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Resilience to stress and social touch
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Modern lifestyle and adversities such as the COVID-19 pandemic pose challenges for our physical and mental health. Hence, it is of the utmost importance to identify mechanisms by which we can improve resilience to stress and quickly adapt to adversity. While there are several factors that improve stress resilience, social behavior—primarily in the form of social touch—is especially vital. This article provides an overview of how the somatosensory system plays a key role in translating the socio-emotional information of social touch into active coping with stress. Important future directions include evaluating in humans whether stress resilience can be modulated through the stimulation of low-threshold C-fiber mechanoreceptors and using this technology in the prevention of stress-related neuropsychiatric disorders such as major depressive disorder.

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Introduction
Modern lifestyle is associated with high levels of stress, especially in the work environment [1]. Additionally, there are social challenges facing humanity today such as the mental health crisis arising from the COVID-19 pandemic. SARS-CoV-2 and its hazardous variants have greatly increased the level of stress in our lives, mainly because they increased perceived threat and social isolation. As a consequence, mental illnesses such as major depressive disorder and anxiety disorders are becoming more prevalent [2]. In order to understand the relationship between resilience to stress and social touch, it is essential to first understand the neurobiological mechanisms of the stress response.

Stress is a concept that was coined by Hans Selye in the last century. This concept describes the way that the physiology of our body continually seeks to conform to environmental demands and adapt to them [3]. Through our sensory systems we perceive these environmental demands or stressors. Sensory information is translated into action potentials in the sensory organs and that information travels to the brain where the danger levels of stressors are evaluated [4]. When the integrity of the organism is threatened, brain structures such as the amygdala and the hypothalamus issue bodily changes and modulate the response to stress [5]. One important component of this response engages the sympathetic nervous system leading, among others, to the release of adrenaline from the medulla of adrenal glands into the blood [6]. Adrenaline increases heart rate and blood circulation, preparing the body to cope with stress [6].

A second system to be recruited is the hypothalamic-pituitary-adrenal (HPA) axis, which is part of the slow response to stress [6]. This neuroendocrine axis translates the neuronal activity associated with stress into a hormonal message that is sent to the adrenal glands [6]. From the cortex of these glands, the main stress hormone (cortisol in humans and corticosterone in rodents), is released into the blood [6]. These hormones increase blood sugar and produce the energy needed to sustain bodily activity under adverse conditions [6]. There are several extra-hypothalamic systems that play a key role both in the generation and regulation of stress responses including, for example, the locus coeruleus and the endocannabinoid system [7].

Stress and the somatosensory system
Sensory systems play a fundamental role in stress since they are responsible for transmitting information from the environment to the amygdala in order to activate and deactivate stress responses. Without that information, the amygdala cannot evaluate the possibility of danger and modulate responses to stress accordingly [8]. Despite the importance of sensory systems in both inducing and inhibiting stress, their therapeutic potential has been left largely untapped.

One potential therapeutic application involves the somatosensory system and, more specifically, the cutaneous mechanoreceptors of the skin — also known as low-threshold mechanoreceptors (LTM) [9]. These LTM contain a class of unmyelinated C fibers that are called CLTM in non-human animals and C-Tactile (CT) afferents in humans [10]. These fibers prefer gentle, dynamically moving stimuli at velocities between 1–10 cm/s and are believed to convey affective touch between

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conspecifics [11]. The socio-emotional information from affective touch reaches an area of the brain called the posterior insula [12]. This area is a multi-sensory hub and well connected with the anterior insula [13] and through this way it connects with other parts of the brain including the prefrontal cortex, anterior cingulate cortex, orbitofrontal cortex, and amygdala [14]. Therefore, affective touch could be a key factor to improve adaptation to stressors and promote resilience to stress. Note, however, that touch targeted towards other LTM s such as A-beta fibers and their projections to primary and secondary somatosensory cortex may also be relevant for stress.

Resilience to stress

Our early life experiences and associated epigenetic changes including, for example glucocorticoid receptor expression, remodel the brain circuits that modulate stress responses [15,16]. Just as a runner improves their performance through daily training, the biological mediators that regulate the response to stress improve their performance each time we are exposed to stressors and adapt to them. In the early 1970s, Norman Garmezy was the first to focus his attention on how humans cope with stress [17]. He observed that some children had an active way of coping with stress and were able to adapt faster [17]. Garmezy coined the term ‘resilience’ to explain this phenomenon and defined it as the ability to adapt quickly to adversity [17]. Resilience to stress is a biological-behavioral process that is continuously generated over time when we are exposed to stress and adapt to it [18]. Simply put, resilience emerges from the learning generated from our life experiences with stress.

The concept of allostasis put forward by Bruce McEween can help elucidate this idea: stress pushes us away from the homeostasis or balance we have with the environment in which we live [5]. Allostasis is related to the process of going back to homeostasis in which the physiological parameters necessary for the mediators of the stress response to function outside of homeostasis are generated [5]. In this scenario, the performance of the sympathetic nervous system and the HPA axis demand an energy cost or allostatic load that allows adaptation to stress [5]. Glucocorticoids, for example, cortisol and corticosterone, by the activation of glucocorticoid receptors, translate the effects of stress into changes in neuronal plasticity in brain structures that control the response to stress, such as the hippocampus and the medial prefrontal cortex [19]. Thus, stress hormones improve the performance of allostatic mediators, such as the sympathetic nervous system and HPA axis, when they are exposed to stressors again [5]. This improves the coping response to stress, in turn, allowing a faster adaptation to stressors [19]. When adaptation to stress does not happen, an allostatic overload is generated which triggers an imbalance both in the release of glucocorticoids, as well as in the ratio between the glucocorticoid receptors and mineralocorticoid receptors [20]. Mineralocorticoid receptors have a 10-fold higher affinity for corticosterone compared to glucocorticoid receptors. At the beginning of the stress response, mineralocorticoid receptors bind glucocorticoids in the cytoplasm of neurons, while glucocorticoid receptors are activated at the end of the stress response and participate in adaptation and recovery [20]. At a pathological level, imbalance in the ratio between the glucocorticoid receptors and mineralocorticoid receptors triggers neuro-inflammatory processes, mainly in the hippocampus, amygdala and medial prefrontal cortex, which impair the ability to cope with stress, increasing the susceptibility to stress-related neuropsychiatric diseases, such as mood disorders and neurodegenerative diseases like Alzheimer’s disease [20]. To prevent these mental illnesses, it is necessary to understand what behavioral factors can improve resilience to stress.

CLTMs and stress resilience

The main biological characteristic of resilience to stress is that the activity of the HPA axis is optimal for generating active coping with stress and achieving adaptation [5]. Lower or higher levels of HPA axis activity generate allostatic overload and trigger the behavioral phenotype susceptible to stress [5]. Research suggests that slow stroking, potentially engaging the CT system, can help to optimize HPA activity. In a recent study, the skin of rats was stroked at a slow (5 cm/s) or fast (30 cm/s) velocity before subjecting them to a chronic unpredictable mild stress paradigm [21]. Interestingly, slow stroking seemed to dampen the HPA axis relative to fast stroking; in the slow stroking condition only, plasma corticosterone levels were similar to those obtained for non-stressed rats [21]. At a behavioral level, slow stroking generated an anxiolytic effect in stressed animals and stimulated active coping, showing an increase in climbing and a decrease in floating behavior in the forced swim test [21]. These results demonstrate that it is possible to modulate stress resilience through the somatosensory system in rats (Figure 1). Moreover, they agree with another study showing that gentle skin stimulation decreases the development of depressive-like behaviors and improves episodic memory in rats that were exposed to the chronic unpredictable mild stress protocol [22].

Resilience and social behavior

Resilient rodents show active coping with social defeat stress, developing a social behavior like non-defeated rodents in social interaction tests [23]. Social behavior can modulate responses to stress via two mechanisms that are activated by social touch. The first mechanism is related to oxytocin; this hormone is synthesized in the paraventricular nucleus of the hypothalamus and social touch stimulates its synthesis and increases its plasma concentration [12,24,25]. Thus one may speculate that CT afferents stimulated by social touch increase oxytocin release through neural connections between the posterior
Conceptual model to explain the cross-talk between social behavior and stress. Social touch is a key element of social behavior in mammals. Gentle stroking touch applied at optimal velocity of cutaneous low-threshold C-fiber mechanoreceptors (CLTM) triggers oxytocin release and attenuates the effects of stress [12*,41], an important phenomenon related with social buffering. An active stress-coping strategy is associated with social buffering and stress resilience [35,36]. Enhances neuronal plasticity in the ventral tegmental area-nucleus accumbens-medial prefrontal cortex (VTA-NAc-mPFC) brain circuit further increasing motivation to develop social behavior in which social touch is included [37]. On the other hand, lack of nurturing touch in early life induces allostatic overload and stress vulnerability [5*,12*]. Increases of neuro-inflammation in brain areas that modulate stress responses triggers neuropsychiatric disorders such as major depression [20].

Social buffering and stress
The tendency of social behavior to mitigate the effects of stress is known as ‘social buffering’ [31] (Figure 1). Thus, affiliative tactile stimulation is an important component of social buffering and through this mechanism can modulate resilience to stress (Figure 1). For example, negative feedback of the HPA axis increases after maternal licking and grooming of pups in animal models [32], while in humans CT stimulating touch reduces sympathetic arousal in preterm infants [33]. The evidence shown above supports the hypothesis that tactile stimulation of the skin can modulate resilience to stress.

Insula and the paraventricular nucleus. There is also cross-talk between the oxytocin system and the HPA axis. Oxytocin can dampen the HPA axis which leads to a decrease in plasma cortisol levels when rodents are exposed to stress [26] (Figure 1). In humans, intranasal administration of oxytocin was found to suppress the cortisol response to psychological stress and to attenuate emotional sensitivity after stress [27]. The second mechanism by which social touch may control stress responses depends on the insula and its complex connections to cortical and subcortical regions. The anterior part of the insular cortex has strong functional connectivity with the hippocampus and the medial prefrontal cortex [14]: when these areas of the brain are activated, they inhibit the HPA axis and the sympathetic nervous system thus decreasing the stress levels [6].

Notably, social touch is not only relevant for stress in mammals. It is also beneficial in fish where tactile stimulation was shown to reduce stress, fear, and aggression [28–30]. Thus, it appears that the mechanism by which touch modulates resilience to stress has been highly conserved in evolution. Indeed, fish apart from making direct physical contact, can detect conspecifics through vibrations of water currents. This sort of remote touch may be a precursor of and functionally similar to direct touch which features in the interactions of land-dwelling species [29].
carry out a social behavior and serves as a natural reinforcer in some mammals [40] (Figure 1). There is a high density of oxytocin receptors expressed in the VTA-NAc-DNA circuit which stimulates its functioning during stress [41, 42] (Figure 1).

Conclusions and future directions
One of the most important evolutionary benefits of social behavior is social buffering, which allows the development of active coping and resilience to stress (Figure 1). In this context, the stimulation of C-LTM s improves resilience to stress in rats. Future experiments in humans are needed to establish a similar causal pathway for CTS and to evaluate their therapeutic potential in stress-related diseases such as major depressive disorder and anxiety disorder. This will become more and more relevant as challenges such as the COVID-19 pandemic increase the level of stress in our lives and heighten its impact on our mental health.

Author contribution
A. D.-S. designed and wrote the article.

Conflict of interest statement
Nothing declared.

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