Title
Neural modulation of social reinforcement learning by intranasal oxytocin in male adults with high-functioning autism spectrum disorder – a randomized trial

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ABSTRACT

Reduced social motivation is a hallmark of individuals with autism spectrum disorders (ASD). Although the exact neural mechanisms are unclear, oxytocin has been shown to enhance motivation and attention to social stimuli, suggesting a potential to augment social reinforcement learning as the central mechanism of behavioral interventions in ASD. We tested how reinforcement learning in social contexts and associated reward prediction error (RPE) signals in the nucleus accumbens (NAcc) were modulated by intranasal oxytocin. Male adults with a childhood diagnosis of ASD (n=15) and healthy controls (n=24; aged 18-26 years) performed a probabilistic reinforcement learning task during functional MRI in a single-center (research center in Germany), randomized double-blind placebo-controlled crossover trial. The interventions were intranasal oxytocin (Syntocinon®, Novartis; 10 puffs = 20 international units (IU) per treatment) and placebo spray. Using computational modeling of behavioral data, trial-by-trial RPE signals were assessed and related to brain activation in NAcc during reinforcing feedback in social and non-social contexts. The order of oxytocin/placebo was randomized for 60 participants. Twenty-one participants were excluded from analyses, leaving 39 for the final analysis. Behaviorally, individuals with ASD showed enhanced learning under oxytocin when the learning target as well as feedback was social as compared to non-social (social vs. non-social target: 87.09% vs. 71.29%, [95% CI 7.28-24.33], p = 0.003; social vs. non-social feedback: 81.00% vs. 71.29%, [95% CI 2.81-16.61], p = 0.027). Correspondingly, oxytocin enhanced the correlation of the RPE signal with NAcc activation during social (versus non-social) feedback in ASD (3.48 vs. -1.12, respectively, [95% CI 2.98-6.22], p = .000), whereas in controls, this effect was found in the placebo condition (2.90 vs. -1.14, respectively, [95% CI 1.07-7.01], p = 0.010). In ASD, a similar pattern emerged when the learning target was social (3.00 vs. -0.64, respectively, [95% CI -0.13-7.41], p = 0.057), whereas controls showed a reduced correlation for social learning targets under oxytocin (-0.70 vs. 2.72, respectively, [95% CI -5.86-0.98], p = 0.008). The current data suggest that intranasal oxytocin has the potential to enhance social reinforcement learning in ASD. Future studies are warranted that investigate whether
oxytocin can potentiate social learning when combined with behavioral therapies, resulting in greater treatment benefits than traditional behavior-only approaches.

**Trial Registration:** The trial is registered at the US National Institutes of Health (ClinicalTrials.gov) #NCT01712464.

**INTRODUCTION**

To date, there is still no approved pharmacological treatment for the core social symptoms of autism spectrum disorders (ASD). The neuropeptide oxytocin (OXT) has been proposed as a promising candidate for treating ASD-related social deficits, as it has been shown to enhance motivation and attention to social stimuli by making them more salient, thereby facilitating social learning and memory [1]. At present, the most effective treatment for improving social functioning in ASD are behavioral therapies which build on the principles of reward-based operant reinforcement learning, such as applied behavior analysis (ABA) or social skills training [2,3]. Such interventions are costly and time consuming but often fail to benefit a substantial number of affected individuals [2]. Thus, a better understanding of the mechanisms for social learning is urgently needed to improve current treatments.

The dopaminergic (DA) system, including the striatum, plays an essential role for reinforcement learning. The ventral striatum, specifically the nucleus accumbens (NAcc), signals reward and the expectation thereof, in order to initiate changes in behavior [4]. Brain activation of the NAcc is closely associated with the processing of reward prediction errors (RPE), that is, the difference between an actual and an expected reward [5]. RPE signals in the NAcc are generated by phasic activity of DA neurons in the ventral tegmental area, reflecting the basic neural mechanism underlying reinforcement learning [4]. In animal studies it could be demonstrated that OXT closely interacts with the DA reward system [6]. For example, OXT modulates social learning, such as establishing social preference and bonding, and acts specifically as a social reinforcement signal within the NAcc [6,7].
humans, Hu and colleagues demonstrated increased learning selectively for social feedback under OXT along with changes in striatal brain activation [8].

Several lines of research have indicated that the OXT system is altered in ASD. Genetic variation of the OXT receptor (OXTR) is significantly associated with ASD [9,10], and baseline plasma OXT may relate to (social) functioning in affected individuals [11]. Animal models of ASD-associated behavior suggest dysfunctions of the OXT system which could be ameliorated with OXT administration (e.g., [12]). Single dosage studies in humans suggest improved prosocial functioning after OXT administration (e.g., [13,14]) and increased brain activation and connectivity in striatal brain regions [15,16]; however, clinical trials with repeated OXT administration have produced mixed findings [17]. Given the proposed link between OXT and DA-mediated social learning, surprisingly few studies have addressed combined effects of behavioral and pharmacological interventions (e.g., [18]), and no study has yet investigated the influence of OXT on social reinforcement learning and the associated neural mechanisms in ASD.

Thus, in the present study we used a probabilistic social reinforcement learning task in young adults with and without ASD during fMRI with intranasal OXT and placebo (PLC) administration in a randomized double-blind within-subjects crossover design. We expected OXT to improve learning in social contexts in ASD, as well as an association with enhanced RPE signals in the NAcc, indicating that OXT in ASD may alleviate social learning deficits through an influence on brain mechanisms mediating reinforcement learning.

MATERIALS AND METHODS

Study design

A single-center, double-blind, placebo-controlled cross-over trial was performed between April 2013 and August 2016. All participants received both oxytocin and placebo treatments in randomized order to compare the modulatory effect of oxytocin on social reinforcement learning and associated RPE signals in the NAcc versus placebo. The study
was approved by the ethical committee of the University Hospital RWTH Aachen, Germany,
and registered at the US National Institutes of Health (ClinicalTrials.gov) # NCT01712464

Participants

Thirty-five healthy male healthy control (HC) participants (aged between 18 and 25
years) and 25 male individuals with ASD (aged between 18 and 26 years) were included into
the randomization procedure (i.e., allocation to treatment of OXT or PLC on the first visit).
Several participants had to be excluded from the analysis for various reasons (dropout on
the second visit, HC n = 1; anatomical brain abnormality, ASD n = 1, HC n = 1; technical and
data quality problems, ASD n = 3, HC n = 6; correct guess of the administered
pharmacological substance, HC n = 1; poor task performance, i.e., acquisition of a “wrong”
association (significant preference [> 60%] of the not reinforced option) either within the
second half or across all trials of at least one condition, ASD n = 2, HC n = 2; medication
status, ASD n = 4). In total, 24 HC (mean age = 22.09, SD = 1.88, mean IQ = 119.10, SD =
9.35) and 15 ASD (mean age = 21.79, SD = 2.60, mean IQ = 113.53, SD = 11.61)
participants were included in the final analysis. They all had normal language function and
were not taking any psychotropic medications at the time of scanning. Because of menstrual
cycle-related changes in plasma OXT [19], no women were included in this study. See Table
1 for demographic information of the final participant samples. A more detailed description of
the trial protocol (including a CONSORT Flow Diagram) and details about exclusion criteria,
screening procedure and other behavioral measures is provided in the supplement.
Individuals with ASD were recruited from a database of participants with ASD from previous
studies at the Departments of Child and Adolescent Psychiatry in Aachen (RWTH Aachen
University) or Frankfurt am Main (Goethe University). All participants with ASD had received a childhood diagnosis by experienced clinicians and reached cut-offs on the Autism Diagnostic Observation Schedule – Generic (ADOS-G) and/or the Autism Diagnostic Interview – Revised (ADI-R). They were screened for other current psychiatric and neurological disorders with a brief clinical interview during the screening procedure. Two participants reported a diagnosis of ADHD and respective medication during childhood, but no current medication or symptoms. Four participants reported a depressive episode in the past, one reported vocal tics during childhood, but no current symptoms. HC participants were recruited from databases of previous studies at the Department of Child and Adolescent Psychiatry in Aachen or via local advertisements. They had no indication of developmental delay or history of any neurological or psychiatric disorder, as assessed by a brief clinical interview. The Beck Depression Inventory-Second Edition (BDI-II) was used to screen for depressive symptoms (ASD: $M = 5.20, SD = 4.75$ vs. HC: $M = 3.13, SD = 3.03$; $p = 0.103$). For a dimensional measurement of reciprocal social behavior, participants filled in the Social Responsiveness Scale (SRS; self rated). The ASD group showed on average moderate deficiencies in social behavior ($T = 68; M = 92.2, SD = 21.7$, missing data $n = 2$), whereas the HC group displayed no deficits ($T = 47; M = 31.6, SD = 26.9$, missing data $n = 8$) (see supplement for further details). The mean score in the ASD group is comparable to reports from other studies investigating adults (e.g., [20-22]). All experimental procedures were conducted at the Research Center Jülich, Germany, with written informed consent of all participants after they had received a complete description of the study.

- Please insert Table 1 about here -

**Procedure**

Participants took part in two sessions on two consecutive days. Each session consisted of 1) intranasal administration of OXT (Syntocinon®, Novartis; 10 puffs = 20
international units (IU) per treatment) or PLC spray, 2) two blood draws, 3) an (f)MRI scan, and 4) neuropsychological assessment and questionnaires. OXT/PLC was administered ~45 minutes (mean 48 minutes, SD = 5.61) before the beginning of the fMRI scan to ensure maximum availability of OXT in the central nervous system [23]. One blood sample was drawn before OXT/PLC administration for baseline, and a second before the beginning of the fMRI measurement for post-hoc validation of OXT-plasma levels during the fMRI measurement. Please refer to the supplement for the OXT-plasma analysis.

**Probabilistic Reinforcement Learning Task**

We employed a modified probabilistic social reinforcement learning task (similar to [24]). Participants were asked to indicate by button press with their left and right index finger whether a learning target would belong to category A or B, followed by probabilistic feedback. They were informed that the categories were arbitrary and had to be learned by means of probabilistic feedback with no underlying rule defining the category. The feedback was either rewarding upon correct choice or neutral upon incorrect choice, both with a probability of 75% (accordingly, a probability of 25% for incongruent, “false” feedback). Three different conditions were used, i.e., SN [social target – non-social feedback], NN [non-social target – non-social feedback], and NS [non-social target – social feedback]. Social learning targets were video clips of a male or female person looking at the participant with a neutral facial expression [SN conditions]. Non-social learning targets were video clips of colored fractals [NN, NS conditions]. Social feedback were video clips of a male or female person smiling at the participant and giving him “thumbs up”, or neutral video clips of this person with eyes closed and snipping fingers as if listening to music [25] [NS condition]. Non-social feedback was provided by videos of a colored fractal where a green checkmark or a blue cross appeared [SN, NN conditions]. To be able to identify potentially differential contributions of targets or feedback being social during reinforcement learning we focused on NS and SN conditions and their comparisons to the NN condition. Due to time
constraints, we were not able to include a condition with social feedback following a social target [SS condition].

**Figure 1.** Illustration and timing of the (A) SN, (B) NN, and (C) NS condition with rewarding feedback.

**Analysis of behavioral data**

Behavioral data were analyzed using the software SPSS 21 (IBM Corporation, Armonk, NY, USA). For each experimental condition and subject, the percentage of correct responses was calculated. To assess learning effects over time and to account for effects
during the initial performance and later stages of the learning phase (i.e., potential floor and ceiling effects), behavioral data were subdivided for each task condition into three intervals with the first two consisting of 3 blocks (8 trials each) and the last one consisting of 2 blocks. General linear model repeated measure analyses (mixed ANOVA) were used to assess main effects and interactions with treatment condition (OXT/PLC), task condition (NN/NS/SN), and interval (1/2/3) as within-subjects factors and group (ASD/HC) as between-subjects factor. For post-hoc tests, Bonferroni’s adjustment procedure was used.

**Computational Model**

Importantly, behavioral choice data were further analyzed using computational modeling of reinforcement learning, according to a basic Q-learning algorithm and a softmax decision function [26]. Learning parameter alpha was estimated using maximum-likelihood estimation and RPE and Q-values for each trial were calculated (see supplement for more details).

**Functional magnetic resonance imaging (fMRI)**

The fMRI protocol and analysis are described in detail in the supplement. In short, scans were acquired on a 3-Tesla head-dedicated MRI system (MagnetomTrioTim, Siemens, Erlangen, Germany) using a T1-weighted 3D magnetization prepared rapid acquisition gradient echo (MPRAGE) sequence and T2*-weighted echo planar imaging scans during task performance. Image preprocessing and analysis were performed using SPM12 (http://www.fil.ion.ucl.ac.uk/spm/software/spm12). Functional images were realigned to the mean image, anatomical scans co-registered to the mean image, segmented and normalized to the Montreal Neurological Institute template. Functional volumes were normalized and smoothed at 6 mm FWHM isotropic Gaussian kernel. The learning target
phase (when subjects performed the category selection) and the feedback phase (when subjects received feedback in response to their choice) of the task were modeled separately using stick functions, convolved with the hemodynamic response function and its first-order temporal derivative. Motion parameters were included as regressors. Feedback events were parametrically modulated by trial-wise individual RPE values. In line with previous findings [8] and our focus on RPE processing in the brain [27], the second level analysis focused on the parametric modulation of feedback events. Beta values representing this modulation were taken to the second level with all conditions modeled separately in a flexible factorial ANOVA, with the within-subjects factors treatment condition (OXT/PLC) and task condition (NN/NS/SN) and the between-subjects factor group (ASD/HC). For the whole brain analysis, only effects above a significance threshold of $p < .05$ (cluster-level corrected, $p < .001$ voxel level) are reported. ROI analyses were thresholded at $p < .05$ (voxel level), FWE-corrected for the ROI. Our analysis focused on brain activation within an anatomical ROI of the NAcc (defined as primary outcome measure before the beginning of recruitment (see https://clinicaltrials.gov/ct2/show/NCT01712464). Exploratory analyses were also performed for the amygdala (see supplement).

RESULTS

Behavioral results

For both participants with ASD ($F (2, 28) = 34.74, p < .001, \eta^2 = .71$) and HC ($F (1.28, 29.49) = 155.35, p < .001, \eta^2 = .87$) we observed a significant main effect of interval on the percentage of correct trials. This effect was evident across and separately for each experimental condition (all $p$s $< .001$; interval 1 vs. 2: 67.11\% vs. 82.95\%, respectively, [95\% CI -18.46 to -13.23], interval 1 vs. 3: 67.11\% vs. 87.13\%, respectively, [95\% CI -23.06 to -16.98], interval 2 vs. 3: 82.95 vs. 87.13, respectively, [95\% CI -5.67 to -2.70]), suggesting successful learning of the stimulus-feedback association during the course of the experiment.
Testing for OXT-induced effects in participants with ASD (according to our a-priori hypothesis) revealed a significant treatment x task interaction ($F(2, 28) = 3.45, p = .046, \eta^2 = .20$), indicating a higher percentage of correct trials for social targets (i.e., SN vs. NN) in the OXT condition (SN: 81.20% vs. NN: 72.07%, [95% CI -15.15 to -3.12], $p = .006, \rho_{corr} = .018$), but not in the PLC condition (SN: 70.87% vs. NN: 73.07, [95% CI -9.53 to 5.13], $p > .10$). Also, a significant task x treatment x interval interaction ($F(4, 56) = 2.72, p = .039, \eta^2 = .16$) revealed that in the second interval ($F(2, 28) = 10.83, p < .001, \rho_{corr} < .001, \eta^2 = .44$), individuals with ASD had a higher mean percentage of correct trials for both social feedback (i.e., NS vs. NN, 81.00% vs. 71.29%, respectively, [95% CI 2.81 to 16.61], $p = .009, \rho_{corr} = .027$) and the social learning target (i.e., SN vs. NN, 87.09% vs. 71.29%, respectively, [95% CI 7.28 to 24.33], $p = .001, \rho_{corr} = .003$) in the OXT condition, but not in the PLC condition (all $p > .10$) (see Figure 2). Similar analyses for the HC group revealed no main effects or interactions (see supplement for further behavioral analyses).

![Figure 2](image.png)

**Figure 2.** Task x treatment x interval interaction in the ASD group.

**Imaging results**

**Whole brain analysis**
Across groups, tasks, and treatment conditions [(HC+ASDOXT,PLC) > baseline], the whole brain analysis revealed a significant correlation of the reward prediction error signal with a broad neural network (Figure 3). Importantly, significant activation was observed within the striatum, including NAcc and putamen [28,29]. See supplement Table S2 for an overview of feedback RPE signals within the brain across ASD and HC participants.

Figure 3. Regions in the fMRI task where activation was associated with learning from feedback across groups, treatment and task conditions.

**ROI analysis**

Using the NAcc as our a-priori defined anatomical ROI, we observed a significant group x social feedback x treatment interaction ([−8 10 -10], Z = 3.40) (Figure 4A & B). The interaction was due to the ASD group showing a higher correlation of the RPE signal with brain activation in the left NAcc for social feedback as compared to non-social feedback in the OXT (NS: 3.48 vs. NN: -1.12, [95% CI 2.98 to 6.22], F (1, 14) = 37.02, p < .001, $\eta^2_p = .73$) but not PLC condition (NS: 0.84 vs. NN: 1.87, [95% CI -3.80 to 1.73], F (1, 14) = .65, p = .435, $\eta^2_p = .04$), whereas the HC participants had a higher correlation for social feedback as compared to non-social feedback in the PLC (NS: 2.90 vs. NN: -1.14, [95% CI 1.07 to 7.01],
\( F(1, 23) = 7.93, \ p = .010, \ \eta^2 = .26) \) but not OXT condition (NS: 1.36 vs. NN: 2.35, [95% CI -3.25 to 1.26], \( F(1, 23) = 0.83, \ p = .371, \ \eta^2 = .04) \). Also, a significant group x social learning target x treatment interaction was found ([8 -8 -10], \( Z = 4.02 \)) (Figure 4C & D). The interaction was due to the ASD group showing a marginally higher correlation of brain activation in the left NAcc for social as compared to non-social learning targets in the OXT (SN: 3.00 vs. NN: -0.64, [95% CI -0.13 to -7.41], \( F(1, 14) = 4.29, \ p = .057, \ \eta^2 = 0.24) \) but not PLC condition (SN: 0.99 vs. NN: 2.23, [95% CI -5.39 to 2.90], \( p = .531) \), whereas HC had a marginally higher correlation for social as compared to non-social learning targets in the PLC (SN: 2.96 vs. NN: -1.35, [95% CI -5.86 to -0.98], \( F(1, 23) = 3.56, \ p = .072, \ \eta^2 = .13) \) condition. Moreover, HC showed a higher correlation for non-social rather than social learning targets in the OXT condition (SN: -0.70 vs. NN: 2.72, [95% CI -5.86 to -0.98], \( F(1, 23) = 8.44, \ p = .008, \ \eta^2 = .27).
**Figure 4.** Neural correlates for the interaction between group x social feedback x treatment ((A) & (B)) and the interaction between group x social learning target x treatment ((C) & (D)).

**Brain-behavior correlations**

We also explored possible correlations between ASD symptom indices, individual characteristics related to reward processing and neural activity in the NAcc during social feedback in the OXT as compared to the PLC condition [OXT<sub>NS</sub> > PLC<sub>NS</sub>], within the ASD group using a multiple regression analysis. We observed a negative correlation between brain activation in the left NAcc ([−12 12 -6], Z = 3.27) and the reward dependence (RD) subscale of the Temperament and Character Inventory-140 (TCI-140), implying that individuals with ASD with lower reward dependence showed more activation in the NAcc during social feedback in the OXT as compared to the PLC condition (see supplement).

**DISCUSSION**

To our knowledge, the present study is the first to demonstrate OXT-induced enhancement of social learning in high-functioning ASD and an associated modulation of the RPE signal in the NAcc, a central neural hub for reinforcement-based learning. Our results suggest that the beneficial effect of OXT on social processing in ASD [13,14] is mediated by an enhancement of the brain’s motivational system, selectively in response to social stimuli, eventually boosting reinforcement learning in social situations. Thus, future studies investigating long-term efficacy of OXT as pharmacotherapy in ASD should consider that OXT might be particularly efficient in concert with behavioral interventions with an emphasis on socially reinforcing context to promote learning.

**Effects of oxytocin on task performance**

Following a single intranasal OXT challenge, we observed enhanced social learning in ASD, but no equivalent effect in HC. Similarly, a previous study demonstrated task performance increases under OXT selectively for participants with low social proficiency
using an incentive delay task with socially rewarding feedback [30]. At first sight, our results
seem at odds with other reports demonstrating that OXT facilitates learning from social
feedback in HC [8,24]. However, we focused exclusively on reinforcing feedback, whereas
these studies included also aversive feedback (i.e., angry faces) upon incorrect choices.
Facilitated learning in such contexts might be mediated by other effects of OXT, e.g.,
reduced threat sensitivity [31], or a decrease in aversiveness of negative stimuli [32]. Future
studies should directly test the differential contributions of reinforcing and aversive feedback
on OXT-induced effects.

**Modulatory influences of OXT on RPE signals in the NAcc**

Using RPE modeling, our task engaged a network of areas typically involved in
reinforcement learning. In particular, NAcc activity is assumed to reflect RPEs, i.e.,
differences between expectation and receipt of reward in order to adjust behavior [27,33] .
We did not find group differences within the general learning network, suggesting no overall
functional disruption of reinforcement learning in ASD. This is consistent with the
observation that behavioral interventions in ASD rely heavily on reinforcement based
learning to successfully modify behavior (e.g., [2]). At the same time, we could demonstrate
a particular sensitivity of NAcc activity for social feedback in HC (but not ASD) for the PLC
condition, supporting prior findings of impaired social reward processing in ASD (e.g.,
[34,35] in line with the social motivation theory of autism [36]. Our finding of an enhancing
effect of OXT for social vs. non-social feedback in ASD suggests that OXT has the potential
to restore the typical preference for social rewards in HC. Thus, learning from reinforcing
feedback during a social situation appears to be important for the effect of OXT in ASD. The
most parsimonious explanation for these results is that this effect is mediated by an OXT
induced increase of DA signaling during social situations, resulting in a targeted
enhancement of social approach motivation. This mechanism may also drive improvements
in social processing (e.g., [13,14]) and modulation in cortico-striatal activation and
connectivity [15,16] as reported in earlier OXT-challenge studies in ASD. These findings are
well in accordance with the social motivation theory of autism [36], suggesting that a
deficient OXT-DA interaction within the NAcc could be an important mechanism to account for reduced social motivation and, ultimately, impaired sociability in ASD.

Exploratory analyses also revealed that individual differences in reward dependence (RD; as measured by the Temperament and Character Inventory-140) was associated with the increase of RPE-correlated activation in the NAcc under OXT in ASD, suggesting stronger OXT effects in individuals with lower sociability and a tendency to learn less from rewarding interpersonal feedback (see supplemental data). This result is also in line with treatment outcomes of social skills trainings, showing that individuals with lowest skills usually benefit most [3].

**Involvement of OXT in dopaminergic modulation of learning**

According to numerous animal studies, OXT and DA interact within the NAcc (see, e.g., [6] for a review) to promote learning from social encounters as a prerequisite for establishing and maintaining social affiliations [37,38]. Similarly, the “social salience hypothesis of oxytocin” [1] suggests that OXT plays an overarching role for regulating the salience of social cues through its interaction with the DA system. Our findings critically add to a growing body of evidence for links between OXT and the DA reward circuitry, including the striatum (e.g., [30]), by providing a plausible mechanistic explanation in the context of social reinforcement learning, i.e., the amplification of striatal RPE signals as one potential mechanism of increased saliency. This notion is also compatible with the view that OXT may primarily amplify approach related behaviors [39]. Here, we found this effect only in individuals with ASD, suggesting that OXT effects may be dependent on individual differences in social functioning [1,30] and that OXT-DA interaction might constitute a central mechanism of reduced social motivation in ASD [41]. Importantly, we observed the effects during exclusively reward-based reinforcement learning (i.e., in the absence of aversive stimuli), suggesting an independent contribution of OXT on DA mediated approach behavior for social functioning.
Accordingly, we did not observe effects of OXT in the amygdala. The amygdala has been associated with salience signaling [1], probably mediating dampening effects on stress and anxiety (e.g., [40]). The occurrence of such effects may be confined to or more pronounced for negative social stimuli (such as threatening or aversive faces). Thus, anxiolytic properties of OXT and their interactions with DA within the amygdala [41,42] do not seem to be essential for a beneficial effect on social reward-based reinforcement learning.

Further studies with a focus on stress and anxiety in combination with reinforcement learning are warranted to elucidate this issue.

Limitations

Although the ASD sample was comparable to typical adult ASD populations in previous studies, we would like to emphasize that the average severity of deficits in reciprocal social behavior was moderate, all participants were male and very high functioning with respect to their cognitive abilities. Thus, the present findings only apply to this subgroup of high-functioning ASD. Given the high heterogeneity within the autism spectrum, generalization to the broader ASD population should be tested in future studies. A replication of our data with larger samples including women and individuals with lower functioning ASD, as well as a focus on children and adolescents is warranted. Furthermore, more research into comorbid conditions is necessary (e.g., ADHD, social anxiety) which also show impaired social reward processing, albeit in a different direction (e.g., hypersensitivity to social rewards in ADHD [43]). Future studies using a similar experimental design probably should include fully social conditions (i.e., social feedback and target). These may yield even stronger effects and are more comparable to real-life situations (e.g., feedback of the therapist during a social skills training, where both the learning target and the feedback are typically social).

Conclusions and future directions
We provide clear evidence for a neurobiological plausible mechanism of OXT-induced behavioral enhancement of social reinforcement learning in high-functioning ASD, i.e., the modulation of RPE signals in the NAcc. Our results implicate that OXT may unfold its therapeutic potential most efficiently in concert with targeted behavioral interventions, which provide opportunities for learning within social contexts along with immediate reinforcement, which is positive and explicitly rewarding (e.g., praise). Single dose administration studies have generally shown positive effects in ASD [13-16], which might be related to an overall enhancing effect on the motivational system in social contexts as demonstrated specifically for reinforcement learning here. In contrast, longer-term treatment studies using multiple dosing per day often failed to demonstrate treatment effects [17] and one important reason might be that these were not designed to provide specific social learning contexts around the times of administration [20], or may have interfered with psychotropic (e.g., in particular dopaminergic) medication [44]. Well-controlled studies which systematically combine social learning opportunities with OXT administration are lacking (but see [18]), but are urgently needed to further elucidate this issue.

Taken together, our findings suggest that it is crucial to further investigate the promising potential of combining OXT with behavioral interventions to inform modifications that might improve current treatment approaches.

Supplementary Information accompanies this paper at (-insert doi-).
ACKNOWLEDGEMENTS

The authors would like to thank Alexander Firk, Marlen Mildebrandt, Hannah Schopf, and Saskia Theune for assisting with participant recruitment, data collection, and data entry. For the discussion of the statistical analyses, the authors are very grateful to Dr. Wolfgang Scharke. The authors would further like to thank all participants who took part in this study and made an indispensable contribution. They also would like to thank their colleagues at the INM3, INM4, and the Child Neuropsychology Section of the University Hospital RWTH Aachen for their continued support during data acquisition and discussions of the data.

FUNDING AND DISCLOSURE

Dr. Freitag has been consultant for Desitin and Roche within the last 3 years. She receives royalties for books and psychotherapy intervention manuals on Autism-Spectrum-Disorder, Attention-Deficit/Hyperactivity Disorder as well as Major Depressive Disorder. Dr. Cholemkery receives royalties for books and psychotherapy intervention manuals on Autism-Spectrum-Disorder. Dr. Konrad has received speaking fees from Shire Pharmaceuticals. The remaining authors declare that they have no conflict of interest. This work was primarily supported by grants to Dr. Schulte-Rüther (German Research Foundation (DFG, SCHU 2493/2-1), Excellence Initiative of the German federal and state
Dr. Freitag was supported by the grant FR2069/2-1 from the German Research Foundation (DFG).

REFERENCES


FIGURE AND TABLE LEGENDS

Table 1. Demographic characteristics of the final participant samples

HC, Healthy Control; ASD, Autism Spectrum Disorder; TCI-140, Temperament and Character Inventory-140; TCI-140 RD, Reward Dependence Scale of the TCI-140

Figure 1. Illustration and timing of the (A) SN, (B) NN, and (C) NS condition with rewarding feedback

Conditions were presented in three separate runs (SN, NN, NS) of approximately 15 minutes, with 64 trials each (including 2 stimuli of each category A or B) resulting in 16 repetitions for each stimulus throughout each run. Each stimulus appeared twice in a learning block of 8 trials with no immediate repetition. Trials were presented in a pseudo-random order and the order of the 3 runs was counterbalanced between subjects. During each target presentation (max 2000ms), participants selected via button-press whether the learning target belonged to category A or B. Upon choice, a fixation cross (3000-5000ms) appeared, followed by a feedback screen (2000ms) with either positive or neutral probabilistic feedback, depending on correct or incorrect category choice, respectively. A complete trial took 13s, resulting in an inter-trial interval of (4000-6000ms).
Figure 2. Task x treatment x interval interaction in the ASD group

Participants with ASD showed better learning with social as compared to non-social targets and feedback in the OXT (A) but not PLC (B) condition during the second learning interval of the probabilistic reinforcement learning task during fMRI.

ASD, autism spectrum disorder; NN, task condition with non-social learning target and non-social feedback; NS, task condition with non-social learning target and social feedback; SN, task condition with social learning target and non-social feedback

Figure 3. Regions in the fMRI task where activation was associated with learning from feedback across groups, treatment and task conditions

Results were significant at p < .05 (cluster-level corrected, p < .001 voxel-level, k = 164 voxels). For illustrative purposes, the uncorrected level is presented here, but results are reported for the cluster level correction.

HC, healthy control; ASD, autism spectrum disorder; OXT, oxytocin; PLC, placebo

Figure 4. Neural correlates for the interaction between group x social feedback x treatment (A & B) and the interaction between group x social learning target x treatment (C & D)
Contrast estimates at the corresponding peak voxel ([-8 10 -10] and [-8 8 -10], respectively) of the NAcc ROI are depicted. Beta values (vertical axis) represent parameter estimates for the degree of the correlation of brain activation with the RPE. ASD: N = 15, HC: N = 24.

* = p < .05; # = p < .10; RPE, reward prediction error; ASD, autism spectrum disorder; HC, healthy control; OXT, oxytocin; PLC, placebo; NN, task condition with non-social learning target and non-social feedback; NS, task condition with non-social learning target and social feedback; SN, task condition with social learning target and non-social feedback
TABLES

Table 1.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>HC (N = 24)</th>
<th>ASD (N = 15)</th>
<th>Analysis by Student's t Test (Two-Tailed)</th>
</tr>
</thead>
<tbody>
<tr>
<td>% Male</td>
<td>100</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>Age in years</td>
<td>22.1 ± 1.88</td>
<td>21.79 ± 2.60</td>
<td>df 23.20, t -0.39, p .700</td>
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<td>IQ (estimated)</td>
<td>119.1 ± 9.35</td>
<td>113.53 ± 11.61</td>
<td>df 37.00, t -1.66, p .106</td>
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<tr>
<td>TCI-140</td>
<td>42.99 ± 4.93</td>
<td>39.39 ± 5.57</td>
<td>df 37.00, t -2.11, p .041</td>
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<tr>
<td>TCI-140 RD</td>
<td>43.21 ± 9.93</td>
<td>32.27 ± 14.21</td>
<td>df 37.00, t -2.83, p .007</td>
</tr>
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