The Associations Between Oxytocin and Trauma in Humans: A Systematic Review

Mariana Fortunata Donadon 1, Rocio Martin-Santos 2,3,4 and Flávia de Lima Osório 1,4*

1 Department of Neuroscience and Behavior, Medical School of Ribeirão Preto, University of São Paulo, São Paulo, Brazil, 2 Department of Psychiatry and Psychology, Hospital Clinic, Institut d’Investigacions Biomèdiques August Pi i Sunyer (IDIBAPS), Centro de Investigación en Red de Salud Mental (CIBERSAM), Barcelona, Spain, 3 Department of Medicine, University of Barcelona, Barcelona, Spain, 4 Technology Institute for Translational Medicine (INCT), National Council for Scientific and Technological Development (CNPq), Brasilia, Brazil

Studies have shown that traumatic experiences may affect hormonal systems mediated by the hypothalamic-pituitary-adrenal (HPA) axis and the oxytocinergic system. This effect is the result of long-term impairments in hypothalamic structures and negative feedback mechanisms within the HPA axis, structures that mediate the response to stress. This deregulation reduces the production and release of cortisol and oxytocin (OXT), which may alter stress responses and lead to increased vulnerability to impairments from stressful experiences. The presence of gene polymorphisms might also have an impact on the vulnerability to psychopathology. We made a systematic review of articles dealing with the relationship between OXT and traumatic emotional experiences in humans. Thirty-five studies were reviewed and significant associations between experiences of emotional trauma (ET) and OXT were found. The main results showed that the presence of ET and post-traumatic stress disorder (PTSD) is strongly associated with reductions in endogenous OXT, and also that the acute effects of OXT administrations in individuals with ET tend to be anxiolytic only in less severe forms. In victims of recent traumatic experiences (RTE), OXT increased the re-experience of traumas and restored the function of different neural networks associated with fear control/extinction in PTSD patients. The results available also suggest that gene receptor polymorphisms may have a protective function in different outcomes after the experience of traumatic events. We conclude that the relationship between ET and OXT is multifaceted, complex, and mediated by contextual and individual factors. Directions for future studies are suggested considering the gaps in the available literature.

Keywords: oxytocin, early trauma, current trauma, PTSD, systematic review, PRISMA, qualitative assessment

INTRODUCTION

Traumatic and stressful experiences throughout life, whether acute or chronic, may lead to changes in different bodily systems that increase the vulnerability to psychopathology (Meewisse et al., 2007; McQuaid et al., 2016). One of the most well-known of such changes concerns hormonal systems, including the hypothalamic-pituitary-adrenal (HPA) axis. The HPA axis plays a fundamental role in responding to both external and internal stimuli, including psychological stressors, and is also believed to be implicated in vulnerability to mental illnesses (Heinrichs et al., 2003; Juruena et al., 2004; Boyce and Ellis, 2005; Neumann and Landgraf, 2012; Olff, 2012; Kuhlman et al., 2015).
When facing stressful situations, especially during the early stages of development, the HPA axis can be either hypo- or hyperactivated, with the possibility of excessive exposure to glucocorticoids and their deleterious effects. These effects can persist throughout the lifespan because as the HPA axis may remain unstable, hypersensitive, or dysfunctional. This contributes to the weakening of the immune system, to increased vulnerability to different physical and mental illnesses, and to the inability to cope with subsequent stressful/traumatic events that may lead to exhaustion of the organism (Mirescu et al., 2004; Smith and Vale, 2006; Faravelli et al., 2012).

Several neurotransmitters and neuropeptides also affect the function of the HPA axis, including oxytocin (OXT). OXT is a neurohormone produced in the hypothalamus by the supraoptic and paraventricular nuclei. It is sent to the posterior pituitary or neurohypophysis and, from there, it is secreted into the bloodstream to produce its effects (Gimpl and Fahrenholz, 2001). OXT has peripheral and central functions and its action in breastfeeding, childbirth and maternal behavior is well established (Ring et al., 2006; Yoshida et al., 2009; Neumann and Landgraf, 2012).

In addition to these functions, OXT has therapeutic potential associated with the promotion of pro-social behaviors such as increased self-confidence, positive social memories, and affiliative behavior. Furthermore, previous studies have investigated the possibility that the administration of OXT may lead to reductions in anxiety and stress levels (Savaskan et al., 2008; Guastella et al., 2009; Rimmle et al., 2009; Ross and Young, 2009; Fischer-Shofaty et al., 2010).

Existing evidence suggests that central OXT release contributes to the modulation and maintenance of cortisol levels that favor the rapid return of the body to its pre-stress baseline state (Amico et al., 2004; Gulpinar and Yegen, 2004; Heinrichs et al., 2004) to minimize the response of the HPA axis to psychologically stressful stimuli. However, stressful experiences might also alter the functioning of the suprachiasmatic nucleus, decreasing the synthesis and release of endogenous OXT (Ozbay et al., 2008; Gonzalez et al., 2009; Nicolson et al., 2010). As result, the negative feedback mechanism of the HPA axis may be affected, leading to hypercortisolism (Yehuda et al., 2000; Brown et al., 2016). On the behavioral level, the decrease in endogenous OXT levels reduces the multiple pro-social functions of this hormone, thereby reducing coping and resiliency responses (Opacka-Juffry and Mohiyeddini, 2012; Frijling and et al., 2015).

In light of the above findings, the release of exogenous administration of OXT can reduce hormonal and subjective responses to stress, reduce cortisol release in the blood stream, and reestablish bodily homeostasis, therefore placing OXT as a potential therapeutic agent (Cardoso et al., 2013, 2014). Another line of research deals with the role that genetic polymorphisms play in OXT receptor genes, which might alter the individual vulnerability to stress since changes in receptor configuration can reduce or prevent the binding of OXT and its consequent effects in the body (Champagne and Curley, 2009; Skuse and Gallagher, 2011; Unternaehrer et al., 2012; Bakermans-Kranenburg and van Ijzendoorn, 2014).

As seen, a number of recent studies have attempted to provide a detailed understanding of the multiple and complex associations between OXT and different traumatic and stressful situations, whether chronic or acute (Myers et al., 2014; Seltzer et al., 2014; Mizushima et al., 2015; Van Zuiden et al., 2017). The objective of the current study was to systematically review this literature to highlight the major contributions of the studies conducted to date dealing with the associations between OXT and traumatic emotional experiences in humans.

**METHODOLOGY**

The guidelines of the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA—Moher et al., 2009) were adopted as the methodological framework of this study. The electronic databases PsycINFO, PubMed, Scielo, Web of Science, and LILACS were searched without limitations in terms of time, language, or publication date (last search conducted on December 3, 2017). A manual search of the reference lists of the selected articles was also performed. The following keywords were used followed by “AND” or “OR:” oxytocin; PTSD; post-traumatic stress; early trauma; childhood maltreatment; emotional trauma; emotional stress; neglect; adversity; sexual abuse; emotional abuse; and physical abuse.

Articles that evaluated associations between OXT and the experience of early trauma (ET), recent traumatic experiences (RTE), and post-traumatic stress disorder (PTSD) in men and women, regardless of age, were eligible for inclusion. ET experiences were considered as those that occurred before 18 years of age (Bremner et al., 2000), and RTEs were considered as those that occurred over the last few days or months (Chatziotis et al., 2014).

Studies of endogenous OXT that did not involve participants with ET, RTE, or PTSD; those investigating OXT and psychiatric disorders excluding PTSD; those involving OXT and social behavior, resilience, social cognition, genetic analysis, and the physiological aspects of the HPA axis among participants without ET, RTE, or PTSD; those involving treatment for ET or PTSD but not involving OXT; and animal studies, letters to the editor, case studies, and reviews unrelated to this topic were excluded.

Two researchers made independent decisions regarding whether to include a study, and divergences in data extraction were discussed until a consensus was reached.

**Figure 1** shows the search results as well the reasons for article exclusion.

The methodological quality of the studies was evaluated using the following references: (a) Strengthening the Reporting of Observational Studies (STROBE; Von Elm et al., 2007); (b) Transparent Reporting of Evaluations with Non-Randomized Designs (TREND; Des Jarlais et al., 2004); and (c) the Revised Recommendations for Improving the Quality of Reports of Parallel-Group Randomized Trials (CONSORT; Moher et al., 2001). The quality percentage of each study was calculated by dividing the number of items scored by the total number of items.
RESULTS

A total of 1,244 articles were found, and 35 articles were selected after applying the inclusion and exclusion criteria (see Figure 1).

Table 1 below presents the main characteristics of the included studies in relation to the samples, outcomes assessment, and methodological quality. For more details, see the Excel dataset in the Supplementary Material (S1).

Importantly, regarding the methodological quality evaluation, all of the studies included in this review had at least 64% of their essential items included in the STROBE (observational studies), TREAD (experimental studies) or CONSORT [experimental or randomized controlled trials (RCTs)].

Depending on their designs or objectives, the studies were divided into four distinct groups: (a) observational studies evaluating endogenous OXT levels; (b) experimental studies related to the reactivity of the oxytocinergic system; (c) RCTs of OXT administration and the experience of either ET, RTE, or PTSD; and (d) observational studies investigating the effect of polymorphisms of the OXT receptor gene.

The major results of each study group are presented below.

(a) Observational studies evaluating endogenous OXT levels.

The association between endogenous OXT levels and ET situations was evaluated by six studies as a major outcome. Four of these studies found significant correlations between endogenous OXT levels and ET, with values ranging from $-0.54$ to $-0.23$ (Heim et al., 2009; Opacca-Juffry and Mohiyeddini, 2012; Chatziotisof et al., 2014; Mohiyeddini et al., 2014). On the contrary, Mizushima et al. (2015) did not find an association between OXT levels and experiencing ET ($p > 0.05; d = 0.19$; insignificant effect size). Importantly, however, OXT secretions
### TABLE 1 | Characteristics of the studies included in this review.

| Author/Year | Study design | Interest sample | Comparison sample | Trauma | (OXT)* | Other variables/Instrument | Trauma tool | Quality assessment** |
|-------------|--------------|-----------------|-------------------|--------|--------|---------------------------|-------------|----------------------|
| Heim et al., 2009 | Cross-sectional | Outpatients Healthy Institution | ET | — | — | CSF/EIA kit | CTQ | 67.0 |
| Opacka-Juffry and Mohiyeddini, 2012 | Cross-sectional | Healthy General Population | ET RTE | — | — | Blood/RIA | ELS-C, ELS-A, RSLEs | 68.0 |
| Chatzittofis et al., 2014 | Cross-sectional | Inpatients Suicide Attempters | ET+ RTE /Non-RTE | — | — | CSF/RIA | KVS, Interview | 74.0 |
| Mohiyeddini et al., 2014 | Cross-sectional | Healthy General Population | ET | — | — | Blood/RIA | ELS-C | 85.0 |
| Muzuki and Fujiwara, 2015 | Cross-sectional | Healthy General Population | ET/Non-ET | — | — | Urine/ASKA kit | CTQ | 77.0 |
| Frijling et al., 2015 | Case-Control | Outpatients/Police Personal Clinic | PTSD + ET/Non-ET | 20M | 22–59 | Police officers RTE + ET/Non-ET | Saliva/RIA | 90.0 |
| Mizushima et al., 2015 | Case-Control | Residential Child Care | ET | 14M | 22-12.7 (2.1) | Healthy General Population | Saliva/Elisa Kit | 64.0 |
| Nishi et al., 2015 | Cohort | Inpatients Intensive care unit | RTE # | — | — | Blood/ELISA Kit | IES-R | 81.0 |
| Reijnen et al., 2017 | Cohort | Military after combat | ET PTSD | — | — | Blood/ELISA Kit | ETIS-SR, DSRI | 80.6 |

### 2 - EXPERIMENTAL STUDIES - STRESS REACTIVITY

| Author/Year | Study design | Interest sample | Comparison sample | Trauma | (OXT)* | Other variables/Instrument | Trauma tool | Quality assessment** |
|-------------|--------------|-----------------|-------------------|--------|--------|---------------------------|-------------|----------------------|
| Pierrehumbert et al., 2010 | Pre/Post | Outpatients/Health Healthy/ Cancer | ET** & ET## | 14M | 20-29.3 (8.50) | Healthy General Population | Non-ET | Blood/RIA | Stress Test/ TSST-C, Interview | 81.8 |
| Munro et al., 2013 | Pre/Post | Survivors Healthy University Students | ET/Non-ET /PTSD | — | — | Blood/RIA | Stress Test / Film Protocol | ELS | 89.0 |
| Seltzer et al., 2014 | Pre/Post | Child Protective Services | ET### | 14M | 8-11.5 | Healthy General Population | No ET | Urine/RIA | Stress Test/ TSST-C | PSC | 92.7 |

### 3 - CLINICAL TRIALS, RANDOMIZED, DOUBLE-BLIND, PLACEBO-CONTROLLED - EXOGENOUS ADMINISTRATION

| Author/Year | Study design | Interest sample | Comparison sample | Trauma | (OXT)* | Other variables/Instrument | Trauma tool | Quality assessment** |
|-------------|--------------|-----------------|-------------------|--------|--------|---------------------------|-------------|----------------------|
| Meinlschmidt and Ham, 2007 | Crossover | Healthy University Students | ET | 10M | 20-28 | Healthy University Students | Non-ET | OXT vs. PLA intranasal (2IU) | [Control]/saliva SSRQ | 61.2 |
| Author/Year            | Study Design    | Interest sample                                      | Comparison sample                                      | Outcomes assessment                                      | Quality assessment** |
|-----------------------|----------------|------------------------------------------------------|--------------------------------------------------------|----------------------------------------------------------|---------------------|
| Grimm et al., 2014    | Crossover      | Healthy General Population                           | Healthy General Population                             | OXT vs. PLA Intranasal (40 UI)                           | CTQ                 |
| Frijling et al., 2016 | Parallel Group | Outpatients Emergency Department                     | Outpatients Emergency Department                      | OXT vs. PLA Intranasal (40 UI)                           | PDS                 |
| Sack et al., 2017     | Crossover      | Outpatients Psychosomatic Clinic                     | Healthy Non-PTSD                                       | OXT vs. PLA Intranasal (24 UI)                           | DSS                 |
| Nawijn et al., 2016   | Crossover      | Outpatients Police Personnel                         | Trauma centers                                         | OXT vs. PLA Intranasal (40 UI)                           | CAPS, SOD           |
| Bradley et al., 2011  | Cross-sectional| Outpatients Primary Care                             | ET/Non-ET                                              | OXTR 5:3576                                             | CTQ, TB             |

4-OBSERVATIONAL STUDIES – ASSOCIATION WITH POLYMORPHISMS OXYTOCIN RECEPTOR GENE

Mean: 76.27%
### TABLE 1

| Author/Year | Study design | Interest sample | Comparison sample | Outcomes assessment | Quality assessment** |
|-------------|--------------|-----------------|-------------------|---------------------|---------------------|
| **Cicchetti and Rogosch, 2012** | Case-Control | Healthy Sumer Camp Program | Healthy Sumer Camp Program | OXTR 53576 | Resilience/PEI Interview 65.0 |
| **Lucas-Thompson and Holman, 2013** | Cohort | 9/11 Attack | PTSD/ RTE Non-PTSD | OXTR 53576 | — PCL 84.0 |
| **McQuaid et al., 2013** | Cross-sectional | Healthy University Students | ET/Non-ET | OXTR 53576 Depression/ BDI CMO 87.0 |
| **Hostinar et al., 2014** | Case-Control | Malpractice Child Protective Services | ET | OXTR 53576 Int/Ext/ YSR Social Support/NRI Interview 81.0 |
| **Myers et al., 2014** | Cross-sectional | Healthy General Population | ET/Non-ET | OXTR 129832701 1113147 Depression/ ELS 77.0 |
| **Dunn et al., 2014** | Cohort | Recent Trauma Hurricane Katrina | PTSD/ RTE Non-PTSD | OXTR 53576/ 2254298 | — PTG, IES-R 71.0 |
| **Dannlowski et al., 2016** | Cross-sectional | Healthy ET/Non-ET | OXTR 53576 fMRI/Brain activation Facial Task/EFMT CTO 67.9 |
| **Tollenaar et al., 2017** | Cohort | Healthy RTE/No PTSD | OXTR 2254298/ Depression and Anxiety/ DSM-IV/ CIDI + Interview 75 |
| **Sippel et al., 2017** | Cohort | War Veterans PTSD | OXTR 53576 Attachment Style Questionnaire DSM-IV: PLC 81.8 |

*Technique Used for Measuring and/or Test Kit And Dose And Via Administration; **Quality Assessment According To STROBE, CONSORT, and TREND; #: Current trauma by motor or vehicle accident; #: ET by cancer survival in childhood; ###: ET by physical abuse; #: APQ, Attachment Prototype Questionnaire; BDI, Beck Depression Inventory; CAN, Canada; CAPS, Clinician Administered; CIDI, Composite Interview Diagnostic Instrument; PTSD Scale; CES, Center for Epidemiological Studies Depression Scale; CMO, Childhood Malpractice Questionnaire; Cross, Cross Sectional; CSF, Cerebrospinal Fluid; CTQ, Childhood Trauma Questionnaire; DSM-IV, Diagnostic and Statistical Manual of Mental Disorders; EDS, Emotional Deregulation Scale; ELS, Early Life Stress Questionnaire; ELS-A, Early Life Stressful Adolescence; ELS-C, Early Life Stressful Experiences In Childhood; ELS-D, Emotional Dysregulation Scale; ENG, England; EPS, Parental Early Separation; ET, Early Trauma; ETI, Early Trauma Inventory; ETI-SF, Early Trauma Inventory Self Report - Short Form; ETI, Early Trauma Inventory; fMRI, Functional Magnetic Resonance Imaging; F, Female; GER, Germany; IES, Impact of Event Scale Revised; IN/ EXT, Internalizing/ Externalizing Symptoms; HR, Heart Rate; ISR, Israel; JAN, Japan; KIVS, The Karolinska Interpersonal Violence Scale; MEG, Magnetoencephalography; MID, Monetary Incentive Delay; MIST, Montreal Image Task; NETH, Netherlands; M, Male; MINI, Mini-International Neuropsychiatric Interview; NRI, Network Relationships Inventory; OXTR, Oxytocin; PCL, Checklist Civilian Version for DSM-IV; PDI, Pentraumatic Distress Inventory; PTSD, Post Traumatic Stress Diagnostic; PEI, Pupil Evaluation Inventory; PG, Parallel Group; PL, Placebo; PSC, Parental Child Conflict Scale; PTG, Post Traumatic Growth; PTSD, Post Traumatic Stress Disorder; R, Recent; RSDS, Dissociative Symptoms Scale; RTE, Recent traumatic experiences; SCD, Clinical Interview And Structured (DSM-IV); SDIS, Social Disability Scale; SSRQ, Standardized Self Report Questionnaire; STAI, State-Trait Anxiety Inventory; SW, Switzerland; TEI, Traumatic Events Inventory; TSCC, Trauma Symptoms Check List For Children; TSQ, Trauma Screening Questionnaire; TSST-C, Trier Social Stress Test, UI, International Units; USA, United States of America; YSR, Youth SelfReport; [OXT], Oxytocin Concentration Endogenous; 9/11 Attack= terrorist attack USA.}
were markedly increased in adolescents who suffered abuse and lived in a stable environment (e.g., a social welfare institution) at the time of the experiment from awakening to bedtime compared with those who had a history of abuse and lived in unstable environments. Mizuki and Fujiwara (2015) showed that only less severe forms of ET were associated with increased OXT levels.

Heim et al. (2009) also performed complementary analyses to evaluate the effect of the recurrence of traumatic events. These authors found that experiencing three or more types of traumatic events during childhood was related with reduced endogenous OXT levels, with a moderate effect size ($n_p^2 = 0.45$). Opacka-Juffry and Mohiyeddini (2012) also conducted complementary analyses and found associations between decreased endogenous OXT levels and increased emotional suppression ($r = -0.30, p < 0.01$) during adulthood.

Studies that evaluated participants with RTEs failed to find any significant correlations. For example, Opacka-Juffry and Mohiyeddini (2012) found that the correlation between OXT and RTE was 0.01 ($p > 0.05$). Chatzitofis et al. (2014) also failed to find an association between endogenous OXT levels and RTEs ($r = -0.30, p = 0.18$); however, these authors found that endogenous OXT levels were lower in individuals with RTEs who also experienced ET (i.e., were re-victimized) than in those who only experienced trauma during childhood ($p = 0.04$).

Nishi et al. (2015) also failed to find an association between PTSD symptoms and endogenous OXT levels ($r = -0.08$ to $-0.00, p > 0.57$). However, these authors showed interesting differences between the genders: In women, endogenous OXT levels were positively correlated with cooperativeness ($r = 0.41, p = 0.01$), whereas in men these levels were negatively correlated with C-reactive protein ($r = -0.22, p < 0.01$), which indicates that OXT plays a role in the coping strategies for PTSD symptoms among women.

The findings of Frijling et al. (2015) also indicated gender differences. Regarding endogenous OXT levels in highly traumatized police officers, only men with PTSD showed lower levels of OXT than those without PTSD ($p < 0.05, d = 0.60$; moderate effect size). No differences were observed in women ($p > 0.05; d = 0.10$; insignificant effect size).

Finally, a recent study by Reijnen et al. (2017) found that pre-deployment OXT levels in soldiers sent to Afghanistan did not predict PTSD development. However, the experience of an ET predicted the development of PTSD, even though no associations were found between OXT levels and the presence/absence of ET in these individuals.

(b) Experimental studies relating to the reactivity of the oxytocinergic system.

Three studies evaluated OXT levels (reactivity patterns) in acute stressful situations, although these results should be interpreted with caution because of their small sample sizes (Table 2).

Girls who experienced physical abuse during childhood showed increased reactivity to stress. In other words, they presented with higher levels of endogenous OXT after undergoing an acute stress test ($p = 0.02$), whereas OXT levels in boys with and without a history of physical abuse did not change after stress induction (Seltzer et al., 2014). On the other hand, adults with a history of sexual abuse (regardless of gender) showed decreasing OXT levels after acute stress (Pierrehumbert et al., 2010).

Munro et al. (2013) also found a decrease in OXT levels after exposure to abandonment scenes ($p = 0.01$) but not after exposure to bonding scenes ($p = 0.39$). However, associations were observed between increased OXT levels and increased dissociative, somatic, and attachment symptoms during bonding scenes as well as between decreased OXT levels and PTSD symptoms during abandonment scenes.

(c) RCTs of OXT administration and trauma.

The results of these trials were grouped by the type of trauma experienced and are shown in Table 3.

### Table 2: Major results of endogenous OXT during experimental studies (reactivity to stress) of participants who experienced trauma ($n = 3$).

| Study            | Type of stress test | [OXT] situation | Reactivity to stress | $r$  | $p$       |
|------------------|---------------------|-----------------|----------------------|------|----------|
| Pierrehumbert et al., 2010 | TSST-C | Pre-Stress [OXT] | ET sexual abuse = ET cancer; childhood = control | $-$ | 0.23     |
| Seltzer et al., 2014 | TSST-C | Post-Stress (+20 min) [OXT] | ET sexual abuse < control | $-$ | 0.06      |
| Munro et al., 2013 | Film Protocol (Abandonment and Bond scenes) | Bonding [OXT] | Dissociation symptoms x higher levels [OXT] | 0.55 | 0.018* |
|                  |                     |                 | Somatization symptoms x higher levels [OXT] | 0.59 | 0.010* |
|                  |                     | Abandonment [OXT] | Basal → Abandonment = ↓ PTSD x lower levels [OXT] | 0.35 | 0.010 |

[OXT]: oxytocin; ET: Early trauma; PTSD: Post-traumatic stress disorder; ♂: Men; ♀: Women; TSST-C: Trier Social Stress Test; <: Minor; >: Major. *Difference statistically significant.
TABLE 3 | Major results of RCTs that administered OXT to participants who experienced early or current trauma (N = 13).

| Study | Outcomes | Treatment | Comparison groups | Main results |
|-------|-----------|-----------|-------------------|--------------|
| (1) EARLY TRAUMA | | | | | |
| Meinlschmidt and Heim, 2007 | [Cortisol] | OXT vs. ET vs. Non-ET Placebo | | Attenuated the ↓ [cortisol] after OTX in subjects with ET |
| | Brain activation fMRI/ 3T | OXT vs. ET vs. Non-ET Placebo | | ↑ Activation left hippocampus and dorsomedial thalamus |
| | Voxel wise | OXT vs. ET vs. Non-ET Placebo | | ↓ [cortisol] after stress task |
| | MIST (Stress task) | OXT vs. ET vs. Non-ET Placebo | | ↑ Activation at right insula, anterior ACC, PCC, left parahippocampal gyrus |
| | [Cortisol] | Placebo | | No effect on the [cortisol] after stress task |
| | | | | ↓ [cortisol] after stress task |
| | | | | ↑ Hormonal limbic reactivity (pgACC, left amygdala, left parahippocampal gyrus, left insula, bilateral putamen and bilateral caudate), during the stress test |
| | | | | ↓ Hormonal limbic reactivity (pgACC, left amygdala, left parahippocampal gyrus, left insula, bilateral putamen and bilateral caudate), during the stress test |
| Fan et al., 2014 | Brain connectivity fMRI/ 3T | Placebo | OXT vs. ET vs. Non-ET Placebo | Resting state ↓ pgACC-amygdala rs/FC |
| | Voxel/wise | Placebo | OXT | It was not significant effect of OXT |
| | Seed/based approach | OXT | | It was no significant effect of OXT & ET interaction |
| | Resting state | Placebo | OXT | Psychosocial Stress Correlated with state anxiety, that correlate with [cortisol] |
| | MIST (Stress task) | Placebo | OXT | Negatively predicted anxiety |
| | | | | Predicted stronger pgACC deactivation during stress |
| | | | | Predicted weaker pgACC deactivation |
| | | | | Attenuated rest-task interaction between pgACC-amygdala rs/FC and pgACC deactivation |
| (2) CURRENT TRAUMA | | | | | |
| Frijling et al., 2016a | Brain connectivity fMR/ 3T | Placebo | OXT vs. Endothelial vs. Non-Endothelial Placebo | Connectivity left amygdala and PFC for trauma images vs. neutral images |
| | Voxel/bold | Placebo | OXT | Connectivity left amygdala and PFC for trauma images vs. neutral images |
| | Accident images vs. Neutral images Task | Placebo | OXT | Amygdala connectivity to the left insula for trauma images vs. neutral images |
| | | | | Amygdala connectivity vmPFC for trauma images vs. neutral images |
| | | | | Flashbacks of memory during accident images |
| | | | | ↑ BL amygdala activation for fearful faces |
| | | | | ♂ Left amygdala to neutral faces |
| Frijling et al., 2016b | Brain activation fMR/ 3T | Placebo | OXT vs. Placebo | Connectivity left amygdala and PFC for trauma images vs. neutral images |
| | Voxel/bold | Placebo | OXT | Connectivity left amygdala and PFC for trauma images vs. neutral images |
| | Facial task | Placebo | OXT | Amygdala connectivity to the left insula for trauma images vs. neutral images |
| Van Zuiden et al., 2017 | PTSD symptoms | OXT vs. Placebo | Placebo | Valence-dependent amygdala reactivity was absent for the left amygdala |
| | | | | Increased reactivity of the left amygdala |
| (3) PTSD | | | | | |
| Eidelman-Rothman et al., 2015 | MEG | Placebo | PTSD vs. Non-PTSD OXT | a resting-state activity in left dPFC, SFG and MFG |
| | Frequency/Hz | OXT | PTSD vs. Non-PTSD OXT | a resting-state activity in left SFG e MFG |
| | Alpha Resting State activation | OXT | PTSD vs. Non-PTSD OXT | a resting-state activity in left SFG and MFG correlated with ex-reperience symptoms |
| Palgi et al., 2016 | Compassion task | Placebo | PTSD | Compassion toward women protagonist |
| | | OXT vs. Placebo | PTSD | No effect on compassion toward masculine protagonist |
| Koch et al., 2016a | Brain activation fMR/ 3T | Placebo | PTSD | Valence-dependent amygdala reactivity was absent for the left amygdala |
| | Voxel/bold | Placebo | PTSD | ↑ Reactivity of amygdala to fearful-angry faces compared with happy -neutral faces |
| | Facial task | Placebo | PTSD | ↓ Reactivity of left amygdala all emotions |
| | | | | ↑ Increased reactivity of the left amygdala |
| | | | | ↓ Activation of left amygdala all emotions |
| | | | | ↓ Activation of left amygdala |

(Continued)
Three studies evaluated participants with just ET. In the first, Meinschmidt and Heim (2007) measured endogenous cortisol and demonstrated that intranasal OXT attenuated the cortisol decrease in participants with ET compared with controls, suggesting the presence of amortization effects related to HPA-axis activities. In the other two studies, participants with ET were assessed in the context of a psychosocial stress situation, and the results indicated that negative outcomes were associated with acute OXT administration.

In one of these studies, Grimm et al. (2014) found that participants with ET, regardless of severity, presented with greater hormonal and limbic reactivity after the use of OXT. However, Fan et al. (2014) found that the activation of the connectivity between the amygdala and the preganual anterior cingulate cortex (pgACC) during stress was only attenuated by OXT in individuals with less severe ET. The use of OXT did not favor this anxiolytic effect in patients with moderate-to-severe ET.

Two studies by Frijling et al. (2016a,b) reported the adverse effects of exogenous OXT administration in individuals with RTEs. In one such study (Frijling et al., 2016a), OXT administration during threatening situations reduced the functional connectivity between the left amygdala and the ventrolateral prefrontal cortex (vlPFC) as well as between the amygdala and the ventromedial prefrontal cortex (vmPFC) circuits responsible for cognitive-emotional regulation and fear extinction. These findings were also accompanied by an increased connectivity between the amygdala and the insula as well as increased episodes of traumatic flashbacks. In the other study (Frijling et al., 2016b), increased amygdala reactivity was elicited by fearful faces, indicating that OXT favors an increase in the processing of fear salience and, consequently, anxiogenic effects.

A single study administered OXT for eight subsequent days and showed that it did not attenuate PTSD symptoms in the short term (45 days). However, only participants with high symptom severity reported improvement after 6 months, which suggests that OXT may have a protective effect in the long term which is mediated by symptom severity (Van Zuiden et al., 2017). In this study, as well as at Fan et al. (2014), it is evident the influence of the severity of experience/symptoms of trauma.

Contrary to the effects observed in subjects with RTE, in those with a PTSD installed, the related effects to acute OXT administration were favorable. Eidelman-Rothman et al. (2015) examined veterans with PTSD and showed that OXT normalized the resting-state brain functioning of these individuals, which was similar to those of controls (i.e., veterans not exposed to trauma). Prior to exogenous OXT administration, an increase in resting-state alpha activity was observed in the left dorsolateral prefrontal cortex (DLPFC) and middle frontol gyrus (MFG). These regions are associated with memory and cognitive control, which are important for emotional control.

Koch et al. (2016a,b) and Nawijn et al. (2016, 2017) examined the same sample of participants (i.e., trauma-exposed police

---

### TABLE 3 | Continued

| Study          | Outcomes                  | Treatment | Comparison groups                      | Main results                                                                 |
|----------------|----------------------------|-----------|----------------------------------------|-------------------------------------------------------------------------------|
| Koch et al., 2016b | Brain connectivity | Placebo   | PTSD vs. PTSD vs. PTSD Non-PTSD         | Connectivity right CeM amygdala to left vmPFC                                  |
|                | fMR/ 3T                   | OXT       | PTSD vs. PTSD vs. PTSD Non-PTSD         | Connectivity right BLA to bilateral dACC                                      |
|                | Voxel/bold                | Placebo   | PTSD vs. PTSD vs. PTSD Non-PTSD         | Connectivity right CeM to left vmPFC                                          |
|                | Resting State             | OXT       | PTSD vs. PTSD vs. PTSD Non-PTSD         | Connectivity right BLA to right dACC                                          |
| Nawijn et al., 2016 | Brain activation       | Placebo   | PTSD vs. Placebo                        | Anxiety and nervousness but not happiness and sadness                        |
|                | fMR/ 3T                   | OXT vs.   | PTSD vs. PTSD vs. PTSD Non-PTSD         | Reaction time on MID task for reward/loss vs. neutral trials                 |
|                | Voxel/bold                | Placebo   | PTSD vs. PTSD vs. PTSD Non-PTSD         | Brain activation at ventral stratum, amygdala, insula, CBF                    |
|                | Task/Monetary             | OXT       | PTSD vs. PTSD vs. PTSD Non-PTSD         | Brain activation at right stratum, dACC, and insula during reward and loss   |
| Sack et al., 2017 | Trauma Script           | OXT       | PTSD vs. PTSD vs. PTSD Non-PTSD         | Activationinsula anterior left during social incentive                        |
|               | Challenge                 | Placebo   | PTSD vs. PTSD vs. PTSD Non-PTSD         | Activation bilateral putamen, right dACC and right insula during social incentive |
| Nawijn et al., 2017 | Brain activation     | Placebo   | PTSD vs. Placebo                        | Activation right Putamen during social incentive                               |
|                | fMR/ 3T                   | OXT       | PTSD vs. PTSD vs. PTSD Non-PTSD         | Activation left stratum; right striatum and insula, and right dorsal ACC      |
|                | Social Task               | Placebo   | PTSD vs. PTSD vs. PTSD Non-PTSD         |                                                                               |

*Early trauma of emotional abuse; α, alpha; ACC, Anterior cingulate cortex; BLA, Amygdala basolateral; BL, Basolateral; CeM, central medial amygdala; CAPS, Clinician-administered PTSD Scale; DMPFC, Dorsomedial pre-frontal cortex; dACC, Dorsal anterior cingulate cortex; dPFC, Dorsolateral pre-frontal cortex; MFG, Middle frontal gyrus; MEG, Magnetoencephalography; MID, Monetary incentive delay task; MIST, Montreal Image Task; PTSD, Post traumatic stress disorder; OXT, Oxytocin; rs/fc, resting state functional connectivity; VmPFC, Ventral medial pre frontal cortex; VPPFC, Ventral lateral pre frontal cortex; SFG, Superior frontal gyrus; MFG, Middle frontal gyrus; ET, Early trauma; [Cortisol], concentration of cortisol; fMRI, Functional magnetic resonance imaging; pgACC, Pregenual cortex cingulate anterior; RTE, Recent traumatic experiences; SPG, superior frontal gyrus; sgACC, Subgenual anterior cingulate cortex.
officers with or without PTSD), in different paradigms outcomes and showed positive effects during acute OXT administration. In a resting state paradigm the acute OXT effects were different between genders. In men with PTSD, OXT decreased subjective anxiety and nervousness as well as restored the connectivity between right amygdala (CeM) and the left vmPFC. In women, OXT restored the connectivity between the right basolateral amygdala (BLA) and the anterior cingulate dorsal cortex (dACC), which decreased the anxiety and fear expression originating from the amygdala (Koch et al., 2016b). In an emotional face-matching task, the acute effects of OXT in subjects with PTSD were the same, regardless of gender: OXT reduced amygdala reactivity to all emotional expressions. In those without RT2E, however, reactivity was increased, which indicates the presence of the anxiolytic effects of OXT only in trauma-exposed individuals who develop PTSD (i.e., an interdependence of inter-individual factors; Koch et al., 2016a). During a monetary task, OXT increases neural responses during anticipation of reward or loss in key regions of the brain’s reward circuit (i.e., the striatum, dACC, and insula) and decreases motivational anhedonia. These effects were positively associated with those of OXT in the ventral striatum (Nawijn et al., 2016). Similarly, during a social incentive delay task, the administration of OXT normalized the aberrant insula response and increased the putamen response, indicating increased neural sensitivity to social reward (Nawijn et al., 2017). Finally, it was observed as positive effects related to OXT, an increase in compassion toward women with PTSD (Palgi et al., 2016) and a decrease in the avoidance symptoms during trauma script exposure (Sack et al., 2012).

(d) Observational studies investigating the effect of the polymorphisms of the OXT receptor gene.

The major polymorphism studied was OXTR rs53576, which was evaluated in 88.8% of the included studies. Table 4 shows the main results found.

Regarding the results linked to the OXTR rs53576 gene polymorphism, participants who exhibited the GG genotype and experienced ET, PTSD, or both reported a series of impairments including emotional dysregulation and problematic attachment (Bradley et al., 2011), PTSD symptoms (Lucas-Thompson and Holman, 2013), symptoms of depression (McQuaid et al., 2013), lower levels of perceived social support, and internalized behavioral problems (Hostinar et al., 2014). Using structural magnetic resonance, Dannlowski et al. (2016) showed that the presence of the G allele (GG/GA) was associated with increased amygdala responsiveness to all emotional facial expressions (negative and positive), constituting a higher vulnerability to alterations in the limbic brain structure in individuals with ET. Furthermore, a negative correlation was found between ventral striatum gray matter volume and participants with ET and the GG genotype.

Cicchetti and Rogosch (2012) showed that the presence of ET and the AA/AG genotype was associated with increased resiliency; therefore, the presence of the AA/AG genotype appears to have a protective function. On the other hand, Tollenaar et al. (2017) indicated that the OXTR rs53576 gene polymorphism does not interact with ET and predict risk factor and/or vulnerability toward the development of depression or anxiety. In addition, Sippel et al. (2017) found that the presence of a minor allele associated with an insecure attachment style was associated with a higher prevalence of PTSD among war veterans.

Other polymorphisms studied included OXTR rs2254298 and rs2268498, but neither Dunn et al. (2014) nor Tollenaar et al. (2017) found changes or association among the presence of ET or PTSD, a single nucleotide polymorphism, and the development of symptoms. In addition, Dunn et al. (2014) investigated the role of rs53576 and did not find a difference between the presence of the AA allele and the AG or GG alleles as risk factors for the development of PTSD or its symptoms.

Finally, one study evaluated the polymorphisms OXTR rs139832701 and rs11131147. This study showed that participants who experienced ET and had these polymorphisms exhibited higher levels of depression and more symptoms of stress (Myers et al., 2014).

DISCUSSION

The present review revealed associations between traumatic experiences in humans and the neuropeptide OXT that involve polymorphisms on OXT receptor genes. These associations are multiple and complex and are mediated by contextual and inter-individual factors. A stronger association with OXT was found with regard to the experience of early or chronic trauma as well as with recurrent, severe, or intense traumatic events. Regarding RTEs, the association was strong among individuals who developed psychopathological conditions (e.g., PTSD).

The findings of the group of studies that evaluated endogenous OXT levels showed a moderate association between reduced OXT levels and the experience of trauma. This finding supports the hypothesis that early adversity persistently, even up to adulthood, alters the functioning of the suprachiasmatic nucleus, which is responsible for the production and release of OXT (Ozbay et al., 2008; Gonzalez et al., 2009; Nicolson et al., 2010; Goldman-Mellor et al., 2012). This change might favor vulnerability to stress during adulthood and impair the social functioning associated with OXT. Moreover, it might reduce coping and resiliency responses (Opacka-Juffry and Mohiyeddini, 2012; Frijling et al., 2015). Interestingly, Munro et al. (2013) showed that an increase in endogenous OXT was associated with symptoms of social detachment, which might not always result in favorable outcomes (Seng, 2010).

However, an increase in endogenous OXT occurred in participants with ET and specific conditions such as less severe forms of ET and social environmental change (from a threatening environment to a protective environment with the establishment of stable social relationships). This finding shows the role of OXT in the development and maintenance of resiliency (Elzinga et al., 2008; Carpenter et al., 2009; Heim et al., 2009). Authors such as Mizuki and Fujiwara (2015) and Mizushima et al. (2015) suggested that an increase in OXT occurs as a response to social stress, thereby promoting the regulation of the oxytocinergic system and an increase in pro-social behaviors under challenging and less unfavorable situations. These findings also reinforce
TABLE 4 | Main results of the association studies of early or current trauma with regard to OXT receptor gene polymorphisms (N = 10).

| Study | Phenotype | Results genetic association with ET, RTE and PTSD | Statistics |
|-------|-----------|-------------------------------------------------|-------------|
| Bradley et al., 2011 | Emotional deregulation | 3 ET or more | GG > AA/AG | <0.001 | $\eta^2 = 0.052$ |
| Ciocchetti and Rogosch, 2012 | Resilience | ET+AA/AG | < | Non-ET +AA/AG | <0.01 | $\eta^2 = 0.083$ |
| Lucas-Thompson and Holman, 2013 | PTSD symptoms | RTE*+GG | > | Non-RTE*+GG | <0.01 |
| Dunn et al., 2014 | PTSD symptoms | Low ET+GG/AA | = | Low ET+AA | 0.07 |
| Dannowski et al., 2016 | Ventral striatum gray matter volume | ET+GG | < | AA | <0.01 |
| Tollenaar et al., 2017 | Depression symptoms | ET/Non-ET | GG=AA/AA | >0.08 | – |
| Sippel et al., 2017 | Insecure attachment style | PTSD | A allele + insecure attachment = risk factor for PTSD | 0.02 |

| Study | Phenotype | Results genetic association with ET, RTE and PTSD | Statistics |
|-------|-----------|-------------------------------------------------|-------------|
| Dunn et al., 2014 | PTSD symptoms | RTE+GG | RTE+AA | RTE+AA | 0.45 |
| Tollenaar et al., 2017 | Depression Symptoms | ET/Non-ET | GG=AA/AA | >0.21 | – |
| Myers et al., 2014 | Depression symptoms | ET >Non-ET | GG=AA/AA | >0.08 | – |

*Recent traumatic experiences by economic stress; **Recent traumatic experiences by negative environment; A, A allele; d, Cohen’s d; ET, early trauma; G, G allele; $r^2$, partial eta squared; PTSD, post-traumatic stress disorder; p, level of significance; Non-ET, absence of early trauma; RTE, Recent traumatic experiences.

Veenema (2012) views concerning the high plasticity of the oxytocinergic system and its dependence on social cues. Studies have indicated the mediating role that other variables play in traumatic situations during adulthood because OXT is reduced only in subjects with ET and/or PTSD, demonstrating decreased resiliency responses. It has also been suggested that stress affects the oxytocinergic system in a sex-dependent manner (see especially Nishi et al., 2015), which has been extensively documented in animals (Ebner et al., 2000; Cameron et al., 2008).

The studies related to stress reactivity have also shown gender dependence because only women with a history of physical abuse show increased OXT secretion under acute stress (Seltzer et al., 2014). When gender was not experimentally controlled, a decrease in OXT secretion was observed (Munro et al., 2013). Previous studies have shown that men and women differ with regard to the release of endogenous OXT following behavioral paradigms. They also differ behaviorally after exogenous OXT administration (Finkelhor et al., 1990; Heim et al., 2000; Seng et al., 2014; Feng et al., 2015; Koch et al., 2016a,b).

One possible explanation of the above gender effects might involve the biochemical differences in the number of available OXT receptors between men and women as well as the binding affinity of these receptors in specific neural networks (Uhl-Bronner et al., 2005; Hoge et al., 2014). Furthermore, the different phases of the menstrual cycle can also influence OXT levels as well as the HPA axis, conferring more or less impairment (Altemus et al., 2001). This effect is seen because of the role of estrogen, which regulates the production of OXT receptors and the possible release of this hormone (Williams et al., 1985; Wigger and Neumann, 1999). In contrast, evidence also shows that androgens inhibit OXT release under stress (Young et al., 1997). According to Taylor et al. (2006), OXT might also favor the
“tend and befriend” response in women and the “fight or flight” response in men.

The results of the studies that focused on exogenous OXT administration varied by the type of stressor. Regarding ET, the effect of this stressor on the functioning of the brain circuits was demonstrated. ET favored functional changes in the brain circuits, especially those associated with limbic regions (e.g., the hypothalamus, pgACC, amygdala, and parahippocampal gyrus), which are HPA axis-modulating regions. In this sense, OXT modulates the neural networks to favor an improved response to stress (Meinschmidt and Heim, 2007).

Other studies that examined severe ET, however, seemed to show a null or differential response to exogenous OXT administration that assumes anxiogenic characteristics. This effect might occur because of previous changes in the oxytocinergic circuit as a consequence of the traumatic experience or because of unregulated interactions between the oxytocinergic system and the other neurotransmitter systems (Fan et al., 2014; Grimm et al., 2014).

The effects of exogenous OXT in individuals with RTE at risk for PTSD were unfavorable because they tended to promote anxiogenic effects and did not reduce the risk of developing different pathologies, including PTSD. The results generally showed that OXT favors fear-related responses; this fact is also an accepted finding in studies conducted with healthy participants (Domes et al., 2010).

The above inconsistency can be understood in light of Heinrichs et al. (2004). These authors argue that OXT can produce an extinction effect on aversive memories among healthy participants, depending on the test type used and the relevance of the stimuli to the evaluated patient. According to their findings and the previous literature (Kirsch et al., 2005; Domes et al., 2007; Gamer et al., 2010), the effects of OXT on amygdala reactivity might differ not only because of inter-individual factors such as gender and level of psychopathology but also because of the context (i.e., the emotional content/valence of the stimulus).

The increased number of flashbacks suggests that the administration of OXT in individuals who experienced recent trauma impedes the functioning of the emotional regulation network in response to exposure to situations reminiscent of the trauma. This finding is contrary to the previous literature (Koch et al., 2016b; Sack et al., 2017) showing the potential role that OXT plays in the extinction of traumatic memories. In previous studies, postpartum women (i.e., those with increased OXT production) experience temporary deficits in memory that contribute to the extinction of the aversive memories associated with childbirth (Brindle et al., 1991; Brett and Baxendale, 2001).

On the other hand, the continuous use of OXT for 8 days had positive effects only for individuals with severe symptoms/traumas, again suggesting an interdependence of inter-individual variables. Thus, the studies of the use of OXT to prevent PTSD suggest the need for caution because of the possible negative/anxiogenic effects on the one hand and the presence of benefits resulting from OXT administration that occur selectively only in highly symptomatic individuals on the other.

Finally, for individuals with PTSD, the effects of acute OXT administration were favorable. Considering that PTSD is associated with hyperactivity in the amygdala and other limbic brain structures when presented with negative emotional stimuli (Brunetti et al., 2010; Frijling, 2017), OXT favored the reestablishment of the functioning of different neural networks associated with fear control and extinction responses, thereby favoring better emotional control and cognitive performance through the reduction of hypervigilance, avoidance, anhedonia, and emotional salience to fear.

Importantly, these studies showed weak effects for the control groups that were composed mostly of participants with RTEs (i.e., those exposed to trauma but without PTSD). This result reinforces the findings of the previous group of studies that indicated the lack of benefits for this specific sample. These data also reinforce the positions of Koch et al. (2016a,b) and Bartz et al. (2010) who argued that OXT is beneficial only for individuals with impaired fear regulation and social functioning. However, Palgi et al. (2016) examined participants exposed to trauma, with or without PTSD, and found that they benefited from the administration of OXT in terms of their compassion response.

Regarding the studies related to the different polymorphisms, the present review suggested that variations and polymorphisms are intrinsically associated with the changes in the stress resilience mechanism and the experience of traumatic experiences (Cicchetti and Rogosch, 2012). One possible explanation is that individuals with polymorphic variations in the OXT receptor gene due to random changes in the position of the amino acids are more vulnerable to the development of disorders resulting from traumatic experiences throughout the lifespan (Feldman et al., 2016).

The genetic polymorphisms that showed more damage were those associated with the GG genotype at the OXT genetic receptors rs53576, rs139832701, and rs11131147. These genetic variations might alter receptor configuration and develop changes at density, alter the number of receptors available in the central nervous system (Champagne and Curley, 2009; Skuse and Gallagher, 2011), and might develop alterations on OXT binding affinity to the receptor, thereby altering the effects/functions of OXT in the body.

Interestingly, Dannlowski et al. (2016) found that the presence of the GG genotype allied with ET might increase the vulnerability to the greater activation of limbic areas during the visualization of positive/negative faces; however, these authors indicated that such a vulnerability would also depend on environmental experiences and the complex interaction between genes and environment. Therefore, stable environments should provide more positive signals and lead to beneficial development, whereas threatening environments should result in detrimental effects on emotional development and lead to vulnerability to psychiatric disorders.

Champagne and Curley (2009) also suggested that participants exposed to stressful environments undergo DNA methylation processes that modify the positions of nitrogenous bases, thereby resulting in the modification of certain gene expression. Therefore, the binding affinity of OXT to the
receptor might be altered, thereby reducing the amount of bound OXT and leading to a reduction in the function of OXT in the organism. This effect would alter the individual's vulnerability to the effects of stress (Jack et al., 2012).

On the other hand, one study (Sippel et al., 2017) indicated that polymorphisms on the OXTR rs53576 receptor gene with the presence of a single A’ allele might contribute to the formation of insecure attachments among individuals with PTSD. This finding is relevant because this attachment style is associated with a reduced response to PTSD treatment. Such dysfunction might translate into a neural mechanism that could predispose individuals to negative assessments of their social environment stimuli, which might compromise the stress-reducing effects of social support and exacerbate the effect of cognitive representations of relationships as threatening, mistrustful, or unstable.

Finally, the other genetic polymorphisms (OXTR rs2254298 and rs2268498) had no effect on PTSD symptoms or their development (Dunn et al., 2014) nor on the depression or anxiety symptoms of participants with ET (Tollenaar et al., 2017). One possible explanation is that the participants were recruited from a specific population of low-income non-Hispanics blacks who were exposed to Hurricane Katrina (Dunn et al. 2014) or healthy participants with ET (Tollenaar et al., 2017). Large samples are required for candidate gene studies.

CONCLUDING REMARKS

This systematic review regarding the relationship between OXT and emotional trauma in humans revealed that reductions in endogenous OXT levels are more strongly associated with the presence of severe and recurrent ET and PTSD.

The results also show that the acute effect of OXT in victims of ET tends to be anxiolytic, but only in those with less severe forms of ET. Among individuals with severe ET, OXT does not seem to have anxiolytic properties and may even increase anxiety. The same holds true for victims of RTE, since OXT has been reported to increase the re-experience of traumatic symptoms. In patients with PTSD, OXT reestablished the function of different neural networks associated with fear control and extinction.

Genetic studies showed that the presence of the GG genotype in the OXTR rs53576 gene polymorphism associated with the experience of ET or PTSD is related to more negative outcomes, whereas the presence of the AA genotype appears to have a protective role.

It should be noted that the relationship between OXT and traumatic situations is mediated by different contextual and inter-individual variables that might predict more or less favorable outcomes. Thus, a stable environment, less severe forms of trauma, and the female gender are associated with resiliency responses.

The studies included in the review have limitations that call for caution in the interpretation and generalization of their findings, including (a) small sample sizes recruited from extremely specific contexts; (b) poor control of confounding variables such as gender and ET severity; (c) highly variable age groups; (d) cross-sectional designs; (e) possible differences between chronic and acute administration of OXT in RCT; and (f) lack of evaluation of both central and peripheral levels of OXT to determine whether differences exist [most of the studies on this topic assessed endogenous levels in peripheral fluids (e.g., blood plasma, urine, saliva) and the methodological validity of these techniques is controversial (Wotjak et al., 1998; Seltzer et al., 2014; Valstad et al., 2017)].

For future studies, we suggest the adoption of longitudinal designs to establish causal relationships; the recruitment of large samples to control for gender, age, and sex hormones; the conduction of RCTs with chronic and/or prolonged administration of OXT; the investigation of other polymorphisms that might interfere with or contribute to increased susceptibility to the harmful effects of traumatic experiences; and the performance of epigenetic investigations on the association between ET, RTE, and PTSD with OXT.

AUTHOR CONTRIBUTIONS

All authors listed have made a substantial, direct and intellectual contribution to the work, and approved it for publication.

ACKNOWLEDGMENTS

The National Council for Technological and Scientific Development (CNPq grant numbers 2015/12548-6 and 301321/2016-7), the Spanish Biomedical Research Networking Center for Mental Health (CIBERSAM), and the Comissionat per a Universitats i Recerca del DIUE, Generalitat de Catalunya, Spain (grant number 2014SGR1435) supported this study.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fphar.2018.00154/full#supplementary-material
Bartz, J. A., Zaki, J., Bolger, N., Hollander, E., Ludwig, N. N., Kolevzon, A., et al. (2010). Oxytocin selectively improves empathic accuracy. *Psychol. Sci.* 21, 1426–1428. doi: 10.1177/0956797610383439

Boyce, W. T., and Ellis, B. (2005). Biological sensitivity to context: I. An evolutionary–developmental theory of the origins and functions of stress reactivity. *Dev. Psychopathol.* 17, 271–301. doi: 10.1017/S095457940500145

Bradley, R., Westen, D., Mercer, K. B., Bender, E. B., Jovanovic, T., Crain, D., et al. (2011). Association between childhood maltreatment and adult emotional dysregulation in a low-income, urban, African American sample: moderation by oxytocin receptor gene. *Dev. Psychopathol.* 23, 439–452. doi: 10.1017/S0954579411000162

Bremner, J. D., Vermetten, E., and Mazure, C. M. (2000). Development and preliminary psychometric properties of an instrument for the measurement of childhood trauma: the early trauma inventory. *Depress. Anxiety* 12, 1–12. doi: 10.1002/(SICI)1098-7316(2000)12:1<1::AID-DAI1>3.0.CO;2-W

Brett, M., and Baxendale, S. (2001). Motherhood and memory: a review. *Psychoneuroendocrinology* 26, 339–362. doi: 10.1016/S0306-4530(00)00008-8

Brunetti, M., Sepede, G., Mingoia, G., Catani, C., Ferretti, A., Merla, A., et al. (2010). Elevated response of human amygdala to neutral stimuli in mild post-traumatic stress disorder: neural correlates of generalized emotional response. *Neuroscience* 168, 670–679. doi: 10.1016/j.neuroscience.2010.04.024

Cameron, N. M., Fish, E. W., and Meaney, M. J. (2008). Maternal influences on the emotional dysregulation in a low-income, urban, African American sample: a prospective analysis of low income adults. *J. Affect. Disord.* 125, 243–249. doi: 10.1016/j.jad.2013.09.018

Ebnert, K., Wotjak, C. T., Landgraf, R., and Engelmann, M. (2000). A single social defeat experience selectively stimulates the release of oxytocin, but not vasopressin, within the septal brain area of male rats. *Brain Res.* 872, 87–92. doi: 10.1016/S0006-8993(00)02464-1

Eidelman-Rothman, M., Goldstein, A., Levy, J., Weisman, O., Schneiderman, I., Mankuta, D., et al. (2015). Oxytocin affects spontaneous neural oscillations in trauma-exposed war veterans. *Front. Behav. Neurosci.* 9:165. doi: 10.3389/fnbeh.2015.00165

Elzinga, B. M., Roelofs, K., Tollenaar, M. S., Bakvis, P., van Pelt, J., and Spinholven, P. (2008). Diminished cortisol responses to psychosocial stress associated with lifetime adverse events: a study among healthy young subjects. *Psychoneuroendocrinology* 33, 227–227. doi: 10.1016/j.psyneuen.2007.11.004

Fan, Y., Herrera-Melendez, A. L., Pestke, K., Feeser, M., Aust, S., Otte, C., et al. (2014). Early life stress modulates amygdala-prefrontal functional connectivity: implications for oxytocin. *Hum. Brain Mapp.* 35, 5328–5339. doi: 10.1002/hbm.22553

Faravelli, C., Sauro, C. L., Godini, L., Lelli, L., Benni, L., Pietrini, F., et al. (2012). Childhood stressful events, HPA axis and anxiety disorders. *World J. Psychiatry* 2, 13–25. doi: 10.5498/wjp.v2.i1.13

Feldman, R., Monakhov, M., Pratt, M., and Ebstein, R. P. (2016). Oxytocin pathway genes: evolutionary ancient system impacting on human affiliation, sociality, and psychopathology. *Biol. Psychiatry* 79, 174–184. doi: 10.1016/j.biopsych.2015.08.008

Feng, C., Hackett, P. D., DeMarco, A. C., Chen, X., Stair, S., Haroon, E., et al. (2015). Oxytocin and vasopressin effects on the neural response to social cooperation are modulated by sex in humans. *Brain Imaging Behav.* 9, 754–764. doi: 10.1007/s11682-014-9333-9

Finkelhor, D., Hotaling, G., Lewis, I., and Smith, C. (1990). Sexual abuse in a national survey of adult men and women: prevalence, characteristics, and risk factors. *Child Abuse Negl.* 14, 19–28. doi: 10.1016/0145-2134(90)90077-7

Fischer- Shofty, M., Shamay-Toory, S. G., Harari, H., and Levkovitz, Y. (2010). The effect of intranasal administration of oxytocin on fear recognition. *Neuropsychologia* 48, 179–184. doi: 10.1016/j.neuropsychologia.2009.09.003

Frijling, J. L. (2017). Preventing PTSD with oxytocin: effects of oxytocin administration on fear neurocircuity and PTSD symptom development in recently trauma-exposed individuals. *Eur. J. Psychotraumatol.* 8:1302652. doi: 10.1080/20080198.2017.1302652

Frijling, J. L., van Zuiden, M., Koch, S. B., Nawijn, L., Veltman, D. J., and Olff, M. (2016a). Intranasal oxytocin affects amygdala functional connectivity after trauma script-driven imagery in distressed recently trauma-exposed individuals. *Neuropsychopharmacology* 41, 1286–1296. doi: 10.1038/npp.2015.278

Frijling, J. L., van Zuiden, M., Koch, S. B., Nawijn, L., Veltman, D. J., and Olff, M. (2016b). Effects of intranasal oxytocin on amygdala reactivity to emotional faces in recently trauma-exposed individuals. *Soc. Cogn. Affect. Neurosci.* 11, 327–336. doi: 10.1093/scan/nsv116

Frijling, J. L., Zuiden, M., Nawijn, L., Koch, S. B. J., Neumann, I. D., Veltman, D. J., et al. (2015). Salivary oxytocin and vasopressin levels in police officers with and without post traumatic stress disorder. *J. Neuroendocrinol.* 27, 743–751. doi: 10.1111/jne.12300

Gamer, M., Zuerbowski, B., and Büchel, C. (2010). Different amygdala subregions mediate valence-related and attentional effects of oxytocin in humans. *Proc. Natl. Acad. Sci. U.S.A.* 107, 9400–9405. doi: 10.1073/pnas.1009985107

Gimpl, G., and Fahrenholz, F. (2001). The oxytocin receptor system: structure, function, and regulation. *Physiol. Rev.* 81, 629–683. doi: 10.1152/physrev.2001.81.2.629

Goldman-Mellor, S., Hamer, M., and Steptoe, A. (2012). Early-life stress and recurrent psychological distress over the lifespan predict divergent cortisol reactivity patterns in adulthood. *Psychoneuroendocrinology* 37, 1755–1768. doi: 10.1016/j.psyneuen.2012.03.010

Gonzalez, A., Jenkins, J. M., Steiner, M., and Fleming, A. S. (2009). The relation between life adversity, cortisol awakening response and diurnal cortisol levels in postpartum women. *Psychoneuroendocrinology* 34, 76–86. doi: 10.1016/j.psyneuen.2008.08.012
Meinschmidt, G., and Heim, C. (2007). Sensitivity to intranasal oxytocin in adult men with early parental separation. *Biol. Psychiatry* 61, 1109–1111. doi: 10.1016/j.biopsych.2006.09.007

Mirescu, C., Peters, J. D., and Gould, E. (2004). Early life experience alters response of adult neurogenesis to stress. *Nat. Neurosci.* 7, 841–846. doi: 10.1038/nn1290

Mizuki, R., and Fujiiwara, T. (2015). Association of oxytocin level and less severe forms of childhood maltreatment history among healthy Japanese adults involved with child care. *Front. Behav. Neurosci.* 9:138. doi: 10.3389/fnbeh.2015.00138

Mizushima, S. G., Fujisawa, T. X., Takiguchi, S., Kumazaki, H., Tanaka, S., and Tomoda, A. (2015). Effect of the nature of subsequent environment on oxytocin and cortisol secretion in maltreated children. *Front. Psychiatry* 6:173. doi: 10.3389/fpsyg.2015.00173

Moher, D., Liberati, A., and Tetzlaff, J. (2009). Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. *PLoS Med.* 6:e1000097. doi: 10.1371/journal.pmed.1000097

Moher, D., Schulz, K. F., and Altman, D. G. (2001). The CONSORT statement: revised recommendations for improving the quality of reports of parallel-group randomised trials. *Lancet* 357, 1191–1194. doi: 10.1016/S0140-6736(00)04337-3

Mohiyeddini, C., Opaka-Jaffry, J., and Gross, J. J. (2014). Emotional suppression explains the link between early life stress and plasma oxytocin. *Anxiety Stress Coping* 27, 466–475. doi: 10.1080/10615834.2014.887696

Munro, M. L., Brown, S. L., Pourniajai-Nazarloo, H., Carter, C. S., Lopez, W. D., and Seng, J. S. (2013). In search of an adult attachment stress provocation to measure effect on the oxytocin system a pilot validation study. *J. Am. Psychiatr. Nurses Assoc.* 19, 180–191. doi: 10.1097/JNA.0b013e31830f3134

Myers, A. J., Williams, L., Gatt, J. M., McAuley-Clark, E. Z., Dobson-Stone, C., Schofield, P. R., et al. (2014). Variation in the oxytocin receptor gene is associated with increased risk for anxiety, stress and depression in individuals with a history of exposure to early life stress. *J. Psychiatr. Res.* 59, 93–100. doi: 10.1016/j.jpsychires.2014.08.021

Nawijn, L., van Zuiden, M., Koch, S. B., Frijling, J. L., Veltman, D. J., and Offl, M. (2016). Intranasal oxytocin enhances neural processing of monetary reward and loss in post-traumatic stress disorder and traumatized controls. *Psychoneuroendocrinology* 66, 228–237. doi: 10.1016/j.psyneuen.2016.01.020

Nawijn, L., van Zuiden, M., Koch, S. B., Frijling, J. L., Veltman, D. J., and Offl, M. (2017). Intranasal oxytocin increases neural responses to social reward in post-traumatic stress disorder. *Soc. Cogn. Affect. Neurosci.* 12, 212–223. doi: 10.1093/scan/nsw123

Neumann, I. D., and Landgraf, R. (2012). Balance of brain oxytocin and vasopressin: implications for anxiety, depression, and social behaviors. *Trends Neurosci.* 35, 649–659. doi: 10.1016/j.tins.2012.08.004

Nicolson, N. A., Davis, M. C., Kruszewski, D., and Zautra, A. J. (2010). Childhood maltreatment and diurnal cortisol patterns in women with chronic pain. *Psychosom. Med.* 72, 471–480. doi: 10.1016/j.psychos.2010.03.009

Nishi, D., Hashimoto, K., Noguchi, H., Kim, Y., and Matsuoka, Y. (2015). Serum oxytocin, posttraumatic coping and C-reactive protein in motor vehicle accident survivors by gender. *Neuropsychobiology* 71, 196–201. doi: 10.1159/000382021

Olff, M. (2012). Bonding after trauma: on the role of social support and the oxytocin system in traumatic stress. *Eur. J. Psychotraumatol.* 3:18597. doi: 10.3402/ejpt.v3i0.18597

Opaka-Jaffry, J., and Mohiyeddini, C. (2012). Experience of stress in childhood negatively correlates with plasma oxytocin concentration in adult men. *Stress* 15, 1–10. doi: 10.3109/10253890.2011.560309

Ozbay, O., Fitterling, H., Charney, D., and Southwick, S. (2008). Social support and resilience to stress across the life span: a neurobiologic framework. *Soc. Cogn. Affect. Neurosci.* 3, 52–60. doi: 10.1093/scan/nsp087

Palgi, S., Klein, E., and Shamay-Tsoory, S. G. (2016). Oxytocin improves compassion toward women among patients with PTSD. *Psychoneuroendocrinology* 64, 143–149. doi: 10.1016/j.psyneuen.2015.11.008

Pierrehumbert, B., Torrisi, R., Lauffer, D., Hallon, Ø., Ansermet, F., and Popovic, M. B. (2010). Oxytocin response to an experimental psychosocial challenge in adults exposed to traumatic experiences during childhood or adolescence. *Neuroscience* 166, 168–177. doi: 10.1016/j.neuroscience.2009.12.016
Reijnen, A., Geuze, E., and Vermetten, E. (2017). Individual variation in plasma oxytocin and vasopressin levels in relation to the development of combat-related PTSD in a large military cohort. *J. Psychiatr. Res.* 94, 88–95. doi: 10.1016/j.jpsychires.2017.06.010

Rimmele, U., Malberg, J. E., Potestio, L., Ping, J., Boikess, S., Luo, B., et al. (2006). Anxiolytic-like activity of oxytocin in male mice: behavioral and autonomic evidence, therapeutic implications. *Psychopharmacology* 185, 218–225. doi: 10.1007/s00213-005-0293-z

Ross, H. E., and Young, L. J. (2009). Oxytocin and the neural mechanisms regulating social cognition and affiliative behavior. *Front. Neuroendocrinol.* 30, 534–547. doi: 10.1016/j.yfrne.2009.05.004

Sack, M., Cillien, M., and Hopper, J. W. (2012). Acute dissociation and cardiac reactivity to script-driven imagery in trauma-related disorders. *Eur. J. Psychotraumatol.* 3:17419. doi: 10.3402/ejpt.v3i0.17419

Sack, M., Spieler, D., Wizelman, L., Epple, G., Stich, J., Zaba, M., et al. (2017). Intranasal oxytocin reduces provoked symptoms in female patients with posttraumatic stress disorder despite exerting sympatimmetic and positive chronotropic effects in a randomized controlled trial. *BMC Med.* 15:40. doi: 10.1186/s12916-017-0801-0

Seng, J. S. (2010). Posttraumatic oxytocin dysregulation: is it a link among posttraumatic self disorders, posttraumatic stress disorder, and pelvic visceral dysregulation conditions in women?. *J. Trauma Dissoc.* 11, 387–406. doi: 10.1080/15299732.2010.496075

Seng, J. S., D’andrea, W., and Ford, J. D. (2014). Complex mental health sequelae of psychological trauma among women in prenatal care. *Psychol. Trauma* 6, 41–49. doi: 10.1037/tpt00013467

Sippel, L. M., Han, S., Watkins, L. E., Harpaz-Rotem, I., Southwick, S. M., Krystal, J. H., et al. (2017). Oxytocin receptor gene polymorphisms, attachment, and PTSD: results from the national health and resilience in veterans study. *J. Psychiatr. Res.* 94, 139–147. doi: 10.1016/j.jpsychires.2017.07.008

Skuse, D. H., and Gallagher, L. (2011). Genetic influences on social cognition. *Pediatr. Res.* 69, 85–91. doi: 10.1203/PDR.0b013e318212f662

Smith, S. M., and Vale, W. W. (2006). The role of the hypothalamic-pituitary-adrenal axis in neuroendocrine responses to stress. *Dialog. Clin. Neurosci.* 8, 383–395.

Taylor, S. E., Gonzaga, G. C., Klein, L. C., Hu, P., Greendale, G. A., and Seeman, T. E. (2006). Relation of oxytocin to psychological stress responses and hypothalamic-pituitary-adrenocortical axis activity in older women. *Psychosom. Med.* 68, 238–245. doi: 10.1097/01.psyc.0000203242.95990.74

Tollenaar, M. S., Molendijk, M. L., Penninx, B. W., Milaneschi, Y., and Antypa, N. (2017). The association of childhood maltreatment with depression and anxiety is not moderated by the oxytocin receptor gene. *Eur. Arch. Psychiatry Clin. Neurosci.* 267, 517–526. doi: 10.1007/s00406-017-0784-z

Uhl-Bornuner, S., Walsiterperger, E., Martinez-Lorenzana, G., Cendes, L. M., and Freund-Mercier, M. J. (2005). Sexually dimorphic expression of oxytocin binding sites in forebrain and spinal cord of the rat. *Neuroscience* 135, 147–154. doi: 10.1016/j.neuroscience.2005.05.025

Unternaehrer, E., Luers, P., Mill, J., Dempster, E., Meyer, A. H., and Staehti, S. (2012). Dynamic changes in DNA methylation of stress-associated genes (OXTR, BDNF) after acute psychosocial stress. *Transl. Psychiatry* 2:e150. doi: 10.1038/tp.2012.77

Valtas, M., Alvares, G. A., Egkoud, M., Matziorinis, A. M., Andreasen, O. A., Westlye, L. T., et al. (2017). The correlation between central and peripheral oxytocin concentrations: a systematic review and meta-analysis. *Neurosci. Biobehav. Rev.* 78, 117–124. doi: 10.1016/j.neubiorev.2017.04.017

Van Zuiden, M., Frijling, J. L., Nawijn, L., Koch, S. B., Goslings, J. C., Luitse, J. S., et al. (2017). Intranasal oxytocin to prevent posttraumatic stress disorder symptoms: a randomized controlled trial in emergency department patients. *Biol. Psychiatry* 81, 1030–1040. doi: 10.1016/j.biopsych.2016.11.012

Veennema, A. H. (2012). Toward understanding how early-life social experiences alter oxytocin-and vasopressin-regulated social behaviors. *Horm. Behav.* 61, 304–312. doi: 10.1016/j.yhbeh.2011.12.002

Von Elm, E., Altman, D. G., Egger, M., Pocock, S. J., Gotzsche, P. C., and Vandenbroucke, J. P. (2007). The Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) statement: guidelines for reporting observational studies. *Prev. Med.* 45, 247–251. doi: 10.1016/j.yjmep.2007.08.012

Wigger, A., and Neumann, I. D. (1999). Periodic maternal deprivation induces gender-dependent alterations in behavioral and neuroendocrine responses to emotional stress in adult rats. *Physiol. Behav.* 66, 293–302. doi: 10.1016/S0306-4522(97)00300-X

Williams, T. D. M., Carter, D. A., and Lightman, S. L. (1985). Sexual dimorphism in the posterior pituitary response to stress in the rat. *Endocrinology* 116, 738–740. doi: 10.1210/endo-116-2-738

Wotjak, C. T., Ganster, J., Kohl, G., Holsober, F., Landgraf, R., and Engelmann, M. (1998). Dissociated central and peripheral release of vasopressin, but not oxytocin, in response to repeated swim stress: new insights into the secretory capacities of peptidergic neurons. *Neuroscience* 85, 1209–1222. doi: 10.1016/S0306-4522(97)00683-0

Yehuda, R., Bierer, L. M., Schmeidler, J., Aferiat, D. H., Breslau, L., and Dolan, S. (2000). Low cortisol and risk for PTSD in adult offspring of Holocaust survivors. *Am. J. Psychiatry* 157, 1252–1259. doi: 10.1176/appi.ajp.157.8.1252

Yoshida, M., Takayanagi, Y., Inoue, K., Kimura, T., Young, L. J., Onaka, T., et al. (2009). Evidence that oxytocin exerts anxiolytic effects via oxytocin receptor expressed in serotonergic neurons in mice. *J. Neurosci.* 29, 2259–2271. doi: 10.1523/JNEUROSCI.5939-08.2009

Young, L. J., Muns, S., Wang, Z., and Insel, T. R. (1997). Changes in oxytocin receptor mRNA in rat brain during pregnancy and the effects of estrogen and interleukin. *J. Neuroendocrinol.* 9, 859–865. doi: 10.1111/j.1365-2826.1997.00654.x

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2018 Donadon, Martín-Santos and Osório. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.