Muscle injections with lidocaine improve resting fatigue and pain in patients with chronic fatigue syndrome

Roland Staud 1
Taylor Kizer 1
Michael E Robinson 2

1Department of Medicine, College of Medicine, 2Department of Clinical and Health Psychology, University of Florida, Gainesville, FL, USA

Objective: Patients with chronic fatigue syndrome (CFS) complain of long-lasting fatigue and pain which are not relieved by rest and worsened by physical exertion. Previous research has implicated metaboreceptors of muscles to play an important role for chronic fatigue and pain. Therefore, we hypothesized that blocking impulse input from deep tissues with intramuscular lidocaine injections would improve not only the pain but also fatigue of CFS patients.

Methods: In a double-blind, placebo-controlled study, 58 CFS patients received 20 mL of 1% lidocaine (200 mg) or normal saline once into both trapezius and gluteal muscles. Study outcomes included clinical fatigue and pain, depression, and anxiety. In addition, mechanical and heat hyperalgesia were assessed and serum levels of lidocaine were obtained after the injections.

Results: Fatigue ratings of CFS patients decreased significantly more after lidocaine compared to saline injections (p = 0.03). In contrast, muscle injections reduced pain, depression, and anxiety (p < 0.001), but these changes were not statistically different between lidocaine and saline (p > 0.05). Lidocaine injections increased mechanical pain thresholds of CFS patients (p = 0.04) but did not affect their heat hyperalgesia. Importantly, mood changes or lidocaine serum levels did not significantly predict fatigue reductions.

Conclusion: These results demonstrate that lidocaine injections reduce clinical fatigue of CFS patients significantly more than placebo, suggesting an important role of peripheral tissues for chronic fatigue. Future investigations will be necessary to evaluate the clinical benefits of such interventions.

Keywords: muscle injections, lidocaine, metaboreceptor, chronic fatigue

Introduction

Chronic fatigue syndrome (CFS) is a disabling disorder whose pathogenesis is only partially understood. It is characterized by unexplained chronic fatigue, exertion intolerance, and sleep and cognitive abnormalities. It shares many clinical features with fatiguing conditions such as auto-immune illnesses, chronic pain syndromes, multiple sclerosis, heart failure, and cancer. Although chronic fatigue has been reported by patients after infections with herpes viruses (EB virus, human herpes virus 6 [HHV-6], and cytomegalovirus [CMV]) and Lyme borreliosis, it remains highly controversial whether patients’ fatigue is caused or maintained by these infectious agents. Current treatment approaches for patients with CFS focus almost entirely on symptom relief and in general rely on graded exercise or cognitive behavioral interventions. Overall, there are no currently accepted CFS treatments directly based on mechanistic understanding of this illness.
Chronic fatigue is not only one of the most important symptoms of CFS but also of other medical conditions including congestive heart failure (CHF). Initially, evaluations of fatigue experienced by such patients have focused on cardiac dysfunction, but measures of heart function only correlate poorly with their clinical fatigue levels. Accumulating evidence has implicated abnormal concentrations of muscle metabolites, including lactate, H+, and ATP for chronic fatigue, and sensitization of intramuscular afferents as important contributors to fatigue in many chronic illnesses including CFS. It has been hypothesized that such afferents become sensitive to metabolic products of skeletal muscle work via metaboreceptors by yet unknown mechanisms. It is well known that stimulation of metaboreceptors during exercise can result in dyspnea and fatigue – the so-called metabo-reflex and early acidosis during exercise appears to be responsible for enhanced metaboreceptors activity in CHF. While increased metaboreceptors activity may assist in the physiological response to exercise in healthy individuals, excessive stimulation of this reflex may be a source of persistent sympathetic over-activity and reduced vagal activity in patients with chronic fatigue. Such abnormalities are characteristic for CHF, chronic pain, and CFS alike. Thus, metaboreceptors activation may play a key role not only in the onset but also in the progression of illness-associated symptoms in chronic pain, CFS, and CHF by maintaining and stimulating compensatory mechanisms, which are deleterious in the long term. Overall, abnormal metaboreceptor activity could be relevant for chronic fatigue in many different conditions including CFS, thus providing a new target for research and treatment with focus on peripheral disease mechanisms.

We hypothesized that blockade of peripheral tissue receptors, including metaboreceptor with lidocaine injections, would reduce their input to the central nervous system and thus decrease or abolish fatigue in chronically fatigued patients, including those with CFS. We elected to inject lidocaine into the trapezius and gluteal muscles because of their large size and accessible location.

Methods
Subjects
All study procedures were approved by the University of Florida Institutional Review Board prior to the start of subject enrollment and the study protocol conformed to the ethical guidelines of the 1975 Declaration of Helsinki. CFS subjects could not have a history of heart disease, COPD, malignancy, or other systemic disorders including psychiatric illnesses that would be exclusionary for a diagnosis of CFS. Prior to testing, all subjects underwent a clinical examination and were excluded from the study if they had abnormal findings unrelated to CFS. Use of analgesics, including non-steroidal anti-inflammatory drugs (NSAIDs) and acetaminophen, was not allowed during the study. All subjects were asked to discontinue analgesics for the duration of five drug half-lives before testing, except narcotics that had to be stopped at least 2 weeks prior to study entry. Low-dose muscle relaxants and/or tricyclic antidepressants (≤10 mg/day) were permissible during the study for the treatment of insomnia.

Inclusion and exclusion criteria
Inclusion criteria for participants were, 1) adults over the age of 18 years; 2) the ability to give informed consent; and 3) subjects had to fulfill the 1993 Center for Disease Control Criteria for CFS which stipulate the presence of chronic, unexplained fatigue for > 6 months that interferes significantly with daily activities or work. In addition, the subjects had to have at least several symptoms, including post-exertional malaise, unrefreshing sleep, impairment of memory or concentration, and muscle/joint pain. Exclusion criteria were, 1) a relevant medical condition besides CFS; 2) current participation in another research protocol that could interfere or influence the outcome measures of this study; 3) the inability to give informed consent; 4) current use of analgesic drugs, hypnotic, or anxiolytic drugs; 5) previous adverse reaction to lidocaine; and 6) previous muscle injections with local anesthetics.

Experimental design
This study used a parallel group, double-blind, placebo-controlled design. The primary outcomes of this study were change of fatigue and pain. All subjects were trained to rate threshold mechanical and suprathreshold heat pulses to the shoulders, arms, back, and legs. The subjects received experimental pain stimuli immediately before and 30 min after muscle injections into both shoulders and gluteal muscles. The order of pressure and heat pain stimuli was counterbalanced to prevent order effects. All injections and experimental pain tests were counterbalanced to avoid order effects. Randomization was performed by our investigational pharmacy using mixed block allocation. Study medications were prepared by nursing staff not directly involved in the study and labeled with subject identifiers only. The subjects were lying comfortably on a table during quantitative sensory testing but were asked to sit up during the muscle injections. The interval between heat and pressure stimuli was always 30 s or until pain after-sensations (AS) were no longer reported by the subjects.
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Ratings of fatigue, pain, and mood
Ratings of overall fatigue, pain, and mood were obtained before and 30 min after the muscle injections. A 15 cm mechanical visual analog scale (mVAS; 0–10) was used for ratings of experimental and clinical fatigue and pain.20 This scale is anchored on the left by “no fatigue/pain at all” and on the right by “the most intense fatigue/pain imaginable.” The same scale was used for ratings of anxiety/depression. This scale, however, was anchored on the left by “no anxious feelings/depressed feelings at all” and on the right by “most anxious feelings/depressed feelings imaginable”.

Muscle injections
All subjects received one muscle injection into the center of each trapezius muscle, midway between the acromion and the cervical spine, and one injection into the upper medial quadrants of each gluteus maximus muscle using a 27 g hypodermic needle for a total of four injections. The order of injections was counterbalanced. The injection depth depended on skin thickness but was at least 1 inch to reach muscle tissue. These injection sites were selected because they represented large muscle groups used for activities of daily living. Study medications were prepared in syringes by nurses not involved in the injections. Each syringe used for muscle injections contained either 5 mL of 1% lidocaine (50 mg, without additives) or 5 mL of normal saline. The injections were conducted while the subjects sat on the examination table connected to an electrocardiograph and blood pressure/pulse monitor. The study physician (RS) provided the following instructions for the participants: “the injections you are going to receive contain either a local anesthetic or an inert substance”. Then he slowly injected the content of each syringe into the designated muscles over 2 min in counterbalanced order.

Thirty min and 60 min after muscle injections, venous blood was drawn for analysis of lidocaine concentrations. The blood samples were immediately sent to the laboratory for analysis.

Expectations of treatment allocation
Immediately after the muscle injections, the participants were asked by forced choice to indicate whether they believed having received either active study medication or an inert substance.

Experimental pain stimuli
Experimental heat stimuli were applied before and ~30 min after the injections to: 1) three marked areas on the volar surface of both forearms 10 cm proximal of the wrist (areas were separated by 3 cm), and 2) the midpoint of the calf 5 cm lateral to the tibia over the anterior tibialis muscle. These areas were selected because they were distal to the injection sites, thus allowing testing of central nervous system effects of tissue injections on pain sensitivity. For each heat stimulus, the technician placed the thermode on each subject’s skin using light pressure while the participant was resting comfortably on the examination table. Similarly, pressure stimuli were applied before and ~30 min after injections to, 1) the midpoint of both trapezius muscles (midway between spinal column and acromion), and 2) the medial upper quadrant of the gluteus maximus muscles. These areas were selected because they were identical to the injections sites, thus allowing testing of local nervous system effects of tissue injections on pain sensitivity. Each heat and pressure stimulus was repeated three times in counterbalanced order. The interval between stimuli was always 30 s or until AS were no longer reported by the subjects. Heat and pressure testing was counterbalanced to avoid order effects.

Heat testing
Thermal probe
Precise heat pulses were generated by a Peltier thermode (Pathways; Medoc Advanced Medical Systems, Ramat Yishai, Israel). The heat probe was composed of a Peltier element that provides fast heating rates of up to 10°C/s and cooling rates of up to 8°C/s. The thermode provided stimulation of a skin area of 3 × 3 cm (surface area: 9 cm²). Special hardware and software allowed precise temperature control. The thermal sensors of the thermode were always calibrated before the experiments.

Heat stimuli
The participants received three 10s ramp and hold heat pulses of 44°C, 45°C, and 46°C each to the middle of the volar forearm and the center of the tibialis anterior muscle (“Experimental pain stimuli” section) on both sides. These temperatures were chosen in preliminary experiments, because they were found to be well tolerated by most participants without eliciting withdrawals. The order of testing was counterbalanced. The temperature of the ramp and hold heat pulses increased from 35°C to the target temperature over 6 s and was maintained at the peak temperature for 4 s. The subjects were asked to rate the sensation intensities after each heat pulse using the mVAS. The average of all heat pulse ratings obtained with the same stimulus intensity at each location was used for statistical analyses.
Mechanical pain threshold testing

Dolorimeter

A calibrated electronic dolorimeter (Wagner Force Measurement, Greenwich, CT, USA) was utilized for 10 s pressure stimuli. The rubber tip of the dolorimeter was 1 cm in diameter.

Mechanical stimuli

For testing of pressure pain thresholds, the electronic dolorimeter was applied to, 1) the center of the shoulders (trapezius muscle), and 2) the upper medial quadrant of the gluteal muscles (“Experimental pain stimuli” section). The order of test locations was counterbalanced to prevent order effects. After the dolorimeter was placed on the target area, pressure was gradually increased by 0.5 kg/s until the subject reported pain for the first time. The average of three threshold ratings obtained at the same location was used for statistical analyses.

Data analysis

Statistical analyses were calculated using SPSS 21.0 software (IBM Corporation, Armonk, NY, USA). Baseline group differences in clinical fatigue, pain, depression, anxiety, and age were analyzed by independent t-tests. The effects of muscle injections on experimental fatigue, pain, depression, and anxiety of CFS subjects were analyzed by repeated measures analysis of variance (ANOVA). Significant main and interaction effects were decomposed with appropriate t-tests. Linear regressions were performed to assess the contributions of lidocaine blood concentrations to behavioral rating changes. Previous results of lidocaine muscle injections on overall pain in similar study subjects have demonstrated moderate effect size (Cohen’s d = 0.7). Therefore, we used Cohen’s power tables estimating that a sample size of 30 subjects would achieve power >0.8 with alpha at 0.05 (two-tailed). Significance was set at alpha 0.05.

Results

Participants

Prior to enrollment, all subjects provided written informed consent and were told about the nature of the study. All subjects met the 1994 Center for Disease Control Criteria for CFS. A total of 30 CFS subjects (average age [SD] 50.8 [12.4] years) were randomized to lidocaine injections and 28 CFS participants (average age; standard deviation [SD] 50.3 [11.1] years) to saline injections. An independent t-test revealed no significant age differences between the groups (t = 0.16, p > 0.05). (see Table 1)

Muscle injections

All participants tolerated the injections well, except two individuals who demonstrated mild sweating and bradycardia after the injection, which rapidly normalized after they were placed in the supine position. No hematomas, pneumothorax, allergic reactions, or other complications were observed.

Effects of muscle injections on fatigue and pain

The average (SD) fatigue and pain ratings of CFS subjects assigned to either lidocaine or normal saline are shown in Table 1.

A repeated measures ANOVA was performed to examine the effect of muscle injections on overall fatigue using time (2) as within and condition (2) as between-subjects factors. This analysis demonstrated a significant main effect of time (F(1,43) = 20.7; p < 0.001; η² = 0.33) and a significant time × condition interaction effect (F(1,43) = 5.2; p = 0.03; η² = 0.11) (Figure 1).

A similar ANOVA with pain ratings as the dependent variable also showed a significant main effect of time (F(1,43) = 26.5; p < 0.001; η² = 0.38) but a non-significant time × condition interaction effect (F(1,43) = 0.13; p > 0.05; η² = 0.003) (Figure 2).

Table 1 Demographics

<table>
<thead>
<tr>
<th></th>
<th>Lidocaine injections</th>
<th>Saline injections</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Females/males</td>
<td>28/2</td>
<td>26/2</td>
<td></td>
</tr>
<tr>
<td>Age (SD), years</td>
<td>50.8 (12.4)</td>
<td>50.3 (11.1)</td>
<td>ns</td>
</tr>
<tr>
<td>Illness duration (SD), years</td>
<td>13.9 (7.1)</td>
<td>15.4 (9.2)</td>
<td>ns</td>
</tr>
</tbody>
</table>

Abbreviations: ns, not significant; SD, standard deviation.

Figure 1 Mean (SD) ratings of overall fatigue of participants with CFS before (black bars) and after (hatched bars) injections with 1% lidocaine or normal saline.

Notes: All subjects received concomitant injections into both shoulders and buttocks. Fatigue ratings were obtained before and 30 min after the muscle injections. Fatigue ratings of CFS participants decreased significantly more after lidocaine compared to normal saline injections (p = 0.03).

Abbreviations: CFS, chronic fatigue syndrome; SD, standard deviation; VAS, visual analog scale.
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Effects of muscle injections on depression and anxiety

The effects of lidocaine or normal saline injections on depression and anxiety are demonstrated in Table 2. Several repeated measures ANOVAs with condition (2) as independent factors and depression or anxiety ratings as dependent factors were performed. All showed significant main effects of time but non-significant time \( \times \) condition interaction effects, indicating that depression and anxiety significantly declined over time but that this effect was not different for lidocaine compared to normal saline injections (\( p > 0.05 \)).

### Table 2 Ratings of fatigue, pain, depression, and anxiety

<table>
<thead>
<tr>
<th></th>
<th>Lidocaine injections</th>
<th>Saline injections</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Before</td>
<td>After</td>
</tr>
<tr>
<td>Fatigue VAS (SD)</td>
<td>6.3 (1.6)</td>
<td>4.1 (2.3)*</td>
</tr>
<tr>
<td>Pain VAS (SD)</td>
<td>4.7 (2.1)</td>
<td>3.1 (2.0)**</td>
</tr>
<tr>
<td>Depression VAS (SD)</td>
<td>1.6 (2.0)</td>
<td>1.2 (1.6)**</td>
</tr>
<tr>
<td>Anxiety VAS (SD)</td>
<td>2.4 (2.2)</td>
<td>2.0 (2.2)**</td>
</tr>
</tbody>
</table>

Notes: ***p < 0.001; **p = 0.003; *p < 0.05.
Abbreviations: SD, standard deviation; VAS, visual analog scale.

When fatigue and pain ratings were simultaneously introduced into the ANOVA analysis, there was a significant interaction effect of time \( \times \) ratings \( \times \) condition noted (\( F(1,43) = 5.6; p = 0.02; \eta^2_p = 0.12 \)). This analysis indicated that lidocaine injections reduced fatigue more effectively than pain without affecting pain ratings more than saline injection.

Effects of muscle injections on mechanical hyperalgesia

**Effect of muscle injections on mechanical hypervalgesia**

Mechanical pain thresholds were tested at both shoulders and gluteal muscles before and 30 min after the injections with either normal saline or lidocaine. After dependent \( t \)-tests did not show significant differences between ratings at either side, the results of both sides were averaged and entered into repeated measures ANOVAs. The effects of muscle injections on mechanical pain thresholds are demonstrated in Figure 3.

Before the injections, the average (SD) mechanical pain threshold at the shoulder was 191.1 (100.2) kPa and 231.4 (118.9) kPa and the mechanical pain threshold at the buttocks was 212.3 (119.4) kPa and 292.4 (110.1) kPa, for subjects randomized to saline and lidocaine, respectively. After the injections, the average (SD) mechanical pain threshold at the shoulders changed to 232.2 (112.8) kPa and 315.6 (154.4) kPa and the mechanical pain threshold at the buttocks changed to 317.4 (148.9) kPa and 358.5 (184.0) kPa, for subjects injected with saline and lidocaine, respectively. Repeated measures ANOVAs demonstrated significant effects of time on mechanical thresholds at both locations (all \( p < 0.001 \)), but time \( \times \) condition interactions were only significant at the shoulders (\( p = 0.03 \)) and not at the buttocks (\( p > 0.05 \)) (Table 3), indicating that lidocaine was only more effective than saline in increasing mechanical pain thresholds at the shoulders but not at any other location.

Effects of muscle injections on heat pain ratings

Heat pain sensitivity was tested with 44°C, 45°C, and 46°C heat pulses at both forearms and legs. Because results obtained with 44°C and 46°C heat stimuli at the arms and legs were similar to those obtained with 45°C stimuli, these findings are not shown here. After dependent \( t \)-test showed no significant differences between sides, the ratings at both sides were averaged and entered into repeated measures ANOVAs. The effects of tissue injections on heat pain ratings of 45°C stimuli at the arms and legs are demonstrated in Figure 4.

Before the injections, the average (SD) ratings of heat pain stimuli at the arms were 3.6 (2.0) VAS units and 4.5 (1.6) VAS units and the ratings of heat pain stimuli at the legs were 3.3 (2.3) VAS units and 4.0 (2.0) VAS units, for subjects randomized to saline and lidocaine, respectively. After the injections, the average (SD) ratings of heat pain stimuli at the arms changed to 3.3 (2.1) VAS units and 4.4 (2.3) VAS units and the ratings of heat pain stimuli at the legs changed to 2.7 (2.2) VAS units and 3.4 (1.8) VAS units, for subjects injected with saline and lidocaine, respectively.
Separate ANOVAs showed small but significant effects of injections on heat pain ratings at the legs (Table 4). All time × condition interactions, however, were non-significant (all \( p > 0.05 \); Table 3). These findings demonstrate that lidocaine injections were not more effective than saline injections for heat pain at both locations.

**Lidocaine serum levels after muscle injections**

Mean (SD) lidocaine serum levels obtained from individuals who received this medication were minimal and measured at 1.19 (0.35) \( \mu \)g/mL and 1.02 (0.27) \( \mu \)g/mL, 30 min and 60 min after the muscle injections, respectively.

**Evaluation of systematic biases related to muscle injections**

Effects of muscle injections on behavioral measures

To evaluate the rating changes of study participants after the muscle injections for systematic biases, Pearson’s product moment correlations of fatigue, pain, depression, and anxiety were performed. Although fatigue changes were significantly correlated with clinical pain changes (\( r = 0.65; p < 0.001 \)), there was no significant correlation detected between fatigue, depression, or anxiety change scores (all \( p > 0.05 \)).

**Effects of lidocaine on behavioral measures**

Serum levels of lidocaine were obtained in all subjects 30 min and 60 min after the muscle injections (“Lidocaine serum levels after muscle injections” section). However, the use of Pearson’s product moment correlation showed no statistically significant relationships between lidocaine serum levels and change scores of fatigue, pain, anxiety, and depression (all \( p > 0.05 \)).

**Treatment allocation concealment**

As problems with treatment allocation concealment can influence the outcome of interventions, all participants were asked (by forced choice) to provide their best guess whether they had received lidocaine or placebo injections. Out of 30 subjects...
randomized to lidocaine, 24 (80%) declared having received the active study drug and six stated having received placebo. Conversely, out of 28 subjects randomized to saline injections, 21 (75%) thought they had received lidocaine injections and seven believed to have received saline injections. A Fisher’s Z test for proportion showed no differences in the proportion of correct guesses between the groups ($z = 0.46; p = 0.65$).

**Discussion**

Previously, a large number of CFS studies have focused on the role of chronic infections/chronic immune activation on chronic fatigue with mixed and often conflicting results.$^{22,23}$ In contrast, very little is known about the contributions of peripheral tissues to chronic fatigue symptoms of CFS patients. In particular, evidence of abnormal concentration of muscle metabolites$^{10}$ and sensitized intramuscular fatigue receptors$^{11}$ has provided the basis for the current hypothesis that fatigue of CFS patients is at least partially affected by peripheral tissue signaling to the central nervous system. This is the first study of CFS patients demonstrating significant reductions of clinical fatigue after muscle injections with 1% lidocaine. Importantly, the interventions occurred at rest and not after a fatiguing mental or physical task, therefore demonstrating relevant effects on baseline fatigue of CFS patients. Although clinical fatigue of CFS patients was reduced by 38% after lidocaine injection, which is clinically relevant, some of this effect is most likely related to non-specific factors (expectations, context, etc.).$^{24}$ Moreover, injections with lidocaine did not result in a significantly greater decrease of clinical pain in CFS patients than saline injections. This finding suggests that the observed improvements in fatigue were not mediated by pain reductions. Importantly, CFS patients were unable to reliably distinguish lidocaine from normal saline injections, thus reducing the influence of bias from the results. Overall, the results of this study provide strong support for our hypothesis that peripheral signal transduction to the central nervous system is mechanistically important for the self-reported fatigue of CFS patients at rest.

**Figure 4** Average heat pain ratings (SD) of 10 s heat stimuli in patients with CFS at 45°C before (black bars) and 30 min after (hatched bars) muscle injections with 1% lidocaine or normal saline into both shoulder and gluteal areas.

Notes: Supra-threshold heat pain stimuli were applied to the (A) forearms and (B) lower legs (over the anterior tibialis muscles). Heat pain ratings at both sites were not significantly different after injections with lidocaine or normal saline ($p > 0.05$).

Abbreviations: CFS, chronic fatigue syndrome; SD, standard deviation.

**Table 4** Time × condition interaction effects of muscle injections on heat pain ratings

<table>
<thead>
<tr>
<th></th>
<th>Heat pain ratings ($F(1,56)$)</th>
<th>$\eta^2$</th>
<th>p-value</th>
<th>Ratings × condition ($F(1,56)$)</th>
<th>$\eta^2$</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arm 45°C</td>
<td>0.995</td>
<td>0.017</td>
<td>&gt;0.05</td>
<td>0.267</td>
<td>0.005</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>Leg 45°C</td>
<td>7.63</td>
<td>0.12</td>
<td>0.008</td>
<td>0.0</td>
<td>0.0</td>
<td>&gt;0.05</td>
</tr>
</tbody>
</table>

Notes: Bold figure indicates the statistically significant result.

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Role of peripheral tissues for fatigue
A large body of evidence supports the important role of peripheral tissue signaling in exercise-related fatigue.25-27 Throughout all muscles, free nerve endings from group III and IV muscle afferents are widely distributed that are usually either silent or demonstrate only low background discharge rates (<1 Hz) at rest. They respond to local mechanical and thermal stimulation and exercise-related muscle metabolites, including lactate, bradykinin, prostaglandins, and potassium ions.28-31 Metabolic activity of muscles is sensed by metaboreceptors associated with Aδ and C-fibers, mediating inputs from alpha- and gamma-motor neurons and their presynaptic modulation. The sensations of fatigue can occur during muscle contractions and are often enhanced after exhausting exercise in normal subjects.32 Conversely, infusions with acid, ATP, and lactate into muscles can cause fatigue in normal individuals.33 However, moderate exercise elicits little long-lasting post-exercise fatigue and no muscle pain in most normal subjects, whereas such symptoms are often exacerbated in CFS patients after mild exercise.34,35

The reason for fatigue exacerbations after often minor exercises in CFS patients is only partially understood and seems to be associated with suboptimal central activation of the neuromuscular system.32 Previous research by our group has demonstrated evidence for significant contributions of peripheral tissue signaling to chronic fatigue exacerbations,11 i.e., when muscle metabolites were allowed to accumulate in the forearms of CFS patients after brief exercise, they reported significantly greater increases of fatigue ratings than during a control condition.11 These results suggested not only that activation of metaboreceptors plays an important role for chronic fatigue but also the presence of sensitized fatigue pathways as the same experiments produced less fatigue in healthy controls.

The results of this study support and extend these findings. By blocking peripheral impulse input from deep tissues with lidocaine, fatigue ratings of CFS patients decreased significantly more than with normal saline, again suggesting that peripheral impulse input plays an important role for CFS. Whether the peripheral tissue contributions to fatigue of CFS patients is dependent on particular muscle groups is unclear at this time as all participants received muscle injections into the same four body areas. Future studies will be necessary to demonstrate whether fatigue reductions of CFS patients depend on specific injection sites or lidocaine dose.

Fatigue and pain
In addition to debilitating fatigue, the majority of patients with CFS experience chronic widespread pain.36,37 Conversely, in many conditions with widespread pain, including fibromyalgia syndrome (FM), most patients also report varying levels of fatigue.38 Some of the hallmarks of chronic widespread pain are varying degrees of central sensitization.39-41 Over the last 10 years, increasing evidence for sensitization of central pain pathways has not only been accumulated in FM38 but also in CFS,39 providing mechanistic explanations for increased pain in both disorders. Although no specific tissue abnormalities have been described in many chronic pain disorders including FM, temporomandibular disorder, low back pain, and CFS, it has been proposed that pain and hyperalgesia of such patients may be at least partially dependent on activation of peripheral receptor systems, specifically metaboreceptors.42 We and others have shown that peripheral impulse input appears to be necessary for maintaining central sensitization in many chronic pain conditions including FM.43-47

We have previously examined the effects of tissue injections with lidocaine on chronic pain of FM patients.21 Similar to this study, four injections into the trapezius and gluteal muscles were performed. Although lidocaine provided significant reductions of local FM pains, there was no superior effect detected for lidocaine on overall pain compared to normal saline injections.21 Similarly, no superior effects of lidocaine compared to saline injections on chronic pain of CFS patients was detected in this study.

At this time, we can only speculate why lidocaine injections were only superior in reducing clinical fatigue but not pain of CFS patients. One important difference between peripheral pain and fatigue signaling may be the different magnitude of metaboreceptor and pain receptor contribution for clinical fatigue and pain in CFS, respectively. Whereas large residual nociceptive signaling of muscle groups outside of the injected areas may have prevented significant pain reductions of CFS patients, the metaboreceptor contributions of the injected muscles to fatigue may be substantial enough so that significant improvements of fatigue became detectable. Future lidocaine injection studies with varying numbers of injections sites will be necessary to more conclusively answer these important questions.

Study limitations
Although unlikely, we cannot exclude a systemic effect of lidocaine injections on chronic fatigue as lidocaine was
detectable in peripheral blood of CFS patients after the injections. The lidocaine levels in peripheral blood, however, were very low and did not significantly correlate with CFS patients’ fatigue reductions.

In general, there were significant effects of muscle injections with either lidocaine or normal saline on fatigue, pain, and mood of CFS patients. This pattern is consistent with placebo effects, suggesting that non-specific treatment effects occurred with both types of injections; nevertheless, there was a small but significant additive effect from lidocaine noted. Furthermore, it is possible that the lidocaine effect on clinical pain failed to reach significance in our study because of insufficient power.

Conclusion
The lidocaine-specific effects of muscle injections provide indirect evidence for the dependence of CFS patients’ resting fatigue on impulse input from peripheral tissues, likely from metaboreceptors. Muscle injections with lidocaine effectively reduced chronic fatigue by 38%, which may be clinically relevant. The significant correlation between fatigue and pain reductions after lidocaine injections suggests the presence of similar afferent pathways for nociception and fatigue in CFS patients. Future studies will be necessary to identify which peripheral tissues play the most important role for chronic fatigue and whether pharmacological therapies can successfully reduce peripheral tissue contributions to CFS symptoms over extended periods of time.

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Disclosure
The authors report no conflicts of interest in this work.

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