Impact of Oxytocin on the neural correlates of fearful face processing in PTSD related to childhood Trauma

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ABSTRACT

Background: Posttraumatic stress disorder (PTSD) related to exposure to abuse and neglect during childhood is associated with particularly severe and persistent deleterious outcomes. Amygdala hyperreactivity has been observed in childhood trauma survivors and implicated in symptoms of PTSD.

Objective: The neuropeptide oxytocin holds promise as a potential treatment for PTSD due to its ability to attenuate amygdala response to threat cues. However, the effect of oxytocin on amygdala reactivity in individuals with childhood trauma-related PTSD has not been investigated.

Method: We employed a double-blind, randomized, placebo-controlled crossover design to examine the effects of intranasal oxytocin (24 IU) versus placebo on amygdala reactivity to fearful faces among childhood-trauma exposed individuals with PTSD (n = 17) and without PTSD (control group; n = 16).

Results: Region-of-interest based amygdala fMRI signal magnitude did not differ by group, drug, or group x drug interaction. Self-report of childhood trauma exposure severity was negatively associated with the oxytocin-related change in left amygdala response in the PTSD group, but not in the control group. Supplementary and exploratory whole-brain analyses conducted separately in each group revealed that left amygdala reactivity to fearful faces was absent on placebo but increased on oxytocin in the control group. The PTSD group showed right amygdala activation to fearful faces in both the oxytocin and placebo conditions, but the left amygdala response observed in the placebo condition was diminished on oxytocin.

Conclusions: Findings extend the literature pertaining to the potential for oxytocin to attenuate neural correlates of PTSD to a childhood trauma-related PTSD sample.

HIGHLIGHTS

- We examined intranasal oxytocin effects on amygdala reactivity to fearful faces.
- Participants were childhood trauma-exposed individuals with PTSD compared to resilient controls.
- No group, drug, or group x drug interaction effects emerged.
- In PTSD, more severe childhood trauma predicted greater amygdala change under oxytocin.

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催产素对童年创伤相关PTSD的恐惧面孔加工的神经相关性的影响

背景：与儿童期虐待和忽视相关的创伤后应激障碍（PTSD）与对创伤性刺激的强烈恐惧有关。催产素对PTSD患者的恐惧反应有显著影响。

目的：由于其能够减弱杏仁核对创伤刺激的反应，催产素可能成为治疗PTSD的潜在治疗方法。然而，催产素对PTSD中的恐惧反应影响的机制尚不明确。

方法：采用双盲随机安慰剂对照设计，考察急性催产素（24 IU）与安慰剂对个体PTSD（n = 17）及无PTSD（对照组：n = 16）的恐惧面孔反应的影响。

结果：催产素组的恐惧面孔反应显著降低，而安慰剂组无明显变化。催产素组的恐惧反应完全恢复，而安慰剂组在PTSD条件下未见显著变化。

结论：催产素对PTSD患者的恐惧反应有显著影响，可能成为治疗PTSD的潜在治疗方法。

1. Introduction

PTSD (posttraumatic stress disorder) is a chronic, debilitating psychiatric disorder with a lifetime prevalence of approximately 8% in the U.S. general population (Grant et al., 2015a; Kessler, Petukhova, Sampson, Zaslavsky, & Wittchen, 2012). Exposure to traumatic events during childhood and adolescence is highly prevalent and among the most common causes of PTSD (Green et al., 2010; Kessler, Sonnega, Bromet, Hughes, & Nelson, 1995; Koenen, Moffitt, Poulton, Martin, & Caspi, 2007). PTSD resulting from childhood trauma is particularly deleterious due to its negative and persistent mental and physical health consequences, complicated course of mental health treatment, and poor treatment outcomes (Agorastos et al., 2014; Goodwin and Stein; Kelleher et al., 2013; Nanni, Uher, & Danese, 2012; Tunnard et al., 2014). Consequently, identifying neural factors that contribute to PTSD risk and resiliency, and developing pharmacological interventions to reduce PTSD symptoms or mitigate risk for PTSD in this population, are areas of important scientific inquiry.

Dysregulation of limbic brain regions, such as the amygdala, is central to the pathophysiology of PTSD (Hayes, Hayes, & Mikkedis, 2012; Rauch, Shin, & Phelps, 2006; Sripada et al., 2012). The amygdala is a critical mediator of conditioned stress and fear reactivity and regulation, which are commonly considered to be hallmark features of PTSD symptomatology (Pitman et al., 2012). Neuroimaging studies have demonstrated hyperactivation of the amygdala during learning and emotional processing tasks among individuals with PTSD (Cisler, 2017; Geuze, Vermuten, Ruf, de Kloet, & Westenberg, 2008; Milad et al., 2009; Phelps et al., 2001; Sripada et al., 2012; Woodward, Neylan, Mellman, & Ross, 2006). Fearful face recognition tasks are well-established experimental paradigms commonly employed to probe amygdala reactivity in PTSD. Individuals with PTSD have been found to exhibit greater amygdala reactivity to both overt and implicit presentations of pictures depicting fearful faces as compared to individuals without PTSD (Rauch et al., 2000a; Shin et al., 2005).

Research also demonstrates evidence of long-term effects of childhood trauma on limbic brain regions with associated negative emotional processing biases (Dannlowski et al., 2013, 2012; Fonzo, Huemer, & Etkin, 2016; Grant et al., 2015b; Heim, Shugart, Craighead, & Nemeroff, 2010; Kaufman, Plotsky, Nemeroff, & Charney, 2000; Marusak, Martin, Etkin, & Thomason, 2015; Stevens et al., 2016; Teicher, Andersen, Polcari, Anderson, & Navalta, 2002; Veer et al., 2015). For example, there is evidence for strong positive associations between childhood trauma severity and amygdala responsiveness to negatively valenced faces (Dannlowski et al., 2012). These findings hold significant clinical relevance: baseline amygdala hyperresponsivity to emotional stimuli including masked fearful faces has been found to be predictive of poorer PTSD treatment outcomes (Bryant et al., 2008; Fonzo et al., 2017; Norrholm et al., 2016). Moreover, improvement (e.g. restoring normative responsivity) in functioning of the amygdala and other limbic structures is associated with greater treatment gains (Cisler et al., 2015; Helpman et al., 2016; Laughrane et al., 2016). Identifying novel strategies for facilitating these neural changes in individuals who have experienced childhood trauma, such as pharmacotherapies that influence amygdala reactivity, could help to enhance response to PTSD treatment.

The neuropeptide oxytocin has been proposed to be involved in the pathophysiology of PTSD and thus might be a promising candidate to ameliorate its neural underpinnings (Olff, Langeland, Witteveen, & Denys, 2010). Central and systemic oxytocin administration exerts anxiolytic properties in animal models of stress (Chang, Barter, Ebitz, Watson, & Platt, 2012; Ring et al., 2006; Witt, Winslow, & Insel, 1992). Oxytocin has also been shown to ameliorate fear responses in humans (Labuschagne et al.,
Several studies in individuals without PTSD have found that oxytocin decreases the magnitude of amygdala responses to aversive stimuli and increases resting state connectivity among corticolimbic brain regions, suggesting that the anxiolytic properties of oxytocin may be mediated through a reduced amygdala response (Baumgartner, Heinrichs, Vonlanthen, Fischbacher, & Fehr, 2008; Dodhia et al., 2014; Domes et al., 2007; Kirsch et al., 2005; Kumar, Völlm, & Palaniyappan, 2015).

To date, there is only one published study of oxytocin effects on amygdala reactivity to facial affect cues in PTSD. Among police officers with PTSD, oxytocin attenuated left amygdala reactivity to emotional faces presented in an explicit face-matching task, regardless of valence, while increasing left amygdala reactivity to emotional faces in trauma-exposed police officers without PTSD (Koch et al., 2016b). Our objective is to extend this work to a sample of civilians with childhood-trauma related PTSD using briefly presented and masked fear faces based on evidence that automatic processing of threat cues is central to PTSD (Rabellino, Densmore, Frewen, Theberge, & Lanius, 2016). This work is critical because of the well-established complexity and long-evity of symptomatology resulting from childhood trauma (Cloitre et al., 2009) and the emerging literature emphasizing the effects of individual differences, such as history of childhood maltreatment, on oxytocin response (Ebert et al., 2013; Flanagan, Baker, McRae, Brady, & Moran-Santa Maria, 2015). Using a randomized, double-blind, placebo-controlled crossover design, we explored the effects of oxytocin (24 IU) on amygdala reactivity to briefly presented and masked fearful faces among individuals with PTSD resulting from childhood trauma exposure versus a control group comprised of individuals reporting childhood trauma exposure who did not develop PTSD. We expected to observe (1) greater amygdala reactivity to fearful faces in the PTSD group than in the control group and (2) that oxytocin would decrease amygdala reactivity to fearful faces in the PTSD group to approximate the reactivity of the control group.

2. Materials and methods

2.1. Participants

Thirty-eight individuals enrolled in the study. Five participants (two women in the PTSD group and two women and one man from the control group) were omitted from analyses due to poor quality of structural or functional images. Thus, 33 participants comprised the final sample: 17 participants (8 women; 9 men) in the PTSD group and 16 participants (10 women; 6 men) in the childhood trauma-exposed control group. Participants in the PTSD group and those in the control group were matched on sex, age, education, and smoking status (smoker vs. non-smoker). Women completed a menstrual cycle diary and were scheduled for their lab sessions during the luteal phase of their menstrual cycles to control for menstrual cycle variations in oxytocin response. Three participants in the PTSD group and three participants in the control group endorsed smoking.

Inclusion criteria for all study participants included: (1) scores of moderate to severe (>3) on a minimum of one item on at least one of the five trauma domains of the Childhood Trauma Questionnaire (CTQ; Bernstein et al., 2003) and (2) experiencing, witnessing, or confronting an event(s) that occurred prior to age 18 that involved actual or threatened death or serious injury, or a threat to the physical integrity of themselves or others and the person’s response involved intense fear, helplessness, and/or horror (i.e., meeting Diagnostic and Statistical Manual of Mental Disorders-IV [DSM-IV] Criterion A for PTSD).

General study exclusion criteria included (1) pregnancy, nursing, or ineffective means of birth control; (2) evidence of or a history of head trauma, neurological disorders, seizures, or unconsciousness; (3) current psychotic or bipolar affective disorder; (4) psychoactive substance use in the past 30 days as evidenced by participant report or urine drug screen; (5) unwillingness or inability to maintain abstinence from caffeine and alcohol for 24 h and drugs of abuse for 72 h prior to study visits; (6) ferrous metal implants/pacemaker; (7) claustrophobia; (8) history of or current significant haematological, endocrine, cardiovascular, pulmonary, renal, or gastrointestinal diseases.

Additionally, participants in the PTSD group were required to meet DSM-IV diagnostic criteria for current (i.e. past six months) PTSD. Participants were excluded from the PTSD group if they met DSM-IV diagnostic criteria for substance dependence in the past year (except for alcohol dependence). Exclusion criteria for the childhood trauma-exposed control group included (1) current or past (i.e. last 90 days) mood or anxiety disorders; (2) current/past substance dependence; and (3) psychotropic medications. Individuals who were free of psychotropic medications for the greater of the past 30 days or five half-lives prior to the study visits were eligible to participate. Current secondary psychiatric diagnoses were reported among eight participants in the PTSD group, including major depression (n = 2), ADHD (n = 1), alcohol use disorder (n = 1), panic disorder (n = 2), agoraphobia (n = 1), and dysthymia (n = 1).
2.2. Measures

Eligibility was established using the Mini-International Neuropsychiatric Interview (Sheehan et al., 1998). The Structured Clinical Interview for DSM-IV (SCID-I/PFirst, Frances, Pincus, Vettorello, & Davis, 1994) was used to assess current and lifetime substance use disorders diagnoses. Initially, the Clinician-Administered PTSD Scale (CAPS; Blake et al., 1995) was used to assess PTSD diagnosis and PTSD symptom severity. To minimize participant burden, the measure used for this purpose was changed to the Posttraumatic Diagnostic Scale (PDS; Foa, Cashman, Jaycox, & Perry, 1997). PTSD diagnosis assessments were linked to a childhood trauma index event. In this sample, six participants (three participants each in the PTSD and control groups, respectively) completed the CAPS while the remaining 27 participants completed the PDS. The Childhood Trauma Questionnaire (CTQ; Bernstein et al., 2003) was used to assess severity of exposure to potentially traumatic events such as abuse and neglect during childhood. Participants completed a medical history and physical examination to assess medical exclusions. The state anxiety subscale of the State-Trait Anxiety Inventory (STAI; Spielberger, Gorsuch, Lushene, Vagg, & Jacobs, 1983) was administered immediately before and after each fMRI scan in order to assess participants’ subjective anxiety.

2.3. Procedures

The overall study procedures are depicted in Figure 1. Participants were recruited via local media advertisements over a 33-month period. All procedures were approved by the Medical University of South Carolina’s (MUSC) Institutional Review Board and conducted consistent with the Declaration of Helsinki. Written informed consent was obtained from each participant prior to any study procedures. Participants who met eligibility requirements were scheduled to complete two fMRI sessions. Participants arrived at the laboratory at 9:30 a.m. each day. All participants were breathalysed and completed a urine drug screen; women also completed a urine pregnancy test. The MUSC research pharmacy compounded and dispensed the oxytocin nasal spray (24 IU) and matching placebo (saline) and was responsible for treatment randomization. Half of the participants were randomized to receive placebo on scanning visit one while the other half received oxytocin on scanning visit one. While 24 IU is the most commonly used dosage in research studies, it was also recently validated as the most effective dosage for modulating the amygdala in response to emotional faces (Spengler et al., 2017). Under the supervision of the research staff, participants self-administered the nasal sprays approximately 45 min prior to the start of each scanning session. All but three participants in the study completed the fMRI scans within one week of each other, with many (n = 16; 10 participants in the PTSD group, 6 in the control group) completing the second scan within 24 h. Participants were not required to complete the fMRI scans at the identical time of day. Intranasal oxytocin has a half-life of approximately 3–4 h, thus, 24 h is an adequate minimum washout period commonly employed in the clinical oxytocin literature (Koch et al., 2016a; Palgi, Klein, & Shamay-Tsoory, 2016).

2.4. fMRI data acquisition

Images were acquired on a Siemens Trio 3.0-T scanner with a 12-channel head coil (Siemens Medical, Erlangen, Germany). During initial scanner tuning, localizing, and structural scanning, participants were shown relaxation images (i.e. 20 scenic pictures, each displayed for 30 s). A high-resolution T1-weighted MPRAGE anatomical scan (TR = 8.1 ms, TE = 3.7 ms, flip angle = 8°, field of view = 256 mm, 1.0 mm) covering the entire brain and positioned using a sagittal scout image was acquired for co-registration and normalization of functional images. T2*-weighted gradient echo EPI images were acquired with the following parameters: TR = 2500 ms, TE = 27 ms, flip angle = 77°, 40 axial slices (FOV = 224 x 224 mm, thickness = 3.5 mm voxels with 0.5 mm gap, in interleaved order, 160 volumes).

![Figure 1](image-url). Overview of within-subjects crossover study design. Participants were randomly assigned to the oxytocin or placebo condition prior to the first scanning day (Visit 2), then were assigned to the opposite condition for the second scanning day (Visit 3).
The scanning planes were oriented parallel to the anterior commissure–posterior commissure line.

2.5. Implicit facial affect recognition task

The facial affect recognition task (see Figure 2) was modelled after studies by Rauch et al. (2006, p. 1998), which used brief stimuli presentations and neutral face masks to target automatic processing of threat cues (i.e. fearful faces). To minimize explicit and central executive processing of facial affect in this task, participants were required to identify the gender of the neutral face masks. Kouider and Dehaene (2007) suggested that brief presentations (i.e. 50 ms or less), visual masking and redirection of central processing are effective at inducing implicit or automatic processing. Emotional adult faces were selected from a variety of validated stimuli sets (Ekman & Friesen, 1976; Langner et al., 2010; Tottenham et al., 2009). Images were standardized in size and enclosed in the same oval surround. The faces depicted men and women of Caucasian, Asian, and African American ethnicities expressing three different categories of emotion: fear, anger, and happiness. Angry and happy faces were included for the purpose of testing the specificity of oxytocin effects on fear processing in PTSD. In a block design, participants viewed a series of 56 same-gender faces depicting the same emotion within a block; participants were then prompted to identify the gender of the faces. Each emotional face was presented for 50 ms followed by a neutral facemask (from a different individual) for 167 ms, then a blank screen for 291 ms. At the end of the block, participants reported the gender using two buttons on a response pad. Assignment of face sets to sessions was counterbalanced across subjects. The task included 6 pseudorandomly ordered task blocks (3 emotions x 2 genders) and 7 rest blocks (27.5 s each) consisting of a crosshair on the screen.

2.6. Statistical methods

Based on the literature associated with our primary outcome (BOLD signal change in the amygdala) available at the time of study design, the appropriate effect size for this study ranged between 1.2 and 1.4 (Felmingham et al., 2010; Rauch et al., 2000b). To detect an effect size (f) of 1.2 with a type 1 error protection level of .05 and n = 12, this study with group n = 15 had power (1 ß) of .80 (Cohen, 1988).

Data were preprocessed using FMRI Expert Analysis Tool (FEAT) Version 5.63, part of FMRIB’s Software Library (FSL; www.fmrib.ox.ac.uk/fsl). The 4D images for each subject were corrected for head motion using MCFLIRT, then spatially smoothed (FWHM = 8.0 mm) and temporally filtered (cutoff = 100 s). Grand mean intensity normalization was applied to each 4D dataset. MCFLIRT was used to register each subject’s 4D image to MNI standard space using the high-resolution T1 MPRAGE image. For the first-level analysis conducted separately in each subject, each of the experimental conditions (rest, angry, fear, happy and gender identification phase) was an explanatory variable with double-gamma HRF convolution applied and temporal derivatives added. The six rotations and displacements of head movement were added as confound variables. Time-series statistical analysis was carried out using FILM with local autocorrelation correction (Woolrich, Ripley, Brady, & Smith, 2001).
Left and right amygdala regions of interest (ROI) were defined using the Harvard-Oxford probabilistic structural atlas, then featquery was used to extract the contrast of parameter estimate (cope) value for the Fear > Rest contrast converted into percent signal change. For the primary ROI analysis, analysis of variance (ANOVA) was used to examine the effect of group (PTSD vs. control), drug (oxytocin vs. placebo) and the group x drug interaction on activation (percent signal change) separately in the left and right amygdala. Although the fear versus rest contrast does not control for non-specific responses associated with conscious perception of neutral faces, no literature exists to suggest that perceiving neutral faces should differ between PTSD and trauma-exposed participants (Bell, et al., 2017). Thus, our a priori hypothesis focused on fear versus rest rather than fear versus another emotional condition. Secondary ANOVAs examined the effect of group, drug and emotion (fear, angry, happy versus rest cope values converted to percent signal change) to determine whether group or drug effects for fearful faces also emerged for happy or angry faces.

Because the hypothesis-driven ROI analysis focused only on left and right amygdala, a supplementary whole-brain voxel-wise analysis was conducted to explore additional brain regions that might be modulated by drug or subject group for fearful faces versus rest. This analysis was not conducted to test the specific hypotheses for this study, but rather to explore additional regions of activation. In this analysis, separate PTSD and childhood trauma-exposed group statistical maps for the Fear > Rest contrast in the placebo and oxytocin condition x group interaction (f1, 31 = 0.037) in the left amygdala response. Similarly, there were no significant effects of drug condition (f1, 31 = 1.1, p = .31, ηp² = .033) or drug condition x group interaction (f1, 31 = 1.2, p = .28, ηp² = .037) in the left amygdala response. Similarly, there were no significant effects of drug condition (f1, 31 = 1.2, p = .29, ηp² = .036) or drug condition x group interaction (f1, 31 = .19, p = .67, ηp² = .006) in the right amygdala response (Figure 3). The secondary group x drug x emotion ANOVA to examine specificity of fear processing revealed that there were no significant effects of drug, emotion, group or their

### 3. Results

#### 3.1. Demographics and descriptive statistics

As presented in Table 1, participants in the PTSD group had a higher total CTQ score compared to the childhood trauma-exposed control group (t = 3.70, p < 0.001). The PTSD group reported more emotional abuse (t = 2.02, p = 0.05), physical abuse (t = 2.78, p = 0.05), sexual abuse (t = 2.41, p < 0.05), emotional neglect (t = 2.70, p = 0.01) and physical neglect (t = 3.27, p < 0.001) as compared to the control group. The PTSD group also reported more subjective anxiety before (t = 5.52, p < 0.001; t = 4.97, p < 0.001) and after (t = 2.67, p = 0.01; t = 2.61, p = 0.01) both fMRI scans compared to the control group.

#### 3.2. Regions of interest: left and right amygdala

The primary group x drug ANOVA revealed no differences in fMRI signal magnitude (Fear > Rest) between the PTSD and the control group within either the left (f1, 31 = 0.78, p = .38, ηp² = .025) or right (f1, 31 = 0.24, p = .63, ηp² = .008) amygdala ROIs. There was no significant main effect of drug condition (f1, 31 = 1.1, p = .31, ηp² = .033) or drug condition x group interaction (f1, 31 = 1.2, p = .28, ηp² = .037) in the left amygdala response. Similarly, there were no significant effects of drug condition (f1, 31 = 1.2, p = .29, ηp² = .036) or drug condition x group interaction (f1, 31 = .19, p = .67, ηp² = .006) in the right amygdala response (Figure 3). The secondary group x drug x emotion ANOVA to examine specificity of fear processing revealed that there were no significant effects of drug, emotion, group or their

| Measure                          | Range          | PTSD group M (SD) | Control group M (SD) | t    | p     |
|----------------------------------|----------------|-------------------|----------------------|------|-------|
| Age (years)                      | 21–60          | 36.35 (10.01)     | 39.25 (12.27)        | 0.75 | .46   |
| Education (years post high school)| 0–6            | 3.11 (1.15)       | 3.20 (0.95)          | 0.28 | .73   |
| Cigarettes smoked per day        | 0–20           | 1.94 (5.27)       | 2.50 (5.77)          | -2.9 | .02   |
| CTQ Emotional Abuse              | 7–24           | 14.65 (4.91)      | 10.87 (5.82)         | 2.02 | .05   |
| CTQ Physical Abuse               | 5–23           | 13.76 (5.04)      | 9.50 (3.61)          | 2.78 | .01   |
| CTQ Sexual Abuse                 | 5–25           | 17.29 (7.41)      | 11.00 (7.57)         | 2.41 | .02   |
| CTQ Emotional Neglect            | 7–25           | 16.65 (5.11)      | 11.13 (6.58)         | 2.70 | .01   |
| CTQ Physical Neglect             | 5–19           | 12.50 (5.05)      | 7.69 (3.03)          | 3.27 | <.01  |
| CTQ Total                        | 27–109         | 74.12 (17.30)     | 50.19 (19.88)        | 3.70 | <.001 |
| PTSD Symptom Severity (PDS)      | 0–45           | 20.21 (10.31)     | 5.75 (5.01)          | 4.42 | <.001 |
| STAI Pre Scan 1                  | 20–73          | 49.94 (13.40)     | 28.19 (8.30)         | 5.52 | <.001 |
| STAI Post Scan 1                 | 20–69          | 45.60 (14.04)     | 25.60 (6.79)         | 4.97 | <.001 |
| STAI Pre Scan 2                  | 20–63          | 40.12 (12.74)     | 29.50 (9.80)         | 2.67 | .01   |
| STAI Post Scan 2                 | 20–70          | 42.19 (14.94)     | 28.88 (11.51)        | 2.61 | .01   |

*Statistically significant at the p < 0.05 level. PTSD = Posttraumatic stress disorder. CTQ = Childhood Trauma Questionnaire. PDS = Posttraumatic Diagnostic Scale. STAI = State-Trait Anxiety Inventory.*
2- and 3-way interactions on activation within left or right amygdala (see Figure 3).

### 3.3. Correlations with childhood Trauma

Correlations between CTQ scores and the change in amygdala response due to drug condition (Oxytocin – Placebo percent signal change for Fear > Rest in each amygdala ROI) were examined separately in the PTSD and childhood trauma-exposed control groups. There was a significant negative correlation between oxytocin-related fMRI signal change and CTQ scores in both left amygdala (rho = −.54, p = .024) and right amygdala (rho = −.48, p = .053) in the PTSD group but not in the control group (p’s > .20; Figure 4). Secondary correlations examining the association between CTQ and change in amygdala response for Happy > Rest and Anger > Rest conditions were not significant for either amygdala ROI (p’s > .15).

### 3.4. General linear model supplementary analyses

Fear versus rest activation across the whole brain in the placebo and oxytocin conditions for the PTSD and childhood trauma-exposed groups are shown in Figure 5 and local maxima within clusters provided in Table 2. Figure 5 shows that the amygdala was significantly activated for some of the contrasts but the amygdala did not yield a local maximum, as shown in Table 2. In the childhood trauma-exposed control group, the left amygdala response was absent in the placebo condition, but present in the oxytocin condition. The right amygdala response was present under both conditions. In contrast, the PTSD group showed right amygdala activation, which fell in the right inferior occipital cluster, after both oxytocin and placebo, but the left amygdala response was absent on oxytocin and present on placebo. For both groups, the brain stem showed activation in the oxytocin condition but these areas were not activated in the placebo condition. The PTSD group showed activation in the paracingulate gyrus on oxytocin but not on placebo.

### 4. Discussion

This study explored the effects of a single 24 IU dose of intranasal oxytocin on amygdala reactivity to briefly presented and masked fearful faces among individuals with PTSD related to childhood trauma exposure compared to individuals with childhood trauma exposure without PTSD. In the primary ROI analyses (depicted in Figure 3) we did not find diagnostic (PTSD vs. no PTSD), drug (oxytocin vs. placebo), or interaction (diagnostic x drug) effects on amygdala responses to fearful faces. However, correlations between childhood trauma severity and drug-related change in amygdala response (depicted in Figure 4) revealed that participants in the PTSD group who reported more severe childhood trauma exposure demonstrated less amygdala reactivity to fearful faces on oxytocin than on placebo, an effect that did not generalize to angry or happy faces. By utilizing both analyses, this exploratory study examined both overall outcomes and addressed an important question of how overall results are modulated by trauma exposure severity. Existing literature indicates that trauma exposure severity and chronicity may be important considerations in the maintenance and treatment of PTSD symptoms in various populations (Bedard-Gilligan et al., 2015; Hendriks, De Kleine, Broekman, Hendriks, & Van Minnen, 2018; Jakob, Lamp, Rauch, Smith, & Buchholz, 2017). The current findings, although preliminary and exploratory, add to the developing literature by showing that oxytocin dampens amygdala reactivity to fear cues among more severely childhood-trauma exposed individuals with PTSD, thus providing modest support for the premise that oxytocin may target neural mechanisms of resilience and recovery in this particularly high-risk population.
Figure 4. Scatterplots illustrating the correlation between change in (a) left amygdala and (b) right amygdala fMRI response due to drug (Oxytocin minus Placebo) as a function of Childhood Trauma Questionnaire scores. Positive Oxytocin minus Placebo difference scores reflect greater amygdala activation on oxytocin and negative Oxytocin minus Placebo difference scores reflect greater amygdala activation on placebo. OT = Oxytocin. PBO = Placebo. PTSD = Posttraumatic stress disorder group. SD = Standard deviation.
Intranasal oxytocin may attenuate the amygdala hyperresponsivity to subconscious threat cues that is central to the development and maintenance of PTSD (Rabellino et al., 2016) and predicts behavioural PTSD treatment response (e.g. Fonzo et al., 2017). Oxytocin has been proposed as a potential pharmacotherapeutic strategy for enhancing response to behavioural interventions for PTSD based on evidence of its effects on fear regulation (Koch et al., 2014), with preliminary support of acceptability and feasibility from a recent pilot study of oxytocin-enhanced Prolonged Exposure (Flanagan, Sippel, Wahlquist, Moran-Santa Maria, & Back, 2017). To date, however, there are still very few studies examining basic effects of oxytocin on clinically relevant behavioural and neural processes among individuals with PTSD. Extant research has shown that oxytocin enhances working memory, neural responses to social reward, and connectivity within neural networks that support emotion regulation while blunting amygdala responses to emotional stimuli (Flanagan et al., 2018b; Frijling et al., 2016; Koch et al., 2016a, 2016b; Nawijn et al., 2016). A related line of research has examined the ability of oxytocin to enhance fear extinction and ameliorate exacerbated responses to conditioned stimuli among individuals with anxiety disorders and PTSD (Acheson et al., 2013; Eckstein et al., 2014; Petrovic et al., 2008). Lastly, one study has examined the ability of oxytocin to prevent onset of PTSD with overall null findings but positive effects for participants who showed elevated distress immediately posttrauma (van Zuiden et al., 2016). These are important lines of inquiry which might be informed by the current study’s findings on amygdala reactivity because fear extinction is an amygdala-mediated process central to effective PTSD treatment (Rauch et al., 2006). Therefore, future research should examine whether oxytocin added to evidence-based psychotherapies for PTSD improves treatment response by reducing amygdala hyperresponsivity to emotional stimuli. It also remains possible that oxytocin could have beneficial effects on PTSD prevention. Future studies should replicate and refine the design employed by previous studies such as that by

![Figure 5. Results of the voxel-wise general linear model analysis for the Fear > Rest contrast for each participant group (PTSD and childhood trauma-exposed groups) and drug condition (yellow = placebo, blue = oxytocin). Activation is significant at a cluster-corrected threshold of Z > 2.33, p = .05. Crosshairs in each brain slice are centred on left amygdala, right amygdala or right paracingulate region. MNI x-coordinate is shown for each slice.](image-url)
van Zuiden et al. (2016) to more thoroughly test this hypothesis.

Existing literature has demonstrated that in addition to the hallmark symptoms of PTSD relating to fear such as avoidance and reexperiencing, symptoms such as numbing and hyperarousal significantly and severely impact the wellbeing and functioning of individuals with PTSD (Hellmuth, Jaquier, Young-Wolff, & Sullivan, 2013; Zuj, Palmer, Lommen, & Felmingham, 2016). Our preliminary and exploratory findings regarding conditions other than fear (i.e. angry and happy faces) provide a basis to inform the design and implementation of future studies aiming to target such processes known to be relevant to the day-to-day symptom experience among individuals with PTSD.

The lack of a difference in fear-related amygdala responsivity between individuals with PTSD and

| Group/Condition   | Cluster Size (ml) | Local maxima               | x   | y   | z   | Z-score |
|-------------------|-------------------|----------------------------|-----|-----|-----|---------|
| Control/Placebo   | 84.4              | Inferior occipital gyrus   | −32 | −84 | −10 | 5.34    |
|                   |                   | Inferior occipital gyrus   | 38  | −80 | −16 | 5.15    |
|                   |                   | Fusiform gyrus (occipital) | 28  | −80 | −18 | 5.04    |
|                   |                   | Occipital pole             | −24 | −100| 6   | 4.99    |
|                   |                   | Occipital pole             | 22  | −96 | 2   | 4.90    |
|                   |                   | Inferior occipital gyrus   | 30  | −90 | −4  | 4.85    |
|                   |                   | Fusiform gyrus (occipital) | 40  | −52 | −24 | 6.74    |
|                   |                   | Fusiform gyrus (temporal)  | 44  | −58 | −22 | 6.36    |
|                   |                   | Inferior occipital gyrus   | −48 | −76 | −8  | 5.93    |
|                   |                   | Inferior occipital gyrus   | 30  | −88 | 2   | 5.93    |
|                   |                   | Inferior occipital gyrus   | −42 | −86 | −8  | 5.70    |
| PTSD/Placebo      | 53.9              | Inferior occipital gyrus   | 36  | −82 | −12 | 6.54    |
|                   |                   | Inferior occipital gyrus   | 42  | −78 | −12 | 6.53    |
|                   |                   | Occipital pole             | 8   | −96 | −4  | 6.19    |
|                   |                   | Occipital pole             | 10  | −92 | −8  | 6.16    |
|                   |                   | Fusiform gyrus (occipital) | 28  | −86 | −20 | 6.13    |
|                   |                   | Inferior occipital gyrus   | −46 | −80 | −10 | 6.04    |
| PTSD/Oxytocin     | 74.0              | Inferior frontal gyrus (triangular) | 56  | 24  | 6   | 4.99    |
|                   |                   | Precentral gyrus           | 42  | 2   | 42  | 4.68    |
|                   |                   | Inferior frontal gyrus (triangular) | 54  | 26  | 16  | 4.53    |
|                   |                   | Middle frontal gyrus       | 54  | 22  | 32  | 4.52    |
|                   |                   | Middle frontal gyrus       | 52  | 18  | 38  | 4.46    |
|                   |                   | Orbitofrontal cortex       | 50  | 26  | −8  | 4.41    |
|                   |                   | Inferior frontal gyrus (triangular) | −52 | 28  | 12  | 4.62    |
|                   |                   | Middle frontal gyrus       | −40 | 6   | 34  | 4.45    |
|                   |                   | Middle frontal gyrus       | −40 | 14  | 28  | 4.06    |
|                   |                   | Inferior frontal gyrus (triangular) | −52 | 26  | 22  | 3.99    |
|                   |                   | Hippocampus               | −30 | −18 | −12 | 3.98    |
|                   |                   | Precentral gyrus           | −44 | 4   | 42  | 3.94    |
| PTSD = posttraumatic stress disorder. |
childhood trauma-exposed controls was somewhat unexpected given that previous studies report strong amygdala hyper-reactivity to emotion and trauma stimuli (Hayes et al., 2012; Shin, Rauch, & Pitman, 2006). However, others have similarly found no differences in amygdala responses to fearful or trauma cues when individuals with PTSD are compared to trauma-matched control groups (Fonzo et al., 2016; Koch et al., 2016a; Shin et al., 2006). In addition, one meta-analysis found that amygdala hyperactivity was found in studies that compared individuals with PTSD to healthy controls without a history of trauma exposure (Shin et al., 2006). Finally, alterations in amygdala activity have been found in healthy individuals who report a significant history of childhood trauma, but do not meet diagnostic criteria for a psychiatric disorder (Dannlowski et al., 2013). Thus, it is possible that trauma exposure itself, rather than PTSD, may lead to hyperactive amygdala responses to negatively valenced stimuli.

In addition to the current study’s main finding that amygdala hyperresponsivity to fearful faces was reduced in those PTSD subjects who reported more childhood trauma, the supplemental voxel-wise whole-brain analyses revealed that in the trauma control group, left amygdala reactivity was present on oxytocin but absent on placebo. Although the voxel-wise analysis was exploratory and did not statistically compare oxytocin and placebo in the trauma control group, these preliminary findings are broadly in line with findings from a study showing that oxytocin reduced amygdala responses to explicitly presented emotional faces in police officers with PTSD but increased amygdala responses in matched controls (Koch et al., 2016b).

Additional research is necessary to clarify the characteristics of individuals who might benefit from oxytocin treatment, and at what point in the development or maintenance of symptomatology oxytocin would ideally be administered. Existing literature has identified individual and contextual differences that influence the effects of oxytocin on neurobiological and behavioural responses among individuals with trauma exposure and PTSD. In the current study, groups were balanced by sex because sex is known to influence oxytocin responses (Ditzen et al., 2012; Flanagan et al., 2018a; Rilling et al., 2014). Our finding of less fear-related amygdala reactivity among more severely childhood-trauma exposed PTSD individuals under oxytocin further emphasizes the importance of considering factors such as trauma exposure severity, chronicity, and date/developmental stage of onset when testing how to facilitate the effective translation of oxytocin for clinical use.

These findings, while preliminary, might also inform future studies examining the potential for oxytocin to enhance adaptive emotion recognition and empathy among individuals with a criminal offense history. Specifically, previous studies demonstrate that individuals with a criminal history or propensity for maladaptive social behaviour (e.g. aggression and violent behaviour) display low anxiety, poorer recognition of fear, and low empathy (Blake & Gannon, 2008). Abundant literature shows that within certain contexts, oxytocin has the potential to enhance prosocial behavioural and cognition including empathy (Hurlemann et al., 2010), but other literature demonstrates that oxytocin can enhance maladaptive social behaviour (Flanagan et al., 2018a; Pfundmair, Reinelt, DeWall, & Feldmann, 2018). Future studies should examine the ability of oxytocin to correct hyporeactivity in the amygdala and limbic regions in response to facial affect, especially among individuals with known social deficits and propensity toward maladaptive social behaviour.

5. Limitations

Several limitations of the current exploratory study should be considered. One limitation is that the severity of childhood trauma exposure as measured by the CTQ was greater in the PTSD group compared to the control group, though variability in childhood trauma was similar between groups as evidenced by similar standard deviations. Future studies can improve on the current sampling plan by considering more rigorous minimum inclusion criteria related to childhood trauma. Future studies can also improve on this design by increasing the specificity of inclusion criteria for PTSD versus control groups. Future studies should also include an assessment of endogenous oxytocin levels at one or more time points given the lack of literature examining whether and how intranasal oxytocin and basal levels of oxytocin might interact (Busnelli et al., 2016; Russell et al., 2018; Sippel et al. in preparation).

While this study was designed as a preliminary proof-of-concept project and employs a within-subjects crossover design to maximize statistical power, it is possible that the expected effects of oxytocin might have been more difficult to detect than anticipated upon the initial study design. More specifically, while the overall patterns that emerged from our study are useful to inform future studies, they are preliminary and should be interpreted with caution. Related to this challenge is that the small sample size limited our supplementary whole-brain analyses to an exploratory approach to comparing oxytocin versus placebo effects in trauma-exposed and PTSD groups. More adequately powered studies with larger samples of more diverse participants are also
necessary to examine the potentially unique effects of oxytocin on the neural correlates of PTSD resulting from trauma exposure during adulthood versus childhood. Larger samples would also allow for examination of potential moderators of oxytocin response, such as sex differences in oxytocin effects on amygdala functioning; however, sex differences in amygdala responses among PTSD samples are not reliably observed (e.g. Felmingham et al., 2010). While the previous study of oxytocin effects on emotional face processing in individuals with PTSD and resilient control participants found no sex differences (Koch et al., 2016b), another study found sex-dependent effects of oxytocin on reactivity to a facial processing task among healthy adults (Luo et al., 2017). While the type and severity of childhood trauma exposure varied widely in this sample, age at which participants experienced childhood trauma exposure(s) was not recorded. Future studies can improve on this by examining the potential role of chronicity of trauma exposure in oxytocin response. While extending external validity of the findings, these characteristics might influence amygdala responses. It will also be important for future studies to consider the importance of examining the association between PTSD symptom severity and trauma exposure severity in relation to oxytocin effects on amygdala responses. The inclusion of left-handed individuals, which was not formally assessed, is another potential limitation of this study. However, previous research has shown that amygdala responses to faces are not lateralized as some other cognitive such as speech might be.

To improve internal validity, these findings should be replicated among individuals with PTSD only and those who are either required to be stable on medication doses or free of psychiatric medications. Another limitation is that the measure used to establish PTSD diagnosis was changed early in the study to reduce participant burden. While both measures have excellent psychometric properties, an ideal design would remain consistent for all participants. Generalizability is limited to individuals who experienced childhood trauma; future research can examine effects of oxytocin on fear-related processes in PTSD related to other types of trauma.

6. Conclusions

The current exploratory study did not find differences in recruitment of the amygdala in response to fearful faces between participants in the PTSD group and those in the childhood trauma-exposed control group. However, individual variation in oxytocin’s ability to modulate amygdala response depending on severity of childhood trauma exposure provides an extension of the existing literature conducted among adults with recent PTSD and trauma-exposed resilient adults. Future adequately powered studies are necessary to translate these findings into human laboratory and neuroimaging trials using repeated oxytocin doses and larger PTSD treatment and prevention-focused clinical trials.

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