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Oxytocin Receptor (OXTR) Methylation and Cognition in Psychotic Disorders

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Key Words Psychosis · Cognition · Oxytocin receptor · Methylation

Abstract

Previous reports have identified an association between cognitive impairment and genetic variation in psychotic disorders. In particular, this association may be related to abnormal regulation of genes responsible for broad cognitive functions such as the oxytocin receptor (OXTR). Within psychotic disorders, it is unknown if OXTR methylation, which can have important implications for gene regulation, is related to cognitive function. The current study examined peripheral blood OXTR methylation and general cognition in people with schizophrenia, schizoaffective disorder, and psychotic disorder not otherwise specified (N = 101). Using hierarchical multiple regression analysis, methylation at the Chr3:8767638 site was significantly associated with composite cognitive performance independent of demographic and medication factors while controlling for multiple testing in this combined diagnostic sample (adjusted p = 0.023).

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Introduction

The neuropeptide oxytocin is associated with broad cognitive processes in humans. Previous findings indicate that both general and social cognitive processes such as working memory and facial emotion recognition, respectively, are related to oxytocin function [1-4]. Within psychotic disorders such as schizophrenia, broad cognitive deficits have been well documented [5-8], and oxytocin abnormalities may contribute to these deficits. Further, the link between social cognition and oxytocin in psychotic disorders has garnered attention due to treatment implications (e.g., intranasal oxytocin may improve facial emotion recognition) [9–11]. Recent evidence suggests that intranasal oxytocin may also improve general cognition in schizophrenia [12–15]. In addition, genetic variants of the oxytocin receptor gene (OXTR), which serves as the binding point for oxytocin action through second messenger systems in the brain, have been linked to a diagnosis of schizophrenia [16] and poorer cognitive performance in psychotic disorders [17, 18]. Thus, the genetic regulation of oxytocin pathways via deoxyribonucleic acid (DNA) variation, and its relationship to cognition in schizophrenia continues to be an active area of research [for review, see 19]. While recent reports have

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8767581	ATGTGATGTAGGCCTTGGGTCCCCAGGGCTGGATGAAGA [CG] GCCCAGCAG	8767630
8767631	T [CG] AAGA [CG] C [CG] TCAGCCACCTCGC <u>GCAGAGAAGATGTGCAC</u> CTG [CG] G	8767680
8767681	[CG] [CG] CTGGCCACCAGGCAGC [CG] AGCCA [CG] TGG [CG] AGCACTGCCAGG [CG] GT	8767730
8767731	[CG] GTG [CG] G [CG] G [CG] CAGCGAGCGCAGCGGCTGGCAGATGGCCAGGCAGCGG	8767780
8767781	TCCAGGGA <u>CATGAGCAGCAGGCAGGTAGGTGGA</u> 8767812	

Fig. 1. Figure 1 shows the *OXTR* amplicon analyzed in the current study. This amplicon is contained within exon 3 of the *OXTR* gene downstream to the translation start site. The sequence is displayed using the GRCh38/Hg38 build coordinates Chr3:8767581 – Chr3: 87677812. The 19-nucleotide forward primer used for the amplicon is single underlined starting at 8767581 and ending at 8767599, and the 24-nucleotide reverse primer is single underlined from

8767789 to 8767812. The reverse primer was ordered as the reverse complimentary primer and was biotinylated to facilitate pyrosequencing. The three sequencing primers are double underlined and found starting at positions 8767599 (19 nucleotides), 8767656 (19 nucleotides), and 8767704 (20 nucleotides). Analyzed CpG methylation sites are shown bracketed (15 total).

identified a relationship between increased *OXTR* methylation and impaired facial emotion recognition in schizophrenia [20], there have been no studies of *OXTR* gene regulation through DNA methylation and general cognitive performance in psychotic disorders. It is possible that oxytocin system abnormalities in psychosis may extend to *OXTR* methylation, and this abnormality may be related to broad cognitive deficits. If true, it would suggest that *OXTR* gene regulation is a potential treatment target for cognitive deficits in people with psychosis.

Therefore, the current study aimed to investigate the relationship of *OXTR* DNA methylation and general cognition in a sample of participants with schizophrenia, schizoaffective disorder, or psychotic disorder not otherwise specified. We hypothesized that site-specific *OXTR* DNA methylation would be associated with cognitive performance across participants.

Methods

Participants

Participants were recruited from the southeastern Michigan area based on the following inclusion criteria for a separate study examining the metabolic side effects of antipsychotic medications in people with a psychotic disorder: (1) Diagnostic and Statistical Manual for Mental Disorders (DSM-IV) [21] Axis I diagnosis of schizophrenia, schizoaffective disorder, schizophreniform disorder, or psychotic disorder not otherwise specified, (2) between the age of 18 and 90, and (3) at least 6 months of stable atypical or typical antipsychotic medication dosage (atypical antipsychotics included risperidone, olanzapine, quetiapine, ziprasidone, aripiprazole, clozapine, paliperidone, iloperidone, asenapine, and lurasidone; typical antipsychotics included haloperidol, fluphenazine, perphenazine, trifluoperazine, chlorpromazine, thioridazine, and thiothixene). Participants were excluded based on the following criteria: (1) diagnosed with type 2 diabetes prior to treatment with antipsychotic medications (criterion required for separate study examining metabolic side effects), and (2) active DSM-IV substance abuse diagnosis. Included participants completed a single visit to undergo a fasting blood draw, comprehensive medication history (relevant to the study of metabolic side effects), diagnostic assessment using the Structured Clinical Interview for DSM-IV (SCID-IV) [22], and cognitive testing. All evaluations were carried out by a single trained research associate. In addition, level of education and smoking history were obtained. Smoking history was assessed by asking the participants if they smoked one or more cigarettes per day. Self-report measures of depression (i.e., Beck Depression Inventory-Revised) [23], psychological stress (i.e., Psychological Stress Index) [24], and state anxiety (i.e., State-Trait Anxiety Inventory) [25] were completed to assess current mood and anxiety. Clinical assessments of current psychotic symptoms were not collected due to time limitations. All participants gave fully informed consent to participate in the protocol, as approved by the University of Michigan Institutional Review Board.

DNA Methylation Analysis

DNA was first extracted using the salt precipitation method [26] and cleaned and concentrated using commercially available kits. Five hundred nanograms of DNA was then bisulfite converted using the EZ DNA Methylation-Gold Kit (Zymo Research, Irvine, Calif., USA). Ten nanograms of bisulfite-converted DNA was placed in a touchdown PCR reaction containing OneTaq Hot Start $2\times$ Master Mix with Standard Buffer (New England BioLabs, Ipswitch, Mass., USA) in order to amplify our chosen *OXTR* region of interest. Specifically, the 232-bp region is contained within the 3rd exon of the *OXTR* gene and covered the coordinates Chr3: 8767580 – 8767812 (using the GRCh38/hg38 build). Methylation within and immediately adjacent to this region has been investigated in several brain-based disease states [27–32]. The studied

Age, years	19 to 71	45.4±10.7
Gender		
Male	_	56 (55)
Female	_	45 (45)
Race		
Caucasian	_	54 (53)
African-American	_	45 (45)
Other	_	5 (5)
Level of education $(n = 91)$	1 to 8	3.9 ± 1.3
DSM-IV Axis I diagnosis		
Schizophrenia	_	32 (32)
Schizoaffective disorder	_	59 (58)
Psychotic disorder NOS	_	10 (10)
Age at diagnosis, years	16 to 64	26.4 ± 9.5
Duration of illness, years	1 to 44	19.0 ± 11.3
Hospitalizations, lifetime	0 to 44	8.9 ± 10.1
Beck Depression Inventory-Revised (n = 94)	0 to 63 total	14.95±11.29
Psychological Stress Index $(n = 94)$	0 to 4 average	2.19 ± 0.71
Social Adjustment Scores-Self-Report ($n = 95$)	0 to 5 average	2.24 ± 0.54
BACS composite z-score $(n = 94)^a$	-4.83 to 0.42	-1.92 ± 1.06
Verbal Memory	-3.85 to 1.44	-1.35 ± 1.09
Digit Sequencing	-5.95 to 1.66	-1.41 ± 1.26
Token Motor Task ($n = 99$)	-5.48 to 1.43	-1.61 ± 1.24
Verbal Fluency	-4.14 to 1.07	-1.06 ± 0.91
Symbol Coding $(n = 98)$	-3.97 to 0.72	-1.56 ± 0.93
Tower of London $(n = 96)$	-4.29 to 1.36	-1.19 ± 1.29
Current cigarette smoker	-	52 (51.5)
Packs of cigarettes per year	37 to 1,095	86.8±123.0
Atypical antipsychotic	_	91 (90)
Typical antipsychotic	_	19 (18.8)
Daily chlorpromazine equivalents, mg	25 to 3,175	530.6 ± 517.9

Table 1.	Demographic,	clinical,	cognitive,	and	physiological	characteristics	of	all	clinical	participants	(i.e.,
n = 101)			•								

Note: data are presented as range, mean \pm SD, or n (%). NOS = Not otherwise specified. ^a Computation of BACS z-scores utilized a healthy control comparison group.

region is contained within a CpG Island of the OXTR gene that spans from approximately 20 to 2,350 bp downstream of the transcription start site. Differential methylation of this CpG region has been shown to have large functional effects on OXTR expression [30, 33]. Within our target amplicon, 15 CpG methylation sites were chosen for analysis by the method of pyrosequencing using three separate pyrosequencing primers for the amplicon [34]. Other CpG methylation sites within the amplicon were not included for the following reasons: (1) the assay design for the region was not included (dictated by the software), and (2) a detectable methylation value was not replicated in \geq 66% of the sample. Methylation calls for individual samples were excluded if the replication methylation value was too variable (defined as a coefficient of variation >2.5%) and thus deemed unreliable. The 15 methylation calls that were included for analysis were normalized using the method of Goodrich et al. [35]. Briefly, this method uses methylation standards (0, 20, 40, 60, 80, and 100% methylated) for each analysis in

order to construct a standardized curve on which each sample is normalized. Figure 1 shows the amplicon region, primers used, and methylation sites analyzed. All primers were designed using the Pyromark AssayDesgin 2.0 software.

Cognition

Cognitive performance was determined using the Brief Assessment of Cognition in Schizophrenia (BACS), a performance-based battery of cognition with six subscale scores, including Verbal Memory, Digit Sequencing (working memory), Token Motor Task (motor speed), Verbal Fluency, Symbol Coding (attention and speed of information processing), and Tower of London (executive functions) [36, 37]. In order to standardize and interpret BACS performance, *z*-scores were computed with reference to a sample of healthy controls who also lived in Southeastern Michigan (n = 63) and are described in Grove et al. [38]. The BACS provides a composite score to describe overall cognitive performance. This

Table 2. Nontransformed methylation of OXTR sites

No.	Site	n	Mean ± SD
1	Chr 3:8767620	91	17.3±8.2
2	Chr 3:8767632	89	23.9±10.8
3	Chr 3:8767638	80	17.3 ± 8.8
4	Chr 3:8767641	77	20.4±13.1
5	Chr 3:8767678	91	21.9 ± 7.7
6	Chr 3:8767681	92	17.6 ± 8.1
7	Chr 3:8767683	88	12.3 ± 7.1
8	Chr 3:8767701	85	21.3 ± 8.7
9	Chr 3:8767708	68	11.1 ± 4.8
10	Chr 3:8767713	67	21.4 ± 9.4
11	Chr 3:8767727	91	20.6±9.9
12	Chr 3:8767731	89	14.6 ± 9.5
13	Chr 3:8767736	90	20.6±13.0
14	Chr 3:8767739	80	19.7±13.2
15	Chr 3:8767742	81	13.0 ± 6.8
Overall methylation	_	92	18.2±5.9

Note: chromosomal locations using Human Genome build 38.

score is analyzed in a 'whole model' approach where, in the current study, a significant relationship between BACS composite score and methylation site warrants further analysis of the subscale scores.

Statistical Analyses

Demographic and clinical characteristics are described by means with standard deviation. Differences between the healthy control sample (used to compute BACS z-scores) and participants with a psychotic disorder were determined using independent samples *t*-tests or χ^2 analyses. The healthy controls did not have methylation data available and were not included in methylation analyses. Pearson correlational analyses were used to identify significant associations between BACS z-scores (composite score first and then followed by subscale scores if necessary) and methylation sites. Hierarchical multiple regression models were used to determine if a specific OXTR methylation site (i.e., identified from significant Pearson correlational analyses) would be associated with BACS composite z-scores independent of variables related to cognition and methylation. The regression models contained two steps. The first step included level of education, with a ranking of 1 (grade 6 or less) to 8 (completed graduate/professional school), as educational attainment has been consistently associated with cognition in schizophrenia [39, 40]. In addition, considering that epigenetics can be mediated by antipsychotic medication type (i.e., atypical or typical) and cigarette smoking, both of which may play an important role in the pathophysiology of schizophrenia [41-43], antipsychotic type and cigarette smoking status (current smoker or nonsmoker) were also included as independent variables in the first step of the models. A methylation site that significantly correlated with BACS composite z-score was entered as an independent variable in the second and final step of the models. Within all models,

standardized betas are reported, along with p values for the amount of variance in the BACS z-score explained by the individual methylation site compared with demographic and medication variables. These values were corrected using Benjamini and Hochberg's False Discovery Rate (threshold of 0.05) [44–46], which has been recommended as an alternative to the conservative Bonferroni correction for clinical studies. In addition, due to the nonnormal distribution of methylation data, all methylation results were log transformed for statistical analyses. While power analysis was not conducted due to the lack of previous studies examining OXTR methylation and cognition in psychotic disorders needed to estimate the effect size, the current sample size is comparable or larger than previous OXTR methylation studies of clinical populations and psychiatric symptoms [28, 29, 47–50]. JMP statistical software (SAS Institute Inc., Cary, N.C., USA) was used for all analyses.

Results

Demographic, Clinical, and Cognitive Characteristics

Demographic, clinical, and cognitive characteristics of the 101 participants with a psychotic disorder are presented in table 1. Compared to the healthy control sample used to compute BACS *z*-scores [n = 63; average age of 43.5 ± 11.3; age range of 19–66; 65% male (n = 41); Bachelor's degree level of education 6.1 ± 1.8, and 70% Caucasian (n = 46)], participants with a psychotic disorder had a significantly lower level of education [t(151) = 8.71, p < 0.0001] and a significantly higher ratio of non-Caucasian participants ($\chi^2 = 9.47$, n = 160, p = 0.002).

Relationships between BACS and OXTR Methylation

The average, nontransformed values of the analyzed *OXTR* methylation sites are listed in table 2. Overall, the investigated sites were 'lowly' methylated (average methylation of sites ~20%), which is in agreement with healthy control studies investigating *OXTR* methylation within and near this region [29, 32].

A significant negative Pearson correlation was observed between Chr 3:8767638 site and BACS composite *z*-score. Neighboring methylation sites (i.e., Chr 3: 8767708, Chr 3:8767681, and Chr 3:8767632), along with the remaining methylation sites, were not significantly correlated with BACS composite *z*-score except for Chr 3:8767742, which was positively correlated (table 3).

Given the significant findings at Chr3:8767638 and Chr3:8767742, we conducted separate hierarchical regression analyses for each site. The multiple regression analyses showed that only *OXTR* methylation site Chr3: 8767638 explained a significant amount of variance in BACS composite *z*-score, along with Verbal Memory, Symbol Coding, and Tower of London *z*-scores, indepen-

OXTR methylation		BACS										
No.	site	Composite	Verbal Memory	Digit Sequencing	Token Motor Task	Verbal Fluency	Symbol Coding	Tower of London				
1	Chr 3:8767620	-0.05	-0.02	-0.11	0.11	0.03	-0.13	-0.10				
2	Chr 3:8767632	-0.13	-0.12	-0.21*	0.11	-0.10	-0.23*	-0.03				
3	Chr 3:8767638	-0.25*	-0.25*	-0.20	0.07	-0.17	-0.24*	-0.27*				
4	Chr 3:8767641	0.08	-0.01	-0.01	0.16	0.11	0.05	-0.02				
5	Chr 3:8767678	-0.06	-0.12	-0.12	0.09	-0.02	-0.16	0.04				
6	Chr 3:8767681	-0.08	-0.03	-0.25**	-0.05	0.04	-0.05	-0.05				
7	Chr 3:8767683	-0.01	-0.03	-0.10	0.16	0.02	-0.10	-0.02				
8	Chr 3:8767701	0.12	0.05	0.02	0.27*	0.00	0.03	0.08				
9	Chr 3:8767708	-0.12	-0.17	-0.24	0.13	-0.11	-0.17	0.01				
10	Chr 3:8767713	-0.02	-0.16	-0.15	0.17	0.02	-0.08	0.10				
11	Chr 3:8767727	-0.05	-0.02	-0.11	0.11	0.03	-0.13	-0.10				
12	Chr 3:8767731	0.02	-0.06	-0.04	0.19	0.07	-0.13	0.02				
13	Chr 3:8767736	-0.03	-0.11	0.05	0.07	0.05	-0.13	-0.06				
14	Chr 3:8767739	0.04	-0.04	0.09	0.13	0.05	-0.07	-0.03				
15	Chr 3:8767742	0.25*	0.12	0.18	0.29**	0.16	0.16	0.10				
Overall methylation	_	-0.02	-0.10	-0.08	0.19	0.03	-0.09	-0.05				

Table 3. Pearson correlations between BACS and log-transformed OXTR methylation

Note: Pearson correlation coefficients between the log-transformed methylation of the *OXTR* gene (rows) and BACS *z*-scores (columns). * p < 0.05, ** p < 0.01.

Table 4.	Results	from	the	hierarchical	regression	models	used t	o predict	the effect	t of	Chr3:8767638	OXTR	methylation	on	BACS
perform	ance (n =	= 72)			-										

BACS	Step 1				Step 2				
	variable sta		model	statistics	variable statistics, β	model			
	level of education	antipsychotic type	current smoker	$\overline{\Delta R^2}$	ΔF	Chr3:8767638	$\overline{\Delta R^2}$	ΔF	FDR- adjusted p value
Composite	0.30*	-0.18	0.17	0.15	4.03*	-0.31*	0.09	7.89*	0.023
Verbal Memory	0.29	-0.21*	0.14	0.15	4.10*	-0.29*	0.08	6.84*	0.023
Digit Sequencing	0.30*	-0.20	0.10	0.15	3.83*	-0.20	0.04	3.20	0.062
Token Motor Task	0.09	-0.14	-0.02	0.03	0.69	0.00	0.00	0.00	0.600
Verbal Fluency	-0.07	-0.01	0.16	0.03	0.75	-0.21	0.04	3.00	0.062
Symbol Coding	0.22	0.00	0.04	0.05	1.12	-0.27*	0.07	5.06*	0.029
Tower of London	0.29*	-0.09	0.25*	0.14	3.72*	-0.27*	0.07	6.04*	0.024

Note: BACS composite and subscale *z*-scores were included as the dependent variable. Level of education, antipsychotic type (atypical or typical), and cigarette smoking status (Step 1) and methylation value (Step 2) were entered as independent variables for each model. The false discovery rate (FDR)-adjusted p values were calculated for the Step 2 variable (Chr3:8767638 site) only. Standardized betas are reported. * p < 0.05.

dent of level of education, antipsychotic type, and smoking status, even after correcting for multiple testing (table 4). The association between Chr3:8767742 and BACS composite *z*-score did not remain significant after accounting for level of education, antipsychotic type, and smoking status ($\Delta R^2 = 0.04$, $\Delta F = 3.19$, $\beta = 0.19$, p = 0.079). Consequently, Chr3:8767742 and BACS subscale *z*-scores were not analyzed with hierarchical regression models.

OXTR and Cognition in Psychosis

Discussion

Oxytocin Methylation and Cognition

Within the current study, a significant negative association was observed between CpG methylation at Chr 3: 8767638 in the OXTR gene and cognitive performance independent of demographic and antipsychotic medication variables in people with schizophrenia, schizoaffective disorder, or psychotic disorder not otherwise specified. Specifically, cognitive deficits in domains such as working memory, attention and speed of information processing, and executive functions may be related to peripheral abnormal OXTR DNA methylation in people with a psychotic disorder.

To our knowledge, this is the first study to investigate OXTR gene methylation and general cognition in people with a psychotic disorder. A previous study has examined facial emotion recognition in people with a psychotic disorder (at site -934 upstream of the OXTR start codon in DNA and the same CpG island as the region in the current study), and found increased OXTR methylation in females, but not males, was associated with greater difficulties identifying angry, sad, and happy faces [20]. In the current study, the association between OXTR methylation and cognition was observed independent of sex. Previous studies have also examined OXTR methylation and social cognition in psychiatrically healthy participants and found that higher degrees of OXTR methylation were associated with greater activation of the superior temporal gyrus/supramarginal gyrus and dorsal anterior cingulate cortex during an animacy task [32]. Further, increased OXTR methylation was associated with higher neural response and decreased functional coupling within brain regions supporting socioemotional processing (i.e., amygdala, fusiform, and insula) [31]. While OXTR methylation in these two studies was assessed in a neighboring region compared to the current study, the methylation regions analyzed in these studies were still contained within the same CpG Island. In conjunction, these findings suggest that decreased OXTR methylation in this CpG Island may contribute to both lower- and higher-level cognitive processing (e.g., working memory and facial emotion recognition, respectively). Yet within psychotic disorders, the relationship between the oxytocin system and cognition is inconsistent. The Psychiatric Genomic Consortium (PGC) did not identify oxytocin (OXT) or OXTR as risk factors for schizophrenia via a genomicwide association analysis [51]. However, this may be due to oxytocin pathway genes not being implicated in

the susceptibility to a psychotic disorder, but rather susceptibility to impaired social behavior (i.e., emotional withdrawal) within the PGC sample [52]. In addition, the role of the oxytocin system in the pathophysiology of positive, negative, and cognitive symptoms continues to be implicated [13, 53].

The mechanism by which abnormal oxytocin methylation influences cognition may be explained by biological models that indicate perinatal stress disrupts the oxytocin system in brain areas associated with numerous cognitive processes (e.g., memory retrieval, decisionmaking, and social cognition) [54, 55]. Specifically, increased methylation of OXTR may lead to abnormal gene expression that disrupts hippocampal and/or medial prefrontal cortex function and causes cognitive impairment (e.g., memory deficits and/or impaired emotion recognition) [see 56 for a review]. This disruption may be exacerbated by psychosocial stressors, which can increase OXTR methylation [28]. Considering that OXTR methylation at Chr 3:8767638 was significantly associated with working memory, processing speed, and executive functions, abnormal activation in associated areas (e.g., prefrontal cortex and related substructures) [57-61] may be linked to decreased peripheral OXTR methylation in people with a psychotic disorder. However, future neuroimaging studies with patients and healthy control populations are needed to determine this potential relationship between OXTR methylation and brain function in psychosis. Thus, while further research is needed to confirm these postulations and resolve discrepant findings, abnormal oxytocin methylation in the periphery and the brain may not only contribute to impaired socioemotional processing in psychosis [20], but also general cognitive deficits in psychotic disorders, making OXTR methylation a potential biomarker or treatment target.

Previous reports also indicate that *OXTR* methylation in overlapping regions of interest is associated with social anxiety disorder [29]. Specifically, elevated Chr3:8767736 methylation and reduced Chr3:8767708 methylation in participants with social anxiety disorder compared with healthy controls was observed [29]. In the current study, no significant correlations were observed between Chr3: 8767708 methylation and BACS composite or subscale scores. However, the previous report highlights the importance of *OXTR* methylation at individual sites, which has apparent variable methylation in association with a given phenotype and may contribute to different disease pathophysiology (i.e., psychotic disorder vs. social anxiety disorder).

Understanding the Influence of Environmental and Lifestyle Factors on Cognition

The potentially positive cognitive effects of nicotine obtained through cigarette smoking in schizophrenia are well documented [62, 63], along with the potential role for nicotinic receptors in cognition [64]. The effect of smoking on DNA methylation has been observed in several populations [65-67]. Additionally, cigarette smoking has been found to affect OXTR gene regulation and transcription in pregnant women [68]. Yet the current findings suggest smoking may not impact the associations between OXTR methylation and cognition in people with a psychotic disorder, as OXTR methylation explained BACS composite variance independent of smoking status. Further, besides the environmental effect of smoking, cardiovascular function plays an important role in cognition [69-71]. Future work may investigate the methylation of known genes that contribute to cardiovascular function in schizophrenia (e.g., methylenetetrahydrofolate reductase) for their role in the relationship between cardiovascular health and cognition in psychotic disorders [72-74].

Limitations

The current study was observational in nature, and only associations can be inferred. Considering the critical need for replication of the current findings, future prospective studies could not only replicate but also expand upon the current findings by assessing coincident changes in gene methylation and cognition in psychotic disorders coupled with brain imaging during disease progression and/or treatment to explore casual relationships. While the current study used a candidate epigenetic approach and investigated a single area of a single gene, along with False Discovery Rate correction to reduce the chance of type 1 errors, few correlations were observed between cognition and gene methylation. Associations between cognition and gene methylation of other genes will need to be tested independently in future studies. Further, consequences of gene methylation through RNA studies and downstream protein levels will need to be assessed to understand the molecular effects of differential gene methylation in cognitive processes. However, as described earlier, differential methylation of the CpG Island that contained the current methylation sites has been linked to significant effects in OXTR expression and oxytocin levels [75]. A limitation of the current study is not assessing DNA methylation at the tissue of interest, which in this case would be the brain. A future study using available postmortem human brain samples could have some

benefits but would not be without significant limitations of its own. Such limitations may include epigenetic changes that could occur from death to tissue processing, accurate phenotyping of subjects at the time of death (e.g., disease status, medication use, lifestyle factors, cognitive status, etc.), and choosing specific brain regions to analyze OXTR DNA methylation. The current study did not use postmortem brain samples to validate the peripheral blood findings due to a lack of sample access. However, the use of peripheral tissues to study brain genetic regulation is an important initial strategy [76-78], and previously reported correlations between OXTR methylation in peripheral blood mononuclear cells and the brain suggest that peripheral methylation may be related to brain function [79]. Additionally, the use of peripheral blood DNA methylation can have significant biomarker value regardless of it not being from the tissue of interest. Nevertheless, preclinical models of cognition in schizophrenia are available [80] and may enable translational studies that can be used to support epigenetic findings in the peripheral blood from humans.

The correlation of DNA methylation across tissues continues to be an active area of research, and correlations between OXTR peripheral blood methylation and the temporal cortex have been reported, suggesting that there may be some degree of similarity [50]. Although likely gene dependent, work in animals and humans continues to support translatability between peripheral blood and brain DNA methylation [81-84]. Nevertheless, correlating site-specific peripheral methylation findings to the brain may be required. As studies continue to assess DNA methylation in the periphery, it is important that similar extraction and assessment methodologies are used in order to test replication of any findings put forth. Finally, although limited in sample size, significant associations remained after correction for multiple testing. However, larger sample sizes with better demographically matched controls than the current study (along with cognition and methylation data for both cases and controls) may allow for replication of these analyses within and across DSM-IV diagnostic groups. Between-group analyses within the current sample are limited by the small number of participants diagnosed with psychotic disorder not otherwise specified (i.e., 10 participants). In addition, the current study did not assess psychotic symptoms, and future studies should explore the relationship between OXTR methylation and the severity of psychosis, as previous reports indicate a relationship between OXTR variants and a diagnosis of schizophrenia [16].

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Conclusion

Methylation levels of the OXTR gene at the Chr 3: 8767638 site are negatively associated with general cognitive performance in people with a psychotic disorder (i.e., schizophrenia, schizoaffective disorder, or psychotic disorder not otherwise specified). This association is independent of demographic and antipsychotic medication type, and remains after adjusting for multiple testing, suggesting that abnormal OXTR methylation may contribute to the cognitive deficits and pathophysiology of psychotic disorders.

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Statement of Ethics

The authors have no ethical conflicts to disclose.

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