Increased Affective Bias Revealed Using Experimental Graded Heat Stimuli in Young Depressed Adults: Evidence of “Emotional Allodynia”

IRINA A. STRIGO, PhD, ALAN N. SIMMONS, PhD, SCOTT C. MATTHEWS, MD, ARTHUR D. (BUD) CRAIG, PhD, AND MARTIN P. PAULUS, MD

Objective: To examine the hypothesis that young adults with major depressive disorder (MDD) would show increased affective bias to painful and nonpainful experimental heat stimuli, as evidenced by an increased responsiveness to warm and hot temperatures. Pain and depression often occur together. Pain is both a sensation and an affective experience. Similarly, depression is associated frequently with somatic symptoms as well as emotional dysphoria. Existing evidence indicates that MDD may be associated with altered pain processing. However, the extent to which alterations in experimentally controlled heat pain sensations are related to increased affective bias in MDD is unknown. Method: Graded nonnoxious and noxious heat stimuli were delivered randomly with a thermode applied to the volar surface of the left arm of 15 unmedicated subjects with current MDD and 15 age- and gender-matched healthy comparison subjects. MDD and non-MDD subjects rated the intensity and unpleasantness of all stimuli. Results: Two main results were observed. First, MDD relative to non-MDD subjects showed decreased heat pain thresholds. Second, a significantly increased affective bias (unpleasantness/intensity) was observed in subjects with MDD, particularly over the range of nonnoxious heat stimuli. This bias was independent of the change in sensory pain thresholds. Conclusion: These findings represent corroborative evidence of abnormal affective heat pain processing in young adults with MDD, and suggest that MDD is associated with “emotional alldony,” a qualitatively altered negative emotional response to normally nonaversive thermal stimuli. Key words: psychophysics, allodynia, MDD, thermode, heat, nociception.

MDD = major depressive disorder.

INTRODUCTION

Pain and depression are common and highly comorbid. Over 75% of patients with a clinical depressive disorder experience chronic or recurring pain, such as headache, stomach pain, neck and back pain, or nonspecific generalized pain (1). Conversely, 30% to 60% of pain patients (i.e., chronic pain, migraine) report significant depressive symptoms (2). The comorbidity between pain and depression contributes significantly to poorer outcomes and increased cost of treatment (3). The high comorbidity between these conditions potentially suggests a common underlying pathophysiology. Specifically, pain and depression may be related clinical manifestations of altered activity within a common neural network. However, the neurobiological basis of pain processing in major depressive disorder (MDD) is incompletely understood.

Pain is a subjective experience, which can be assessed along two dimensions, i.e., sensory pain intensity, which describes the discrimination of the stimulus intensity, and affective pain unpleasantness, which describes the emotional impact of the stimulus (4,5). During brief experimental noxious stimuli, these dimensions are highly correlated (6–8), and both of these dimensions are greatly influenced by the cognitive, emotional, and physiological state of the individual (9–11). For example, pleasant emotions and cognitions (i.e., expectation of pain relief) can attenuate pain perception, and negative emotions or cognitions, such as catastrophizing (i.e., thinking that pain will never end), can amplify perceived pain intensity and unpleasantness (9). Therefore, increased attention to one’s negative feeling state, which is often observed in depressed patients (12), may fundamentally alter the pain experience and underlying brain circuitry in MDD. This conceptualization is consistent with recent evidence of increased perception and neuronal activation in the brain areas encoding the affective, but not the sensory-discriminative aspect of pain processing in subjects with depressed mood (13,14) and in patients with comorbid chronic pain and depression (15). Heightened emotional reactivity and emotional biasing of daily self-reports of subjective pain experience are consistently observed in chronic pain patients, especially in those with a history of depression and increased daily depressive moods (16–18).

Several prior studies examined sensitivity to experimental pain in depressed versus nondepressed patients, with discordant results. Decreased pain sensitivity in depressed individuals was observed in several studies, as evidenced by increased pain thresholds (19–24). Other studies reported increased pain sensitivity in depression, as evidenced by decreased pain thresholds (25) and decreased pain tolerance (26–29). Finally, one study reported no difference in pain sensitivity in mild depression (30). Several methodological differences could be responsible for these discrepancies. First, some studies that found decreased or no change in pain perception in depression did not control for the confounding effects of antidepressant use (20,27,28) and/or rule out comorbid clinical pain complaints (19,24), which are both known to interfere with pain sensitivity (31,32). Second, other studies looked at a nonhomogeneous patient sample including patients with a range of psychiatric disorders as a primary diagnosis (e.g., bipolar, psychosis) that are known to influence differently the perception of pain (33). Third, the majority of earlier studies on experimental pain did not examine separately the sensory and affective dimensions of the experimental pain experience,
MDD = major depressive disorder; BDI-II = Beck Depression Inventory, Second Edition; DSM-IV = Diagnostic and Statistical Manual of Mental Disorders, 4th Edition; PTSD = posttraumatic stress disorder; GAD = general anxiety disorder; PD = panic disorder.

TABLE 1. Patient Characteristics

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MDD = major depressive disorder; BDI-II = Beck Depression Inventory, Second Edition; DSM-IV = Diagnostic and Statistical Manual of Mental Disorders, 4th Edition; PTSD = posttraumatic stress disorder; GAD = general anxiety disorder; PD = panic disorder.

“Lifetime, not current diagnosis.

which is crucial for understanding pain processing in depression (13–15). In the present study, we controlled for all of the above discrepancies to identify how MDD affects the sensory and affective perception of experimental heat pain.

The aim of the current experiment was to examine whether unmedicated young adults with MDD, but without comorbid pain conditions or other medical problems, would show altered sensitivity to brief heat stimuli. We hypothesized that MDD relative to non-MDD subjects would demonstrate a bias toward the affective component of brief heat stimuli. Support for this hypothesis would contribute to a more comprehensive understanding of the pathophysiology of pain and depression as well as suggest a possible mechanism that could explain the emotional bias of daily self-reports of pain in chronic pain patients.

MATERIALS AND METHODS

Subjects

Fifteen unmedicated (>30 days) subjects with current MDD (three males, mean age 24.5 ± 5.5 years) gave written informed consent to participate in this study, which was approved by the University of California, San Diego Human Protection Program (Institutional Review Board protocol #050511). The data were collected from February 2004 to August 2005. Subjects completed the structured clinical interview for Diagnostic and Statistical Manual of Mental Disorders, 4th Edition (DSM-IV) (34) and met the DSM-IV criteria for current MDD (Table 1), based on the diagnosis of a board-certified psychiatrist (S.C.M.). Depressive symptoms severity was measured with the Beck Depression Inventory, Second Edition (BDI-II) (35). The method of constant stimuli was used to measure subjects’ sensitivity to heat. Heat stimulation started from a baseline of 32°C and rose linearly at a rate of 1.5°C to one of five predetermined temperatures (40°, 42°, 44°, 46°, 48°C). The duration of each stimulus was 5 seconds, excluding the rise/fall time. Each temperature was presented twice, and the order of presentation was randomized.

Education 15.7 1.7 15.4 2.0 0.4
BDI-II 27.8 7.6 1.3 1.7 13.1
MFSI-SF 43.5 15.1 8.1 8.6 11.3
Neuroticism 70.5 8.3 41.4 9.8 8.4
Heat pain threshold 41.7 3.3 34.9 1.2 2.4

MDD = major depressive disorder; M = males; F = females; BDI-II = Beck Depression Inventory, Second Edition; MFSI-SF = Multidimensional Fatigue Symptoms Inventory, Short Form.

“p < .01; b p < .05; c NS.

TABLE 2. Demographics and Behavioral Measures

<table>
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<th></th>
<th>MDD (3 M, 12 F)</th>
<th>Comparison Group (3 M, 12 F)</th>
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<td>MFSI-SF</td>
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<td>8.1 8.6</td>
<td>11.3</td>
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<td>Neuroticism</td>
<td>70.5 8.3</td>
<td>41.4 9.8</td>
<td>8.4</td>
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<td>Heat pain threshold</td>
<td>41.7 3.3</td>
<td>34.9 1.2</td>
<td>2.4</td>
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</table>

MDD = major depressive disorder; BDI-II = Beck Depression Inventory, Second Edition; MFSI-SF = Multidimensional Fatigue Symptoms Inventory, Short Form.

“p < .01; b p < .05; c NS.

**Temperature Sensitivity**

The method of constant stimuli was used to measure subjects’ sensitivity to heat. Heat stimulation started from a baseline of 32°C and rose linearly at a rate of 1.5°C to one of five predetermined temperatures (40°, 42°, 44°, 46°, 48°C). The duration of each stimulus was 5 seconds, excluding the rise/fall time. Each temperature was presented twice, and the order of presentation was randomized and unknown to the subjects and the experimenter. A 9-cm² thermode (Medoc TSA-II, Ramat-Yishai, Israel) was applied to each subject’s alcohol or substance dependence; b) met DSM-IV criteria for alcohol or substance abuse within the past 30 days; c) were experiencing active suicidal ideation; d) had a history of bipolar or psychotic disorder; e) had an active medical problem; f) had a history of chronic pain. In addition to the BDI-II, all subjects completed a) the Multidimensional Fatigue Symptom Inventory, Short Form (MFSI-SF) (a self-report 30-item questionnaire which assesses general, physical, emotional, and mental manifestations of fatigue as well as vigor—an estimate of energy level) (36); and b) the neuroticism module of the NEO Five-Factor Inventory (NEO-FFI) (a self-report 60-item version of NEO personality inventory that assesses five domains of adult personality, including neuroticism, extraversion, openness to experience, agreeableness, and conscientiousness) (37).

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vo lar forearm, and the site of stimulation on the forearm skin was varied slightly to avoid sensitization. The skin under the thermode was adapted to the baseline thermode temperature before the start of stimulation. The interval between successive stimuli was at least 30 seconds, and the minimum interval between stimulation of the same skin site was at least 1 minute. Subjects were asked to rate the intensity and unpleasantness of each stimulus, using three validated visual scales (6). After each temperature stimulus, subjects were asked to rate the maximum sensation of pain, using a scale that ranged from 0 (“no pain sensation”) to 10 (“extremely intense pain sensation”). If the stimulus produced no painful sensation, subjects were asked to rate how far from painful the sensation was using a scale that ranged from 0 (“extremely warm”—the pain threshold) to −10 (“no sensation”). Furthermore, subjects rated the maximum unpleasantness evoked by each temperature stimulus, even if it was perceived as nonpainful, using a scale that ranged from 0 (“not at all unpleasant”) to 10 (“extremely unpleasant”). Each subject’s pain threshold was estimated by fitting linear functions to the nonpainful/painful intensity response curves (7,38). To specifically evaluate affective bias, all unpleasantness ratings were weighted by the heat intensity ratings at that temperature (affective bias = unpleasantness rating/intensity rating). This transformation was used to control for possible group differences in subjects’ response bias (39). Intensity ratings were rescaled from −10 to 10 to 0 to 10 before affective bias was calculated.

Statistical Analyses

We used Pearson correlations to explore how the affective bias measure related to other common psychophysiological variables. Specifically, we tested the relationship between affective bias (averaged across all stimulus temperatures), BDI, MFSI-SF, and the neuroticism score of the NEO-FFI (significance level was adjusted based on Bonferroni correction for multiple comparison). These measures were selected because MDD has been associated with high neuroticism and fatigue (40,41). To specifically test the hypothesis that MDD relative to non-MDD subjects show bias toward the affective component of brief heat stimuli, we ran repeated-measures analyses of variance (ANOVA) on the affective bias ratings (group as between-subject factor; temperature as within-subject factor). To determine the individual contrasts that powered the group by temperature interaction effects, we did exploratory post hoc t tests at each temperature (40°, 42°, 44°, 46°, and 48°C). The slopes and intercepts of the individual affective bias curves were determined by linear regression and the coefficients were compared between groups after correcting for multiple comparisons. Differences in slopes reflected between-group differences in discrimination, whereas differences in the intercept of the affective bias curve reflected between-group differences in the initial reactivity to temperature stimuli. The relationship between affective bias (averaged across all stimulus temperatures) and sensory pain thresholds was tested, using Pearson correlations. The between-group difference in the strength of these correlations was tested by contrasting the Fisher Z-transform of the correlation values in each group. In addition, we ran two exploratory follow-up repeated-measures ANOVAs for the intensity and unpleasantness ratings separately to provide further details about how depression affects these variables. Finally, to explore which behavioral measure, i.e., intensity, unpleasantness, or affective bias, explained the greatest degree of variance between the groups at each temperature level (40°, 42°, 44°, 46°, and 48°C), we performed five exploratory stepwise linear regressions where all three measures were entered simultaneously into the model. All statistical analyses were done with SPSS 12.0 (SPSS Inc., Chicago, Illinois).

RESULTS

Demographics and Psychophysiological Variables

Subjects with MDD exhibited significantly higher levels of a) depression, as measured by the BDI-II; b) fatigue, as measured by the MFSI-SF; and c) neuroticism, as measured by the NEO-FFI (Table 2). In addition, sensory pain thresholds were significantly lower in the MDD relative to the non-MDD subjects (t = 2.4; p < .05) (Table 2).

Best Measure to Differentiate the Groups

The results of exploratory stepwise linear regression analysis with group as a dependent variable and three predicting factors (subjective stimulus intensity, unpleasantness, and affective bias ratings, i.e., unpleasantness rating/intensity rating) are shown in Table 3 for each stimulus temperature. Group membership was significantly predicted by the affective bias measure at all temperatures tested, except for 48°C. At all temperature stimuli, except for 48°C, the affective bias measure explained most of the between-group variance; adding unpleasantness and intensity ratings did not improve the model. It is important to note that although intensity and unpleasantness ratings were used to calculate the affective bias ratio, it was the ratio that best explained the variance between MDD and non-MDD patients whereas the linear combination of the intensity and unpleasantness ratings did not. At 48°C, all factors were needed to predict the group differences.

Table 3

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>AB</th>
<th>AB and Unpl</th>
<th>AB and Unpl and Int</th>
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<tbody>
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<td>.431*</td>
<td>.452</td>
<td>.535</td>
</tr>
<tr>
<td>42</td>
<td>.536*</td>
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<td>48</td>
<td>.296</td>
<td>.383</td>
<td>.522</td>
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</table>

* R-coefficients are shown; *p < .01; *p < .05.
Stepwise linear regression model with group as dependent variable and three predicting factors: affective bias (AB), unpleasantness (Unpl), and intensity (Int).

Table 4

<table>
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<tr>
<th>Measure</th>
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<td>Neuroticism</td>
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<td>.457*</td>
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</table>

* Correlation coefficients after covarying for sensory pain thresholds; *p < .01; *p < .05 (Bonferroni correction for multiple comparisons); * from NEO-FFI (NEO Five-Factor Personality Inventory); * missing data on 2/30 subjects.
AB = affective bias; BDI-II = Beck Depression Inventory, Second Edition; MFSI-SF = Multidimensional Fatigue Symptom Inventory, Short Form.

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controlling for sensory pain thresholds in the combined group (r = >.4; p < .05) (Table 4).

**Increased Pain Affect in MDD**

Figure 1A shows ratings of the perceived intensity and unpleasantness to brief heat stimuli in MDD and non-MDD subjects. Repeated-measures ANOVAs revealed significant main effects of both group (p < .01) and temperature (p < .01) on the perceived intensity (p < .01) and unpleasantness (p < .01) rating to brief temperature stimuli. Additionally, a significant group by temperature interaction on the perceived unpleasantness rating was observed (p < .01). Post hoc analyses showed that subjects with MDD rated the heat stimuli as more intense at 42°C (p < .05), 44°C (p < .01), and 46°C (p < .01), but not at 40°C (p = .06) or 48°C (p = .2) (Figure 1A, left). Similarly, subjects with MDD rated the stimuli as more unpleasant at 40°C (p < .05), 42°C (p < .01), 44°C (p < .01), and 46°C (p < .01), but not at 48°C (p = .07) (Figure 1A, right). Figure 1B shows ratings of the affective bias to brief heat stimuli. Repeated-measures ANOVAs showed significant main effects of group (p < .01) and temperature (p < .01) and a significant group × temperature interaction (p < .01) on the affective bias rating. For both groups, as intensity ratings increased, unpleasantness ratings also increased. Post hoc analyses showed a greater affective bias rating in the MDD relative to the non-MDD group at 40°C (p = .05), 42°C (p < .01), 44°C (p < .01), and 46°C (p < .01), but not at 48°C (p = .1)—the temperature at which the affective bias ratings did not differ between the groups (Figure 1B). There were no significant differences in the slopes of the stimulus-response functions related to the different temperature stimuli (t = −1.0; p = .3), suggesting that there were no differences in scaling of heat stimuli. However, a significant difference between the intercepts of the stimulus-response functions was observed between the groups (MDD = 41.7 ± 3.3; CON = 43.9 ± 1.1, p < .05, t (28) = 2.4).

![Figure 1. Psychometric functions. A. Intensity (left) and unpleasantness (right) ratings to temperature stimuli in major depressive disorder (MDD) (●) and comparison (CON) (□) subjects. Repeated-measures analysis of variance (ANOVA) showed significant effect of group and temperature for both measures, and significant group × temperature interaction for the unpleasantness measure (p < .01). B. Affective bias ratings during temperature stimuli in MDD (●) and comparison (CON) (□) subjects. Affective bias to heat stimulation was calculated by dividing the individual subject’s unpleasantness scores by the corresponding intensity scores. Repeated-measures ANOVA showed significant main effect of group, temperature, and significant group × temperature interaction (p < .01). Arrows indicate heat pain intensity thresholds for MDD (solid) and CON (broken) group (MDD = 41.7 ± 3.3; CON = 43.9 ± 1.1, p < .05, t (28) = 2.4).](image-url)
AFFECTIVE BIAS DURING HEAT IN MDD

Divergence of Subjective Pain Experiences in MDD

Heat pain thresholds were slightly lower in the MDD compared with the non-MDD subjects ($t = 2.4; p < .05$) (Table 2). We investigated whether increased affective bias in MDD was predicted by the decreased sensory pain by examining the correlation between the two measures. In healthy comparison subjects, the average affective bias showed an almost perfect negative correlation with sensory pain thresholds ($r = -9; p < .01; t = -6.5$), i.e., subjects with higher affective bias scores experienced pain at lower stimulus temperatures. In comparison, there was no significant correlation between the mean affective bias and the sensory pain thresholds in the MDD group ($r = -3; p = .2; t = -1.2$). Furthermore, there was a significant difference between the groups in the strengths of correlation coefficients ($p < .01$; Fisher Z-transform). In summary, whereas pain thresholds were related to the degree of affective bias in healthy comparison subjects, no such relationship was observed in individuals with MDD. This supports the notion that physiological thresholds are at least partially disconnected from the psychological assessment in these subjects.

DISCUSSION

This investigation tested the hypothesis that young adults with MDD (without comorbid pain conditions) have altered affective processing that can be revealed by their responses to experimental nonnoxious and noxious heat stimuli. Our results indicate that individuals with MDD, when compared with age- and gender-matched healthy comparison subjects, perceive heat stimuli as both more intense and more unpleasant, suggesting that MDD is associated with hypersensitivity in both the sensory and the affective dimensions of heat pain. Interestingly, after controlling for the sensory dimension, individuals with MDD still displayed an increased bias in the affective dimension of heat stimuli that was particularly apparent in the nonnoxious heat range. Furthermore, in our data, the decreased sensory thresholds to pain did not account for the increased affective bias in these patients. These results are in line with previous reports that associated depressive symptoms with alterations in the affective aspect of self-reported pain experience (14,16,25,42), and they suggest independent neural pain-processing networks of the affective and sensory pain dimensions in MDD (15).

Our results indicate that the increased affective bias revealed by heat stimuli (i.e., the affective-motivational divided by the sensory processing of pain) in MDD is not due to between-group differences in the ability to distinguish heat stimuli because there were no differences between the slopes of the stimulus-response functions. Instead, our results suggest that this increased affective bias in MDD is due to a leftward shift of this function, which suggests that MDD is associated with earlier engagement of the affective-motivational processing system. The affective bias for the subjects with MDD at 40°C (a heat stimulus that is normally well below pain threshold) is as large as that at 46°C (a stimulus that is clearly suprathreshold for pain). Both groups showed similar affective biases at the highest temperature. The latter observation suggests that both MDD and non-MDD subjects showed a strong affective bias when there is a real notion that potential damage to the body is taking place (43). However, in individuals with MDD, this motivational component seems to be already active at stimulation intensities well below normal pain threshold, and strikingly, even below the heat intensity that they report as distinctly painful.

The fact that subjects with MDD showed increased affective bias in the range of nonpainful stimuli is reminiscent of the concept of sensory allodynia, which refers to the condition in which ordinarily nonpainful stimuli evoke the sensation of pain. Allodynia in pain sensation is commonly observed after acute (e.g., sunburn) or chronic (e.g., fibromyalgia) tissue injury. People with such injuries are hypersensitive to light touch, warmth, cold, or light pressure stimuli (44). Because the difference in affective bias that we observed between MDD and comparison subjects is strongest in the nonnoxious range, we would like to suggest that the concept of allodynia can be extended to the affective realm; thus, “emotional allodynia” is the abnormal elicitation by subthreshold stimuli of the affective-motivational component associated with the perception of pain. Emotional allodynia in depressed individuals seems to be due to increased baseline negative affect (45,46) or higher reactivity to potentially aversive stimuli (47,48). Like sensory allodynia, emotional allodynia may be a consequence of psychological injury or the presence of a psychiatric condition. This notion is useful because it suggests particular processing abnormalities and specific differences in neural activation patterns that one may expect to find in individuals with MDD. Moreover, it provides a conceptual bridge between disorders that have been described within the pain field and those within psychiatry.

Prior studies that examined experimental pain perception in MDD reported conflicting results (19–28,30–33). For example, based on findings of higher heat pain and electrical pain thresholds, it was suggested that depressed individuals do not feel pain, and that they experienced what was described as “perceptual unresponsiveness” and “affective indifference” (23). Others, however, found no differences in electrical pain tolerance or found decreased cold-pressor tolerance in depression (26,27,33). Recent studies suggested that depressed individuals might have different reactions to “superficial” (i.e., cutaneous) versus “deep” (i.e., muscle, joint, viscera) pain (28,29,49), with decreased sensitivity to the former and increased sensitivity to the latter. That finding is particularly interesting because one of the main psychophysical differences between “deep” and “superficial” pain is the greater unpleasantness of the former (6,7).

Neurophysiologically, it is the different amounts of signaling via unmyelinated small diameter fibers (C-fibers) that may distinguish different sensations arising from “deep” or “superficial” tissues. Both visceral afferents (e.g., stomach ache) and muscle afferents (e.g., ischemic pain) have a high proportion of C-fibers (50,51), which convey a strong affective component of pain in these areas. Consistent with this idea, investi-
gators concluded from a recent study of ischemic pain that sensitivity to the afferent activation of C-fibers is especially enhanced during depression (49). The small-diameter sensory C-fibers can be viewed as homeostatic, based on the idea that these afferents constantly relay information about the tissue status (52). Thus, one might speculatively associate depression with a homeostatic dysfunction (53) characterized by abnormal cortical interpretation of the homeostatic afferent C-fiber activity, resulting in a heightened baseline affective bias, or emotional allodynia. Deficits in nocturnal thermoregulation have been observed in MDD, which may underlie abnormal sleep patterns in this disorder (54). Furthermore, C-fibers play a critical role in central sensitization, which is one of the mechanisms responsible for sensory allodynia (55).

Our results showing increased affective bias to experimental heat stimuli in young adults with current MDD and no comorbid chronic pain conditions have direct clinical relevance. Individuals with chronic pain are more likely to emotionally bias their daily self-reports of pain if they have a history of clinical depression (16–18). Although our findings cannot determine whether affective biasing is a predisposition to develop chronic pain, longitudinal studies on depressed patients like the ones examined in this study will be able answer this question. Likewise, measuring affective bias to experimental pain in healthy controls may decipher the relationship between affective biasing and predisposition to depression.

We acknowledge that our findings are based on a small sample size. Although we observed large statistical differences between MDD and non-MDD groups, further studies confirming our results would aid in generalizing present findings. Another possible limitation of our study is the relatively young age of our participants, which might question the applicability of our findings to older patients with MDD. Because chronic pain conditions are common in older patients with depressive disorders (1,56), identifying older patients with MDD and without chronic pain comorbidities is more challenging. However, it is imperative to select individuals who are not currently comorbid with chronic pain to examine the alterations of pain processing in this population that are specifically related to MDD. Future studies examining responses to brief heat stimuli in older MDD patients without chronic pain as well as in patients with comorbid chronic pain and MDD conditions would aid in clarifying the relationship between pain and depression.

One interesting observation in our study that merits further investigation is the role of personality traits, and neuroticism in particular, in affective pain processing. It is widely accepted that people with high levels of neuroticism have a general tendency to experience negative affect and to report more somatic complaints (57,58) as well as show a heightened suffering component of chronic pain (59). Future studies should examine whether, in a nonclinical population, neuroticism affectively biases responses to experimental pain or leads to the expression of emotional allodynia.

To summarize, unmedicated young adults with MDD showed increased affective bias toward brief heat stimuli, which was independent of sensory pain thresholds in these patients. These results extend previous findings of increased sensitivity to deep pain in depressed patients, and suggest that increased pain sensitivity in MDD may be influenced by increased affective bias (or emotional allodynia). The increased affective bias is particularly striking during cutaneous heat stimuli at temperatures that are below pain threshold, which patients with MDD abnormally report as unpleasant. To our knowledge, this is the first investigation to link abnormalities in experimental pain processing with abnormalities of affect processing in young adults with clinical depression without comorbid chronic pain condition. Future investigations may provide new insights that could improve treatments for both chronic pain and depression.

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AUTHOR QUERIES

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AQ1— Please provide department for these authors.