Neuroepigenetic impact on mentalizing in childhood

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ARTICLE INFO
Keywords:
Oxytocin
Middle childhood
FMRI
Mentalizing
Epigenetics

ABSTRACT
Mentalizing, or the ability to understand the mental states and intentions of others, is an essential social cognitive function that children learn and continue to cultivate into adolescence. While most typically developing children acquire sufficient mentalizing skills, individual differences in mentalizing persist throughout childhood and are likely influenced by a combination of cognitive functioning, the social environment, and biological factors. DNA methylation of the oxytocin receptor gene (OXTRm) impacts gene expression and is associated with increased brain activity in mentalizing regions during displays of animacy in healthy young adults. The establishment, fine-tuning, and implications of such associations in the context of broader social functioning remain unclear. Using a developmental neuroimaging epigenetic approach, we investigated the contributions of OXTRm to individual variability in brain function during animate motion perception in middle childhood. We find that higher levels of OXTRm are associated with increased neural responses in the left temporo-parietal junction and inferior frontal gyrus. We also find a positive association between neural activity in LTPJ and social skills. These findings provide evidence of epigenetic influence on the developing child brain and demonstrate that variability in neural social perception in childhood is multifaceted with contributions from individual social experience and the endogenous oxytocin system.

1. Introduction
Middle childhood, often defined as the period between 6 and 12 years of age, is a time of extensive cognitive, social, and emotional development (Skuse, 2017; Del Giudice, 2014). During this time, children develop complex reasoning skills and display substantial changes in self-regulation, executive function, and peer interactions (Bigelow, 1977). Understanding of mental states, referred to as a theory of mind, improves into middle childhood and is essential for successful navigation of the complex social world (Miller, 2009; Miller et al., 2018). Individual differences in theory of mind in preschool have been associated with variability in socioeconomic status (Hughes and Ensor, 2005), cognitive skills (Carlson et al., 2002; Lecce et al., 2017) and language (Hughes and Ensor, 2007); these differences in preschool are well documented, and are predictive of teacher-rated social competence in elementary school (Devine et al., 2016). While less studied at older ages, theory of mind continues to develop in middle and late childhood, with evidence from cross-sectional and longitudinal studies demonstrating significant individual differences in the skill of mental attribution, or mentalizing. Variability in the magnitude of neural activity within key mentalizing regions during theory of mind tasks has also been linked to social cognition behaviors in children (Mukerji et al., 2019). Taken together, individual differences in mentalizing ability that persist through middle childhood are likely impacted by complex interactions between cognitive skills, the social environment, and biological influence that help shape the development of a child’s ability to effectively reason about the mental states of others (Lecce et al., 2017; Devine et al., 2016; Lecce et al., 2011; Banerjee et al., 2011; Tucker-Drob and Briley, 2014).

One important biological modulator of social and emotional processing is the hormone and neuromodulator oxytocin. Intranasal delivery of oxytocin enhances activity of neural circuits associated with social reward and social attention in children with Autism Spectrum Disorder (ASD) (Gordon, 2016, 2013), improving their social cognitive function and promoting sociality (Anagnostou, 2014). Intranasal oxytocin administration also improves mentalizing and emotion
的认识和在个体差异中的表现。在本研究中，我们假设OXTR在社会知觉功能中扮演一个重要的角色，特别是在发展早期。

此外，我们使用了单独收集在发展时期的数据来支持这个假设。在目前的研究中，我们假设在5至11岁的儿童中，OXTR和功能脑成像数据的相关性会增强。

2. Materials and methods

2.1. Experimental design

我们选择了67名儿童（32名男性）作为研究对象。每个参与者在第一次访问时会接受15分钟的假扫描，以便适应假扫描环境。参与者在第二次访问时会被安排进行真正的MRI扫描。我们使用了儿童报告的父系和行为数据来支持这个假设。在当前的研究中，我们假设在儿童中，OXTR和功能脑成像数据的相关性会增强。
to indicate the highest degree completed by the child’s caregiver(s) and in all instances degree completion of two caregivers were provided. For each participant, parent education was scored from 0 (8th grade) to 5 (PhD or other terminal degree) and summed for both caregivers to generate a parent education value ($M = 6.55$, $SD = 2.09$, range $= 2–10$). Between the two visits, caregivers were emailed a link to complete the Social Skills Improvement System rating scale (SSIS) (Gresham and Elliott, 2008) via Qualtrics (Qualtrics, 2019). The SSIS is a 79-item parent-report questionnaire designed to assess social skills and problem behaviors in children 3–18 years of age. Each item is rated on a 4-point Likert-type scale from 1 (never) to 4 (almost always). Seven caregivers of participants included in the MRI analysis did not complete the questionnaires.

2.3. Saliva Collection and Epigenotyping Procedures

Saliva was collected using CS-2 sponges in OG-250 collection kits from DNA Genotek (Ottawa, Canada) and stored at room temperature. Prior to DNA isolation, samples were incubated at 50 °C for 1 h and centrifuged at 1000 RPM for 10 min to recover all liquid from sponges. DNA was isolated following the manual purification protocol from DNA Genotek, resuspended in Hydration Solution (Qiagen, Hilden, Germany) and quantified using NanoDrop.

Two hundred nanograms of DNA were subjected to bisulfite treatment (Kit MECOV50, Invitrogen, Carlsbad, CA), which converts non-methylated cytosines to uracil and leaves methylated cytosines unmodified for downstream sequencing. A 116-base pair region of OXTR containing CpG Site – 924 was amplified via polymerase chain reaction (PCR) using 20 nanograms of bisulfite-converted DNA as a template, 0.2 μM primers TSL101F (5′-TTAGTTTGGATTATAAGATT-3′) TSL101R (5′-bionin-AATAAATTACCTCCCA TCTCATTTCCTAA-3′) and using Pyromark PCR kits (Qiagen, Hilden, Germany). Samples were amplified in triplicate with the following cycling conditions: Step 1: (95 °C/15 min)/1 cycle, Step 2: (94 °C/30 s, 56 °C/30 s, 72 °C/30 s)/50 cycles, Step 3: (72 °C/10 min)/1 cycle, Step 4: 4 °C hold. Each PCR plate contained methylation standards (0% and 100% methylated), a positive control, and negative controls from bisulfite conversion and PCR. PCR amplification of a 116 bp product was confirmed by gel electrophoresis. DNA methylation for each sample was quantified by pyrosequencing (PyroMark Q24, Qiagen) using primer TSL101S (5′-AGAATTTATTATAATTTTT-3′) and Pyromark Gold Q24 Reagents (Qiagen, Hilden, Germany). DNA collection for three samples was insufficient to obtain methylation levels. Interquartile outlier detection was used to identify three samples with high replicate variability. These samples were rerun to estimate deviant values and replicates that caused deviation for these samples were removed. Average mean deviation within replicates was ± 1.71%. Reported epigenotypes are an average of the three replicates.

2.4. Animacy stimuli

Stimuli were presented with PsychoPy (Peirce, 2009) using an LCD AVOTEC projector onto a screen located behind the subject’s head and viewed through an integrated head-coil mirror. As in Jack et al. (2012) participants passively viewed modified Heider and Simmel clips that engage a network of structures that support reasoning about others (Jack et al., 2012; Castelli et al., 2000). Eight ‘Animate’ animations and eight ‘Control’ animations were presented to subjects during scanning. Each animation featured three white geometric shapes (triangle, diamond, and circle) on a black background. The Animate condition involved goal-directed behavior such as playing hide and seek or dancing with one another while the Control animations showed the shapes moving around the screen in a random fashion with the same velocity and overall motion as the Animate condition. Each animation was approximately 16 s and Animate and Control animations were presented in alternating block design. Participants were instructed to simply observe the shapes as they moved along the screen.

2.5. Imaging procedures

Scanning was performed at the UVA Fontaine Research Park on a 3 T Siemens Prisma scanner with a 32 channel headcoil. Cushions were placed around the participants’ ears and forehead to minimize head movement during the scans. An experimenter stood by the participants’ feet to ensure that the participants remained awake and comfortable. If the experimenter noticed the participant move, they gently placed their hand on the participant’s leg to remind the participant to stay still.

T1-weighted high-resolution structural images were acquired using Siemens’ magnetization-prepared rapid-acquired gradient echo (MPRAGE) pulse sequence with the following specifications: echo time (TE) = 2.98 ms; repetition time (TR) = 2300 ms; flip angle (FA) a 9°; image matrix = 240 mm × 256 mm; slice thickness = 1 mm; 208 slices. Whole-brain functional images were acquired using the same parameters as the Adolescent Brain Cognitive Development protocol (Casey, 2018). The specifications for the T2 * weighted echo planar imaging (EPI) sequence sensitive to BOLD contrast were with the following specifications: TE = 30 ms; TR = 800 ms; FA = 52°; image matrix = 90 mm × 90 mm; slice thickness = 2.4 mm; slice gap = 2.4 mm; 332 slices.

2.6. fMRI preprocessing

Anatomical images were skull stripped using AFNI (Cox, 1996) and the results were manually corrected. Subsequent data preprocessing and analysis was carried out using FEAT (FMRi Expert Analysis Tool) Version 6.0, part of FSL (FMRIB’s Software Library) (Smith, 2004). The following pre-statistics processing was applied: motion correction using MCFLIRT (Jenkinson et al., 2002); spatial smoothing using a Gaussian kernel of full width half maximum 5 mm; grand-mean intensity normalization of the entire 4D dataset by a single multiplicative factor; highpass temporal filtering (Gaussian-weighted least-squares straight line fitting, with sigma = 50.0 s). Registration to the Montreal Neurologic Institute (MNI) Template standard space image was carried out using fMRIB’s software library linear registration tool (FLIRT) (Jenkinson et al., 2002). Registration from high resolution structural to standard space was then further refined using FSL’s nonlinear registration, FNIRT (Andersson et al., 2007). Head motion was quantified by an algorithm that quantifies the root mean square of the relative frame-wise displacement (FDRMS) between each volume of functional data (Smith, 2004). Participants with mean FDRMS greater than 0.15 mm were excluded from all analyses.

At the subject level, time-series statistical analysis was carried out using FSL’s improved linear model (FILM) with motion parameters added to the model and local autocorrelation correction (Woolrich et al., 2001). Regressors for Animate and Control conditions were modeled by convolving the time course with a double-gamma hemodynamic response function (HRF) and with temporal filtering applied. An Animate > Control contrast was conducted and the contrast of parameter estimates (COPE) from this analysis for each individual was carried forward to higher-level analysis.

2.7. fMRI analysis

Group-level analysis of the data was conducted with FSL’s local analysis of mixed effects (FLAME) stage 1 and 2 (Beckmann et al., 2003; Woolrich et al., 2004). Z (Gaussianised T/F) statistic images were thresholded using clusters determined by $Z > 2.3$ ($p < 0.01$) and corrected cluster significance threshold of $p < 0.05$ (Worsley, 2012). We first conducted an exploratory ROI analysis targeting regions involved in mentalizing to identify regions in which Animate > Control activation was significantly associated with OXTRm. The mentalizing ROI mask was defined by a meta-analytic approach using Neurosynth.org (Tor D, 2011) feature set keyword “mentalizing”. Three regressors were
included in the model: a group mean regressor, mean-centered OXTRm, and mean-centered age in months. Age was specifically included in this model as a control variable because of the wide age range in this study and well documented associations between mentalizing and age. Contrasts were computed testing for positive and negative linear associations between OXTRm and Animate > Control activation and between age and Animate > Control activation. Significant clusters associated with OXTRm regressor were registered to each participant’s native space and average Animate > Control Z statistic values for each individual were extracted. We next used the average Animate > Control BOLD activation in the significant ROI cluster from the OXTRm regressor for each individual to test for associations between parent reported social skills and BOLD activation. Stepwise linear regression analysis was conducted in R (Crawley, 2007; R Development Core Team, 2011) and model comparison was used to determine the best fitting model to predict BOLD activation in the significant cluster including OXTRm, social skills, age, sex, and parent education. Stepwise linear regression allows for identification of specific and unique variables that allows for a model that is flexible yet tailored enough to describe sample or the population well. An additional ROI model was conducted in FSL using OXTRm and all covariates, a group mean regressor, mean-centered OXTRm, mean-centered age in months, biological sex, and mean-centered parent education to confirm the primary association with OXTRm with the smaller sample size. Finally, we conducted an exploratory whole-brain analysis to identify any other regions in which Animate > Control activation was significantly associated with OXTRm while controlling for age.

3. Results

Sixty-seven typically developing Caucasian children aged 5–11 years old participated in this study. OXTRm was analyzed at CpG site – 924, a conserved CpG site previously shown to impact gene expression in a DNA methylation dependent manner (Perkeybile, 2019; Kusui, 2001). Saliva-derived DNA methylation at this site is highly correlated with blood-derived methylation and has been associated with individual differences in infant brain response to emotional faces at 7 months (Krol et al., 2019a) and behavioral temperament at 18 months (Krol et al., 2019b). At this site, OXTRm ranged from 44% to 77% methylated (M = 63.46, SD = 5.49) and a Shapiro-Wilk test showed no significant departure from normality (W = 0.97, p = 0.065). There were no significant associations of OXTRm with demographic control variables including child age (r(65) = 0.07, p = 0.57), sex (r(65) = 0.54, p = 0.595), or parent education (r(58) = 0.22, p = 0.089).

Sixty caregivers completed SSIS which assesses Social Skills and Problem Behaviors (Gresham and Elliott, 2008). Social Skills scores ranged from 59 to 129 (M=97.05, SD=14.69), and Problem Behavior scores ranged from 2 to 44 (M=19.68, SD=10.30); in neither domain was there a significant departure from normality (p > 0.05). There was no significant association between OXTRm and caregiver rated Social Skills (r(58) = 0.12, p = 0.351). There were also no significant associations of Social Skills with child age (r(58) = −0.01, p = 0.945), sex (t (58) = 1.88, p = 0.066), or parent education (r(58) = 0.02, p = 0.899). Data is similar for Problem Behaviors (all p-values > 0.05). See Fig. S2 for histograms of primary study constructs.

The main effect of Animate > Control condition was generally consistent with prior research with greater activity in left occipital regions, bilateral posterior superior temporal sulcus, and fusiform gyrus. Left lateral frontal regions including the inferior frontal gyrus and orbitofrontal cortex were also activated (See Fig. 1 and Table 1). Multiple comparisons were corrected for using the false discovery rate (FDR) q < 0.05 voxel significance level and spatial extent threshold (k) > 10 contiguous voxels. For exploratory analyses, OXTRm was used as a regressor to predict individual variability in the processing of animate clips while controlling for age.

3.1. OXTR DNA methylation impacts Left Temporo-Parietal Junction response to animacy

We conducted an exploratory functional ROI analysis to assess the effect of OXTRm on activity within regions involved in mentalizing as defined by a meta-analytic approach using the Neurosynth.org feature set keyword “mentalizing”. This functional ROI analysis revealed a significant positive main effect of OXTRm on Animate > Control activity in the left temporo-parietal junction (TPJ) such that increased levels of methylation predicted increased BOLD activity (Fig. 2). Peak activity (Z = 3.21) for this cluster of voxels (k = 151) occurred at x = −46, y = −60, z = 24. The result from the functional ROI analysis including additional covariates of sex and parent education with the smaller sample size is similar and shown in S3.

| Anatomical Region | Hem | x   | y   | z   | Z  |
|-------------------|-----|-----|-----|-----|----|
| Superior Temporal Sulcus | R   | 48  | -40 | 10  | 7.57|
| Inferior Frontal Gyrus | R   | 46  | 34  | 4   | 7.48|
| Supramarginal Gyrus | R   | 60  | -26 | 40  | 7.06|
| Parietal Operculum Cortex | R   | 54  | -22 | 22  | 7.05|
| Precentral Gyrus | R   | 46  | 10  | 24  | 6.91|
| Fusiform Gyrus | R   | 46  | 10  | 24  | 6.90|
| Inferior Frontal Gyrus | L   | -46 | 20  | 8   | 3.88|
| Inferior Frontal Gyrus | L   | -42 | 24  | 12  | 3.70|
| Inferior Frontal Gyrus | L   | -50 | 26  | 12  | 3.57|
| Orbitofrontal Cortex | L   | -28 | 30  | -12 | 3.56|
| Inferior Frontal Gyrus | L   | -46 | 30  | 2   | 3.54|
| Orbitofrontal Cortex | L   | -38 | 26  | -14 | 3.26|

Fig. 1. Response intensity of brain regions demonstrating significantly greater activity in the Animate > Control condition.
3.2. Social skills predict the BOLD response in the Left TPJ

In order to better understand the association between OXTRm and left TPJ activity response to animacy, we investigated the possible association of child social skills within the left TPJ. We first conducted an exploratory analysis whereby the TPJ cluster was registered to subject space and the mean Z statistic value was extracted for each participant and correlated with scores from the SSIS social skills summary scale. Recall that caregivers of sixty children completed the questionnaire. Pearson correlation indicated that increased TPJ activity was related to higher social skills scores on the SSIS (r(58) = 0.29, p = 0.026). Demographic variables were tested as potentially confounding factors and revealed no significant effect or improvement to model fit.

Given that social skills and OXTRm both were positively associated with increased TPJ activity in the perception of animacy but not associated with one another, we conducted stepwise linear regression analysis to determine the best fitting model for these data. Results indicate that the statistical association between OXTRm and social skills as predictors for TPJ activity increased when both were included in the model, F(2, 57) = 6.92, p = 0.002, adjusted R² = 0.17. Additionally, we stepwise tested for demographic variables of age, sex, and parent education as confounding variables and results revealed parent education, but not age or sex, as a significant covariate of TPJ activity and improved model fit F(3, 56) = 7.06, p < 0.001, adjusted R² = 0.24. The final predictive model showed the TPJ activity was positive relative to OXTRm (B = 0.055, p = 0.001) and social skills score (B = 0.012, p = 0.038), but negative relative to parent education (B = −0.103, p = 0.017). There was no evidence of interactions or mediation in the model. See Fig. 3 for partial correlation matrix of all variables and Table S1 for stepwise linear regression table.

3.3. Exploratory whole brain analysis indicates OXTR DNA methylation influences the Left Inferior Frontal Gyrus

Finally, we performed an exploratory whole-brain analysis to identify potential associations between OXTRm and brain activity in regions not included in the mentalizing mask. This analysis revealed a significant positive main effect of OXTRm in the left inferior frontal gyrus (Fig. 4). Peak activity (Z = 3.36) for this cluster of voxels (k = 310) occurred at x = −56, y = 18, z = 16. While there was no linear association between age and Animate > Control activation in the ITG cluster, visual inspection of the data led us to test a non-linear inverted u-shape trend which was not significant.

4. Discussion

We examined the association between the neural response to animate motion in regions of the brain involved in mentalizing and individual differences in the endogenous oxytocinergic system in middle childhood. Controlling for chronological age, we found that higher levels of OXTRm were associated with greater activity in the left temporal parietal junction, similar to findings in young adults (Jack et al., 2012). Our multiple regression demonstrated that in addition to OXTRm, children’s social skills were also a positive significant predictor of the BOLD response in the left TPJ, providing evidence that behavior may account for unique variance in individual differences in neural response to animate motion. These results suggest a complex interplay of social skills and biological influence in the development of children’s mentalizing abilities.

Our group has previously shown that OXTRm is positively associated with BOLD response to animate motion in healthy young adults in the left STG extending to the TPJ and dorsal ACC (Jack et al., 2012). However, the young adult study lacked additional phenotyping measures to characterize how the observed correlations were meaningful in the context of individual differences in overt social behavior. The primary motivation of the current study was to probe associations between OXTRm and neural systems supporting social perception during a developmental period characterized by significant changes in social cognition and behavior. By taking a developmental approach, we aimed to maximize the amount of variability due to differences in maturation
and leverage this variance to explain the directionality of associations we found in young adults. In our child sample, we find a positive association of OXTRm and activity in the left TPJ that is similar to the findings in young adults. The consistency of this association in the left tempo-parietal areas in childhood and adulthood suggests that this association is established by middle childhood. While both the left TPJ and STG are consistently recruited in mentalizing and perspective taking tasks (Gallagher, 2000; Völlm, 2006; Jackson et al., 2006), the shift in the association with OXTRm from the TPJ in childhood to the STG in adults may reflect a fine-tuning of the system across development.

In our exploratory whole brain analysis, we also found that OXTRm is positively correlated with activity in the left IFG in children. While the left IFG is robustly significant in the main effect of Animate > Control condition in both children and adults, the association with OXTRm in the left IFG is a novel finding in childhood that has not previously been shown in adults. Across the lifespan the left IFG is associated with a wide variety of cognitive functions ranging from inhibitory control (Swick et al., 2006) to empathy (Chakrabarti et al., 2006) and is specifically well known to be involved in semantic processing (Binder et al., 2005; Perani, 1999; Noppeney and Price, 2004; Jessen, 2000). It is also proposed to be a region involved in differentiating abstract from concrete concepts (Wang et al., 2010) and interpreting communicative intent (Wang et al., 2006). The left IFG has also been shown to be more active in children and adolescents as compared to adults in nonverbal theory of mind tasks (Kobayashi et al., 2007). There is some evidence to suggest that activity in frontal regions, including the IFG, follows an inverted U-shape from childhood to adulthood as mentalizing becomes more automatic (Wang et al., 2006; Blakemore, 2008). It is possible that the observed association with OXTRm and IFG activity may reflect a mediating factor in a developmental shift in mentalizing abilities during perception of animate motion. Future work should investigate the role of OXTRm on developmental trajectories of mentalizing in a longitudinal sample.

When considering how variability in DNA methylation and social skills might relate to individual differences in neural response during animate motion, we hypothesized that social skills and OXTRm would be negatively correlated with one another and inversely related to TPJ activity. We hypothesized that individuals with higher levels of methylation, and presumably less sensitivity to endogenous oxytocin, recruit brain areas involved in mentalizing to a greater extent possibly due to increased resource intensive processing of social contingencies (Jack et al., 2012). It would follow that those individuals with higher methylation levels would also exhibit lesser social skills and greater social skills would be associated with reduced neural activity. Instead, we find no correlation between OXTRm and parent-reported social skills. Moreover, we find a significant positive association between social skills and left TPJ activity such that individuals with more social skills exhibit greater TPJ activity. Perhaps this is because at these ages, children are developing social skills concurrently with that activity; as the skills mature, greater efficiency may tamp the neural activity. A longitudinal investigation of change in neural activity across childhood may ultimately shed light on this trajectory.

The results of the stepwise multiple regression revealed that incorporating social skills in addition to OXTRm significantly improved the model fit. This model provides evidence that the variance in TPJ activity explained by OXTRm and the variance explained by social skills are unique, with no support for a mediation model. Parent education was also a significant variable but only in the full model including OXTRm and social skills, and importantly we find no evidence of an interaction or mediation for this term either. These data highlight that individual differences in the neural response of social perception is multidimensional and dependent on the environment, biological contributions, and social behaviors, and likely comprised of a combination of these influences.

4.1. Limitations and future directions

The present study focuses on individual differences in the endogenous oxytocin system and neural substrates of social perception and social behavior from a development perspective. While our previous young adult research is informative, we cannot make direct developmental comparisons between children and young adults as these were independent samples. Another important difference is the use of different tissues in children and adults. Working with developmental populations required us to use saliva as a non-invasive peripheral tissue to isolate DNA and derive OXTRm levels while working with young adults allowed for the use of an intravenous blood draw. We have previously shown that blood-derived and brain-derived DNA methylation at our CpG site of interest are correlated in animal models and that both reliably assess ASD phenotype in humans (Gregory, 2009). More recently we have shown that OXTRm derived from saliva and from whole blood and peripheral blood mononuclear cells are highly correlated in a large sample of adults (Krol et al., 2019a) and that methylation does not vary across different cell types in the blood (Puglia et al., 2018). Taken together this suggests that cell type frequency is less likely to impact relationships we identified. Even so, the use of saliva derived DNA methylation levels in developing children warrants consideration (Holbrook et al., 2017; Bakulski et al., 2016). Because the CpG sites we study are not on the Illumina array, testing for cell type differences using saliva cell type reference panels (Middleton, 2021) would require us to generate both sets of data to test our sites of interest and control for cell type differences. This limitation should be considered for future studies. These experiments provide compelling evidence for the feasibility of saliva as a non-invasive peripheral marker of OXTRm but do not allow us...
to make direct comparisons between children and adults regarding methylation values or the association with neural response. Lastly, because methylation levels have been shown to differ as a function of ethnic groups, we restricted our sample to self-identified Caucasian Americans. Future work should explore the impact of epigenetic differences within and between ethnic groups.

Another important limitation to the present study is the use of a cross-sectional design. Our lab has shown that saliva derived OXTRm is stable in adulthood but dynamic in infancy (Krol et al., 2019b), and in human infants and animal models changes in methylation values at these CpG sites are related to the social environment (Perkeybile, 2019; Krol et al., 2019b). It is still unknown if OXTRm at this CpG site is dynamic through childhood and the present study is only informative of OXTRm and neural activity at one time point in development for each individual. We do not observe any chronological age-related associations with variability in the oxytocinergic system itself, or any presence of chronological age confounding the observed associations with neural activity to animate motion. But middle childhood is characterized by sweeping developmental shifts in mentalizing and social interactions and there is robust evidence demonstrating developmental changes in neural processing during middle childhood and adolescence (Blake more, 2008). There is likely within-individual variability in the trajectories of the establishment and fine tuning of the associations between OXTRm and neural substrates of social perception that cannot be captured without a longitudinal design.

The current study aimed to investigate how individual variability in the oxytocinergic system and neural response to animate social perception inform the development of social skills and behaviors in typically developing children. The results of the multiple regression analysis emphasize that individual variability in social perception is scaffolded on more than DNA methylation alone and indicate the need for future studies to directly probe these complex associations between biological markers, social environment, and social behaviors. The current study assessed children’s social skills only from parent report to obtain a global measure of social skills across several domains of social behavior. Including direct assessments of children’s social behavior may provide more robust information to untangle the correlation between biological and neural associations of variability in social behaviors. Finally, a longitudinal design is needed to investigate these associations and how shifts in the oxytocinergic system, environment, social perception processing, and social skills relate to the formation of social support systems and successful social integration.

4.2. Conclusion

In summary, we find that similar to adults, increased OXTRm is associated with increased activity in the left tempo-parietal area during the perception of animate motion in children. We also find a novel association in left IFG activity and OXTRm in children. When considering other factors that might contribute to individual differences in mentalizing we find that in addition to OXTRm, social skills contribute to individual differences in TPJ activity regardless of chronological age. While we find no evidence of an association between OXTRm and parent-reported social skills, the full model highlights the complexity of social skills, social environment, and biological influence in the development of children’s mentalizing abilities. Nonetheless, these data provide the first evidence in children of the impact of the epigenome on the developing social brain and point to a clear role for the oxytocin system in understanding individual differences in social developmental trajectories.

Funding

This research was supported by the University of Virginia and a UVA Brain Institute Transformative Neuroscience Pilot Grant “Individual variability in the oxytocinergic system and the development of human sociality” (AL, JC, and JM).

CRediT authorship contribution statement

Amalia Skyberg: Conceptualization, Methodology, Software, Validation, Formal analysis, Investigation, Data Curation, Writing – Original Draft, Visualization, Project administration. Stef en Beeler-Duden: Formal analysis, Investigation, Data Curation, Writing – Review & Editing. Alison Goldstein: Investigation, Data Curation, Project administration, Writing – Review & Editing. Christina Gancayco: Software, Data Curation, Writing – Review & Editing. Angeline Lillard: Conceptualization, Resources, Supervision, Funding, Writing – Review & Editing. Jessica Connelly: Conceptualization, Methodology, Resources, Supervision, Funding acquisition, Validation, Writing – Review & Editing. James Morris: Conceptualization, Methodology, Validation, Resources, Supervision, Funding acquisition, Writing – Review & Editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data statement

This data will be made available by request on OSF.

Acknowledgments

We thank Morgan Lynch, Samuel Wilson, Arpitha Shenoy, Jamie Levin, and Michelle Miller for their assistance with data collection and all the families who participated. We thank Kevin Tarczon and Zoe Bell for their assistance with preparation of the saliva samples. We also thank Drs. Kathleen Krol and Krassimira Botcheva for helpful comments on drafts of this manuscript.

Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.dcn.2022.101080.

References

Anagnostou, E., 2014. Intranasal oxytocin in the treatment of autism spectrum disorders: a review of literature and early safety and efficacy data in youth. Brain Res. https://doi.org/10.1016/j.brainres.2014.01.049.

Andersson, J.L.R., Jenkinson, M., Smith, S., 2007. Non-linear Registration Aka Spatial Normalisation. FMRIB Technical Report TR07JA2.

Bakulski, K.M., Halladay, A., Hu, V.W., Mill, J., Fallin, M.D., 2016. Epigenetic research in neuropsychiatric disorders: the "issue issue". Curr. Behav. Neurosci. Rep. 3, 264.

Banerjee, R., Watling, D., Caputi, M., 2011. Peer relations and the understanding of faux pas: longitudinal evidence for bidirectional associations. Child Dev. https://doi.org/10.1111/j.1467-8624.2011.01669.x.

Beckmann, C.F., Jenkinson, M., Smith, S.M., 2003. General multilevel linear modeling for group analysis in FMRI. Neuroimage. https://doi.org/10.1016/S1053-8119(03)00475-X.

Bigelow, B.J., 1977. Children’s friendship expectations: a cognitive-developmental study. Child Dev. https://doi.org/10.2307/1128905.

Binder, J.R., Westbury, C.F., McKiernan, K.A., Possing, E.T., Medler, D.A., 2005. Distinct brain systems for processing concrete and abstract concepts. J. Cogn. Neurosci. https://doi.org/10.1162/0898929054021192.

Blake more, S.J., 2008. The social brain in adolescence. Nat. Rev. Neurosci. https://doi.org/10.1038/nrn2353.

Carlson, S.M., Moses, L.J., Breton, C., 2002. How specific is the relation between executive function and theory of mind? Contributions of inhibitory control and working memory. Infant Child Dev. https://doi.org/10.1002/icd.298.

Casey, B.J., et al., 2018. The Adolescent Brain Cognitive Development (ABCD) study: imaging acquisition across 21 sites. Dev. Cogn. Neurosci. https://doi.org/10.1016/j.dcn.2018.01.001.
