl, Pointer, & Schmid, 1990).

Our data show a significant increase of bandwidth in the 1st order motion and 2nd order contrast conditions. The CSF bandwidth is recognized as a range of spatial frequencies detectable at a given contrast and it has been shown to be a reliable index of spatial vision. Therefore, this finding is not surprising for the 1st order motion condition where TBI participants also showed an increased sensitivity. However, this result is unexpected for the 2nd order contrast condition in light of the decreased maximum gain for this condition. This finding indicates that despite the decreased sensitivity, the range of detectable envelope spatial frequencies is broader to contrast-modulated 2nd order stimuli.

While we did not specifically test patients for visual symptoms, some patients did report visual disturbances. Interestingly, these reports were uncorrelated with any of our measures, suggesting that contrast sensitivity changes are a separate potential concern, independent of the common vision complaints after concussion. The discrepancy is not necessarily surprising—our qCSF measures probe very low-level pattern perception functions, while the common visual complaints of patients are typically related to “high-level” tasks such as reading or computer use. Thus it is possible that in some patients, there is either different or extended brain injury that combines with the low-level losses, and it is these additional losses related to high-level deficits that trigger the visual complaints.

Using a statistically-optimized method such as the qCSF imposes a risk of generating misleading estimates due to incorrect selection of the prior for each parameter—this may be particularly concerning in a neurologically abnormal population. We do not think this is a critical factor in our study. Firstly, the qCSF method was thoroughly validated in normal population for both 1st and 2nd order stimuli (Lesmes et al., 2010; Reynaud et al., 2014) and secondly, using the qCSF with the same priors has been successfully used with a clinical population with amblyopia (Gao et al., 2014). Another possible limitation is that the qCSF estimate may not be sensitive to possible losses of contrast sensitivity in particular bands of the spatial frequency, deficits that are well-described in brain-lesioned population (Bodis-Wollner & Diamond, 1976; Hess et al., 1990).

In summary, this study brings a unique dataset that provides a comprehensive summary of TBI-related effects on fundamental aspects of low-level visual processing in TBI patients. Our approach utilizing the qCSF allowed us to evaluate—for the first time—the whole CSF curves for five different types of 1st and 2nd order visual stimuli. Comparing the data to a large normative dataset provided psychophysical evidence of increased sensitivity to 1st order motion stimuli, and decreased sensitivity to orientation- and contrast-defined stimuli following TBI. These findings are in general agreement with the clinical reports (Kapoor & Cuffreda, 2002), however do not correlate with the complaints within our sample. The underlying neural causes for the alterations reported here require an integrated approach combining precise psychophysical characterization of the full CSF, in conjunction with neurophysiological measurements as well as temporary neuromodulation.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.visres.2016.03.004.

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