Noxious or Non-noxious Inputs to Oxytocin Neurons: Possible Roles in the Control of Behaviors

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Oxytocin plays an essential role in milk ejection and parturition in mammals. Oxytocin has also been shown to be involved in the control of various behaviors, including anxiety-related behaviors, food intake and affiliative behaviors.

We previously showed that noxious stimuli or stimuli previously paired with noxious stimuli (conditioned fear stimuli) activate hypothalamic oxytocin neurons via activation of brainstem catecholaminergic/prolactin-releasing peptide (PrRP)-positive neurons. Oxytocin neurons are activated not only by noxious stimuli but also by non-noxious touch stimuli. Social contact has been suggested to activate oxytocin neurons. Non-noxious tactile stimuli induce 50-kHz ultrasonic vocalization, an index of positive states in rats, and activate hypothalamic oxytocin neurons, suggesting that pleasant tactile stimuli activate oxytocin neurons.

Physiological roles of oxytocin released during noxious or non-noxious tactile stimuli remain to be clarified. Noxious stimuli increase anxiety-related behavior, while pleasant sensory stimuli have pro-social actions. We have shown that endogenous oxytocin reduces anxiety-related behaviors, induces a decrease in amounts of food intake per meal, and facilitates social recognition via distinct neural pathways. Roles of oxytocin released during sensory stimuli may be dependent upon the sensory stimuli used, and oxytocin may contribute to the prevention of overreactions to noxious stimuli or mediate pro-social or anxiolytic actions of pleasant tactile stimuli.

KEYWORDS: oxytocin, noxious stimuli, conditioned fear, pleasant touch, ultrasonic vocalization

1. Introduction

In mammals, oxytocin is mainly synthesized in the hypothalamus and released from the neurohypophysis, and it facilitates parturition or lactation in females. However, oxytocin is synthesized and released into the peripheral circulation not only in females but also in males. Furthermore, oxytocin is an evolutionarily conserved peptide, and oxytocin or its homologs are expressed not only in mammals but also in non-mammal vertebrates including agnathans, fishes and amphibians [18, 37]. Some kinds of insects, snails and nematodes also express oxytocin homologs. It is thus easy to speculate that oxytocin has functions other than parturition or lactation. In fact, C. elegans, which has a long evolutionary distance from mammals, expresses the homolog of oxytocin named nematocin [4, 22]. Nematocin has been shown to be required for normal sexual behaviors of males and for salt-chemotaxis-related behaviors [5]. These findings suggest that sexual behavior and salt homeostasis are fundamental roles of oxytocin and its related peptides.

In mammals, the neurohypophysis releases oxytocin and its closely related peptide vasopressin. Oxytocin and vasopressin differ by only two amino acids. Genes for these two peptides have the same structures (three exons and two introns) and are located adjacent on the same chromosome in opposite transcriptional directions. Thus, the oxytocin and vasopressin genes are considered to have derived from a gene duplication that occurred approximately 450 million years ago. Oxytocin and vasopressin are synthesized in distinct neurons located in the magnocellular regions of the paraventricular and supraoptic nuclei in the hypothalamus and are released into the peripheral blood stream from the neurohypophysis, to which oxytocin- or vasopressin-synthesizing neurons project. Oxytocin is also synthesized in neurons located in the paraventricular region of the paraventricular nucleus of the hypothalamus or in the bed nucleus of stria terminals. These oxytocin neurons project to other brain regions [36, 37, 49].

Considering that an oxytocin homolog plays an important role, across the evolutionarily diverse animal species, in normal reproductive behaviors, which are regarded as prototypes of social behaviors, it is not surprising that in mammals that construct a complex society, oxytocin is involved not only in reproduction-related behaviors, including the formation of pair-bonding in monogamous species, sexual behavior, and parental behavior, but also in various other social behaviors such as social bonding among consociates. In mammals oxytocin has been shown to facilitate social memory, increase recognition of emotional expression, and mediate social buffering that reduces stress responses by

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being accompanied by mothers, partners or cage mates. Thus, it is possible that oxytocin facilitates not only finding and forming appropriate pairs, giving birth and nursing children but also establishing friendship and building and maintaining appropriate social relationships [10, 17, 34, 59, 68, 76]. Oxytocin has been implicated in disorders associated with dysfunctional social behaviors, including autism spectrum disorders [27, 41, 47].

Oxytocin has also been reported to have various actions in mammals [24, 50, 56, 65], including human-dog bonding [48], anxiolytic effects [49, 79], modulation of stress-induced activation of the hypothalamo-pituitary-adrenal axis [20, 30, 62], modulation of (emotional) memory [45], induction of analgesia [25], termination of food intake [77], suppression of salt appetite [66], an increase in energy consumption [6, 67], control of body temperature [33], anti-inflammatory actions [11], facilitation of wound healing [26, 73], restoration of ovariectomy-induced osteoporosis or obesity [12], protection of cardiomyocytes [28], recovery of age-dependent muscle mass decline [19], protective actions on neuronal damage [71], and control of neuronal development [35, 82].

Oxytocin actions appear to be mediated mainly by the oxytocin receptor [24]. However, oxytocin has also been reported to act on vasopressin receptors and the GABA receptor. Oxytocin can induce analgesia via activation of vasopressin V1A receptor [61]. Social deficits observed in oxytocin receptor-deficient mice are restored by oxytocin application, indicating that oxytocin acts on receptors other than the oxytocin receptor to facilitate social behavior [60]. Oxytocin has also been proposed to act on the GABA receptor containing a delta subunit to block ethanol-induced reduction in locomotion [7].

Here, we review recent findings with focus on the activation and functions of oxytocin neurons in response to somatosensory inputs.

2. Sensory inputs to oxytocin-synthesizing neurons

2.1 Noxious inputs

Hypothalamic oxytocin neurons receive somatosensory information. Hypothalamic neurosecretory neurons in the supraoptic nucleus, where oxytocin neurons are localized, have been shown to receive nociceptive information from peripheral organs bilaterally [29]. Plasma oxytocin concentrations are increased following noxious stimuli. Nociceptive information is conveyed by Aδ or C fibers to dorsal horn neurons, which project ascending fibers to the medulla oblongata (such as the ventrolateral medulla, nucleus tractus solitarius), lateral parabrachial nucleus, periaqueductal gray, thalamus, and hypothalamus [23, 69]. Noxious stimuli activate noradrenergic neurons in the medulla oblongata, facilitate noradrenaline release within the hypothalamus [57] and activate oxytocin neurons in the hypothalamus [53].

Oxytocin release in response to noxious stimuli is impaired by noradrenaline depletion or by an α1 adrenergic receptor antagonist [52]. The medulla oblongata contains A2 noradrenergic and A1 noradrenergic cells. Local application of a noradrenaline neuron-selective neurotoxin into the A1 noradrenergic region of the ventrolateral medulla but not total ablation of the dorsomedial medulla, which contains A2 noradrenergic neurons, impairs noxious stimuli-induced oxytocin release [54]. All these findings suggest that noxious stimuli activate hypothalamic oxytocin neurons via an activation of medullary A1 noradrenaline neurons projecting to the hypothalamus (Fig. 1).

There are direct projections from the spinal cord to the hypothalamus, including the paraventricular nucleus [14, 23]. Nociceptive afferents also project to the periaqueductal gray matter or parabrachial nucleus [69], from where some neurons project to the hypothalamic paraventricular nucleus [30]. The roles of these projections in sensory transmission to oxytocin neurons in the hypothalamus remain to be clarified.

Not only noxious stimuli but also stimuli previously paired with noxious stimuli, conditioned fear stimuli, activate noradrenergic neurons in the medulla oblongata and oxytocin neurons in the hypothalamus [56]. Hypothalamus-projecting medullary noradrenergic neurons that are activated in response to conditioned fear stimuli are suggested to co-express prolactin releasing peptide (PrRP) and are mainly located in the A2 region of the nucleus tractus solitaries [52, 56, 83]. PrRP is expressed mainly in the A2 noradrenergic neurons, modestly in the A1 noradrenergic neurons and slightly in the dorsomedial hypothalamus [16, 55]. Although conditioned fear stimuli activate PrRP neurons located in these three regions, the majority of the hypothalamus-projecting PrRP neurons which are activated by conditioned fear stimuli are found in the A2 region [84]. Conditioned fear stimuli-induced activation of oxytocin neurons in the hypothalamus is impaired by PrRP deficiency [80] and by local destruction of noradrenergic projections to the hypothalamus [83]. All of these findings suggest that conditioned fear stimuli activate hypothalamic oxytocin neurons via activation of the medullary A2 PrRP/noradrenaline neurons projecting to the hypothalamus. The upstream region of the A2 PrRP/noradrenaline neurons has been shown to be the medial amygdala [80].

Physiological roles of oxytocin released in response to noxious stimuli or conditioned fear stimuli remain to be clarified. Oxytocin has analgesic [61] or anti-fear [36] actions, consistent with a view that oxytocin attenuates noxious or conditioned fear responses and contributes to the prevention of overreactions of the body in response to noxious stimuli or conditioned fear stimuli.

2.2 Non-noxious inputs

Hypothalamic oxytocin neurons receive information not only from nociceptive receptors but also from non-nociceptive sensory receptors.
2.2.1 Suckling inputs

In lactating females, activation of somatosensory receptors by infant suckling stimulates oxytocin-synthesizing neurosecretory neurons in the hypothalamus. Oxytocin released into the bloodstream contracts myoepithelial cells surrounding the mammary alveoli and induces milk ejection. Suckling stimuli induce burst firing of oxytocin neurons. The burst firing of oxytocin neurons is synchronized bilaterally among the whole paraventricular and supraoptical nuclei, which is caused by bilateral and intrahypothalamic communications [8].

Massaging or licking of the nipple activates ipsilateral dorsal horn neurons of the spinal cord via C-type afferent fibers derived from the skin around the nipple. Studies with lesions and stimulation have shown that the sensory information from the dorsal horn ascends within the ipsilateral lateral funiculus of the spinal cord, relaying in the lateral cervical nucleus in the cervical segment of the cord, indicating the involvement of the spinocervicothalamic tract. The dorsal column and spinothalamic pathways may not be important for the milk-ejection reflex, since sections of these pathways do not block the milk-ejection reflex. The afferents conveying suckling information from the lateral cervical nucleus cross the midline in the lower brainstem, project to the mesencephalic lateral tegmentum and, after synaptic relays such as in the zona incerta and in the field of Forel, finally reach hypothalamic oxytocin neurons [75]. Involvement of noradrenergic pathways from the medulla oblongata to the hypothalamus has also been suggested in the control of oxytocin release during suckling [75].
Not only suckling stimuli but also visual or auditory stimuli previously associated with suckling have been suggested to trigger milk ejection. The sight or sound of the infant facilitates or triggers the reflex in rats, sheep, and cows [15, 21, 58].

2.2.2 Sexual sensory inputs

Oxytocin-synthesizing neurons in the hypothalamus have been shown to be activated during sexual behavior [75]. Plasma oxytocin concentrations are increased at ejaculation and orgasm. In the pre-copulatory phase, plasma oxytocin concentrations start rising, suggesting sexual arousal or sensory inputs as well as vaginocervical stimulation by intromissions activate oxytocin neurons. Tactile stimulation of the penis activates oxytocin neurons in the hypothalamus [78]. Centrally and peripherally released oxytocin plays a role in ejaculation in males, in lordosis in females, and in pair bond formation [32, 72].

2.2.3 Non-sexual sensory inputs

Tactile stimuli that are not apparently related to reproductive behaviors have also been shown to increase plasma oxytocin concentrations [64, 70]. Mutual contact and grooming are associated with increased urinary oxytocin concentrations in female tamarins [63]. In chimpanzees, urinary oxytocin concentrations increase following grooming [13]. In humans, skin-to-skin contact with the baby and the massage-like hand movements by the baby induce oxytocin release in the mother [44]. Recently, we found in rats that non-noxious stroking stimuli that induce emission of 50-kHz ultrasonic vocalization, an index of a positive emotion in rats, activate hypothalamic oxytocin neurons and increase plasma oxytocin concentrations, suggesting that pleasant touch stimuli activate hypothalamic oxytocin neurons [51].

Innocuous tactile stimuli activate low-threshold mechanoreceptors of cutaneous sensory neurons. The touch information is conveyed via the large-diameter fast-conducting myelinated Aβ afferents or small-diameter slow-conducting unmyelinated C fibers. The Aβ fibers mediate a discriminative touch sense of the glabrous or hairy skin, while the unmyelinated C fibers mediate a sense of affective touch of the hairy skin [1, 46, 85]. C fibers with low-threshold mechanoreceptors are lacking in glabrous skin. Affective touch sense is induced by gentle stroking with an appropriate velocity. The relationship between stroking velocity and firing rate of C fibers shows an inverted U-shape. Similarly, the relationship between stroking velocity and pleasure sense shows an inverted U-shape. Brushing at a speed of 1-10 cm/s, which induces a pleasant sensation, activates C tactile fibers most effectively. Brushing at a lower or higher speed is less effective. In contrast, firing rates in myelinated Aβ fibers in response to stroking increase as the stroking velocity increases. However, Aβ afferents may also contribute to affective feeling, since gentle touch on the palm, where C tactile fibers are lacking, can cause a pleasant sensation [38].

There appear to be, at least, two subtypes of C fibers with low-threshold mechanoreceptors that contribute to pleasant touch: first, neurons that express vesicular glutamate transporter, tyrosine hydroxylase [40], a chemokine-like protein TAFA4 and the Runx1-dependent transcription factor Zfp521 [42, 43], and second, neurons that express the Mas-related G-protein-coupled receptor MrgrpB4 [74].

C fibers conveying low-threshold mechanoreceptor inputs project to lamina I neurons of the spinal dorsal horn. The dorsal horn neurons project to the contralateral parabrachial nucleus [3]. Clinical observations that surgical cutting of the anterolateral spinotential tract for treatment of intractable pain impairs the sense of affective touch suggest that the anterolateral spinotential pathway signals both affective touch and noiception [46]. Roles of the spinocervicalthalamic pathway remain to be clarified. Detailed studies with animal experiments concerning neural pathways should be performed.

Gentle stroking activates somatosensory S1 and S2 cortices and also the posterior insular cortex, posterior superior temporal sulcus, medial prefrontal cortex, dorsal anterior cingulate cortex, and medial orbitofrontal cortex, where affective or social processing is performed [46].

Gentle touch reduces anxiety and decreases pain sensation and has pro-social actions. Affective touches also influence the development of neural connections controlling social behavior [31, 46, 70]. Oxytocin has been suggested to have similar actions. It is thus tempting to speculate that oxytocin mediates social or affective actions of gentle touch.

3. Conclusion

Noxious stimuli activate dorsal horn neurons via Aδ and C fibers. The nociceptive information ascends in the contralateral spinotential tract and enters the hypothalamus and stimulates oxytocin neurons via activation of brainstem reticular formation including noradrenergic neurons in the medulla oblongata. On the other hand, affective touch activates dorsal horn neurons via C fibers expressing VGLUT3, tyrosine hydroxylase and TAFA4 or MrgrpB4 and stimulates oxytocin neurons. Oxytocin induces anxiolytic and analgesic actions and facilitates pro-social behaviors. Thus, oxytocin may attenuate actions of noxious stimuli and mediate prosocial, anxiolytic or developmental actions by affective touch.

Oxytocin application has been reported to induce various actions including pro-social [76], metabolic or anti-ageing effects and has drawn much attention. However, some actions of oxytocin application are controversial and the
mechanisms underlying the majority of oxytocin actions remain to be clarified [39, 81]. In conclusion, somatosensory stimuli activate oxytocin neurons in the brain and thus affect physical or psychological health.

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