Associations Among Posttraumatic Stress Disorder Symptoms, Substance Use, and Affective Attentional...
Associations Among Posttraumatic Stress Disorder Symptoms, Substance Use, and Affective Attentional Processing in OEF/OIF/OND Veterans

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Abstract: The majority of research examining affective attentional bias in posttraumatic stress disorder (PTSD) has not examined the influence of co-occurring psychiatric disorders. This study examined the individual and interactive effects of PTSD symptoms and substance use disorders (SUDs) on affective attentional processing among 323 veterans deployed to Iraq or Afghanistan. Participants were divided into those with SUD (SUD+; n = 46) and those without (SUD−). Substance use disorder was determined using the Structured Clinical Interview for DSM-IV. Posttraumatic stress disorder was measured using the Clinician Administered PTSD Scale. A computerized go/no-go task (Robbins et al., 1994, Robbins et al., 1998) assessed affective attentional processing. Relative to those without SUD, those with SUD showed a significant association between PTSD symptoms and increased omission and commission accuracy rates and decreased d prime. No effects of valence were found. Findings suggest the need to consider co-occurring SUD when investigating the effects of PTSD on attentional control.

Key Words: PTSD, substance use disorder, attention, affect, inhibition, veterans (J Nerv Ment Dis 2017;00: 00–00)

Posttraumatic stress disorder (PTSD) and substance use disorders (SUDs) are relatively common military-related psychiatric illnesses, which are both marked by disrupted attentional control (Drobes et al., 2006; Harvey et al., 1996). Individuals with PTSD are hypothesized to display both an attentional bias toward and difficulty disengaging from information with a threatening or general negative valence (Naim et al., 2014; Pmeles et al., 2007). Despite research showing the effects of SUD on attentional bias/inhibitory control (Field and Cox, 2008; Field et al., 2009; Lambe et al., 2014), it is currently unknown if PTSD-related affective attentional bias might be exacerbated when the commonly co-occurring condition of SUD is present (Seal et al., 2011). In this study, we examine the relative and combined contributions of these disorders in the critical domain of affective attentional bias/inhibitory control.

Attentional bias toward threat-related information has been implicated in both the onset and maintenance of PTSD (Armstrong et al., 2013; Kimberle et al., 2010). This threat-related bias may have an impact on both information processing and subsequent behavior, as it functions as a gating mechanism that directs attention based on stimulus valence (Constans, 2005). For example, studies using the Stroop paradigm have demonstrated that, compared with individuals without PTSD, those with PTSD demonstrate slower reaction times when asked to name the color of the ink for trauma- and threat-related, relative to neutral, words (Cisler et al., 2011; Constans et al., 2004). It is possible that the additional effort involved in inhibiting prepotent response tendencies in the context of information with a threat-related valence accounts for these slowed response times (for review see Buckley et al., 2000). In contrast, research using visual attention tasks has shown that affective attentional bias can facilitate performance. Specifically, relative to those without PTSD, those with PTSD more quickly identify visual targets when the stimuli consist of negatively valenced words (Bryant and Harvey, 1997) or threatening faces (Fanti et al., 2012). Overall, the literature examining affective attentional bias in PTSD remains inconsistent, with several studies failing to find an association between PTSD and attentional bias for negatively valenced stimuli (Ashley et al., 2013; Elsesser et al., 2005). Subsequent research has highlighted that these inconsistent findings may be due to variability in task methods and/or stimuli type (Cisler et al., 2011; Olatunji et al., 2015). Moreover, it remains unclear whether alterations in affective attentional processing in PTSD result from enhanced threat detection, difficulty disengaging from threat-related stimuli, and/or deficits in inhibitory control (Aupperle et al., 2012; Esterman et al., 2013; DeCuir et al., 2015).

The majority of research examining affective attentional bias among those with PTSD has neither controlled for nor examined the influence of co-occurring psychiatric disorders (Bryant and Harvey, 1997; Constans et al., 2004), although there are several exceptions. For example, two recent studies have explored the impact of co-occurring depression on attentional bias in PTSD (Wittekind et al., 2015; Hauschildt et al., 2013). This research, using tasks with visual affective cues, supports the effects of depression as opposed to PTSD, on affective attentional bias after trauma exposure. In addition, Amick et al. (2013) found a moderating effect of mild traumatic brain injury (mTBI) on the association between PTSD and affective-related attentional control. Specifically, among individuals with mTBI, increasing PTSD symptom severity was associated with a more liberal response pattern to positively versus negatively valenced stimuli. However, this association was not evident among individuals without TBI. Cumulatively, these preliminary findings underscore the need to investigate the influence of co-occurring deployment-related conditions to more precisely characterize the association between PTSD symptoms and affective attentional processing, and clarify inconsistencies in this literature.

Similar to those with PTSD, individuals with SUDs demonstrate alterations in attentional control. Findings show differential attentional bias toward substance-related cues (Field and Cox, 2008; Weinstein and Hauschildt, 2006), which has been proposed as a key predictor of addictive behaviors (Cox et al., 2002; Marissen et al., 2006). Experimental studies using modified Stroop tasks have demonstrated support for this. Drug-related attentional bias was found in a variety of SUDs, including nicotine, alcohol, cocaine, and heroin use (Cox et al., 2002; Heather et al., 2006). Notably, one study found that drug-related attentional bias was
exacerbated after trauma cue exposure among cocaine-dependent inpatients with PTSD, suggesting interactive effects of PTSD and SUD on attention (Tull et al., 2011).

Although the majority of findings have highlighted an attentional bias for substance-related cues among those with SUDs, recent research has extended this to examine the presence of affective attentional bias related to positively and negatively valenced cues more broadly. Although studies are limited, preliminary findings support the influence of affective stimuli (i.e., both positive and negative valence versus neutral) on attentional/inhibitory control performance among individuals with alcohol (Lambe et al., 2014), as well as opiate use disorders (Dunning et al., 2011; Lubman et al., 2008).

Given the high rates of comorbidity between PTSD and SUD (Seal et al., 2011) and the independent effects of each disorder on attentional bias (e.g., Cisler et al., 2011; Hester et al., 2006), it is important to investigate the relative and interactive influence of each disorder on attentional bias, as this information is critical to a more thorough understanding of the mechanisms that contribute to impaired attentional control in this population. Thus, this study examined the effects of PTSD and co-occurring SUD on performance using a validated affective go/no-go task (Robbins et al., 1998; Robbins et al., 1994) that has been shown to be sensitive to alterations in affective attentional bias in other related clinical samples (Amick et al., 2013; Kaplan et al., 2006; Seymour et al., 2015; Steiner et al., 2013). Task performance was examined among a sample of Operation Enduring Freedom, Operation Iraqi Freedom, and Operation New Dawn (OEF/OIF/OND) Veterans recruited from a large metropolitan area in the Northeastern United States. This study investigated the effects of PTSD and SUD, as well as their interaction, on several different indices of attentional performance, including commission and omission accuracy, reaction time, and signal detection indices of discriminability and response bias. We predicted that errors of omission and commission would be significantly greater in response to negative versus positive stimuli. Consistent with existing literature, we predicted that, relative to those without PTSD, individuals with PTSD would show significantly more errors of commission. In addition, we hypothesized that there would be a significant PTSD × SUD interaction, such that PTSD symptom severity would be more strongly associated with decrements in omission and commission accuracy in response to negative affective stimuli among those with SUD than among those without this comorbidity.

METHODS

Recruitment

An initial sample of 359 consecutively enrolled veterans was recruited from the longitudinal cohort study of the VA Rehabilitation Research and Development-supported Traumatic Brain Injury National Network Research Center: The Translational Research Center for TBI and Stress-Related Disorders (TRACTS; McGlinchey et al., 2017). Inclusion criteria for this larger study were OEF/OIF/OND deployment and the age of 18 to 65 years. Exclusion criteria included the following: a) history of neurological illness or seizures unrelated to head injury, b) current diagnosis of psychotic disorder, c) current diagnosis of bipolar disorder, d) cognitive disorder due a medical condition other than TBI, and e) psychiatrically instability (i.e., homicidal/suicidal ideation that required immediate crisis intervention or current, unstable psychological diagnosis that would interfere with accurate data collection).

Participants

From this initial sample of 359 individuals, 25 participants were excluded for poor performance on effort measures and 11 were excluded for a history of moderate or severe TBI. The final sample was composed of 323 participants: 46 (42 men) individuals with a recent substance abuse/dependence diagnosis (SUD+ group) and 277 (248 men) participants without current substance abuse/dependence (SUD− group). Before completing any of the experimental procedures, all participants provided written informed consent. The local institutional review board approved all study procedures. Participants were recruited from the Boston Metropolitan and surrounding areas by a full-time recruitment specialist via events involving Army and Air National Guard, Marine and Marine Reserves, and Army and Army Reserve Units.

Materials and Procedure

For the majority of participants, study measures were collected in one-day sessions with a standardized order of test administration. Data in the analyses described in this article are a subset of the larger TRACTS database. After obtaining informed consent, a doctoral-level psychologist assessed TBI, PTSD, SUD, and other DSM-IV Axis I diagnoses using structured interviews. A consensus diagnosis for TBI, PTSD, SUD, and other Axis I disorders was determined via case review by at least three psychologists and/or psychiatrists. Diagnostic assessment was followed by administration of neuropsychological tests including the go/no-go task and self-report measures.

Assessments

Substance Abuse/Dependence

The Structured Clinical Interview for DSM-IV Axis I Disorders (SCID-I/NP; First et al., 2002) is a semistructured interview that includes modules designed to assess either the lifetime or current (past-month) experience of DSM-IV Axis I psychiatric disorders (http://www.scdi4.org/psychometric/). All participants received the SCID-I/NP to determine eligibility and to characterize the individual's psychological history. The SCID-I/NP was used to diagnose participants with current alcohol/substance abuse or dependence. Participants meeting criteria for current alcohol or substance abuse or dependence were included in the SUD+ group; those who did not meet current criteria were included in the SUD− group.

Posttraumatic Stress Disorder

The Clinician-Administered PTSD Scale (CAPS-IV; Blake et al., 1995) was used to assess the presence and severity of PTSD symptoms as well as PTSD diagnostic status. The CAPS is a well-validated and reliable structured clinical interview used to evaluate the DSM-IV-TR re-experiencing (criterion B), avoidance/numbing (criterion C), and hyperarousal (criterion D) symptoms of PTSD (Blake et al., 1993; Blake et al., 1995). Participants are queried about the intensity (0–4) and frequency (0–4) of each of the 17 DSM-IV-TR PTSD symptoms, from which a total score is derived (minimum score = 0, maximum = 136), reflecting overall symptom severity. The CAPS total score was used for primary analyses to examine the dynamic interaction of PTSD symptom severity and SUD status upon behavioral measures of inhibitory control. The CAPS also was used to determine what percentage of participants met full DSM-IV-TR diagnostic criteria for PTSD; participants considered to have PTSD if they endorsed at least one B, three C, and two D symptoms (frequency rating ≥ 1 and intensity rating ≥ 2 were required to be counted as a symptom).

Effort

To ensure that all participants included in the analyses were adequately motivated to perform the tasks of interest, we administered the Green's Medical Symptom Validity Test. Adequate effort was determined using established cutoffs (Green, 2003).

TBI Diagnosis

The Boston Assessment of TBI-Lifetime (BAT-L; Fortier et al., 2013) was used to assess potential brain injury during three
lifetime periods: premilitary, military, and postmilitary. Preliminary validation in a subsample of participants (n = 131) demonstrated excellent correspondence between the BAT-L and the Ohio State TBI Assessment Method (Kendall's tau b = 0.95), a validated method for TBI identification (Fortier et al., 2013). Traumatic brain injury criteria including altered mental state, posttraumatic amnesia, and loss of consciousness were evaluated through open-ended questioning. Participants were excluded from analyses if they had any history of a moderate or severe TBI, as these injuries can have lasting impairments on cognitive performance.

**Affective Go/No-Go**

Participants performed the affective Go/No-Go task (Robbins et al., 1998; Robbins et al., 1994) after the clinical interviews. The affective go/no-go task is a continuous performance task. A series of stimulus words with either positive or negative valence was presented on the center of a monitor for 300 milliseconds with a 900-millisecond interstimulus interval. Participants completed 10 blocks. Each block contained 18 words (9 positive and 9 negative valence). For each block, target word valence (positive or negative) was constant and switched every two blocks. Valence of the target and distractor words was nonspecific (e.g., unrelated to deployment trauma). Order of presentation was counterbalanced across participants. The first two blocks of the task served as practice trials. The critical data set consisted of the remaining eight blocks. Within each block, there were nine words that were consistent with the target valence (“go” target words) and nine words that were inconsistent with the target valence (“no-go” distractor words).

At the start of each block, the participant was informed of the targeted valence for that block (either positive or negative). The participant’s task was to determine if the valence of the presented word matched (go condition) or did not match (no-go condition) the targeted valence. The participant was instructed to press the spacebar when the valence of the stimulus word matched the targeted valence and to withhold pressing the spacebar when the valence of the stimulus word did not match the targeted valence. Participants were asked to press the button as quickly and accurately as possible. Reaction times and errors of omission and commission were recorded for each trial.

**Affective Go/No-Go Task Outcome Measures**

Dependent measures included indices of omission errors (i.e., failure to press the space bar for a word that matched the targeted valence), commission errors (i.e., pressing the space bar for a nonmatched word for a targeted valence), and reaction time (RT; in milliseconds). Omission accuracy is an index of the percent of correct responses on potential omission trials and was calculated as \[1 - (\text{omission errors/total possible omission errors}) \times 100.\] Similarly, commission accuracy was calculated as \[1 - (\text{commission errors/total possible commission errors}) \times 100.\]

In addition, d prime (d’) and criterion were examined as measures of discrimination accuracy and response bias during the task. d’ is an index of overall stimulus discriminability, or sensitivity, and was calculated separately for the positive and negative blocks. d’ was calculated according to the following formula: \[d' = Z(\text{hit rate}) - Z(\text{false alarm rate}).\] Criterion is a measure of response bias (to respond or withhold), which is calculated according to the following formula: criterion = \[-Z(\text{false alarm rate}) + Z(\text{hit rate})/2\]. Negative criterion values indicate a liberal response bias (resulting in more hits and also more false alarms). A positive criterion value indicates a more conservative response bias, with fewer hits and fewer false alarms. If criterion equals zero, then the subject’s criterion is neutral, showing no decision bias toward either response type (i.e., go or no-go). For analysis, univariate outliers (i.e., participant task outcomes that were greater or less than 3 standard deviations from the sample mean) were identified and removed.

**Statistical Analyses**

Statistical analyses were performed using SPSS 19.0 software. Participant characteristics were compared using \(\chi^2\) analyses for categorical variables and t tests and analyses of variance for continuous variables.

Analysis of go/no-go task data was performed using a general linear model (GLM) with repeated measures. General linear model was performed for each of the dependent measures with substance abuse/dependence group (SUD group) as the between subjects factor and block valence (positive and negative) as the repeated within subjects factor. Each GLM model included total CAPS scores, and a CAPS score × SUD group interaction as covariate terms. SPSS examines the effects of constant covariates and covariate interactions as between-subjects factors, allowing us to examine our primary question of interest, namely, whether the association between PTSD symptom severity and behavioral performance differed as a function of substance abuse/dependence status. The total CAPS score, rather than PTSD presence or absence, was examined because the critical question to be examined was if the association between PTSD symptom severity and performance on the go/no-go task varied in the SUD+ compared with the SUD− group. To control for any potential effects of age, education, current depressive disorder, and lifetime mTBI (y/n), these variables also were included as covariates in these models as all of these variables are known to influence performance on tasks of executive functioning. Our previous work did not find a main effect of mTBI upon go/no-go performance (Amick et al., 2013). However, as our prior work found an interaction between mTBI and PTSD, and research shows effects of mTBI in related areas of cognitive performance (e.g., Karr et al., 2014), we included mTBI as a covariate in all analyses. To clarify the nature of significant PTSD × SUD interactions, Pearson correlations were conducted to examine the relationship between PTSD symptom severity and performance on the go/no-go task separately for each SUD group.

**RESULTS**

**Participant Characteristics**

The SUD+ and SUD− groups did not differ with respect to age, education, sex, number of lifetime mTBIs, or the presence of anxiety or mood disorders. As expected, the groups did differ with respect to PTSD symptom severity and number of individuals with a PTSD diagnosis. That is, among those in the SUD+ group, PTSD symptom severity was significantly higher and significantly more individuals were diagnosed with PTSD, relative to those in the SUD− group (see Table 1).

**Affective Go/No-Go**

**Reaction Times**

For the repeated measures GLM involving RTs, there was no effect of block valence, SUD group, CAPS score, or their interaction (block valence × SUD group or CAPS × SUD group). None of the covariate terms were significant (\(P > 0.05\)).

**Commission Accuracy**

The main effect of valence and all of the interactions involving valence were not significant (all \(P s > 0.32\). The main effect of SUD group \([F(1,315) = 1.22, p = 0.27; \text{partial eta square} = 0.004]\) was not significant. The main effect of CAPS score \([F(1,315) = 5.10, p = 0.025; \text{partial eta square} = 0.016]\) and the interaction between SUD group and CAPS score \([F(1,315) = 4.34, p = 0.038; \text{partial eta square} = 0.014]\) were significant. As shown in Figure 1A, within the SUD+ group, commission accuracy decreased with increasing PTSD
TABLE 1. Demographic and Psychiatric Characteristics of the Sample as a Function of Substance Use Disorder Diagnosis

<table>
<thead>
<tr>
<th></th>
<th>SUD+ Group (n = 46)</th>
<th>SUD− Group (n = 277)</th>
<th>t Value</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, yr</td>
<td>30.76 (7.73)</td>
<td>32.06 (8.59)</td>
<td>-0.96</td>
<td>0.34</td>
</tr>
<tr>
<td>Sex (male-female)</td>
<td>42:4</td>
<td>248:29</td>
<td>0.14</td>
<td>0.71</td>
</tr>
<tr>
<td>Education, yrs</td>
<td>13.54 (1.74)</td>
<td>13.98 (1.96)</td>
<td>-1.42</td>
<td>0.16</td>
</tr>
<tr>
<td>No. lifetime mTBI</td>
<td>1.85 (2.97)</td>
<td>1.52 (2.07)</td>
<td>0.94</td>
<td>0.35</td>
</tr>
<tr>
<td>Current anxiety disorder DX (Y:N)</td>
<td>12:34</td>
<td>54:223</td>
<td>1.05</td>
<td>0.30</td>
</tr>
<tr>
<td>Current mood disorder DX (Y:N)</td>
<td>15:31</td>
<td>69:208</td>
<td>1.22</td>
<td>0.27</td>
</tr>
<tr>
<td>Current PTSD DX</td>
<td>37:9</td>
<td>160:117</td>
<td>8.52</td>
<td>0.004</td>
</tr>
<tr>
<td>CAPS total score</td>
<td>60.02 (28.96)</td>
<td>48.31 (28.95)</td>
<td>2.54</td>
<td>0.012</td>
</tr>
</tbody>
</table>

SUD+, participants who met criteria for a diagnosis of a substance use disorder; SUD−, participants who did not meet criteria for a substance use disorder diagnosis; No. Lifetime mTBI, number of lifetime mild traumatic brain injuries; Current anxiety disorder DX (Y:N), current anxiety disorder diagnosis; Current mood disorder DX (Y:N), current mood disorder diagnosis; current PTSD DX, current posttraumatic stress disorder diagnosis; CAPS total score, total score on the clinician-administered PTSD scale; yr, year.

symptom severity (CAPS score, r = -0.33, p = 0.026), but the SUD− group showed no association between commission accuracy and CAPS scores (r = -0.09, p = 0.12). There was a main effect of education [F(1,315) = 7.15, p = 0.008, partial eta square = 0.022], but not age, current major depressive disorder, or lifetime history of mTBI (all p’s > 0.4).

Omission Accuracy
Like the commission errors, there was no main effect of valence, and all of the interactions involving valence were also not significant (all p’s > 0.23). The main effects of SUD group [F(1,315) = 4.80, p = 0.03; partial eta square = 0.02] and CAPS scores [F(1,315) = 5.11, p = 0.02; partial eta square = 0.016] and the interaction between SUD group and CAPS scores [F(1,315) = 7.560, p = 0.006, partial eta square = 0.023] were significant. As shown in Figure 1B, within the SUD+ group, omission accuracy decreased with increasing PTSD symptom severity (CAPS score, r = -0.39, p = 0.007), whereas in the SUD− group, there was no association between omission errors and CAPS scores (r = 0.01, p = 0.81). There was a main effect of education [F(1,316) = 9.75, p = 0.002, partial eta square = 0.029], but not age, current depressive disorder, or history of lifetime mTBI (all p’s > 0.25).

d’ Prime
There was no main effect of valence, and all of the interactions involving valence were not significant (all p’s > 0.28). The main effect of SUD Group [F(1,315) = 2.47, p = 0.03, partial eta square = 0.008] was not significant. CAPS scores [F(1,315) = 5.7, p < 0.02, partial eta square = 0.02] and the interaction between SUD group and CAPS scores [F(1,315) = 6.00, p = 0.015; partial eta square = 0.019] were significant. Within the SUD+ group, d’ (an index of overall discriminability) decreased with increasing PTSD symptom severity (CAPS scores, r = -0.38, p = 0.009), whereas in the SUD− group, there was no association between overall stimulus discriminability and CAPS scores (r = -0.05, p > 0.32). There was a main effect of education [F(1,315) = 12.45, p < 0.001, partial eta square = 0.038], but not age, current depressive disorder, or lifetime number of mTBI (p’s > 0.5).

Criterion
There was no main effect of valence or any of the interactions involving valence (all p’s > 0.25). The main effect of SUD group [F(1,315) = 2.17, p = 0.14, partial eta square < 0.01] or CAPS scores [F(1,315) = 0.12, p = 0.73, partial eta square < 0.001] and the interaction of SUD group and CAPS scores [F(1,315) = 1.77, p = 0.19; partial eta square = 0.006] were not significant. None of the covariate terms were significant (all p’s > 0.18).

DISCUSSION
This study investigated the independent and interactive effects of PTSD symptom severity and SUD on affective attentional processing. Veterans with and without SUD who were deployed to Iraq or Afghanistan demonstrated differential associations between PTSD symptoms and responding using an affective go/no-go task (Robbins et al., 1994; Robbins et al., 1998). Among SUD+ individuals, commission and omission accuracy on the go/no-go task decreased as PTSD symptom severity increased. Similarly, among SUD+ participants, go/no-go task discriminability (as indexed by d’ score) decreased with more severe PTSD symptoms. Conversely, there was no significant association between PTSD symptom severity and commission accuracy, omission accuracy, or discriminability in the SUD− group.

Indices of attentional/inhibitory control were negatively associated with PTSD severity. Consistent with prior research demonstrating cognitive dysfunction in PTSD (e.g., Gillie and Thayer, 2014; DeGutis et al., 2015; Esterman et al., 2013; Falconer et al., 2008; Leskin and White, 2007; Vasterling et al., 1998), there was a main effect of PTSD symptom severity on errors of commission and omission. Furthermore, for both types of attentional errors, accuracy declined with increasing PTSD symptom severity among the SUD+ group. In terms of omission accuracy, this could indicate lapses in attention or vigilance with increasing PTSD severity, which may then lead to lack of responding among individuals with SUD (e.g., Wright et al., 2014). In terms of commission accuracy, there was a significant negative association between PTSD symptom severity and commission accuracy in the SUD+, but not the SUD− group. Taken together, these findings do suggest an enhanced relation between PTSD symptoms and attentional/inhibitory control deficits when SUD, a disorder marked by impaired inhibitory control (Smith et al., 2014), co-occurs.

Within the SUD+ group, discriminability declined with increasing PTSD severity suggesting that, regardless of stimulus valence, the PTSD × SUD interaction resulted in disrupted attentional control or greater difficulty with identification/discrimination of emotional stimuli. This latter interpretation is supported by findings showing deficits in emotion recognition among individuals with SUD (e.g., Castellano et al., 2015) and may suggest that deficits in this domain contribute to observed PTSD × SUD interactions in both omission and commission accuracy.

Contrary to hypotheses, stimulus valence did not alter performance on any index of the affective go/no-go task, suggesting that...
effects may be due to general deficits in cognitive control and inhibitory function, rather than affective attentional bias among individuals with PTSD and SUD (Aupperle et al., 2012; Conrod and Nikolaou, 2016). The lack of an association between PTSD symptom severity and response to negatively valenced targets in this study adds to inconsistent findings in this literature, offering further support for studies that have failed to find evidence of affective attentional bias in PTSD (Bryant and Harvey, 1997; Cisler et al., 2011; Elsesser et al., 2005; Wittekind et al., 2015). Of note, using the identical task, we previously did not observe an association between affective attentional control and PTSD symptom severity, when mTBI was not comorbid (Amick et al., 2013).

Overall, study results extend previous literature showing individual effects of PTSD and SUD symptoms on disrupted behavioral responding using tasks requiring attentional control (e.g., Field and Cox, 2008; Lambe et al., 2014; Esterman et al., 2013; DeGutis et al., 2015). It is possible that the affective go/no-go task, with its specific demands upon inhibitory processes in addition to affective information processing may be more sensitive to the subtle effects of co-occurring SUD on attention and inhibitory control. Thus, findings may point to clinically relevant alterations in inhibitory control and attention function in response to emotionally valenced stimuli that may be unique to individuals with both disorders.

Despite the novel and significant contributions of the current investigation, several limitations should be noted. This study did not include biochemical verification of substance abstinence and instead relied on self-report and clinician assessment of substance free status before testing. In addition, in this preliminary research, all SUDs were collapsed into a single category (SUD+ versus SUD−) rather than examining the individual or dimensional contributions of each type of SUD. Future studies with larger sample sizes should investigate the effects of different SUDs on the relation between PTSD and attentional processing. This may be particularly important given results suggesting variable effects of SUD on attentional processing among individuals with alcohol, nicotine, heroin, and cocaine use disorders (e.g., Rzetelny et al., 2008; Lambe et al., 2014). In addition, future research should explore more complex, three-way interactions among PTSD and co-occurring conditions with established effects on attentional bias (e.g., mTBI; Amick et al., 2013), such as depression, anxiety, and sleep disorders. The study sample was primarily composed of male veterans. Thus, caution should be used when generalizing findings to nonveteran or female populations. Lastly, future investigations should incorporate different attentional bias/inhibitory control tasks to better elucidate the influence of PTSD and SUD on affective attentional bias specifically as opposed to attention and inhibitory control more broadly. These investigations should include investigation of varied stimulus type (verbal versus visual affective stimuli) and task design/methods to more comprehensively characterize the nature of interactive effects of this comorbidity on affective attentional bias.

In conclusion, existing research supports individual effects of SUD and PTSD on affective attentional processing and inhibitory function (e.g., Aupperle et al., 2012; Buckley et al., 2000; Conrod and Nikolaou, 2016; Drobes et al., 2006; Pineles et al., 2006). Results from this initial investigation highlight the need to consider the impact of SUD and other commonly co-occurring disorders when examining relationships between PTSD and affective and nonaffective attentional processing, cognitive control, and behavioral responding to more accurately identify the nature of each disorders’ unique and interactive effects on these processes (Amick et al., 2013; Samuelson et al., 2006).

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DISCLOSURE

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