Sex differences in emotional insight after traumatic brain injury

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*Study was performed at authors’ prior institutions:
Neumann: Carolinas Rehabilitation in the Physical Medicine and Rehabilitation
Zupan: Brock University, Department of Applied Linguistics.

Disclosure: Dawn Neumann was one of the co-creators of a publicly available electronic application that was designed to aid in treating certain aspects of alexithymia. Dr. Neumann does not receive any direct royalties from this App.

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ABSTRACT

Objective: To compare sex differences in alexithymia (poor emotional processing) in males and females with traumatic brain injury (TBI) and uninjured controls.

Setting: TBI rehabilitation facility in the USA and a University in Canada.

Participants: Sixty adults with moderate to severe TBI (62% males) and 60 uninjured controls (63% males).

Design: Cross-sectional.

Main Measures: Toronto Alexithymia Scale-20 (TAS-20).

Results: Uninjured males had significantly higher (worse) alexithymia scores than uninjured female participants on the TAS-20 (p=.007), whereas, no sex differences were found in the TBI group (p=.698). Males and females with TBI had significantly higher alexithymia compared to uninjured same-sex controls (both ps<.001). The prevalence of participants with scores exceeding alexithymia sex-based norms for males and females with TBI was 37.8% and 47.8% respectively, compared to 7.9% and 0% for male and females without TBI.

Conclusions: Contrary to the majority of findings in the general population, males with TBI were not more alexthymic than their female counterparts with TBI. Both males and females with TBI have more severe alexithymia than their uninjured same-sex peers, and moreover, both are equally at risk for elevated alexithymia compared to norms. Alexithymia should be evaluated and treated after TBI regardless of patient sex.

Key Words: emotions, brain injury, alexithymia, affective symptoms, sex differences
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Abbreviations:

DDF: Difficulty Describing Feelings

DIF: Difficulty Identifying Feelings

EOT: Externally Oriented Thinking

ES: Effect Size

GCS: Glasgow Coma Score

LOC: Loss of Consciousness

PTA: Posttraumatic amnesia

SD: Standard Deviation

TAS-20: Toronto Alexithymia Scale-20

TBI: Traumatic Brain Injury

GCS: Glasgow Coma Score

LOC: Loss of Consciousness

PTA: Posttraumatic amnesia

SD: Standard Deviation

TAS-20: Toronto Alexithymia Scale-20

TBI: Traumatic Brain Injury
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Processing emotions is fundamental to psychosocial functioning and overall well-being. Alexithymia is an emotional processing deficit that can interfere with the ability to access, recognize, label, differentiate, express, and think about emotions. One common etiology of alexithymia is structural brain damage, which is believed to disrupt areas of the brain involved in processing emotions. This is referred to as “organic alexithymia” and includes traumatic brain injury (TBI). Non-organic etiologies of alexithymia include childhood or adult psychological trauma, being raised in a culture that discouraged emotional expression, or medical illnesses/disorders.

Studies show that alexithymia is more common in people with TBI (31.5% to 63.9%) compared to people without TBI (3.3% to 15.4%). Research in participants with TBI shows that alexithymia is linked to poorer emotional functioning (i.e., anxiety, depression, aggression), unhealthy coping (i.e., avoidant coping, suicide ideation), and reduced quality of life. Given the high rates of alexithymia after TBI and its associations with a variety of negative outcomes, it is important to identify characteristics that might increase risk for alexithymic deficits.

In the general population, one characteristic that has been extensively explored in relation to alexithymia is sex. Findings typically indicate males are more alexithymic than females. A meta-analysis of 41 studies (32 nonclinical and 9 clinical populations) revealed that on average, alexithymia was significantly higher in men than women participants (effect size [E.S] nonclinical sample=.234; clinical sample ES=.163). The Toronto Alexithymia Scale (TAS-20) was used as the outcome measure in 85% of these studies. This sex discrepancy reported in the general population is most commonly explained by the “Normative Male Hypothesis.” This theory
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focuses on the effects of gender socialization, postulating that during childhood, men are more often than women discouraged to feel, show, and communicate their emotions, which increases their risk for alexithymia. The authors expected this gender-based theory would only apply to a subset of male participants, thus they anticipated the small effect size found for sex differences in their meta-analysis.

To our knowledge, no study to-date has compared alexithymia differences between males and females with TBI. There was, however, one study conducted in Finland that examined alexithymia in males and females with mild to severe TBI in comparison to uninjured peer controls of the same sex, but not to each other. That study reported that males with TBI had significantly higher (worse) alexithymia scores than males without TBI, whereas alexithymia scores in females with TBI were similar to their uninjured counterparts. This finding would suggest that women with TBI may not experience alexithymic deficits. However, it was an isolated study that included a sample in which there were twice as many men compared to women. Thus, results for the women (n=18) may have been underpowered. There is a clear need for more research to further explore and potentially replicate this finding.

Currently it remains unknown whether alexithymia differs between men and women within the TBI population. Since alexithymia in persons with TBI is more likely to be the result of the neurological injury (as opposed to socialization), the typical sex differences observed in the general population may not apply to this population. The primary objective of the current study was to compare alexithymia between male and female participants within TBI and non-TBI samples. The secondary objective was to examine how alexithymia in males and females with TBI differs from their uninjured
counterparts. Although a similar comparison has been explored in another study (described above), it is unknown if their findings would generalize to a sample that differed in injury severity and geographical location. Within these two study objectives, the prevalence of alexithymia problems will also be explored. However, the traditional TAS-20 cut-off score for severe alexithymia (>60) will not be used for two main reasons. The >60 cut-off was a preliminary recommendation based on a small sample of patients referred to a behavioral medicine clinic (14 males, 25 females). Second, that recommendation applies the same cut-off score for men and women, despite the fact there are sex-based differences in the norms reported for this measure. To overcome these limitations, alexithymia prevalence in this paper will reflect the proportion of participants with alexithymia scores that exceed sex-based norms provided for the TAS-20.

METHODS

Study Design and Setting.

This cross-sectional study was performed at a rehabilitation hospital in North Carolina, USA and a University in Ontario, Canada. Each site received approval from their ethics committees and all participants provided informed consent before participation. Participants with TBI were recruited via letters and flyers distributed to current and former patients at partnering brain injury facilities, and to local TBI support groups. Peer controls were recruited through flyer advertisements posted at universities and other various community establishments.

This study was conducted as part of a larger research project on emotional
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processing after TBI, which resulted in publications that are distinct from the focus and findings of the current manuscript.\textsuperscript{7,17-19}

**Participants**

This study included 60 participants with TBI (62\% male) and 60 uninjured controls (63\% male) who were frequency matched for age and sex. Participants with TBI met at least one of the Mayo Classification criteria\textsuperscript{20} for moderate to severe TBI (i.e., Glasgow Coma Scale score <13 at the time of injury; posttraumatic amnesia $\geq$ 24 hours; or loss of consciousness for $\geq$ 30 minutes). Uninjured controls were excluded for any history of TBI regardless of severity. All participants were excluded for developmental disorders that were affective in nature (e.g., Autism Spectrum Disorder), non-traumatic neurological disorder (e.g., stroke), major psychiatric disorder, and uncorrected vision or hearing impairments that would impede task participation.

On average, male and female participants with and without TBI were near 40 years of age and had 14-15 years of education. Although the sample was predominantly white, 17.4\% (n=4) of females with TBI were Black/ African American. The majority of participants with TBI were single. For uninjured controls, the majority of females were married, and the males were evenly split between married and single. Participants with TBI were predominantly injured from a motor vehicle accident, and on average, were 12-14 post-injury (range: 6 months-37 years). Median loss of consciousness was 21 days and 14 days for males and females, respectively. More detail regarding demographics and injury characteristics (TBI only) are presented in Table 1.

Power analyses were based on the primary objectives of the larger study, which examined group differences in emotion recognition on the Diagnostic Analysis of
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Nonverbal Accuracy 2-Adult Faces.\textsuperscript{21} The sample size calculation indicated that 120 participants were sufficient to detect medium effect sizes, with 80\% power for independent sample \textit{t}-tests using two tails. Although affect recognition is not the same as alexithymia, they are related constructs.\textsuperscript{7}

\textbf{Measures}

\textit{Toronto Alexithymia Scale-20 (TAS-20)}.\textsuperscript{2,11,12} The TAS-20 is comprised of 20 statements that evaluate three alexithymia components: Difficulty Identifying Feelings (DIF); Difficulty Describing Feelings (DDF); and Externally Oriented Thinking (EOT). Participants rate agreement using a Likert scale from 1 (strongly disagree) to 5 (strongly agree). The total TAS-20 scale ranges from 20-100 points. Higher scores indicate a greater degree of alexithymia. Based on TAS-20 norms, average male scores are 47.30 (S.D.=11.32) and average female scores are 44.15 (S.D.=11.19).

\textbf{Data Analyses}

Two-tailed independent samples \textit{t}-tests were conducted to examine between group differences on demographic variables. A Chi-square test was used to examine if there was a significant difference in the number of males and females in the TBI and uninjured control groups. The relationship of alexithymia outcomes with demographic and injury-related variables were calculated with Pearson correlations. A series of two-tailed independent samples \textit{t}-tests were conducted to compare mean alexithymia sex differences within the TBI and uninjured control groups (TAS-20 total and 3 subscales)
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(\(\alpha = .0125\)), as well as for between group comparisons of males with and without TBI and females with and without TBI (\(\alpha = .0125\)). Additionally, counts and percentages were calculated to indicate prevalence of participants who met the criteria of having a score greater than one standard deviation above the sex-based norms. This equated to a score of \(\geq 59\) for males and \(\geq 56\) for females. The relationship between alexithymia classifications with group and sex were explored using Fisher’s Exact tests due to Chi-Square violations. SPSS Version 25 was used for all calculations.

Results

Demographics and Injury Related Variables

The TBI and uninjured control groups did not significantly differ in age (\(t=.146, p=.884\)), sex (\(X^2=.036, p=.850\)), or race (\(X^2=3.748, p=.290\)), but did differ by marital status (\(X^2=17.13, p=.002\)). Peer controls had significantly more years of education than participants with TBI (\(t=-3.06, p=.003\)). Alexithymia was not significantly correlated with age (TBI: \(r=-.139, p=.291\); uninjured controls: \(r=.037, p=.777\)), race (TBI: \(\rho=-.171, p=.193\); HC: \(\rho=.061, p=.642\)), or study site (TBI: \(\rho=-.093, p=.482\); uninjured controls: \(\rho=.029, p=.823\)) for either group, nor was it correlated with time post-injury (\(r=.049, p=.711\)) or PTA (\(r=-.053, p=.775\)) for participants with TBI. However, the TAS-20 was significantly correlated with TBI years of education (\(r=-.395, p=.002\)) and LOC (\(r=.335, p=.035\)). Since significant correlations were relatively weak and because this pilot study was not powered to control for these variables, the following analyses do not factor in these variables. This is addressed in more detail in our discussion of limitations.
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Demographics were also examined for sex differences within each group. Males and females within the TBI sample did not significantly differ in age ($t = -.181, p = .857$), education ($t = .647, p = .521$), or race ($X^2 = 2.960, p = .228$). Similarly, uninjured males and uninjured females did not differ in age ($t = -.776, p = .441$), education ($t = .931, p = .357$), or race ($X^2 = .610, p = .737$). Male and female participants with TBI did not significantly differ in time-post injury ($t = .910, p = .367$), PTA ($t = -.228, p = .821$), or LOC ($t = .720, p = .476$).

Comparisons of alexithymia in males versus females

Statistical values are reported in Table 2 and the proportion of participants with above average alexithymia are shown in Figure 1. Within the TBI sample, mean alexithymia scores for males and females did not significantly differ from one another in the overall TAS-20 score, nor on any of the TAS-20 subconstructs. Effect sizes were also small. In contrast, uninjured male participants had significantly higher scores compared to uninjured female participants on TAS-20 total, DDF, and EOT, with large effect sizes. DIF did not differ between uninjured male and female participants. Fisher’s Exact tests indicated that the proportion of males and females who had above average alexithymia did not statistically differ for the TBI group ($p = .591$) or the Uninjured controls group ($p = .292$).

Comparisons of Alexithymia in Participants with TBI versus Uninjured Controls by Sex

Compared to uninjured males, male participants with TBI had significantly higher (worse) total alexithymia scores, as well as DIF and DDF subconstructs, but not on EOT. Compared to uninjured female participants, female participants with TBI had significantly higher alexithymia overall and significantly higher scores on all three alexithymia subconstructs. Aside from EOT for males, all effect sizes were large. The
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Proportion of individuals who had greater than average alexithymia was significantly larger for males and females with TBI ($p=.002$ and $p<.001$, respectively) compared to their same-sex uninjured peers.

[Insert Table 2 and Figure 1]

Discussion

Alexithymia is common after TBI and frequently associated with negative psychosocial outcomes.  
This is the first study to examine sex differences in alexithymia within a TBI population, and the first to use sex-based norms for identifying alexithymia in a clinical population. Findings from this study revealed that a similarly large proportion of males and females with TBI had alexithymia scores that exceeded sex-based norms (37.8% to 47.8%, respectively). Further, men and women with TBI had similar alexithymia scores, on average. This suggests alexithymia is a problem for both men and women with TBI. This contrasted findings for the uninjured control group. While proportions of alexithymia in men and women without TBI did not differ (7.9% and 0%, respectively), TAS-20 scores for uninjured males were significantly higher than uninjured females for all components but DIF. The sex differences found in our uninjured controls are consistent with prior studies in the general population. We suspect that our TBI group did not conform to the typical alexithymia sex differences due to the different etiology of alexithymia in this population (i.e., neurological damage versus gender socialization/ “Normative Male Hypothesis”). Even if the Normative Male Hypothesis principle was relevant to some participants with TBI prior to their injury, the neurological insult likely overshadows the influence of socialization on
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Alexithymia differences between males and females with TBI were also compared to uninjured controls of the same sex. Compared to their uninjured counterparts, male and female participants with TBI had significantly higher alexithymia scores for overall alexithymia and all subconstructs, except EOT, where males with and without TBI did not differ. This finding regarding EOT may suggest that regardless of injury, males are less likely to focus on inner emotional states, as opposed to external factors. Moreover, our results further indicated that a greater proportion of male and female participants with TBI had elevated alexithymia compared to uninjured controls (males: 37.8% vs 7.9%; females: 47.8% vs 0%, respectively). These findings contradict results from the Finland study described earlier, which found that females with TBI did not differ from their uninjured peers, neither by means scores or prevalence of alexithymia. It is possible our findings differ from the Finland study due to cultural or injury severity differences (we did not include participants with mild TBI). Their female participants may have had milder injuries leading to less of an impact on alexithymia traits. Our females with TBI had higher mean alexithymia scores than theirs (53.39 vs 47.4, respectively). Additionally, the Finland study used the “high” alexithymia cut-off (>60) to determine prevalence of alexithymia for all participants, whereas our study used sex-based norms which may partially explain the discrepancy in identification of women with alexithymia problems between our study (47.8%) and theirs (22%). Since women have lower alexithymia norms than men, reaching a universal cut-off would require a bigger deviation from their own norms, than it would for men, to reach “high
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alexithymia” levels. Consequently, it is not a fair method for conducting sex comparisons.

Research on sex differences after TBI has become a growing focus of interest in TBI outcomes, including other aspects of social cognition.\textsuperscript{3,29-31} It is widely acknowledged that after a TBI, social cognition is frequently impaired\textsuperscript{32-34} and these deficits are associated with worse social outcomes.\textsuperscript{33} The most relevant to alexithymia, is the research on empathy. Past research shows an association between alexithymia and empathy in the TBI population.\textsuperscript{3,7} Consistent with our alexithymia findings, a study examining sex differences in empathy\textsuperscript{31} found that mean empathy scores did not differ between men and women with TBI but a larger proportion of women with TBI (44%) fell below emotional empathy norms than men with TBI (17%).\textsuperscript{31} Although in our study the proportions did not differ statistically, the percentage of females who had alexithymia scores that exceeded the norms was larger than that for males (47.8% vs 37.8%). Not all sex difference outcomes in the social cognition arena are consistent with our alexithymia findings.\textsuperscript{3,29,30} For example, another study on empathy found men with TBI were more impaired than females.\textsuperscript{3} While it is suspected that cultural differences and the use of different empathy measures could potentially account for these conflicting results, it is evident that more work is needed to expand knowledge and resolve the inconsistency in findings regarding sex differences in social cognition after TBI.

Findings from this study should convince researchers and rehabilitation professionals that it is important to take sex and gender-based differences into consideration when working with individuals with TBI to address alexithymia and related social cognition deficits. Clinicians working with people with TBI should evaluate
alexithymia and other emotional processing functions in their patients regardless of sex, and be cognizant of the potential psychosocial impact that deficits in these areas may have on their women patients. Gender expectations for women presume women are emotional, empathic individuals. However, emotional experiences or expressions are often blunted with alexithymia, and related to lower empathy, and are thus discrepant with these expectations preventing females with alexithymia from fitting this stereotypical mold. For men, society often expects them to be less emotionally expressive. Thus, an alexithymic deficit may be more detrimental to women with a TBI than men. It may be helpful for clinicians to determine how much their patients who have alexithymia relate to gender norms relevant to emotional expression, and seek information about family expectations as well. The more discrepancy there is, the more important it will be to educate the patient and families about potential changes associated with alexithymia in how they experience and/or express their emotions. This process could help to prioritize rehabilitation goals. Early evidence suggests alexithymia is treatable after TBI. Moreover, targeted treatment can potentially reduce related emotion dysregulation deficits.

**Limitations and Future Directions**

The sample size was driven by a related study objective on emotion perception and was not specifically powered to examine within group alexithymia sex differences nor to control for potential covariates (e.g., years of education). As is typical with preliminary studies with modest sample sizes, results should be interpreted with caution and used to determine appropriate sample sizes needed for future research. Future studies should explore contributions of potential covariates to alexithymia outcomes,
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such as education, executive functioning, coping mechanisms, depression, anxiety, and posttraumatic stress disorder (PTSD), which may also vary by sex. Although this study was not heavily unbalanced with regards to sex (60% male), future research focused on sex differences should strive for equal distribution.

It is also a limitation that we did not collect information about participants’ gender identity. While sex and gender are interrelated, there are important distinctions. Sex is primarily defined by biological factors, whereas gender is tied to characteristics that society deems to be more or less stereotypically male or female. Participants may have identified with a gender different from their biological sex, which could have impacted study outcomes. For instance, some females with TBI could possibly have identified with more male-like characteristics, which could have eradicated the sex differences. Future work should collect both sex and gender-related data to better characterize the sample.

Another study limitation is that there are mixed reports in the literature regarding the psychometric properties of the TAS-20, and it has been recommended to be administered in conjunction with other measures. To the authors’ knowledge, the closest objective measure is the Levels of Emotional Awareness Scale (LEAS), a time-consuming assessment that is not a direct measure of alexithymia nor one that addresses all of the alexithymia constructs. Despite its potential limitation, the TAS-20 was largely selected due to being the most widely used instrument for measuring alexithymia, which facilitates comparison to other studies. Further, it is the only measure specific to alexithymia that has been used in the TBI population. The TAS-20 includes statements that people with severe TBI can evaluate as
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characteristic of themselves or not. Future studies could use the LEAS as an objective measure to complement the TAS-20. A final limitation is that it is unknown if the alexithymia identified in the participants with TBI was due strictly to neurological damage or if it was present prior to their injury (i.e., primary and/or secondary alexithymia). Unfortunately, current measures do not have the capacity to make this distinction.

Conclusions

Study findings suggest that a similarly large proportion of males and females with TBI exceed the alexithymia norms reported for their sex, and in general, have a similar degree of alexithymia. This is in contrast to the general population which typically shows males are more alexithymic. Etiological differences (i.e., organic alexithymia) may be a possible explanation for this difference. Rehabilitation professionals should be aware of these potential emotional deficits in their patients for males and females alike, and should evaluate, educate, and treat accordingly.

Conflicts of Interest/Grant Funding:

Dr. Dawn Neumann contributed to the development of an electronic application that is designed to aid in the treatment of alexithymia. She does not receive royalties for the App, but Indiana University does.

Dr. Barbra Zupan has no conflicts of interest.

Source of funding: This work was supported by Cannon Research Center at Carolinas Rehabilitation in Charlotte, North Carolina, and the Humanities Research Institute at Brock University.

Related Presentations:

References


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**Figures Legends:**

492 Figure 1. Percentage of Participants with Above Average Alexithymia (>1SD above sex-based norms)
**Figure 1**

Legend:
Figure 1. Percentage of Participants with Above Average Alexithymia (>1SD above sex-based norms)
<table>
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<tr>
<th></th>
<th>TBI Males (n=37)</th>
<th>TBI Females (n=23)</th>
<th>Uninjured Controls Males (n=38)</th>
<th>Uninjured Controls Females (n=22)</th>
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<td>87%</td>
<td>4.3%</td>
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</tr>
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</table>

| Time post-injury (years) | 14.65 (10.96) | 12.10 (9.86) |
|                         | Range: 1-37   | Range: 0.6-27 |

| LOC (days) (n=40) | 48.04 (57.68) | 36.30 (40.08) |
|                   | Median: 21    | Median: 14    |
|                   | Range: 0.5-180| Range: 1-120  |
| 25th%: 3          |              | 25th%: 6.5    |
| 50th%: 21         |              | 50th%: 14     |
| 75th%: 90         |              | 75th%: 60.1   |

| PTA (days) (n=31) | 18.56 (41.79) | 21.77 (34.06) |
|                  | Median: 7     | Median: 7     |
|                  | Range: 0.5-180| Range: 0.5-120|
| 25th%: 6          |              | 25th%: 3.5    |
| 50th%: 7          |              | 50th%: 7      |
| 75th%: 7          |              | 75th%: 30     |

Note: LOC=Loss of consciousness; PTA=Post-traumatic amnesia
<table>
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<tr>
<th></th>
<th>TBI Mean (S.D.)</th>
<th>Uninjured Controls Mean (S.D.)</th>
<th>Male vs Female p value (ES)</th>
<th>Male vs Female p value (ES)</th>
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<th>TBI vs Ctrl p value (ES)</th>
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