How Oxytocin Modulates Distress and Support-Seeking in Adolescence According to Attachment Style? An Exploratory Eye-Tracking and Neurophysiological Study

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Introduction

There is a growing literature about oxytocin (OT), the paradigmatic "attachment hormone" that represents the biological aspect of attachment [1]. Oxytocin, a hypothalamic neuropeptide, acting both as a neuromodulator in the central nervous system and as a hormone in the periphery [2] appears more and more to be a potential preventive intervention or aid in psychiatric treatment, due to its prosocial effects [3]. It is a key molecule implicated in the regulation of several reproductive and social functions; for example, administering exogenous OT via a nasal spray increases trust and empathy [4,5]. Few studies have explored how oxytocin influences attachment behaviors and representations, although attachment is among the factors influencing the effectiveness of emotion regulation (ER) [6].

Attachment is an innate psychobiological behavioral system, activated in times of perceived distress [7] inciting children to seek proximity and comfort from their attachment figures [8,9]. The system is deactivated once a perceived sense of security and safety is reestablished. The quality of early attachment interactions with an attachment figure leaves an enduring mark on the developing person: attachment representation (secure or insecure; withdrawn or anxious) is likely to be associated with the way stress is perceived and dealt with across the lifespan [10-12]. Individuals with secure attachment have successfully mastered the process of seeking proximity to others for relief from distress, leading to a capacity to experience, express, and tolerate temporally distressing events. When attachment security is not
attained, the use of alternative, insecure attachment strategies of avoidance or anxiety may be triggered. Insecure withdrawn adolescents tend to suppress emotional expression and comfort-seeking strategies [13]. Insecure anxious adolescents, who tend to fear rejection and separation, express a greater desire for company, and present cortisol dysregulation secretion [14].

These studies, among others, show that insecure attachment is a potential factor of emotional dysregulation and the development of psychopathology in adolescence and adulthood [15,16]. Prior research indicates that intranasal oxytocin (INOT) yields different effects depending upon the participant’s attachment security, in randomized controlled studies (INOT versus placebo, PB) among adult populations [17]. Other effects of INOT on stress reactivity have been found. The conjunction of INOT administration and of social support from best friends had a psychosocial stress-buffering effect on hypothalamic-pituitary-adrenal (HPA) activation [18] among 37 healthy men exposed to the Trier Social Stress Test. Similarly, INOT reduced skin conductance response (SCR) in relation to fear-associated stimuli [19]. To date, only one study has explored INOT response in adolescents (healthy controls versus inpatients) regarding attachment-related trust [20]. There, INOT was found to increase the level of trust of inpatients to a “healthy control level” in a trust game over internet with their mother and a stranger. Among healthy adolescents, attachment security moderated the effects of INOT.

Together, these studies provide indications that a single dose of INOT can affect attachment representation, behavior, and physiological responses in humans. However, despite the importance of INOT treatment outcomes in attachment and socio-emotional contexts, the effects of OT on social support-seeking under stress in insecure adolescents are still insufficiently studied. To extend this line of work, this pilot study investigated the effects of a single dose of INOT on insecure male adolescents, through a randomized, double-blind, placebo-controlled trial, as part of a larger ER study [21,22].

In this study, we employed a distress-then-comfort-seeking paradigm divided into two phases. First, the adolescent’s attachment system was activated by the visualization of pictures of distress. Second, to determine how adolescents deactivate their attachment system, three pictures were presented simultaneously (comfort, joy-complicity, and neutral), corresponding to the phase of comfort-seeking. A multimodal approach (gaze parameters and SCR) was used to objectively assess INOT effects on this dynamic attachment-related ER. We hypothesized that INOT would modify behavioral and/or physiological reactions of insecure adolescents toward more secure strategies. During distress exposure, INOT should reduce stress reactivity and insecure strategies. During the support-seeking phase, INOT should enhance support-seeking toward comfort pictures.

Materials and Method

Participants

As part of a bigger study among 81 adolescents recruited in secondary schools in Besançon, France, twenty-five insecure male volunteers participated in a double-blind, placebo-controlled, parallel-treatment-with-INOT, randomized trial. A group of secure male volunteers (N=12), who participated in the study without treatment, was used as a reference group [22]. The reference group was concomitant with the study groups but was not randomized. It was used to estimate and characterize values in secure patients. All participants had normal or corrected-to-normal vision. Exclusion criteria were medical or psychiatric illness, use of medication, substance abuse, or smoking (more than 20 cigarettes a day). Participants were instructed to abstain from food and caffeine-based drinks for 2 hours before INOT administration. An informed consent form was signed by the participants, and their parents (for minors). An open-loop gift card with a value of 20 € was given to each adolescent for participation. The study was approved by the French Agency for the Safety of Health Products and the local ethics committee, and registered in the Clinical Trials Register (NCT02301715).

Study procedures

Two visits were necessary for the assessment protocol. During the first visit, participants were screened for inclusion and exclusion criteria by a psychiatrist; their psychiatric and attachment profiles were also assessed. During the second visit, an eye-tracking task combined with SCR measurement was performed 45 min after the treatment with INOT or PB. For more details, see the previously published study protocol [21].

Attachment assessment

All eligible participants responded to the Attachment Style Interview [23], in order to determine their attachment style. The ASI is a semi-structured interview, validated in French, and adapted for use with adolescents. It assesses attachment style, based on the ability to make and maintain supportive relationships with one parent and two “very close others”, who can be friends or family members. Attitudes about closeness to/distance from others, autonomy, and fear/anger in relationships were also assessed. Overall attachment style was categorized in terms of secure attachment or insecure attachment, the latter comprising the standard typologies of anxious and withdrawn. When two profiles of insecurity were found, the attachment style was coded dual. In order to control for individual psychometric dimensions, several psychometric questionnaires were used, including the Beck Depression Inventory (BDI) [24], the Spielberger State-Trait Anxiety (STAI-YA, STAI-YB) [25], and the Toronto Alexithymia Scale-20 (TAS-20) [26].
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Treatment

Each participant received either an active INOT (Novartis, Austria) (N = 12) or PB (N = 13) spray. Age-dependent dosage was employed: younger participants (aged 13 to 15) received 16 international units (IU), and older participants (aged 16 to 20) received 24 IU [27]. The placebo was custom-designed by the hospital pharmacy to match drug minus the active ingredient. Either INOT or PB was administered to each participant, 45 minutes before the experimental phase, which took place between 2 p.m. and 4 p.m., to take into account diurnal OT variation.

Pictures

The visual stimuli consisted of pictures from the Besançon Affective Picture Set-Adolescents (BAPS-Ado) [28], representing distress, comfort, joy-complicity, and a neutral state. Pictures of distress represented faces expressing sadness, anguish, or scenes of loss and separation. Comfort pictures represented scenarios of a parent comforting an infant or an adolescent after an episode of distress. Pictures of joy-complicity represented joyful moments. Finally, neutral scenes represented persons in the distance, walking along a street, or in the subway. The picture sequences were organized to "activate" and "deactivate" the attachment system. During the activation phase, a distress picture (N = 20) was presented for 10 sec. During the deactivation phase, 3 pictures (comfort N = 20, joy-complicity N = 20, and neutral N = 20) were presented together for 20 sec.

Eye-tracking measurement

Eye movements were recorded using the Remote Eye-Tracking Device (RED 500, Senso Motoric Instruments, SMI, Teltow, Germany) at a rate of 250 Hz. Details relating to the apparatus can be found in the previously published protocol [21]. Prior to detailed statistical analysis, the whole of each picture was treated as a single area of interest (AOI) for the 4 categories: distress, comfort, joy-complicity, and neutral. A gaze fixation should last for 80 ms on a surface of 100 pixels to be classified as a fixation [29]. When a single distress picture was presented, dwell time, which corresponds to the time that the gaze was within the distress picture, was measured. The mean of dwell time percentage, expressed as a percentage of the total duration of a trial, was determined as follows: dwell time (ms) / (end time - start time). It represents the salience or visual attractive power of the picture and reflects engagement patterns.

When three pictures were presented together (comfort, joy-complicity, and neutral), two other variables were measured: glance count and entry time (ms). Glance count corresponds to the number of glances at a target within a certain period, with saccades coming from outside. It was calculated by averaging all glances inside the AOI per trial. It reflects attentional focus. Entry time (ms) corresponds to the average time from stimulus onset to the first fixation on the AOI, in milliseconds per trial [29]. This parameter reflects the time needed to detect emotional visual stimuli and indicates the first gaze orientation toward each of the three pictures [30].

Neurophysiological measurement

The SCR was recorded using Biopac© MP 36 (Biopac © Systems, Inc., Santa Barbara, CA, USA). A pair of Ag/AgCl electrodes filled with isotonic electrode gel was attached to the distal phalanges of digits II and III on the non-dominant hand. The 3-minute baseline responses were recorded, with the Biopac system, without stimuli. The neurophysiological monitoring AcqKnowledge 4.3 software was synchronized with Experiment Center 3.0 software by event markers representing the beginning of each picture. The AcqKnowledge application was used to extract latency, amplitude of skin conductance responses (SCR), and skin conductance level. The SCR was defined as the maximum change in conductance (μSiemens) in the 0.1 to 6-second window after stimulus onset.

Intranasal oxytocin administration side-effects

We used questionnaires to detect any side-effects of INOT administration. All adolescents were asked whether they had experienced the most commonly reported side-effects immediately after the task, and then 30 days later, when adolescents answered the same questions by phone interview.

Statistical analyses

Analyses were performed using SAS, version 9.4 (SAS Institute, Cary, NC). To investigate the effects of INOT on the variables studied, a Wilcoxon test was used. Continuous variables are represented as: (Median [Q1; Q3]). Effects are considered significant if the p-value is below .05 for dwell time and SCR latency for distress pictures, and for glance count (Gaze parameters) for each picture category during the support-seeking phase. Exploratory secondary analyses will be performed with an adjusted level of statistical significance of .001 (for multiple criteria: entry time, dwell time, SCR amplitude, etc. and different picture categories). The statistical analysis was only performed between INOT and PB groups. Data from the reference group were used in a descriptive analysis, and no statistical comparison was performed with this group.

Results

Preliminary analyses

Data were first analyzed for outliers. Outliers for each eye-tracking variable were deleted from the analyses, using z-score with a mean threshold of +/- 3.29. After threshold application, almost all data were kept (99.9%).

Population characteristics

The study population was composed of 25 insecure male adolescents and 12 secure male adolescents. Both groups presented similar psychological dimensions (see Table 1 for population characteristics).

Table 1: Demographic and psychometric characteristics of adolescents in function of treatment.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Insecure OT (N = 12)</th>
<th>Insecure PB (N = 13)</th>
<th>Reference Group Secure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age Mean (SD)</td>
<td>17.67 (1.97)</td>
<td>17.31 (1.18)</td>
<td>15.83 (1.47)</td>
</tr>
<tr>
<td>TAS-20 Median (Q1, Q3)</td>
<td>45.00 (36.00, 55.00)</td>
<td>53.00 (44.00, 56.00)</td>
<td>46 (39.00, 54.00)</td>
</tr>
<tr>
<td>STAI-YB Median (Q1, Q3)</td>
<td>31.00 (30.00, 37.00)</td>
<td>38.00 (36.00, 41.00)</td>
<td>33.20 (29.00, 36.00)</td>
</tr>
<tr>
<td>STAI-YA Median (Q1, Q3)</td>
<td>28.00 (23.00, 32.00)</td>
<td>31.00 (26.00, 35.00)</td>
<td>29.70 (25.00, 35.00)</td>
</tr>
<tr>
<td>BDI Median (Q1, Q3)</td>
<td>1.00 (0.00, 5.00)</td>
<td>4.00 (2.00, 5.00)</td>
<td>2.70 (1.00,4.00)</td>
</tr>
</tbody>
</table>

Distress exposure

Effect of INOT on gaze parameters

Intranasal OT tended to reduce dwell time percentage for distress pictures \([W(1) = 3.4201, p = 0.0644] \); PB \((92.25 \ [88.08; 94.99]) \) INOT \((87.05 \ [80.26; 89.80]) \).

Effect of INOT on neurophysiological parameters

At a physiological level, it was hypothesized that INOT would attenuate neurophysiological arousal in response to distress pictures. As expected, the Wilcoxon test indicated a significant main effect of INOT on SCR latency (sec) \((W(1) = 4.2722, p = 0.0387) \). Intranasal OT increased SCR latency (sec) \((2.68 \ [2.40; 2.87]) \) compared to PB \((2.43 \ [2.23; 2.49]) \). For the reference group, SCR latency (sec) was 2.33 \((2.24; 3.12) \).

Intranasal OT did not influence SCR amplitude.

Support-seeking phase

Effect of INOT on gaze parameters

Following our hypothesis, INOT should enhance comfort-seeking. The Wilcoxon test indicated a significant main effect of INOT on increasing glance count for comfort \((W(1) = 4.8565, p = 0.027) \) and for neutral \((W(1) = 4.502 ; p = 0.0339) \). Other comparisons were not significant with an adjusted alpha level (Table 2).

Table 2: Gaze parameters for comfort, joy-complicity, and neutral pictures. Continuous variables are represented as: (Median [Q1; Q3]). Reference group will be described only.

<table>
<thead>
<tr>
<th>Picture Category</th>
<th>Gaze Parameters</th>
<th>Study Groups</th>
<th>Reference Group</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Entry time (ms) 1303.75 [959.23; 1939.38]</td>
<td>1877.13 [1186.40; 1991.60]</td>
<td>1292.67 [823.74; 2662.55]</td>
</tr>
<tr>
<td></td>
<td>Dwell time (ms) 4438.11 [3643.53; 6018.29]</td>
<td>5224.91 [4339.60; 6105.30]</td>
<td>5529.16 [3773.50; 6336.23]</td>
</tr>
<tr>
<td>Joy-Complicity</td>
<td>Glance count 4.92 [3.42; 6.58]</td>
<td>3.74 [3.00; 4.79]</td>
<td>3.4 [2.45; 4.58]</td>
</tr>
<tr>
<td></td>
<td>Entry time (ms) 974.15 [781.67; 1415.12]</td>
<td>1860.52 [934.45; 1967.53]</td>
<td>1013.05 [613.28;1494.67]</td>
</tr>
<tr>
<td></td>
<td>Dwell time (ms) 4506.62 [3679.71; 5995.52]</td>
<td>5644.77 [4898.54; 6354.23]</td>
<td>5729.16 [5028.26; 6298.41]</td>
</tr>
<tr>
<td>Neutral</td>
<td>Glance count 5.08 [4.13; 6.59] (^*)</td>
<td>3.65 [2.85;4.50] (^*)</td>
<td>3.33 [2.70;3.98]</td>
</tr>
<tr>
<td></td>
<td>Entry time (ms) 1013.37 [665.78; 2203.50]</td>
<td>2308.23 [1590.25; 2972.14]</td>
<td>2142.27 [1443.24;2716.27]</td>
</tr>
<tr>
<td></td>
<td>Dwell time (ms) 4795.81 [3793.75; 6305.74]</td>
<td>5294.77 [5092.53; 6844.56]</td>
<td>5290.21 [4382.77; 5889.95]</td>
</tr>
</tbody>
</table>

\(^*\)Significant statistical difference between the 2 study groups \((p-value<0.05) \) for glance count on each picture categories or \(p-value <0.001 \) for secondary criteria.

Discussion

This pilot study examined the effects of INOT versus PB on emotion regulation (ER) in 25 insecure male adolescents (12 INOT, 13 PB) under conditions of a distress-then-comfort-seeking paradigm. During the first phase, “distress exposure”, our results indicate that a single dose of INOT tended to reduce dwell time on distress pictures and decreased physiological arousal in response to them. This result might suggest that INOT reduces engagement patterns on distress and thus reduces the stress in response to distress. This behavior is supported by two studies on adult populations: one study indicating that INOT decreased eye contact in response to negative facial expressions [31], and another indicating that INOT decreased amygdala activity and cortisol secretion in response to negative emotional stimuli [18,32].

An important question in our study was the effect of INOT on comfort-seeking in insecure adolescents. As expected, insecure adolescents explored comfort pictures more under INOT than PB. This effect is corroborated by previous research highlighting the role of INOT in social support-seeking in distressed women [33]. Nevertheless, little is known about the effects of INOT on more specific attachment processes, such as support-seeking, among insecurely attached people [34,35]. In our study, we differentiate comfort from joy-complicity. Interestingly, the effect of INOT on
glance count was only significant for comfort pictures. During this support-seeking phase, another hypothesis of the study was that INOT would increase the feeling of security, and allow insecure adolescents to process emotionally not only attachment-related, but also non-attachment-related information (i.e. neutral pictures) as secure adolescents do [22]. Our results indicate that INOT enhanced glance count for comfort and neutral pictures in insecure adolescents compared to PB. In our previous study using the same paradigm, secure adolescents displayed a similar pattern: exploring the comfort and joy-complicity pictures, and then exploring the neutral pictures, which contain more details (e.g. streets and subways) [22]. In agreement with other studies suggesting that the exploratory system is optimal only when the attachment system is deactivated, and when the adolescent is soothed [7,36], our study suggests that insecure adolescents felt more secure to explore non-attachment-related information [6,8,37].

Despite the robustness of a multi-level approach, several limitations to our pilot study deserve consideration. The size of our sample does not allow us to comment on the generalization of our results. Future work is needed to determine INOT effects on emotion regulation in insecure adolescents in relation to attachment style, especially as the literature shows that the styles of insecure attachment [20], and its intensity [12,35,38-40], can both influence OT effects. We did not include females in this study as we were not able to control for pregnancy risk. This limitation is general in the literature.

### Conclusion

Insecure attachment is a potential factor of emotional dysregulation and the development of psychopathology in adolescence. Oxytocin, the paradigmatic “attachment hormone”, could potentially promote social bonding, and social support-seeking under stress. Our pilot study is one of the first double-blind randomized controlled trials to examine INOT effects on ER in insecure adolescents, in the specific context of attachment. Our findings show that INOT reduced neurophysiological arousal to distress, and increased comfort-seeking and exploration. We found that INOT produced no consistent side-effects or adverse outcomes, at single doses of 16-24 IU, assessed at two time points: about 90 min after nasal spray administration, and 30 days after administration. Future research could consider using INOT on a larger sample, over a longer period, and in different contexts, as INOT may have a nuanced range of effects that depend not only on attachment style but also on contextual factors.

### References


