Oxytocin Receptor Gene (OXTR) and Father Support Interact to Predict Depressive Symptoms Postpartum

Parambir Bhatti  
*Chapman University*

Taylor Delaney  
*Chapman University*

Michael Poulin  
*University at Buffalo*

Jennifer Hahn-Holbrook  
*Chapman University*, hahnholb@chapman.edu

Follow this and additional works at: [https://digitalcommons.chapman.edu/psychology_articles](https://digitalcommons.chapman.edu/psychology_articles)

Part of the Maternal and Child Health Commons, Mental Disorders Commons, Other Psychiatry and Psychology Commons, Psychological Phenomena and Processes Commons, and the Women's Health Commons

**Recommended Citation**

[https://doi.org/10.1016/j.biopsycho.2019.03.015](https://doi.org/10.1016/j.biopsycho.2019.03.015)

This Article is brought to you for free and open access by the Psychology at Chapman University Digital Commons. It has been accepted for inclusion in Psychology Faculty Articles and Research by an authorized administrator of Chapman University Digital Commons. For more information, please contact laughtin@chapman.edu.
Oxytocin Receptor Gene (OXTR) and Father Support Interact to Predict Depressive Symptoms Postpartum

Comments
NOTICE: this is the author's version of a work that was accepted for publication in Biological Psychology. Changes resulting from the publishing process, such as peer review, editing, corrections, structural formatting, and other quality control mechanisms may not be reflected in this document. Changes may have been made to this work since it was submitted for publication. A definitive version was subsequently published in Biological Psychology, volume 147, in 2019. DOI: 10.1016/j.biopsycho.2019.03.015

The Creative Commons license below applies only to this version of the article.

Creative Commons License

This work is licensed under a Creative Commons Attribution-Noncommercial-No Derivative Works 4.0 License.

Copyright
Elsevier

This article is available at Chapman University Digital Commons: https://digitalcommons.chapman.edu/psychology_articles/147
Accepted Manuscript

Title: Oxytocin Receptor Gene (OXTR) and Father Support Interact to Predict Depressive Symptoms Postpartum

Authors: Parambir Bhatti, Taylor Delaney, Michael Poulin, Jennifer Hahn-Holbrook

PII: S0301-0511(18)30122-4
DOI: https://doi.org/10.1016/j.biopsycho.2019.03.015
Reference: BIOPSY 7686

To appear in:

Received date: 12 February 2018
Revised date: 26 March 2019
Accepted date: 26 March 2019

Please cite this article as: Bhatti P, Delaney T, Poulin M, Hahn-Holbrook J, Oxytocin Receptor Gene (OXTR) and Father Support Interact to Predict Depressive Symptoms Postpartum, Biological Psychology (2019), https://doi.org/10.1016/j.biopsycho.2019.03.015

This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.
OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION

Oxytocin Receptor Gene (OXTR) and Father Support Interact to Predict Depressive Symptoms Postpartum

Parambir Bhatti\textsuperscript{a,c}, Taylor Delaney\textsuperscript{a,c}, Michael Poulin\textsuperscript{b}, Jennifer Hahn-Holbrook \textsuperscript{a,d}

\textsuperscript{a} Chapman University, Department of Psychology, One University Drive, Orange, California 92866, USA.
\textsuperscript{b} University at Buffalo, Department of Psychology, Park Hall 206, Buffalo, NY 14260, USA.
\textsuperscript{c} A.T. Still University, School of Osteopathic Medicine, 5850 East Still Circle, Mesa, Arizona 85206, USA.
\textsuperscript{d} University of California, Merced, Department of Psychology, 5200 Lake Road, Merced, California 95340, USA.

*Correspondence regarding this manuscript can be addressed to J. Hahn-Holbrook at: 5200 N Lake Road, Merced, CA 95323. Email: jhahn-holbrook@ucmerced.edu

Highlights

- \textit{OXTR} rs53576 SNP moderated the effect of father support on PPD symptoms
- Father support was most protective for mothers with the GG genotype
- Father support was a stronger predictor of PPD symptoms than mother or family support
- The stress-buffering effect of father support was only found in GG genotype mothers

Abstract

Postpartum depression (PPD) is a debilitating mental illness affecting approximately 13% of mothers after birth. Both genetic and psychosocial factors contribute to PPD risk, but very little is known about how these factors interact. We tested whether the rs53576 polymorphism in the oxytocin receptor (\textit{OXTR}) gene accounts for variation in the impact of low social support...
OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION

as a risk factor for depression among mothers during the perinatal period. New mothers ($N = 220$) provided saliva or blood DNA samples and completed surveys assessing PPD symptoms and perceived social support. In a significant interaction, social support from the baby’s father predicted PPD symptoms to a greater extent among mothers with the GG compared to AG and AA genotypes. These results add to converging evidence that variation in OXTR rs53576 moderates the impact of the social environment on PPD.

**Keywords:** Postpartum depression, Social support, Oxytocin receptor gene, rs53576

1. Introduction

Postpartum depression (PPD) is a prevalent mental health disorder suffered by approximately 13% of women in the United States and 17% of women globally (Hahn-Holbrook, Cornwell-Hinrichs & Anaya, 2018; O'Hara et al., 1996). The majority of cases of PPD go undiagnosed (Dennis & Chung-Lee, 2006) and even mild or moderate symptoms of depression are deleterious for mothers and potentially harmful to the infant and the family (Brummelte & Galea, 2016; Netsi et al., 2018). PPD is diagnosed through clinical interviews, although self-report questionnaires are more commonly used to measure depressive symptoms (Yim et al., 2015). Given the consequences of PPD and its symptoms, hundreds of studies have examined biological and psychological risk factors.

Poor social support numbers among the most consistent risk factors for PPD symptoms (for a review, see, Yim et al., 2015). Indeed, individuals who report higher levels of social support live longer and are at reduced risk of depression, among many other stress-related
maladies such as diabetes, cardiac illness, or infection (for review, see Holt-Lunstad et al., 2010). Social support is thought to convey health benefits both directly, through its ability to elevate mood, and indirectly by buffering individuals against stress-induced wear and tear on the body (Cohen & Wills, 1985; for a review, see Uchino, 2006). Social support appears particularly vital during the postpartum period, as a meta-analysis of 59 studies of 12,810 postpartum women identified poor social support as one of the strongest predictors of PPD symptoms (O'Hara et al., 1996), with comparable results obtained in a larger and more recent meta-analysis of over 70 studies encompassing approximately 25,000 postpartum mothers (Robertson et al., 2004).

Subjective perceptions of oneself as the beneficiary of social support, distinct from objective metrics, have been found particularly protective against PPD symptoms (Yim et al., 2015). Although the perceived support of family and friends also appears beneficial, feeling supported by the baby’s father may hold special significance. For example, in a study of 12,361 postpartum Australian women, Milgrom and colleagues (2008) found that social support from the mother’s partner protected against PPD symptoms to a greater extent than support from other family members. In consideration of these prior findings, and of the clear functional relevance of partner support to successful parenting, we anticipated that any observed relationships between oxytocin mechanisms and social support would be particularly evident with regard to father support.

The neuropeptide hormone oxytocin facilitates social relationships in humans and other mammals (Carter & Keverne, 2002; Insel, 2010; McCall & Dinger, 2012), attuning the brain to the social world (Bartz, 2016) and helping to biologically mediate the attenuating effects of affiliation on anxiety (e.g., Labuschagne et al., 2010; Windle, Shanks, Lightman, & Ingram, 1997). Oxytocin administration can bolster trust, empathy and interpersonal communication in affiliative contexts (Kosfeld et al., 2005; Domes et al., 2007; Ditzen et al., 2009). Conversely,
oxytocin appears to heighten feelings of exclusion when social support is perceived to be absent (Bartz, 2016).

Individual differences in social functioning have been linked to genetic variation in the oxytocin receptor system (Thompson, Parker, Hallmayer, Waugh, & Gotlib, 2011). Single nucleotide polymorphisms (SNPs) in the OXTR gene modulate the activity and peptide binding abilities of circulating levels of oxytocin in the body. A growing body of research has focused on one SNP in particular—rs53576. Polymorphisms in rs53576 involve a guanine (G) to adenine (A) substitution in the third intron of the OXTR gene. Because people inherit one copy of this SNP from each parent, an individual’s genotype can be homozygous for the A allele (AA), heterozygous (GA), or homozygous for the G allele (GG). Variations in rs53576 have been found to correlate with the volume of the hypothalamus and the functional and structural connectivity of the hypothalamus to the amygdala and the dorsal anterior cingulate cortex (Tost et al., 2010), regions implicated in PPD-relevant outcomes such as social distress and stress reactivity (see Saphire-Bernstein et al., 2011, for a review).

Consistent with the hypothesized role of oxytocin, variation in rs53576 has been linked to key social cognitive functions (for reviews, see Gong et al., 2017; Li et al., 2015). For example, the number of G alleles (0 to 2) someone carries positively correlates with parental sensitivity (Bakersman-Kranenburg and van Ijzendoorn, 2008), attachment security (Chen et al., 2011), emotion detection (Rodrigues et al., 2009), prosocial behavior (Poulin, Holman, & Buffone, 2012), optimism and self-esteem (Saphire-Bernstein et al., 2011, but see Cornelis et al., 2012 for a null result). However, in adverse social contexts, rs53576 G allele carriers may be vulnerable to adverse outcomes. For instance, in a study of 288 university students, McQuaid and colleagues (2013) found that maltreatment in early life resulted in higher levels of depressive
OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION

symptoms among G allele carriers (GG/GA genotypes), whereas AA genotypes exposed to comparable maltreatment evinced no such association, and Bradley et al. (2013) obtained similar results for G carriers in a study of 971 African-American adults living in unstable homes with low levels of familial support. Together, these overall findings suggest that new mothers with rs53576 G alleles may be more attuned to the social world, and hence either relatively protected or more susceptible to PPD symptoms contingent on their social circumstances.

Here, we assessed whether rs53576 moderates the association between low social support (particularly from fathers) and PPD symptoms. We predicted that the presence of G alleles would predict a new mother’s sensitivity to the quality of social support she receives. In light of prior research, we further predicted that father support would be more protective against PPD symptoms than family support (Milgrom et al., 2008). We tested the number of G alleles (rather than dichotomously contrasting ‘GG versus other’ or ‘AA versus other’) to reduce errors associated with erroneously assuming allele dominance (e.g., Bradley et al., 2011; Ludmer et al., 2015; 2017) that appear to have contributed to problems in past candidate gene research (Duncan & Keller, 2011). In a complementary prediction, we also tested whether individuals homozygous for the A allele in rs53576 would report higher levels of depressive symptoms in comparison to those carrying the G allele given a previous report (see Saphire-Bernstein et al., 2011). Although our a priori predictions concern only rs53576, we also explored the effects of another OXTR polymorphism, rs2268498, linked to pregnancy and difficult labor (Algovik, Kivinen, Peterson, Westgren, & Kere, 2010).

2. Methods

2.1 Participants
A total of 224 perinatal mothers were recruited from farmers’ markets, shopping malls, and mother-infant classes in Orange County, California. To be eligible for the study, participants had to be English- or Spanish-speaking, at least 18 years of age, have an infant under the age of 12 months, not be taking any steroid medications, and not have any previous or current diagnosis of polycystic ovarian syndrome. Four mothers failed to complete the survey packet, leaving a final sample of 220 (53% White, 28% Latina, 11% Asian, 1% African-American/Black, 7% Other or More than one). Seven participants had mothers who were deceased and so did not fill out the maternal support scale, leaving a sample of 213 for analyses involving perceived mother support. Ninety-six percent of the sample was married or living with a partner at the time of data collection.

2.2 Experimental procedure

Once signed informed consent was obtained, participants were provided with a series of questionnaires assessing demographic information (including age, marital status, and income; see Table 1), levels of perceived support, and postpartum depressive symptoms. For the majority of mothers in the study, survey data and saliva samples for DNA genotyping were collected at the field recruitment site. A subset of the sample (N = 23) recruited to participate in an unrelated pilot laboratory study filled out the survey materials online prior to their laboratory session, where they provided blood samples for DNA genotyping. Participant data obtained in the field versus the lab did not differ in terms of social support, depressive symptoms, OXTR G allele frequency, or demographic factors, and are therefore combined for all analysis. Upon completion of the study, participants were debriefed and compensated with $5 cash or a $10 gift certificate to a local baby boutique. All procedures were approved by the Chapman University Institutional Review Board.
OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION

2.3 Measures

**Perceived social support.** Separate seven-item scales were employed to assess the degree of perceived social support from family, the mother’s mother, and the baby’s father, respectively (Turner et al., 1990). The scales each consisted of statements such as: “No matter what happens, I know that my [family/mother/baby’s father] will always be there for me if I need them”; “Sometimes I’m not sure if I can completely rely on the [family/mother/baby’s father] [reverse coded]”; “I know my [family/mother/baby’s father] will always stand by me.” These items were rated on a 4-point Likert scale ranging from 1 (*strongly disagree*) to 4 (*strongly agree*); ratings for each scale were calculated by summing the items, yielding a potential range of 7 to 28, with higher scores indicating greater perceived social support (mother support *α* = .89, family support *α* = .87, father support *α* = .91). This social support measure has been validated for use with pregnant and postpartum mothers, and was previously found to correlate with PPD symptoms (Turner et al., 1983; Collins et al., 1993; Feldman et al., 2000; Hahn-Holbrook et al., 2013). Maternal, father and family social support measures were assessed as individual predictors; a composite measure was also created by z-scoring these three scores and averaging them to create a measure of overall social support (Cronbach’s alpha *α* = .90).

**Postpartum depressive symptoms.** The ten-item Edinburgh Postnatal Depression Scale (EPDS; Cox et al., 1987; Wisner et al., 2002) was utilized to assess participants’ degree of depressive symptoms with regard to mood, anxiety, guilt, sleep issues, and thoughts of self-harm during the past week according to 4-point (0 to 3) scales worded to suit each item, with higher scores indicate greater symptoms of depression. These ratings are summed, yielding a sum score ranging from 0 to 30 (*α* = .84) which may be used as a continuous variable to assess the severity of depressive symptoms (Cox et al., 1987).²
**Perceived stress.** Perceived general life stress was measured according to the Perceived Stress Scale (PSS; Cohen et al., 1983) as a potential confounding variable because perceived stress correlates with both social support and PPD symptoms (see Yim et al., 2015, for a review). This ten-item measure utilizes a 5-point scale (0 to 4); all ratings are summed, yielding a potential range of 0 to 40 ($\alpha = .89$), with higher scores indicating more perceived stress.

**Genotyping.** DNA was collected via both saliva and blood, yielding similar results (James, Panford-Walsh, Birnboim, & Iwasiow, 2013). Genomic DNA was extracted from saliva using Sequenom/Agena iPLEX genotyping procedures; genomic DNA was extracted from whole blood samples using the QIAamp 96 DNA Blood kit according to the manufacturer’s instructions, save that the initial sample of blood was scaled up to 300 µL (Qiagen). The $OXTR$ rs53576 and rs2268498 SNPs were genotyped using MassARRAY genotyping technology (Sequenom/Agena), resulting in the genotypes AA, AG, and GG for rs53576 and CC, CT, and TT for rs2268498.

### 2.4 Analytic Strategy

**Covariate analysis.** Preliminary analyses were carried out to identify potential confounding variables. Previous research, for example, has demonstrated that peoples of Asian descent are less likely to carry the rs53576 G allele than people of European or Hispanic descent (Butovskaya et al., 2016; Kim et al., 2010), that Asians living in the US report lower levels of psychosocial resources, and that psychological reactions to social support can vary by race and ethnicity (Kim et al., 2010; Mojaverian & Kim, 2013). Accordingly, we sought to rule out the possibility that any apparent effects of the rs53576 polymorphism might owe to race/ethnicity. We likewise tested whether demographic factors, levels of social support, or PPD symptoms differed as a function of $OXTR$ genotype. Variables that differed at the level of $p < .10$ as a
function of *OXTR* variants, or that significantly correlated with social support, were included as covariates in the moderation analysis. Linear regressions were used in analyses involving continuous outcome variables and binary logistic regressions were used for categorical variables.

**Social support analysis.** Pearson’s correlations were utilized to test whether support from family, the mother, and/or the baby’s father were associated with PPD symptoms individually. Next, to explore whether each form of social support was a unique predictor of PPD symptoms, all three sources of social support were included in a multivariate linear regression model.

**Moderation analysis.** Multivariate linear regression was used to assess whether the rs53576 genotype moderated the relationship between social support and depressive symptoms postpartum. Similar exploratory analyses were conducted for rs2268498, however, we had no *a priori* predictions for this polymorphism. Cross products were created between rs53576 and rs2268498 genotypes (coded as number of G or T alleles carried, respectively; i.e. 0, 1 or 2) and each of the social support measures. These cross products were then regressed on PPD symptoms in combination with the target *OXTR* polymorphism and the corresponding social support variable. A moderation model for the composite support measure was conducted first, followed by separate models for father, mother, and family support. All variables were checked for normality of distribution before analysis, revealing that social support from the baby’s father and family were both positively skewed. Accordingly, these variables were reverse log-transformed before analysis to improve the normality of these distributions. (Follow-up analyses were also run using the non-transformed variables. The pattern of results using the raw data did not differ from log-transformed models.)
3. Results

Preliminary Analyses

Table 1 reports demographic information on this sample and levels of support and PPD symptoms for each rs53576 and rs2268498 genotype. Analyses indicated that these genotypes were in Hardy-Weinberg equilibrium. Levels of PPD symptoms did not differ as a function of rs53576 ($p = .11$) or rs2268498 ($p = .22$) genotype. Mothers with a CT genotype on rs2268498 reported more perceived family support than mothers with a CC genotype. Women who were married to the infant’s father reported higher levels of perceived support from family and the baby’s father compared to those who were not. There were too few Black mothers to perform racial and ethnic statistical comparisons; however, these mothers were included in all other analysis. White participants had more G alleles on rs53576 than did Hispanic or Asian participants ($p = .01$). Asian participants had more A alleles on rs2268498 than White ($p < .01$) or Hispanic mothers ($p < .05$). Furthermore, White participants reported significantly higher levels of support from the baby’s father than did Hispanic or Asian participants, $ps < .03$. White participants likewise reported more family support than Hispanic participants, $p = .02$, but not Asian participants. Consistent with the well-established relationship between stress and social support, perceived stress was inversely correlated with all three support variables. Given the potentially confounding differences with regard to PPD symptom outcomes orthogonal to rs53576 status, ethnicity/race (White as the contrast reference group), marital status, and perceived stress were included as covariates in all subsequent moderation analysis.

Social Support Analyses

As predicted, and consistent with previous research, social support from family, the mother, and the baby’s father were all inversely correlated with PPD symptoms (see Table 2). To
ascertain which form of social support was most protective, we entered the three sources of social support into a multivariate linear regression model. As anticipated, only father support was a unique predictor of PPD symptoms ($B = -1.97, SE = 0.89, \beta = -0.17, p = .03, 95\% CI [-3.72, -0.21]$) when included alongside mother ($B = -1.82, SE = 1.41, \beta = -0.10, p = .20, 95\% CI [-4.60, 0.96]$) and family support ($B = -1.66, SE = 1.00, \beta = -0.14, p = .10, 95\% CI [-3.63, 0.31]$) in the regression model.

**OXTR rs53576 Moderation Analyses**

**Total Support.** We tested whether OXTR rs53576 moderated the relationship between the composite social support measure (comprised of the average support from father, mother, and family) and depressive symptoms in a regression model. There was a non-significant interaction ($p = .07$) between the composite support measure and OXTR rs53576 ($B = 0.73, SE = 0.41, \beta = -0.24, 95\% CI [-0.07, 1.54], r^2 \text{ change} = 0.01$). Nonetheless, given the appearance of a trend, we explored the simple slopes association between total support for each genotype. Total support predicted 10.5% of the variance in PPD symptoms for GG carriers ($B = -1.96, SE = 0.65, \beta = -0.32, p = .003, 95\% CI [-3.25, -0.67]$), 12.0% for AG carriers ($B = -2.06, SE = 0.56, \beta = -0.35, p < .001, 95\% CI [-3.17, -0.95]$), and only 0.02% for AA carriers ($B = -0.27, SE = 0.92, \beta = -0.05, p = .77, 95\% CI [-2.13, 1.59]$). Categorical slopes comparisons confirmed that the relationship between total support and PPD symptoms was significantly stronger in the GG group than the AA group ($p = .019$), whereas the AG group did not significantly differ from either the AA ($p < .076$) or the GG group ($p = .916$).

**Father Support.** We next assessed our primary moderation prediction regarding perceived support from the baby’s father. As anticipated, the rs53576 polymorphism significantly moderated the association between baby’s father support and PPD symptoms
OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION

(interaction term: B = 2.32, SE = 0.79, β = 0.52, p = .004, 95% CI [0.77, 3.88], r² change=.02; see Figure 1). Simple slopes analyses indicated that, for GG genotypes, perceived support from the baby’s father predicted approximately 10.3% of the variance in PPD symptoms (B = -3.67, SE = 1.27, β = -.33, p = .005, 95% CI [-6.20, -1.15]). In AG genotypes, father support predicted 6.1% of the variance in PPD symptoms (B = -3.03, SE = 1.23, β = -.24, p = .02, 95% CI [-5.48, -0.58]). In AA genotypes, father support was not significantly related to PPD symptoms (B = -1.09, SE = 1.90, β = -.10, p = .57, 95% CI [-4.95, 2.77], r² = .01). Categorical slopes comparisons showed that the relationship between father support and PPD symptoms was significantly stronger in the GG group than the AG (p = .039) or the AA groups (p = .002), while the AG and AA groups did not significantly differ from each other (p = .297).

*Three-way interaction between stress, father support, and OXTR rs53576.* In order to better understand the role of stress in our results, we examined the possibility that the interactive effects of social support and rs53576 would be strongest when mothers reported high (compared to low) stress, consistent with the stress-buffering hypothesis (Cohen & Willis, 1985; Uchino, 2006). In an exploratory analysis, we tested the potential three-way interaction between perceived stress, father support and OXTR genotype. We z-scored father support and perceived stress, and then included stress, father support, rs53576 and a three-way cross-product interaction term (along with the four possible two-way cross-products) in a regression model predicting PPD symptoms. We detected a significant three-way interaction between perceived stress, father support and rs53576 (B = 0.83, SE = 0.31, β = .35, p = .008, 95% CI [.22, 1.44], r² change=.014). In GG genotypes, the inverse relationship between father support and PPD symptoms was indeed stronger when women reported higher (compared to lower) perceived stress (B = -.651, SE = 0.25, β = -.17, p = .010, 95% CI [-1.14, -0.16], r² change= 0.03). No such buffering
interactions between perceived stress and father support were observed in the AG (B = .11, SE = 0.39, β = .03, p = .760, 95% CI [-0.65, 0.88], $r^2$ change= -0.006) or AA genotype groups (B = 1.16, SE = 0.59, β = .23, p = .061, 95% CI [-0.058, 2.37], $r^2$ change= .04).

**Mother Support.** We examined interactions between mother support and OXTR rs53576. The rs53576 genotype did not significantly moderate the effect of maternal support on PPD symptoms (B = .20, SE = 0.31, β = .09, p = .506, 95% CI [-.40, .80], $r^2$ change = -.001).

**Family Support.** We tested the interaction between family support and OXTR rs53576. The rs53576 genotype did not significantly moderate the effect of family support on PPD symptoms (B = -.017, SE = 0.32, β = -.01, p = .958, 95% CI [-.65, .62], $r^2$ change = -.002).

**rs2268498 Analyses**

Participants’ rs2268498 genotype did not moderate the effect of social support (either the composite measure [p = .366] nor father [p = .543], mother [p = .158], or family support [p = .612], individually) on PPD symptoms.

4. Discussion

The present results indicate that OXTR genotype moderates the strength of the association between social support from the baby’s father and PPD symptoms. Specifically, we found that PPD symptoms were more tightly linked to the social support mothers received from the baby’s father in the GG genotype group relative to mothers with AG or AA genotypes. Consistent with the hypothesized role of oxytocin in heightening the impact of both social affiliation and exclusion on mental health outcomes, GG homozygotes (and to a lesser extent, AG heterozygotes) displayed more PPD symptoms when father support was perceived to be low, whereas no such association obtained for AA homozygotes.
The general consensus in the *OXTR* literature is that individuals carrying one or more G alleles for rs53576 are more sensitive to their social environments than those carrying A alleles (Bradley et al., 2013; Chen et al., 2011; Hostinar et al., 2014; McQuaid et al., 2013; Riem et al., 2011; Sturge-Apple et al., 2012). For example, individuals with one or two copies of the G allele of rs53576 have been found to evince lower cortisol responses to stress after social support when compared to individuals with the same genotype who received no social support (Chen et al., 2011). Relatedly, Sturge-Apple and colleagues studied 201 mothers with young toddlers and found that inter-parental conflict was more likely to ‘spill over’ and result in decreased maternal sensitivity and increased punitive parenting in GG homozygotes than in AA/AG genotypes.

Our multiple regression including maternal and family support revealed that support from the baby’s father drove the inverse correlation between perceived social support and low PPD symptoms as well as uniquely interacting with rs53576. Perceived deficiencies in the social support provided by the baby’s father may be particularly critical predictors of maternal distress in the postpartum period given the shared responsibility and vital need for cooperation between parents (Milgrom et al., 2008). The transition to parenthood is often associated with sudden deterioration in marital relationship quality (Doss, Rhoades, Stanley et al., 2009). Our research suggests that such declines could be especially deleterious for women who are more socially attuned as a function of their *OXTR* gene variants. In turn, the partners of mothers possessing such variants may be at greater risk of depressive symptoms themselves as an indirect consequence, given that partners’ mental health tracks the postpartum mental health of mothers, with partners of mothers with PPD 30% more likely to be depressed themselves (Iles et al., 2011). Future research might explore whether partners of mothers with more socially sensitive
OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION

OXTR variants do indeed report greater depressive symptoms under contexts of relationship discord.

Consistent with the stress-buffering hypotheses, exploratory analysis in this study revealed that the association between father support and rs53576 was further moderated by perceived stress. Namely, in mothers with GG genotypes, social support was a more potent buffer against PPD symptoms when mothers reported higher (compared to lower) levels of perceived stress. There was no evidence of such a stress buffering effect in the larger group of AG mothers or the smaller group of AA mothers. However, these exploratory analyses should be interpreted with caution given that studies testing stress buffering models typically utilize more objective measures of stress exposure (e.g., major life event inventories), rather than the perceived stress scale used here. Our perceived stress measure shared approximately half of its variance with PPD symptoms ($r^2 = .51$), which is slightly higher than meta-analytic estimates of the shared variance using life events inventories ($r^2 = .38-.40$; Beck, 2001). We controlled for perceived stress in all of our moderation analyses because stress not only covaries with PPD symptoms, but also with social support and demographic variables relevant to OXTR variation (e.g., race/ethnicity), and hence introduces noise likely to obscure relationships between social support and OXTR. Perceived stress is characterized by feelings of inability to cope with life’s challenges, but not necessarily with affective states associated with depression such as low mood and anhedonia. As such, controlling for perceived stress in our moderation analyses may have particularly highlighted the interaction between OXTR and support with regard to aspects of PPD symptoms which are distinct from perceived stress.

The results of this study should be interpreted cautiously in light of several limitations. First, our sample size of 220, while sufficient to detect the relationships reported, is smaller than
OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION

the ideal participant pool for genetic studies (Duncan & Keller, 2011). A larger sample may have detected the significant relationships between the OXTR genotype and PPD symptoms reported in previous studies (Saphire-Bernstein et al., 2011). Second, we administered a self-report continuous scale of depressive symptoms rather than assessing mothers clinically diagnosed with PPD, diminishing the generalizability of our findings to clinical populations. Third, we have not elucidated the potential mechanisms linking OXTR variation with PPD symptom outcomes. OXTR SNPs may influence PPD symptoms via a number of processes, including downstream effects on oxytocin receptor expression (Mizumoto, Kimura, & Ivell, 1997), or covarying linkages with other SNPs not assessed here (Lin, Vance, Pericak-Vance, & Martin, 2007). Highly powered studies might explore whether other OXTR SNPs (for example, rs2254298; Thompson et al., 2011) or other genetic variants (such as those related to the release of oxytocin) moderate the relationship between perceived social support and maternal mental health. Future work directed toward identifying mediating mechanisms might also explore the extent to which variations in OXTR determine the extent to which natural shifts in oxytocin associated with childrearing impact postpartum depression symptoms and maternal behavior. For example, if mothers who carry G alleles for OXTR 53576 are more sensitive to the actions of oxytocin, then events associated with exposure to natural doses of oxytocin such as vaginal childbirth (compared to C-section) or breastfeeding (compared to formula-feeding) may be more important predictors of PPD symptoms or maternal bonding in mothers who carry G alleles.

To the best of our knowledge, the present study is the first to explore how social and genetic factors interact to predict PPD symptoms. Looking ahead, understanding of the etiology of PPD symptoms may be advanced by similar efforts to synthesize literatures across disciplinary boundaries to create biopsychosocial models of PPD (for a review, see Yim et al.,
By determining which varieties of social support affect which sorts of postpartum mothers, we may one day be able to craft personalized interventions to prevent and ameliorate PPD.

Acknowledgments

We are grateful to our community partners, Granola Babies Boutique and MOMS of Orange County, as well as to our research assistants, Shiva Amanat, Mariel Barojas, Maddisen Espeseth, Samantha Halela, Nikki Shahbazi, and Bonnie Truong. We are also grateful to Chapman University’s Office of Undergraduate Research and the Biology Honor Society (TriBeta) for providing grant support for this project.

Declaration of Conflicting Interests

The authors have no conflicts of interest to declare.
OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION

Footnotes

1 For the purposes of this paper, we reserve the term PPD for clinically diagnosed cases; otherwise, in regard to studies that rely on self-report screeners, we refer to PPD symptoms.

2 We also included a single-item dichotomous (Yes/No) self-report measure of prior history of depression, "Have you ever experienced depression during your lifetime?" as a potentially relevant covariate. Follow-up tests confirmed that including history of self-reported depression did not change the statistical significance of any of the analyses reported here, although depression history did predict PPD symptoms ($B = 2.14, SE = 0.64, \beta = .23, p = .001$, $95\% \text{ CI} [.87, 3.40], r^2 = .05$).

3 We conducted follow-up tests to assess whether father support might exert a different impact on PPD symptoms according to race/ethnicity. Tests revealed that the race/ethnicity x father support interaction term was not statistically significant, nor did including this term change our pattern of results with regard to rs53576 moderating the effects of father support on PPD symptoms. Moreover, removing the race/ethnicity covariates from the models altogether also did not change the pattern of results.
References


**OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION**


OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION


OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION


OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION


OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION


**OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION**

Table 1

Associations between demographic factors, **OXTR** genotype, social support and postpartum depression symptoms

<table>
<thead>
<tr>
<th>Variable</th>
<th>EPDS Score</th>
<th>Father Support</th>
<th>Mother Support</th>
<th>Family Support</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infant Age (months)</td>
<td>6.01 (0.22)</td>
<td>-0.03</td>
<td>-0.06</td>
<td>-0.09</td>
</tr>
<tr>
<td>Maternal Age (years)</td>
<td>33.14 (0.37)</td>
<td>-0.06</td>
<td>0.07</td>
<td>-0.15*</td>
</tr>
<tr>
<td>Income</td>
<td>4.50 (0.15)</td>
<td>-0.08</td>
<td>0.16*</td>
<td>0.16*</td>
</tr>
<tr>
<td>Married</td>
<td>81%</td>
<td>-0.04</td>
<td>0.31***</td>
<td>0.09</td>
</tr>
<tr>
<td>Perceived Stress</td>
<td>14.64 (6.44)</td>
<td>0.733***</td>
<td>-0.21**</td>
<td>-0.27***</td>
</tr>
</tbody>
</table>

rs53576 Genotype

<table>
<thead>
<tr>
<th></th>
<th>β / %</th>
<th>β / %</th>
<th>β / %</th>
<th>β / %</th>
</tr>
</thead>
<tbody>
<tr>
<td>AA</td>
<td>6.15 (4.15)</td>
<td>25.56 (3.91)</td>
<td>24.16 (4.60)</td>
<td>25.46 (2.52)</td>
</tr>
<tr>
<td>AG</td>
<td>7.60 (4.57)</td>
<td>25.69 (3.27)</td>
<td>24.67 (4.20)</td>
<td>24.90 (3.41)</td>
</tr>
<tr>
<td>GG</td>
<td>6.25 (4.41)</td>
<td>25.86 (4.05)</td>
<td>24.62 (4.85)</td>
<td>25.21 (4.05)</td>
</tr>
</tbody>
</table>

rs2268498 Genotype

<table>
<thead>
<tr>
<th></th>
<th>β / %</th>
<th>β / %</th>
<th>β / %</th>
<th>β / %</th>
</tr>
</thead>
<tbody>
<tr>
<td>CC</td>
<td>6.06 (4.06)</td>
<td>24.92 (5.23)</td>
<td>25.68 (3.46)</td>
<td>26.16* (2.38)</td>
</tr>
<tr>
<td>CT</td>
<td>7.44 (4.48)</td>
<td>26.20 (2.36)</td>
<td>24.05 (4.77)</td>
<td>24.62b (3.77)</td>
</tr>
<tr>
<td>TT</td>
<td>6.61 (4.42)</td>
<td>25.53 (4.32)</td>
<td>24.72 (4.62)</td>
<td>25.40 (3.57)</td>
</tr>
</tbody>
</table>

Note. *p < .10; *p < .05; **p < .01, ***p < .001; Values with different superscripts within the same row are significantly different from one another (p < .05); household income is coded as 1 = less than $10,000; 2 = $10,000-$49,999; 3 = $50,000-$74,999; 4 = $75,000-$99,999; 5 = $100,000-$149,999; 6 = $150,000-$199,999; 7 = $200,000-$299,999; 8 = $300,000-$399,999; 9 = $400,000-$499,999; 10 = over $500,000.
Table 2

Correlation Matrix of the Association between Sources of Social Support and Postpartum Depression Symptoms

<table>
<thead>
<tr>
<th></th>
<th>Total Support</th>
<th>Mother Support</th>
<th>Family Support</th>
<th>Baby’s Father Support</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pearson’s r</td>
<td>Pearson’s r</td>
<td>Pearson’s r</td>
<td>Pearson’s r</td>
</tr>
<tr>
<td>EPDS Score</td>
<td>-0.292 (.000)</td>
<td>-0.177 (.010)</td>
<td>-0.257 (.000)</td>
<td>-0.239 (.000)</td>
</tr>
<tr>
<td>Mother Support</td>
<td>-</td>
<td>-</td>
<td>.450 (.000)</td>
<td>.115 (.095)</td>
</tr>
<tr>
<td>Family Support</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>.456 (.000)</td>
</tr>
</tbody>
</table>
OXTR, SOCIAL SUPPORT & POSTPARTUM DEPRESSION

**Figure 1.** Association between depressive symptoms and father support, as moderated by OXTR rs53576 genotype. OXTR rs53576 genotype significantly moderated the effect of father support on postpartum depression risk. Father support was a more potent predictor of depressive symptoms in mothers with more G alleles. Simple effects tests showed that father support and depressive symptoms were significantly inversely correlated amongst GG and AG genotypes; whereas there was not a significant relationship in AA genotypes. ($r^2$ values and correlations represent raw data without covariate adjustment. Covariate adjusted coefficients and effect sizes are reported in the results section.)

![rs53576 Genotype scatter plot](image.png)