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Unpredictable stress delays recovery from exercise-induced muscle pain: contribution of the sympathoadrenal axis

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Abstract
Introduction: Although stress is a well-established risk factor for the development of chronic musculoskeletal pain, the underlying mechanisms, specifically the contribution of neuroendocrine stress axes, remain poorly understood.
Objective: To evaluate the hypothesis that psychological stress-induced activation of the sympathoadrenal stress axis prolongs the muscle pain observed after strenuous exercise.
Methods: Adult male Sprague-Dawley rats were exposed to unpredictable sound stress and eccentric exercise. The involvement of the sympathoadrenal stress axis was evaluated by means of surgical interventions, systemic administration of epinephrine, and intrathecal β2-adrenergic receptor antisense.
Results: Although sound stress alone did not modify nociceptive threshold, it prolonged eccentric exercise-induced mechanical hyperalgesia. Adrenal medullectomy (ADMdX) attenuated, and administration of stress levels of epinephrine to ADMdX rats mimicked this effect of sound stress. Knockdown of β2-adrenergic receptors by intrathecal antisense also attenuated sound stress-induced prolongation of eccentric exercise-induced hyperalgesia.
Conclusion: Together, these results indicate that sympathoadrenal activation, by unpredictable sound stress, disrupts the capacity of nociceptors to sense recovery from eccentric exercise, leading to the prolongation of muscle hyperalgesia. This prolonged recovery from ergonomic pain is due, at least in part, to the activation of β2-adrenergic receptors on muscle nociceptors.

Keywords: Delayed-onset muscle soreness, β2-adrenergic receptor, Adrenal medulla, Nociceptor, Rat

1. Introduction
Chronic musculoskeletal pain, a leading cause of physical disability and decreased quality of life, produces huge direct and indirect economic losses.24 Although the pathophysiology of musculoskeletal pain is still poorly understood, a sizeable body of evidence indicates that psychological stress plays an important role in its induction and aggravation.9,16,17,54 For instance, epidemiological studies have shown a tight relationship between psychological stress and musculoskeletal pain in diverse populations.10,36,38,49 And, clinical studies have emphasized the role of persistent stress in the induction of chronic widespread musculoskeletal pain, identifying a key role of nociceptive afferent inputs in the maintenance of primary and secondary muscle hyperalgesia.35–58 Preclinical evidence has also shown a contribution of nociceptor sensitization to stress-induced muscle pain, revealed by increased excitability of muscle nociceptors after exposure to psychological stress.11,26 However, the mechanism by which stress induces musculoskeletal pain persistence remains to be established.

Single nucleotide polymorphisms in the β2-adrenergic receptor16,17,52 and enzymes mediating catecholamine metabolism16,17,42 determine increased susceptibility to develop persistent musculoskeletal pain after exposure to stress and physical injury. In rodent models of stress-induced muscle hyperalgesia, there is a concomitant persistent increase in plasma epinephrine.32,33 This stress-induced hyperalgesia is prevented by excision of the adrenal medulla,3,32,33 whereas systemic administration of stress levels of epinephrine in adrenal-medullectomized rats recapitulates the pain phenotype observed in rats exposed to unpredictable sound stress.32,33 Finally, systemic administration of stress levels of epinephrine, alone, aggravates muscle hyperalgesia produced in rats previously submitted to neonatal stress.3 Although these observations
support the suggestion that the sympathoadrenal axis is involved in stress-induced muscle pain, the underlying mechanism remains unknown.

We have recently demonstrated that physical resilience, the intrinsic capacity to recover from tissue injury, exhibits a cytokine signature, which is detected by muscle nociceptors. Disruption of a nociceptor’s capacity to detect this signature prevents the recovery of baseline nociceptive threshold. The fact that nociceptors also respond to the sympathoadrenal mediator epinephrine suggests that stress can also modulate the nociceptors also respond to the sympathoadrenal mediator epinephrine. Therefore, we hypothesized that stress-induced activation of sympathoadrenal stress axis disrupts the nociceptor capacity to act as sensors of physical resilience, delaying the recovery of normal muscle nociceptive threshold.

2. Materials and methods

2.1. Animals

Experiments were performed on adult male Sprague-Dawley rats (Crl:CD), weighing 250 to 400 g (approximately 8–12 weeks old), obtained from Charles River (Holister, CA). Rats were housed 3 per cage in an AAALAC International accredited animal facility, the Laboratory Animal Resource Center of the University of California, San Francisco, under a 12-hour light/dark cycle (lights on 7 AM–7 PM; room temperature 21–23˚C), with water and food available ad libitum. On completion of experiments, rats were euthanized using CO2-induced asphyxia followed by bilateral thoracotomy. Animal care and use conformed to NIH guidelines. The University of California, San Francisco Institutional Animal Care and Use Committee approved all experimental protocols. Concerted effort was made to reduce the suffering and number of animals used.

2.2. Eccentric exercise

The method used to eccentrically exercise the rat hind limb has been described previously. Briefly, rats were anesthetized with 2.5% isoflurane in oxygen, and the incision site infiltrated with 0.25% bupivacaine (Marcaine; Hospira, Lake Forest, IL). Adrenal glands were then located through bilateral incisions in the abdominal wall, their capsules incised, and the medullae excised; the fascia and skin were then sutured separately. Rats were provided with 0.45% saline to drink (ad libitum) for the first 7 days after surgery and received an additional injection of meloxicam 24 hours after surgery. To allow for maximum recovery of hypothalamic–pituitary–adrenal axis function, adrenal medullectomy was performed at least 5 weeks before additional experimental procedures.

2.3. Sound stress protocol

Exposure to sound stress occurred over 4 days, as described previously. Animals were placed 3 per cage and the cage placed 25 cm from a speaker that emitted 4 pure tones (5, 11, 15, and 19 kHz), whose amplitudes varied randomly from 20 to 110 dB sound pressure level at random times each minute, lasting 5 or 10 seconds. Sham stressed animals were placed in the sound chamber for 30 minutes, but without exposure to the sound stimulus. After sound stress or sham treatment, rats were returned to their home cages, in the animal care facility. Animals were exposed to the stressor on days 1, 3, and 4.

2.4. Adrenal medullectomy

Rats were injected with meloxicam 2 mg/kg s.c. (Metacam; Boehringer Ingelheim Vetmedica, St. Joseph, MO), anesthetized with 2.5% isoflurane in oxygen, and the incision site infiltrated with 0.25% bupivacaine (Marcaine; Hospira, Lake Forest, IL). Adrenal glands were then located through bilateral incisions in the abdominal wall, their capsules incised, and the medullae excised; the fascia and skin were then sutured separately. Rats were provided with 0.45% saline to drink (ad libitum) for the first 7 days after surgery and received an additional injection of meloxicam 24 hours after surgery. To allow for maximum recovery of hypothalamic–pituitary–adrenal axis function, adrenal medullectomy was performed at least 5 weeks before additional experimental procedures.

2.5. Chronic administration of epinephrine

Chronic administration of epinephrine was performed by osmotic minipumps (ALZET model 2004; DURECT Corporation, Cupertino, CA) implanted in the interscapular region. Rats were injected with meloxicam (2 mg/kg s.c.), anesthetized with 2.5% isoflurane in oxygen, and the incision site infiltrated with 0.25% bupivacaine. Stress levels of epinephrine were induced by filling the osmotic minipumps, so that they delivered 5.4 μg/0.25 μL/h of epinephrine. Nociceptive threshold was measured in epinephrine-implanted, adrenal-medullectomized (ADMedX) rats, 14 days after surgery.

2.6. Intrathecal antisense oligodeoxynucleotides

To evaluate the role of nociceptor β2-adrenergic receptors (β2-AR) in stress-induced prolongation eccentric exercise-induced muscle hyperalgesia, we attenuated its expression by means of intrathecal (i.t.) injections of antisense oligodeoxynucleotides (AS ODN) directed against β2-AR mRNA. The AS ODN sequence, 5′-AAA GGC AGA AGG ATG TGC-3′, was designed against a unique sequence of the rat β2-AR mRNA (GeneBank accession number NM_012492). This AS ODN sequence has been previously reported to produce specific β2-AR protein knockdown in the peripheral nervous system and behavioral changes consistent with such a decrease in protein expression in the rat. The mismatch (MM) ODN sequence, 5′-ATA GCC TGA TGG AAG TCC-3′, was designed by mismatching 6 bases (denoted by bold typeface) of the AS sequence.

Oligodeoxynucleotides were synthesized by Invitrogen (San Francisco, CA), dissolved in sterile 0.9% NaCl (B. Braun Medical, Inc, Irvine, CA) to a concentration of 10 μg/μL, and stored in 20-μL aliquots at −20˚C until use. Immediately before injection, aliquots of ODNs were further diluted in sterile 0.9% NaCl to a concentration of 2 μg/μL, and injected i.t. (40 μg/20 μL) daily for 3 consecutive days under brief anesthesia with 2.5% isoflurane as previously reported. Thereafter, rats were injected with ODN every other day, to ensure sustained knockdown of nociceptor β2-AR. Tail flick was systematically checked to ensure proper i.t. injections.
2.7. Measurement of muscle mechanical nociceptive threshold

Mechanical nociceptive threshold in the right gastrocnemius muscle was quantified using a digital force transducer (Chatillon DF2; Amtek, Inc, Largo, FL) with a 7 mm-diameter probe.2-6 Rats were lightly restrained in cylindrical acrylic holders with lateral slats that allow for easy access to the hind limb for application of the force transducer probe to the belly of the gastrocnemius muscle. The nociceptive threshold was defined as the force, in millinewtons, required to produce a flexion reflex in the hind limb. The results are expressed as percentage change from baseline mechanical nociceptive threshold. All behavioral testing was conducted between 10 AM and 4 PM. Behavioral data collection was performed unblinded to treatment condition.

2.8. Statistics

Group data are expressed as mean ± SEM of n independent observations. Statistical comparisons were made using two-way repeated-measures analysis of variance followed by Bonferroni multiple-comparisons post hoc test. Statistical significance was set at P < 0.05. Statistical software Prism 8.0 (GraphPad Software, Inc, La Jolla, CA) was used to perform data analysis and graph plotting.

3. Results

3.1. Sound stress prolongs eccentric exercise-induced muscle hyperalgesia

Two protocols of unpredictable sound stress followed by eccentric exercise were studied (Fig. 1A). As previously reported,2,4 naive (control) rats submitted to eccentric exercise exhibited a significant decrease in nociceptive threshold (i.e., mechanical hyperalgesia) in the gastrocnemius muscle. Hyperalgesia reached a peak 48 hours after eccentric exercise (n = 6/group, P < 0.05; Fig. 1B) and fully recovered by day 5 (n = 6/group, P < 0.05; Fig. 1B). By contrast, rats submitted to unpredictable sound stress either 1 or 14 days before displayed muscle hyperalgesia that persisted for up to 20 days (n = 6/group, P < 0.05; Fig. 1B). Exposure to unpredictable sound stress 14 days before eccentric exercise produced a small, albeit significant, increase in peak hyperalgesia compared with naive rats (n = 6/group, P < 0.05; Fig. 1B). Given that rats submitted to sound stress with the last exposure either 1 or 14 days before eccentric exercise displayed similar duration of muscle hyperalgesia (Fig. 1B), we performed subsequent experiments using the eccentric exercise one day after sound stress.

3.2. Adrenal medullectomy attenuates stress-induced prolongation of exercise-induced muscle hyperalgesia

To explore the contribution of the sympathoadrenal neuroendocrine stress axis to prolongation of eccentric exercise-induced muscle hyperalgesia produced by sound stress, the effect of ADMdX or a sham procedure was studied (Fig. 2A, B). ADMdX alone did not produce a change in nociceptive threshold, compared with control (sham) rats, at baseline (n = 6/group, P > 0.05; Fig. 2B), or 1 day after the last sound stress (n = 6/group, P > 0.05; Fig. 2B). Importantly, ADMdX alone (no sound stress) had no effect on eccentric exercise-induced muscle hyperalgesia (n = 6/group, P > 0.05; Fig. 2B). The ADMdX plus sound stress rats, however, experienced significantly less muscle hyperalgesia compared with sham controls, on days 1 to 20, after eccentric exercise (n = 6/group, P < 0.05; Fig. 2B).

3.3. Sustained administration of stress levels of epinephrine prolongs eccentric exercise-induced muscle hyperalgesia

To determine whether epinephrine contributes to the prolongation of eccentric exercise-induced muscle hyperalgesia produced by sound stress, we studied its sustained administration in ADMdX rats (Fig. 3A). Administration of saline or epinephrine at stress levels to ADMdX rats, alone or in combination with sound stress, did not modify the nociceptive threshold compared with baseline (n = 6/group, P > 0.05; Fig. 3B). However, compared with saline, epinephrine treatment prolonged eccentric exercise-induced muscle hyperalgesia, producing a significant decrease in nociceptive threshold from day 3 to day 20 after eccentric exercise (n = 6/group, P < 0.05; Fig. 3B). There were no significant differences in the mechanical hyperalgesia of rats receiving epinephrine at stress levels and rats receiving epinephrine plus sound stress (n = 6/group, P > 0.05; Fig. 3B).
to be elucidated. Furthermore, it is unclear how exposure to stressful psychological stimuli leads to muscle nociceptor mechanical sensitization.11,26 Therefore, the finding that the sympatheticadrenal mediator epinephrine, acting at 2 adrenergic receptors on nociceptors, plays a central role in the prolongation of exercise-induced muscle pain produced by unpredictable stress provides insight into how stress can aggravate musculoskeletal pain.

### 4. Discussion

Although it is well established that psychological stress plays an important role in the induction and aggravation of chronic musculoskeletal pain,41,54 the underlying mechanism remains

#### 3.4. Knockdown of β2-AR attenuates sound stress-induced prolongation of exercise-induced muscle hyperalgesia

The role of nociceptor β2-AR in the prolongation of eccentric exercise-induced muscle hyperalgesia by preexposure to sound stress was evaluated by means of intrathecal AS ODN for β2-AR mRNA. The ODN treatment did not produce significant changes in nociceptive threshold, when measured in the ipsilateral gastrocnemius muscle (time point 1), rats were submitted to unpredictable sound stress on days 1, 3, and 4. One day after the last sound stress session, they were submitted to eccentric exercise and the nociceptive threshold measured up to day 20 after eccentric exercise. Of note, nociceptive responses of ADMdX rats are similar to those displayed by naive (control) rats

### Figure 3. Administration of stress levels of epinephrine prolongs eccentric exercise-induced muscle hyperalgesia.

(A) Experimental protocol used to study muscle nociceptive threshold after the administration of stress levels of epinephrine or saline, alone or in combination with unpredictable sound stress (SS), in ADMdX rats. Two weeks after osmotic pumps containing epinephrine or saline were implanted, ADMdX rats were submitted to sound stress (SS + Epinephrine; SS + Saline) or sham procedure (Epinephrine) on days 1, 3, and 4. One day after the last sound stress session, they were submitted to eccentric exercise and the nociceptive threshold measured in the ipsilateral gastrocnemius muscle up to day 20 after eccentric exercise. (B) Two-way ANOVA showed significant effects for time (F 8,120 = 394.7, P < 0.001), treatment (F 2,15 = 34.11, P < 0.001), or treatment by time interaction (F 16,120 = 10.30, P = 0.7497). Post hoc analysis revealed significant differences between control rats receiving saline and rats receiving epinephrine alone (Epinephrine) from day 3 to day 20 after eccentric exercise (P < 0.001) and between control rats (SS + Saline) and rats receiving epinephrine plus sound stress (SS + Epinephrine) from day 3 to day 15 after eccentric exercise (P < 0.001). No significant differences between rats receiving epinephrine alone (Epinephrine) and rats receiving epinephrine plus sound stress (SS + Epinephrine) were observed at any time point after eccentric exercise (P > 0.05). **P < 0.01; ***P < 0.001 (SS + Saline vs Epinephrine); #P < 0.05; ##P < 0.01, ###P < 0.001 (Sham + SS vs ADMdX + SS). ANOVA, analysis of variance.
rather limited to its recovery phase. Indeed, the main difference in muscle pain between rats submitted to sound stress and control rats started after the peak of hyperalgesia, and persisted for over 2 weeks after eccentric exercise, indicating that sound stress selectively affects the capacity to recover from the hyperalgesia induced by eccentric exercise, i.e., stress affects physical resilience. Of note, sound stress does not, by itself, affect nociceptive threshold in the gastrocnemius muscle. Of note, sound stress does not, by itself, affect nociceptive threshold in the gastrocnemius muscle. Of note, sound stress does not, by itself, affect nociceptive threshold in the gastrocnemius muscle. Of note, sound stress does not, by itself, affect nociceptive threshold in the gastrocnemius muscle. Of note, sound stress does not, by itself, affect nociceptive threshold in the gastrocnemius muscle. Of note, sound stress does not, by itself, affect nociceptive threshold in the gastrocnemius muscle. Of note, sound stress does not, by itself, affect nociceptive threshold in the gastrocnemius muscle.

4.2. Stress level of epinephrine is necessary and sufficient to prolong muscle hyperalgesia after eccentric exercise

Adrenal medullectomy attenuated the sound stress-induced prolongation of muscle hyperalgesia, indicating that sympathoadrenal response is necessary for stress-induced impairment of recovery from eccentric exercise-induced hyperalgesia. And, the administration of epinephrine at stress levels mimicked the effect of sound stress, confirming that adrenal medulla-derived epinephrine is sufficient to produce the effect of sound stress. Indeed, systemic epinephrine induced 15% to 25% decrease in mechanical nociceptive threshold between days 5 and 20 after eccentric exercise, whereas complete return to baseline nociceptive threshold occurs at day 5 in normal control rats.

Our observations are in line with previous finding that sound stress produces persistently increased levels of sympathoadrenal catecholamines, including epinephrine. Of note, alone the excision of the adrenal medulla does not modify muscle, or cutaneous mechanical nociceptive threshold or, as shown by the present results, the response to eccentric exercise. These observations indicate that the effect of adrenal medullectomy is not due to a change in baseline nociceptive threshold but due to its enhancement by sound stress. Although we did not explore the source of the stress-induced prolongation of hyperalgesia that is resistant to adrenal medullectomy, the contribution of extra-adrenal sources of catecholamines might play a role. Indeed, sympathetic postganglionic neurons contribute to sound-stress-induced hyperalgesia, as revealed by its attenuation after chemical or surgical sympatheticctomy, or systemic administration of an α - 1 adrenergic receptor antagonist.

Besides its direct actions on nociceptors, catecholamines enhance the release of a number of proalgesic cytokines from mast cells, lymphocytes, cell types that play a central role in the induction and recovery of skeletal muscle injury. Indeed, a transient rise in catecholamine is observed after strenuous exercise, believed to act on the β2-AR expressed by skeletal muscle fibers, contributing to tissue regeneration. Furthermore, ex vivo exposure of skeletal muscle tissue to epinephrine induces the release interleukin-6, a well-established proalgesic mediator, which contributes to eccentric exercise-induced muscle pain. Thus, although our results suggest that nociceptors are prominent targets of sympathoadrenal catecholamines (see below), we cannot rule out the contribution of other players in the prolongation of muscle pain induced by sound stress.

4.3. Nociceptor β2-AR receptors are necessary for prolonged muscle hyperalgesia after eccentric exercise

Consistent with a role of nociceptor β2-AR, its knockdown by intrathecal antisense attenuated the prolonged muscle hyperalgesia after eccentric exercise. We have previously shown that this antisense sequence, administered intrathecal, produced a marked decrease in β2-AR in peripheral nerves and antinociceptive effects in stress-dependent pain. Because only sensory neurons were exposed to both antisense treatment and circulating catecholamines, nociceptor β2-ARs likely play a central role in the prolongation of muscle pain induced by sound stress. Sensory neurons express β2-AR and epinephrine induces nociceptor excitation and sensitization. The involvement of nociceptor β2-AR in visceral mechanical hyperalgesia after exposure to stressful stimuli has also been
reported. These observations, and the fact that catecholamines do not cross the blood–brain barrier, further support the fact that the proalgesic effect of epinephrine observed here depends on peripheral targets such as nociceptors.

The finding that β2-AR antisense did not completely abolish the prolongation of exercise-induced hyperalgesia may be due to a number of factors. For instance, it has been observed in many physiological settings that glucocorticoids enhance the β2-AR expression. Indeed, in vitro exposure of colonic nociceptors to epinephrine and corticosterone produced upregulation of β2-AR, concomitant with enhanced nociceptor excitability. Given that the pronociceptive effect of exposure to unpredictable sound stress also involves the activation of the hypothalamic–pituitary–adrenal axis and the release of corticosterone, enhanced nociceptor expression of β2-AR in this model is possible, which could have limited the efficacy of antisense. Finally, the involvement of peripheral β1- and β2-AR has also been reported in preclinical studies where pain is associated with increased levels of catecholamines.

4.4. Nonnociceptor mechanisms in the prolongation of exercise-induced muscle pain by sound stress

Although our findings point to a contribution of nociceptors in the prolongation of eccentric exercise-induced pain by sound stress, additional mechanisms may also be involved. For instance, a contribution of dysfunctional descending pain modulatory controls cannot be excluded. Indeed, we and others have demonstrated that exposure to repeated stress markedly attenuates the antinoceptive capacity of endogenous inhibitory pain circuits. Furthermore, repeated stress activates GABAergic neurons of the rostral ventromedial medulla that inhibit opioidergic/GABAergic neurons of the dorsal horn, acting as descending pain facilitatory (ie, pronociceptive) controls, selectively enhancing mechanical nociception. Moreover, clinical studies have shown attenuated diffuse noxious inhibitory controls in fibromyalgia patients who experience chronic widespread pain. And, peripheral sensitization induced by muscle contraction interacts with abnormal descending pain modulation leading to pain facilitation in fibromyalgia patients. Together, these observations support the suggestion that repeated stress also contributes to the prolongation of hyperalgesia induced by eccentric exercise, by disrupting normal function of supraspinal modulatory pain controls.

4.5. Stress disrupts the capacity of nociceptors to sense physical resilience status

Nociceptors are sensitized by signals generated in damaged tissue, observed as hyperalgesia, providing a motivational drive for adaptive/protective behaviors. We have demonstrated active suppression of the hyperalgesia induced by eccentric exercise, well before completion of structural recovery, allowing for resumption of physical activity. Thus, muscle nociceptors are posited to act as sensors of functional recovery after strenuous exercise. Indeed, a sizeable amount of evidence shows that nociceptors are sensitized by psychological stress. Importantly, nociceptors express a number of receptors that allow for detection of neuroendocrine stress axes signals, including by glucocorticoid and β2-adrenergic receptors. Also, as mentioned above, catecholamines can act on a number of cell types involved in skeletal muscle damage and repair, modulating their capacity to release cytokines that could modify nociceptor excitability. Thus, consistent with our previous observations, we postulate that persistent sympathoadrenal axis activation, by unpredictable sound stress, induces modifications in nociceptor excitability, and in the cytokine signature of the damaged tissue, likely disrupting the role of nociceptors as sensors of the status of functional recovery, potentially contributing to pain chronification.

4.6. Clinical considerations

Although pharmacological treatments provide only modest relief for chronic pain, including chronic musculoskeletal pain, therapeutic exercise has shown some promise. Aggravation of ongoing muscle pain by exercise can, however, limit its clinical usefulness. Our findings indicate that stress is likely detrimental to the success of exercise-based therapy. Thus, cognitive and behavioral treatments might be beneficial to minimize the pain induced by therapeutic exercise, underscoring the well-established importance of a multidisciplinary approach for the treatment of chronic musculoskeletal pain.

Disclosures

The authors have no conflict of interest to declare.

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The authors thank Dr Oliver Bogen for technical help. Author contributions: P. Alvarez designed and performed experiments, analyzed data, and wrote the manuscript; P.G. Green performed experiments and analyzed data; J.D. Levine analyzed experiments and supervised the study. All authors read and approved the manuscript.

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