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Title: The Role of Oxytocin in Maternal-Fetal Bonding & Social Interaction

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Abstract

Oxytocin is a small peptide usually associated with its effects in the reproductive system such as induction of labor and lactation. However, recent evidence has indicated that oxytocin plays an important role in social behavior in mammals, including humans. This review article outlines the basics of the role of oxytocin in parental-fetal bond formation and pair-bonding. Social recognition forms the basis of all social interaction and therefore the role of oxytocin in recognition is also discussed. Most studies done as of yet have been conducted of mammals such as prairie voles and rats. However, there has been some evidence which shows that similar mechanisms occur in humans. Pair-bonding and parent-fetal bond formation have very similar mechanisms and oxytocin plays an important role in both. For both processes to occur, there needs to be social recognition and memory of the infant or the partner, both of which are aided by oxytocin and other neurotransmitters, especially arginine vasopressin (AVP) and dopamine. Moreover, once the bond has been established, oxytocin plays a role in persistence of the bond even in the absence of the fetus/parent or the partner. There are similarities between the behavior and mechanisms of social bonding and addiction, therefore oxytocin could play a future role in treatment of social diseases, such as autism and addiction.

Introduction

Oxytocin (OXT) is a small nine-amino acid peptide produced by the supraoptic (SON) and paraventricular (PVN) nuclei of the hypothalamus. It is then primarily released by exocytosis

from the neurohypophysis and nerve terminals as a result of physiological stimuli. OXT binds to oxytocin receptors (OXTR) which causes a release of Ca²⁺. This calcium release can be blocked using OXTR antagonists such as d(CH2)5,Tyr(Me)2,Orn8]-vasotocin¹. OXT has long been known for its effects in inducing parturition and lactation. However, recent studies have implicated this peptide in other aspects of mammal and human life, such as the modification of social behavior, including pair-bonding between partners and parental bonding with the offspring². Social bonding is a vital process for the survival of species since it increases the chances of reproduction and protection of the young against predators³. Furthermore, social interaction is essential since isolation results in physical and mental disorders, and ultimately even death in a number of animal and human models¹.

Social Recognition

Social recognition is an important aspect and vital first step in the formation of social bonds in mammals (including humans), for example pair bonding and the formation of a relationship between parents and their offspring². In order for social recognition to occur, the mammal must go through steps, which include social approach, investigation, sensory learning, processing, and memory⁴. Olfaction, a way to calculate the degree of olfactory input (amygdala), and the reward circuitry are all essential parts of social recognition. This process utilizes steroid hormones, neuropeptides, and neurotransmitters such as norepinephrine and DOPA⁴. Oxytocin (OXT) also plays a vital part in social recognition².

Social Recognition in Different Parts of the Brain

Social recognition in rodents can be induced by OXT administration in the lateral septum, medial preoptic areas and olfactory bulb⁵. The lateral septum (LS) plays an important role in social recognition since rats administered OXT or AVP site-specifically spend more time with familiar rats compared to unfamiliar ones². Vasopressin works with OXT in social recognition, but it acts mainly only in the LS. In fact, when vasopressin in administered in the LS, social recognition improves, while administration of vasopressin receptor antagonist in the LS inhibits social recognition⁴. When OXT is specifically administered to the medial amygdala, it reverses the social memory deficit usually found in OXT-knockout mice. Therefore, the medial amygdala is also involved in social recognition².

There is an elevated noradrenergic stimulation in the olfactory bulb initiated by OXT or vasopressin. This decreases the inhibition on the olfactory bulb and there is an increase in the activity of the main output cells of the bulb. This results in plasticity in the olfactory bulb. Difference in the plasticity encodes for the difference between behavior towards strangers later in life⁴. Rodents primarily use olfactory stimuli for social recognition⁵ and can retain facial memory for up to two hours following recognition through olfactory cues⁵. When treated with OXT, their social memory is prolonged⁴. The recognition and learning of olfactory stimulus can be reproduced by norepinephrine administration in the locus coeruleus, indicating that norepinephrine is essential for the social recognition process⁴.

Social recognition in animals

Social recognition is important since it allows an animal to distinguish between its partner and a stranger². Social recognition increases when rats are administered with both peripheral and central OXT administration². In fact, there is an increase in retention time when the social discrimination test is administered². OXT-knockout mice display a lack of social memory. The lack of social recognition found in OXT knockout mice can be reversed by an OXT intracere-broventricular (ICV) injection given before the first exposure. However, an OXT injection administered after the initial exposure has no effect on social recognition⁵. This shows that OXT plays a role in the formation of social recognition, but not in recognizing the specimen after the first encounter⁵.

Pair bond formation is the rewarding mating experience associated with the olfactory sensory cue of the partner⁶. Social recognition and memory are essential for partner preference formation in monogamous mammals, such as prairie voles. In fact, lesions in the olfactory bulb of male prairie voles decrease partner preference formation⁴. Certain studies show that OXT in the mouse brain is not used to discriminate between strangers and familiar animals, but rather discrimination between specific mice to recognize a certain mouse⁵. This corresponds to the formation of a pair bond to a specific mouse⁵.

Social recognition in the maternal bond induced by OXT is observed by a variety of mammals including sheep, goats, Southern pig-tailed macaques, rhesus macaques and rodents². In preweaning rate, administration of OXTR antagonist prior to the formation of the mother-infant bond inhibits the rat from making the association between the mother's social cues and the odor⁴. Newborn rat pups associate the maternal care they receive, such as licking, with stimuli of the mother, especially the odor⁴. In fact, when newborn pups are "licked" with a paintbrush with the odor of peppermint, the pups show a preference towards the smell of peppermint throughout their life⁴. Mice with defective OXT systems are unable to recognize familiar animals using particular social cues, such as facial features, odor or walk⁵. However, they respond normally to other odors, such as lemon⁵.

AVP also plays a role in social recognition since administration of AVP in rats increases their social memory and treatment with AVP receptor antagonist centrally and peripherally inhibits social recognition². CD38-knockout mice show a deficiency in social recognition. CD38 is a transmembrane protein needed for the translocation of Ca²⁺ ions inside the cells by activation cADP ribose. The Ca²⁺ ions induce release of OXT and therefore, lack of CD38 stops social recognition². Total removal of OXT, OXTRs or CD38 in genetically-engineered mice eliminates any type of social memory, even a few seconds after seeing another animal⁵. However, there is no change in non-social memory and learning, indicating that OXT has no effect on memory outside social purposes⁵.

Social Recognition in Humans

In humans OXT in used to distinguish between emotions by looking at the faces of familiar people by subtle facial cues. Intranasal OXT administration increases this ability⁶. Administration of OXT increases both the short- and long-term memory in facial recognition of males and

females². When ten people were treated with intranasal OXT and another ten were given a placebo, and they were all shown faces of strangers 45 minutes after administration. They were then asked to choose the faces they saw in the pictures, and the people treated with OXT recognized more people compared to those with the placebo⁷. Therefore, OXT may be used therapeutically in patients with social deficits, although more research is needed⁵.

Parental Care

Affectional bonds are a primary human need, especially the one between mother and infant⁸. This bond increases the infant's chances of surviving beyond reproductive age⁹. This relationship is different in rodents and herding animals. While rodents display maternal behaviors universally towards infant rodents, herding animals such as sheep are selective towards their own infants and develop a stronger mother-infant bond¹⁰. In cases where a mother-infant bond is impossible, such as in rabbits, the infants tend to share their early environment with siblings¹¹. In prairie voles, both virgin males and females tend to avoid and, in certain cases, even attack infants, but parturient mothers are attracted to the infants and find them irresistible¹⁰. Furthermore, blood transfusions from a parturient rat to a virgin female rat show an increase in the parental response of the virgin rat¹⁰.

The oxytocinergic and dopaminergic systems have been associated with the bond formed between parents and their infants. Apart from these systems, other factors such as stress during the pregnancy, early caregiving experience and relationships throughout the life also have an effect on this bond⁹. Increased plasma OXT concentrations in the first six months of the infant's life have shown a positive parental behavior in both parents¹².

Although most studies have been done on other mammals, similar mechanisms support animal and human parenting mechanisms¹⁰. Apart from its importance in forming the parent-infant bond, OXT is also essential to protect the parents from the stress and demands of the perinatal period¹³.

Maternal Behavior

Apart from its well-known effects regarding labor and milk ejection, OXT also increases maternal behavior¹⁴. The main biological mechanism involved in the onset and maintenance of maternal behavior is the oxytocinergic and dopaminergic systems which induce motivation and reward circuits¹⁵. OXT has projections to the medial preoptic area (MPOA), the bed nucleus of the stria terminalis, the ventral striatum including the nucleus accumbens and the ventral tegmental area, which are all involved in positive maternal behavior⁹. In fact, the nucleus accumbens of biparental prairie voles contains more OXTRs (OXTR) than non-monogamous species which to not display biparental care¹⁶.

DOPA signals start in the ventral tegmental area and the substantia nigra and project into the ventral striatum, dorsal striatum, prefrontal cortex and anterior cingulate cortex⁹. Onset of reward mechanisms is through the activation of the mesocorticolimbic pathways which involves the ventral striatum and the medial prefrontal cortex and temporal prediction errors activate the nigrostriatal pathway which involves the dorsal striatum and the dorsal prefrontal cortex⁹.

The OXT and DOPA systems connect at the mesocotricolimbic pathway system which is involved with motivation and reward in both mother and infant⁹. The mesocotricolimbic pathway induces the onset of maternal care by inhibiting avoidance and violence towards the infant¹⁰.

Recognition of the infant is an essential part of bond formation. This can be selective or nonselective. Nonselective recognition occurs in mothers that give birth to multiple young ones, for example most rodents, including rats. In nonselective recognition, maternal care is focused toward a general stimulus on the infant rather than a specific one, and if another young has the same stimulus the mother will care for it during the postpartum period. Selective recognition usually occurs in species where the mother gives birth to a small number of infants (usually one), such as sheep and most primates. In selective recognition, maternal care is directed toward the specific infant birthed by the mother and other offspring are rejected¹⁷.

Pregnancy & Parturition

During pregnancy, the medial prefrontal cortex (MPOA) enhances the mesocorticolimbic pathways by increasing its effect on the amygdala ¹⁰. The amygdala is involved with learning and memory ¹⁸. Although not a direct part of the mesocorticolimbic pathway, it is linked with reward since memory of previous rewards following certain actions create motivation for similar action by the mesocorticolimbic pathway ¹⁸. The MPOA also monitors the concentration of steroid hormones during pregnancy ¹⁸. During pregnancy there is an increase in plasma progesterone and estrogen which are secreted by the ovaries. This is followed by a sudden drop in progesterone which signals the start of parturition. When the MPOA receives these signals, there is an increase in brain sensitivity to OXT and prolactin induced by an increased production of their receptors ¹⁰.

At pregnancy and lactation, there is an increase in gene expression of OXT in brain regions associated with an increase in maternal behavior, such as dopaminergic substantia nigra. There is also neural modification of the hippocampus, which is associated with facilitated learning, spatial memory and emotion processing of facial cues, all of which are essential to form a close mother-infant bond⁹.

During parturition, the MPOA activates the ventral tegmental area (VTA) in two ways; firstly, through direct projection, and also indirectly through the PVN of the hypothalamus. This leads to an increased DOPA concentration in the nucleus accumbens, which activates DOPA D1 receptors. This stops the inhibition of the nucleus accumbens on the ventral pallidum, activating the nucleus pallidum. This makes the mother responsive to infant cues via projections of the ventral pallidum to the thalamus, and cortical and mesencephalic motor nuclei, which are all involved in the display of maternal nurturing towards her pup¹⁰.

Infancy

OXT following parturition and throughout infancy is important since it is involved in motor output response towards infant cues¹⁰, own infant recognition⁹ and maternal aggression towards intruders¹⁹. Postnatal stimuli from the infant, such as crying, facial expressions and suckling stimulation aid in reshaping the maternal brain in the period immediately following

parturition, which is a time of high neural plasticity in the mother⁹. OXT is essential since an intracisternal injection of OXT in a rat mother few days following the birth of a pup induces grooming four months later¹⁶.

Several parts of the brain are involved in forming a bond with the infant following parturition ¹⁸. The MPOA is essential for positive maternal behavior. Destruction of the MPOA stops all signs of maternal care in rats and administration of OXT, estrogen, DOPA or prolactin in the MPOA of virgin female rats induce maternal behaviors. There is stronger activation of the nucleus accumbens after seeing pictures of their children in mothers who display more positive behavior and less intrusiveness. Infant cries also activate the anterior insula and prefrontal cortex. The anterior insula is associated with feelings of empathy, which are essential as a motivator for maternal care¹⁰. The prefrontal cortex is involved in cognition, motivation, decision making and emotion, all of which are needed for positive maternal behavior¹⁸. Activation of the prefrontal cortex is also linked with maternal sensitivity towards pups, decreased stress responses to separation from her infants and secure attachment following reunion. There is an increased activation of the amygdala of infant when seeing pictures of their mothers compared to pictures of strangers. This increases saliency towards the mother¹⁰.

Recognition of own infant is an important aspect of mother-infant bonds. Sheep mothers start recognizing their offspring after a few hours following parturition¹¹. When viewing their own infants, mothers have an increased activation of the mesocorticolimbic pathway and nigrostriatal pathway, which are associated with positive maternal care⁹.

In humans, the mesocorticolimbic dopaminergic reward regions and the PVN and SON of the hypothalamus are activated when secure mothers are shown pictures of their children. Furthermore, activation of reward centers, such as the NAcc, is correlated with an increase in peripheral OXT levels in the mother²⁰. Conversely, depressed mothers who have lower OXT levels do not have activation of these brain centers and therefore, do not experience reward in maternal behavior, finding it difficult to bond with their child.

Mothers have reduced reactivity to stressors through decreases corticotrophin releasing factor, adrenocorticotropin releasing factor and cortisol¹³. However, mothers have a more aggressive response towards strangers, used to protect the infant, elicited by OXT²¹. In fact, OXT-knock-out mice showed very little aggression towards strangers²¹. An increase in OXT activates the amygdala in humans¹⁹. This induces fear and aggression, increasing the chances of maternal survival and infant protection¹⁹.

OXT is well known for its role in milk ejection during breastfeeding. However, recent studies have shown that mothers who breastfeed have more activity in the brain areas which are related to maternal behavior, such as the striatal reward regions¹⁴. Breastfeeding has been shown to improve depressive symptoms²². Firstly, mothers who breastfeed improve their feeling of self-efficacy, which is inversely associated with the feeling of helplessness felt by mothers suffering from post-partum depression. Furthermore, breastfeeding increases the bond between the mother and the infant. Breastfed children also sleep more when compared to children who are not breastfed. This allows a more stable sleeping pattern for the mothers,

which can improve her mood²². There is also an increase in OXT and DOPA concentrations in the substantia nigra during suckling which explains why breastfeeding is linked with a stronger mother-infant bond compared to mothers who do not breastfeed⁹. Similarly, in mothers who deliver vaginally, there is a greater activation of the hypothalamus when hearing own infant cries, leading to an OXT surge and an increased DOPA release compared to mothers who deliver via Caesarean section⁹.

Later Life

OXT is responsible not only for the onset of maternal care, but also for the maintenance of it⁹. The mother-child bond is one which lasts a whole lifetime, especially in humans. Long-term persistence of the mother-infant bond necessitates both maternal and fetal memory, which occurs through synaptic plasticity within the neural circuit regulating maternal behavior. It is hypothesized that the action of dopamine and OXT on the nucleus accumbens is vital for memory formation in the maternal-infant bond (similar to pair bonding). During a study, rats are allowed an hour of bonding postpartum and then the offspring is removed. Then, dopamine agonists, OXT antagonists, and control solutions are injected into the mother's nucleus accumbens. Ten days later, the mothers are exposed once again to offspring. The study shows that both by blocking both D1 and D2 receptors, maternal memory is stopped and there is no maternal-infant bond formation. The OXT antagonists used were relatively non-selective and also block V1a vasopressin receptors, and therefore, the results regarding OXT are not reliable and with regards to this study, these receptors may or may not be involved in maternal memory formation.

Most studies are done on rodents since it is difficult to measure central OXT concentrations and there is no evidence that central and peripheral OXT concentrations are correlated²³. However, certain studies have been done on humans in the past few years to show that even in humans, OXT plays an important role in maternal behavior¹⁴. Another study shows that mothers who have a secure bond with their child, who display an understanding of emotions and empathy experience a rise in central OXT levels when interacting with their child. This change is not seen in mothers who do not have a secure bond with their child, such as depressed mothers¹⁴.

Furthermore, the rise in OXT levels in secure mothers is correlated with a rise in OXT in their infants, showing the dual effect that a healthy relationship between the mother and the infant has on both parties²⁴. Therefore, the quality of maternal care in the early life of the infant affects the OXT system development in the child, and has an effect on the parental behavior of the offspring when they become a parent⁹.

Measuring maternal behavior is very difficult, but two methods for doing this are used. These are the Adult Attachment Interview (AAI) and the Strange Situation Procedure (SSP)⁹. In the SSP, a young child (<2 years) and the mother are put in a strange or challenging situation to assess whether the infant chooses the mother over a stranger in such situations²⁵. Using this method, it has been found that the development of secure attachment between mother and infant is affected by the mother's attachment history and stress throughout her life⁹. AAI uses an interview, made with both the mother and the child, which is analyzed for specific markers

(words or phrases) and patterns. The relationship between mother and child is then sorted as "secure", "insecure/ dismissing" or "insecure/preoccupied". During an AAI a difference is made between "cognitive" attentiveness and "affective" attentiveness. Cognitive information is information given during the conversation regardless of emotions felt by the mother and affective information is information regarding the emotions and fears of the mother. "Insecure/dismissing" mothers tend to focus on the cognitive information and "insecure/preoccupied" mothers focus on the affective information. "Secure" mothers coordinate well between the two types of information. OXT levels in the "insecure" mothers tend to be less than in "secure" mothers, indicating that OXT has an important role in providing secure attachment between mother and child.

Maternal Neglect

Neglect is most often associated with the mother, since she is normally considered the primary caregiver, even though mothers go through neuroendocrine changes during pregnancy, birth and lactation, which prepares them for caregiving⁹. There are two types of neglect; physical and emotional neglect⁹. While physical neglect is when the mother fails to provide materialistic needs, such as food, hygiene, clothing, or education, emotional neglect is when the caregiving lacks physical affection and emotional warmth⁹.

Physical neglect is associated with loss of "cognitive" understanding, while emotional neglect usually results in lack of "affective" understanding⁹. Emotional neglect, unlike physical neglect, produces negative long-term effects on the child's social and emotional behavior⁹. Reduced maternal care is linked with lack of OXT, since depressed mothers have decreased levels of salivary OXT¹⁰. Also, in rhesus monkeys, there was a decrease in OXT levels of the cerebrospinal fluid (CSF) in monkeys who had non-maternal rearing⁹. Stress throughout pregnancy is associated with decreased OXTR binding in brain areas associated with positive maternal care⁹. This increases anxiety in the mother and reduces licking and grooming, therefore decreasing OXTR binding in the offspring, producing long term effects in the maternal behavior of the offspring⁹. Reduced levels of OXT in late pregnancy are associated with increased symptoms of depression postpartum²⁶.

Depressed mothers have a lower nucleus accumbens activation which causes reduced caregiving motivation and decreased anterior insula activation. Therefore the mother has reduced empathy towards infant cues such as cries or facial expressions 10 . Empathic overarousal also has adverse effects on maternal behavior since it can interfere with effective parenting and can lead to intrusive parenting 10 . A reduced level of licking and grooming in rat pups is an indication of maternal neglect. These pups show a higher level of DNA methylation of estrogen receptor - α gene promoter, resulting in an inhibition of the development of the OXT system 9 .

Drug abuse in mothers is also associated with negative maternal care⁹. Cocaine abuse decreases OXT levels in the hypothalamus, inhibiting the mesocorticolimbic pathway, which is essential for good maternal behavior⁹. Furthermore, cocaine abuse reduces OXTR levels in the brain, therefore decreasing maternal behavior towards pups²⁷. The pups fail to develop the

oxytocinergic system properly, making them more susceptible to future cocaine abuse themselves²⁷. A socially rich environment later in life, may compensate for the maternal neglect, but it cannot reverse the effects caused earlier in life⁹.

Paternal Behavior

Most studies focus on maternal behavior since the mother is usually considered the primary caregiver. However, OXT also has a role in positive paternal care¹⁰. When OXT is administered intra-nasally, there is an increase in father-infant touch¹⁰. This increases the duration of infant gaze towards the father and salivary OXT of the infant¹⁰. OXT levels in the father correlate with proprioceptive tough and exploratory play between the father and the infant, creating a stronger father-child bond¹⁰. Fathers who are involved in caregiving have stronger activation of the VTA and the anterior insula following infant stimuli¹⁰. When anterior insula activation is too high or too low, this interferes with paternal behavior since the father experiences too much or too little empathy towards the infant¹⁰.

Biparental care, such as that found in prairie voles, is associated with a decrease in the father's testosterone levels²⁸. Men with a higher testosterone level show decreased sympathy towards the cries of an unknown infant²⁸. Men with lower testosterone levels, show an increased involvement in fatherhood at the expense of sexual interaction²⁸. They display increased empathy, frustration tolerance, but decreased sexual motivation²⁸. This is also important in positive paternal care since it prevents sexual motivation from "competing" with parenting effort¹⁰. Paradoxically, testosterone is beneficial in paternal behavior since it is converted to estrogen in some species¹⁰. Estrogen induces the development of the oxytocinergic system¹⁰

Size of testes also affects paternal care¹⁰. Men with smaller testes usually show more involvement since testes size has an inverse relationship with VTA activation which is involved in increased paternal behavior¹⁰. As of yet, there is not enough evidence to correlate oxytocin and testosterone levels in fatherhood, although several studies have shown that an increase in oxytocin levels alters testosterone levels and has an effect on paternal behavior²⁹³⁰.

Alloparental Behavior

Alloparental care is parental behavior by someone other than the biological parent³¹. OXT treatment increases female involvement in virgin rats to pups³² as opposed to aversive behaviors which are seen when there is no administration of OXT⁹. In male prairie voles, administration of OXT antagonist decreases alloparental behavior transiently¹⁶. At 21 days postpartum, male prairie voles treated with OXT antagonist show a decrease in parental care and in time spent with pups, but at 60 days postpartum, none of the above effects are seen¹⁶.

Baby "schema" (cuteness) plays an important role in producing parental care in non-parents¹². There is stronger ventral striatum activation in nulliparous women when viewing pictures of "cute" babies¹². There is also an increased in activation of the nucleus accumbens and VTA when non-parents see pictures of "cute" babies¹⁰. Children who were raised in orphanages and then adopted show an increase in amygdala activation, increasing their trust¹⁰. Therefore, they tend to approach unfamiliar adults, putting themselves at risk¹⁰.

Pair Bonding

Pair bond formation is when a male and a female animal develop a monogamous relationship beyond reproductive purposes¹⁸. Monogamous mammals, following the formation of a pair bond, share a territory and defend their nest together with their partner¹⁸. They also both participate in the rearing of their young¹⁸ and guard each other². Most mammals display non-monogamous relationships, unlike prairie voles and humans, which develop monogamous pair bonds¹⁸. In fact, pair bond formation is only present in less than 5% of mammalian species².

The mechanisms which drive partner preference formation are very similar to those which cause maternal-infant bonding. As with the mother-infant bond, once the bond has occurred, it endures even in the case of absence of partner stimulation. Therefore, pair bonding also involves neural plasticity in selective neural pathways as a result of specific social stimuli from the partner, such as olfactory stimuli (similar to maternal bond in sheep). A noticeable difference is that both male and female prairie voles develop pair bonding following mating, as opposed to parental bond, which is focused mostly on the female¹⁷.

Compared to more promiscuous animals, prairie voles display higher OXTR concentrations in the nucleus accumbens, amygdala, bed nucleus of the stria terminalis and prefrontal cortex¹⁸. This shows that OXT plays a role in the formation of a pair bond. There are similarities between the mechanisms of being in love and substance addiction³³. Subjects who are in love experience stress-induced relapse, lack of regard for consequences, are unable to quit, and lose track of time³³. They also lose their ability to make rational decisions about personal risks involving their choices³³. All of these behaviors are also seen in people who are suffering from substance abuse³³. Also, both mechanisms involve OXT stimulation in the brain of the subject³³.

Partner Preference Formation

Partner preference formation differs between the field and the laboratory⁵. In the wild, there is abundant mate choice and monogamy is observed when a male and a female pair are seen together frequently and share their nest. In the laboratory, there is no mate choice and pairs are selected randomly by the experimenter⁵. Sexually naive animals are put together for a period of cohabitation. After the cohabitation period, the animals are tested in a three-hour partner preference test, where the animal is put in a testing chamber consisting of three cages. One contains the partner they spent their cohabitation period with; one contains a stranger animal; the other is empty. The animal can move freely between the three chambers⁵. Monogamous prairie voles show preference for their partner since they spend twice as much time in their cage than in the other two⁵. However, non-monogamous montane voles spend most of the time in the neutral cage⁵. The success of pair bond formation increases when the cohabitation period is longer⁵. In fact, the amount of OXTRs in the *nucleus accumbens* of female prairie voles is directly related to the speed of partner preference formation¹⁸. When the voles were administered with OXTR antagonist after the cohabitation period, partner preference formation

was reduced, but sexual activity was not affected⁵.

Both monogamous and non-monogamous animals have genes encoding for OXT and OXTRs, since both are needed for lactation⁵. Therefore the difference in pair bonding behaviors may be due to OXTR distribution patterns. For example, the *nucleus accumbens* contains higher concentrations of receptors in monogamous mammals compared with non-monogamous species⁵. Moreover, OXT administration induces partner preference formation in both male and female prairie voles, indicating that OXT has a role in both sexes, even though this effect is much less potent in the male³⁴. In fact, a higher dose of OXT is needed in male prairie voles for partner preference formation³⁴.

The difference of the effect of OXT in pair bond formation between male and female voles may be due to the OXTR concentration difference in male and female prairie voles³⁴. While in male voles, there is a high concentration of OXTRs in the LS, in the female, there is a high concentration of OXTRs in the prelimbic cortex and the nucleus accumbens³⁴. Another difference between partner preference formation of the male and female prairie voles is the importance of mating for this process to occur¹⁸. In female rats, OXT-induced partner preference formation is present even when mating does not occur². However, in males, prior to mating, OXT treatment does not induce partner preference formation and OXTR antagonist has no effect on pair bond formation². Therefore, mating is essential for the formation of a pair bond in the male². Once the pair bond has been formed, male prairie voles showed aggression towards unfamiliar females¹⁸.

In prairie voles that have not formed a pair bond, OXT or AVP administration, increases social behavior such as side-by-side contact and decreased aggression. In male rats, chronic OXT administration increased social interaction with females². In monogamous marmosets, administration of OXT induces more contact, decreases partner approach latency and increases food sharing. OXTR antagonist inhibits these actions².

Social recognition is an important aspect of pair bond formation since it allows the mammal to distinguish between the partner and a stranger. OXT plays an important role in social recognition². OXT-knockout mice have reduced social memory which is restored by treatment with OXT before the first social encounter with a particular partner³³.

While the role of OXT in parturition explains its increase during the formation of the maternal-fetal bond, a study showed that similar vaginocervical stimulation during mating induces OXT release in the NA to facilitate the formation of partner preferences in females³⁵.

The Interaction of OXT with Other Neurotransmitters & Hormones

One of the chemicals in the brain which interacts with OXT in the formation of all social bonds, including the parental bond and pair bond formation, is DOPA¹⁸. In fact, administration of haloperidol, which blocks the activity of DOPA, before mating, stops partner preference formation in male prairie voles¹⁸.

Both oxytocinergic and dopaminergic systems work together in the formation of a pair bond³⁴. In fact, formation of pair bond stimulated by DOPA receptors in the nucleus accumbens of the female prairie vole is inhibited by OXTR antagonist and partner preference formation stimulated by OXT is inhibited by treatment with DOPA receptor antagonist in the nucleus accumbens of the female prairie vole³⁴. Pair bonding is affected by DOPA stimulation in the nucleus accumbens¹⁸. However, there is also dopaminergic activity in other regions of the mesocorticolimbic pathways, such as the prefrontal cortex¹⁸.

Two DOPA receptors are involved in the regulation of partner preference formation; D1R and D2R¹⁸. These two receptors produce opposing effects on pair bond formation when activated¹⁸. Administration of D2R agonists establishes partner preference formation in situations where there should be no partner preference formation¹⁸. However, when D1R are activated, pair bond formation is blocked¹⁸. In fact, there is a lower concentration of D1-receptors and a higher concentration of D2-receptors in monogamous prairie voles as compared to non-monogamous meadow voles¹⁸. Following the formation of a pair bond, activation of D1R in the nucleus accumbens induces aggressive behavior towards strangers, therefore inhibiting the formation of a second pair bond¹⁸.

Cyclic adenosine monophosphate (cAMP) works with both the oxytocinergic and dopaminergic systems to regulate partner preference formation⁶. D2 receptor stimulation decreases cAMP concentration and D1 receptor stimulation increases cAMP levels⁶. In fact, an increase in cAMP signaling induced by a decrease in PKA inhibits pair bond formation, while a decrease in cAMP levels in the nucleus accumbens increases partner preference formation⁶.

AVP, which is structurally very similar to OXT produces effects similar to OXT during the formation of a pair bond³⁴. However, its effects are more potent in males, whereas females are more responsive to OXT³⁴.

Corticosterone and OXT work together in the formation of pair bonds³⁴. In sexually naive females, exposure to males increases central OXT release and decreases corticosterone release³⁴. OXT administration decreases corticosterone concentration and therefore OXT seems to interact with the HPA axis to regulate partner preference formation³⁴. When female prairie voles are administered corticosterone, they show an increased preference to strangers compared with controls. This shows that corticosterone in females has the opposite effect of OXT, since administration of OXT increases preference to the romantic partner³⁶.

Pair Bonding in Humans

Most studies and experiments done so far study pair bonding in animals, mostly prairie voles, however, some research shows that the mechanisms observed in prairie voles are similar to those in humans². In humans, MRI scanning showed that when seeing a romantic partner, there was more activation of the ventral tegmental area and caudate when compared to a familiar person¹⁸.

A male and female participant are treated with OXT and shown a picture of an additional male and female. A day later, the participants are shown pictures of an additional man or woman and

are asked to choose which one they would like to get to know better. The participants show an increased interest in the person they were shown the previous day. However, the participants are only asked to choose the person they want to spend time with, not have a romantic relationship with and therefore, this experiment does not provide reliable proof for the role of OXT in pair bond formation¹⁸.

When the Relationship Closeness Induction Task is conducted, administration of OXT increases intimacy in a conversation in females, but does not have the same effect in males². Hugging, touching, massage, nipple stimulation and orgasm induce release of OXT and promote trust and increased eye contact³³. A couple were asked to talk about their first date and they were observed for social cues³⁷. The subjects showed nonverbal displays such as nods, smiles and leaning towards partner when recalling experiences of romance³⁷. This was positively correlated with an increase in plasma OXT levels in the subjects³⁷.

Certain variations of the OXTR gene, including rs13316193, rs2254298, rs1042778, rs226849413 and rs226849 are associated with a decrease in empathy during a romantic relationship². Rs7632287 is associated with increased pair bonding in females². Interestingly, females who have experienced abuse or neglect early in their childhood have decreased levels of OXT⁶. This affects their ability to form social relationships, such as pair bonds⁶.

Initial Stages of a Romantic Relationship

There is a heightened level of OXT in the plasma of new couples. These increased OXT levels are correlated with more interaction between couples and can predict which relationships are likely to have survived six months later¹⁸.

When a person is in love, every encounter with the partner causes OXT release, which enters the mesocorticolimbic pathway, increasing the importance of social cues and memory. Therefore, the person pays more attention towards certain characteristics of their partner such as sights, sounds, odors, behaviors and specific characteristics³³. During the first phase of partner addiction, high levels of sensory information is gathered about the partner such as looks, touches, words, smells, shape of body and face. This induces positive reward stimulation by releasing DOPA in the nucleus accumbens and activating opioid receptors. The oxytocinergic, dopaminergic and opiate systems work together to form a positive feedback loop, whereas the experiences with the partner create reward outcomes and the reward increases the feelings towards the partner³³.

Long-Term Relationships

In substance addiction, the subject develops tolerance towards the drug, producing negative symptoms when the subject is not under the effect of the substance. In very much the same way, when a person is away from their partner, especially in the initial stages of the relationship, they feel symptoms similar to those of tolerance. OXT seems to play a role in mitigating these symptoms³³.

The positive feedback loop accumulates and ultimately the subject adapts³³. Therefore, DOPA signaling is altered in favor of D1R, leading to a reduction in reward and an increase in negative of aggressive behavior, producing one of the following three possible outcomes³³.

Once the euphoria at the start of the relationship ends, it is replaced by a sense of constant happiness as opposed to extreme reward felt when the mesocorticolimbic pathway is at its highest activity. This is similar to tolerance in substance abuse³³. Another possibility is that once the initial euphoria subsides, encounters more frequently contain conflict and the relationship has a high amount of negative emotions³³. However, the couple increase the amount of encounters despite the negative effects. This is similar to dependence-induced escalation of consumption³³. The last possibility is that DOPA release continues but receptor signaling favors D1receptors, which are associated with aggressive behavior³³. Therefore, the subject displays an aggressive behavior when they try to defend their "territory" (jealousy in a relationship). This is similar to compulsive and escalating drug abuse³³. OXT administration increases good communication behavior in relation to negative communication behavior during a conflict². Positive marital relationships are correlated with increased life expectancy; better immune function; cardiovascular health².

The End of a Romantic Relationship

The loss of a partner puts a human at a higher risk of both mental and physical health problems. This has been studied in detail using monogamous prairie voles, which showed signs of depression following bond loss³⁸. A study conducted in 2014 uses male prairie voles and allows them to form a pair-bond with a female for 24 hours. Then, some are allowed to remain with their partner while another group are separated for four weeks. It is shown that the male prairie voles which are separated from their partner show a significantly higher level of anxiety in both the elevated plus maze and the light-dark box³⁹. Although human studies are limited, similar anxiety and depression symptoms are often seen following the end of a romantic relationship in humans.

Reunion after a break-up is similar to relapse in substance abuse³³. In both initial stages of sobriety and after a breakup, there are increased levels of dynorphin which induces a negative effect³³. OXT mitigates the negative effects and decreases the chances of reunion/relapse³³. Consolation and social encounters after a break up increase release of OXT, which further decreases the symptoms of withdrawal³³.

Conclusion

Although the role of oxytocin in human social behavior and its implications on pharmacotherapy are still not well known, current studies have shown that this neurotransmitter plays an essential role in these processes. Oxytocin nasal spray has been shown to increase trust towards strangers during a game. It also reduces amygdala and caudate nucleus regions of the brain which cause fear, conditioning, and social conditioning⁴⁰. Such studies help reveal the mechanisms involved in normal human interaction and bonding, and can increase the understanding of social disorders. Furthermore, oxytocin could play a vital role in future

therapeutics battling social disorders such as autism⁴¹, and has also been implicated in treating addiction⁴².

References

- 1. Viero C, Shibuya I, Kitamura N, Verkhratsky A, Fujihara H, Katoh A, *et al.* (2010) *Oxytocin: Crossing the Bridge between Basic Science and Pharmacotherapy*. CNS Neurosci Ther. Oct;16(5):e138-56.
- 2. Lieberwirth C, Wang Z. (2014) *Social bonding: regulation by neuropeptides*. Front Neurosci. Jun 24;8:171.
- 3. Neumann ID. (2009) *The advantage of social living: brain neuropeptides mediate the beneficial consequences of sex and motherhood.* Front Neuroendocrinol. Oct;30(4):483-96.
- 4. Modi ME, Young LJ. (2012) *The oxytocin system in drug discovery for autism: animal models and novel therapeutic strategies.* Horm Behav. Mar;61(3):340-50.
- 5. Hammock EA, Young LJ. (2006) Oxytocin, vasopressin and pair bonding: implications for autism. Philos Trans R Soc Lond B Biol Sci. Dec 29;361(1476):2187-98.
- 6. Ross HE, Young LJ. (2009) Oxytocin and the Neural Mechanisms Regulating Social Cognition and Affiliative Behavior. Front Neuroendocrinol. Oct;30(4):534-47.
- 7. DeGutis JM, Chiu C, Grosso ME, Cohan S. (2014) *Face processing improvements in prosopagnosia: successes and failures over the last 50 years.* Front Human Neurosci. 8:10.3389/fnhum.2014.00561.
- 8. Bowlby J. (1969) Attachment and Loss. 2nd ed. New York: Basic Books;.
- 9. Strathearn L. (2011) *Maternal neglect: oxytocin, dopamine and the neurobiology of attachment*. J Neuroendocrinol. Nov;23(11):1054-65.
- 10. Rilling JK, Young LJ. (2014) *The biology of mammalian parenting and its effect on offspring social development.* Science. Aug 15;345(6198):771-6.
- 11. Anacker AM, Beery AK. (2013) *Life in groups: the roles of oxytocin in mammalian sociality.* Front Behav Neurosci. Dec 11;7:185.
- 12. Luo L, Ma X, Zheng X, Zhao W, Xu L, Becker B, et al. (2015) Neural systems and hormones mediating attraction to infant and child faces. Front Psychol. Jul 17;6:970.
- 13. Bell AF, Erickson EN, Carter CS. (2014) *Beyond labor: the role of natural and synthetic oxytocin in the transition to motherhood.* J Midwifery Womens Health. Jan-Feb;59(1):35,42: quiz 108.
- 14. Kim S, Soeken TA, Cromer SJ, Martinez SR, Hardy LR, Strathearn L. (2014) *Oxytocin and Postpartum Depression:* Delivering on What's Known and What's Not. Brain Res. Sep 11;0:219-32.

- 15. Swain JE, Kim P, Spicer J, Ho SS, Dayton CJ, Elmadih A, et al. (2014) Approaching the biology of human parental attachment: brain imaging, oxytocin and coordinated assessments of mothers and fathers. Brain Res. Sep 11;1580:78-101.
- 16. Miller TV, Caldwell HK. (2015) Oxytocin during Development: Possible Organizational Effects on Behavior. Front Endocrinol (Lausanne). May 19;6:76.
- 17. Numan M, Young LJ. (2016) *Neural mechanisms of mother-infant bonding and pair bonding: Similarities, differences, and broader implications*. Horm Behav. Jan;77:98-112.
- 18. Love TM. (2014) *Oxytocin, Motivation and the Role of Dopamine*. Pharmacol Biochem Behav. Apr;0:49-60.
- 19. Campbell A. (2013) *The evolutionary psychology of women's aggression*. Philos Trans R Soc Lond B Biol Sci. Oct 28;368(1631):20130078.
- 20. Strathearn L, Fonagy P, Amico J, Montague PR. (2009) *Adult attachment predicts maternal brain and oxytocin response to infant cues*. Neuropsychopharmacology. Dec;34(13):2655-66.
- 21. Bosch OJ. (2013) *Maternal aggression in rodents: brain oxytocin and vasopressin mediate pup defence*. Philos Trans R Soc Lond B Biol Sci. Dec 5;368(1631):20130085. doi:10.1098/rstb.2013.0085.
- 22. Figueiredo B, Dias CC, Brandao S, Canario C, Nunes-Costa R. (2013) *Breastfeeding and postpartum depression: state of the art review*. J Pediatr (Rio J). Jul-Aug;89(4):332-8.
- 23. Gimpl G, Fahrenholz F. (2001) *The oxytocin receptor system: structure, function, and regulation.* Physiol Rev. Apr;81(2):629-83.
- 24. Feldman R, Gordon I, Zagoory-Sharon O. (2010) *The cross-generation transmission of oxytocin in humans*. Horm Behav. Sep;58(4):669-76.
- 25. Rehn T, McGowan RTS, Keeling LJ. (2013) *Evaluating the Strange Situation Procedure (SSP) to Assess the Bond between Dogs and Humans*. PLoS One.8(2):e56938. doi:10.1371/journal.pone.0056938.
- 26. Skrundz M, Bolten M, Nast I, Hellhammer DH, Meinlschmidt G. (2011) *Plasma Oxytocin Concentration during Pregnancy is associated with Development of Postpartum Depression*. Neuropsychopharmacology. Aug;36(9):1886-93.
- 27. Williams S, Johns J. (2014) Prenatal and gestational cocaine exposure: Effects on the oxytocin system and social behavior with implications for addiction. Pharmacol Biochem Behav. Apr;0:10-21.
- 28. Mascaro JS, Hackett PD, Rilling JK. (2014) *Differential neural responses to child and sexual stimuli in human fathers and non-fathers and their hormonal correlates.*Psychoneuroendocrinology. Aug;46:153-63.
- 29. Weisman O, Zagoory-Sharon O, Feldman R. (2014) *Oxytocin administration, salivary testosterone, and father-infant social behavior*. Prog Neuropsychopharmacol Biol PsychiatryMar 3;49:47-52.
- 30. Bales KL, Saltzman W. (2016) *Fathering in rodents: Neurobiological substrates and consequences for offspring.* Horm Behav. Jan;77:249-59.

- 31. Perkeybile AM, Delaney-Busch N, Hartman S, Grimm KJ, Bales KL.(2015) *Intergenerational transmission of alloparental behavior and oxytocin and vasopressin receptor distribution in the prairie vole.* Front Behav Neurosci. 9:10.3389/fnbeh.2015.00191.
- 32. Saltzman W, Maestripieri D. (2011) *The neuroendocrinology of primate maternal behavior*. Prog Neuropsychopharmacol Biol Psychiatry. Jul 1;35(5):1192-204.
- 33. Burkett JP, Young LJ. (2012) *The behavioral, anatomical and pharmacological parallels between social attachment, love and addiction*. Psychopharmacology (Berl). Nov;224(1):1-26.
- 34. Young KA, Gobrogge KL, Liu Y, Wang Z. (2011) *The Neurobiology of Pair Bonding: Insights from a Socially Monogamous Rodent*. Front Neuroendocrinol. Jan;32(1):53-69.
- 35. Waldherr M, Neumann ID. (2007) *Centrally released oxytocin mediates mating-induced anxiolysis in male rats*. Proc Natl Acad Sci U S A. Oct 16;104(42):16681-4.
- 36. Aragona BJ, Wang Z. (2004) *The prairie vole (Microtus ochrogaster): an animal model for behavioral neuroendocrine research on pair bonding*. ILAR J. 45(1):35-45.
- 37. Debiec J. (2007) From affiliative behaviors to romantic feelings: a role of nanopeptides. FEBS Lett. Jun 12;581(14):2580-6.
- 38. Gobrogge K, Wang Z. (2015) Neuropeptidergic Regulation of Pair-bonding and Stress Buffering: Lessons from Voles. Horm Behav. Nov;76:91-105.
- 39. Sun P, Smith AS, Lei K, Liu Y, Wang Z. (2014) *Breaking bonds in male prairie vole: long-term effects on emotional and social behavior, physiology, and neurochemistry*. Behav Brain Res. May 15;265:22-31.
- 40. Kosfeld M, Heinrichs M, Zak PJ, Fischbacher U, Fehr E. (2005) Oxytocin increases trust in humans. Nature. Jun 2;435(7042):673-6.
- 41. Opar A. (2008) Search for potential autism treatments turns to 'trust hormone'. Nat Med. Apr;14(4):353-.
- 42. Kovacs GL, Sarnyai Z, Szabo G. (1998) Psychoneuroendocrinology. Nov;23(8):945-62.