The effects of slow-paced versus mechanically assisted breathing on autonomic function in fibromyalgia patients


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Abstract: Paced breathing has shown efficacy in fibromyalgia (FM), but the mechanisms associated with symptom change are largely unknown. We investigated whether changes in respiratory rate (RR) alone resulted in autonomic changes during normal, paced, and mechanically assisted breathing in untrained FM patients and controls. Participants included 20 FM patients and 14 controls matched for age and body mass index. During a single visit, participants completed three 15-minute breathing sessions: 1) normal breathing, 2) slow-paced breathing, and 3) mechanically assisted breathing (continuous positive airway pressure) while supine. Continuous blood pressure and electrocardiogram were recorded, and measures of heart rate variability (HRV) and spontaneous baroreceptor sensitivity (sBRS) were calculated. During normal breathing, FM patients had higher heart rate (HR), but lower HRV and sBRS variables compared to controls with no difference in RR. Compared to the paced breathing condition, FM patients had significantly lower HR with higher HRV and sBRS variables during mechanically assisted breathing, despite no significant change in RR. Mechanically assisted breathing provided greater benefits in autonomic function than paced breathing in untrained FM patients. Future research will be needed to elucidate the central pathways involved in these autonomic changes and whether training in paced breathing can eventually replicate the results seen in mechanically assisted patients.

Keywords: fibromyalgia, heart rate variability, baroreceptor sensitivity, paced breathing, autonomic activity, mechanically assisted breathing

Introduction

Fibromyalgia (FM) is a disorder characterized by chronic widespread pain. Individuals diagnosed with FM often report significant symptoms of comorbid fatigue, sleep dysfunction, and limited physical activity. Psychologically, they frequently present with symptoms of depression, anxiety, and catastrophizing about the impact pain has on daily routine, work, and social activities. FM is typically assessed and treated within the biopsychosocial model framework, and multidisciplinary interventions that include medications and behavioral pain management strategies have demonstrated the most improvement in pain severity and daily functioning. Several recently published studies have demonstrated improvements in pain and other FM-related symptoms in patients after training in a breathing-focused intervention. Specifically, heart rate variability (HRV) biofeedback paired with slow-paced breathing has demonstrated efficacy in the improvement of pain severity and functioning in FM. However, the underlying physiological changes associated with the efficacy of these interventions are not well understood.
Physiologic dynamics among cardiac, respiratory, and blood pressure regulation in humans have been explored under various conditions, including normal breathing, paced breathing, yogic breathing, and Zen Ranzai practice.\textsuperscript{17,19,34} In healthy subjects, slow-paced breathing increases the amplitude of respiratory sinus arrhythmia (RSA), which refers to the cyclical fluctuations in heart rate (HR) that correlate with respiratory patterns. RSA is assessed using high-frequency HRV, is vagally mediated, and is used as a noninvasive measure of parasympathetic function.\textsuperscript{5,6} Respiratory rate (RR) also stimulates carotid baroreceptors through oscillatory changes in stroke volume and arterial blood pressure. Spontaneous baroreceptor sensitivity (sBRS) is a measure of the increase in the interval between heartbeats per mmHg arterial blood pressure rise. Lehrer et al have explored the interactions among HRV, respiratory activity, and baroreflexes through an HRV biofeedback protocol.\textsuperscript{25,26} At a personally unique frequency, HR and respiration are in phase and result in higher RSA through the interaction of respiratory activity and baroreflex response. However, these studies reported on patients who were trained in slow-paced breathing or HRV biofeedback. The primary aim of this study was to explore whether different breathing conditions were associated with changes in HRV and sBRS in untrained FM patients. Study participants were assessed during normal, slow-paced, and mechanically assisted breathing conditions if RR was the same.

**Patients and methods**

**Participants**

Study procedures were approved by the Mayo Clinic Institutional Review Board, and all participants provided written informed consent. Participants were recruited from the Mayo Clinic Department of Medicine Fibromyalgia Treatment Program or from the local community (matched healthy controls). Inclusion criteria for the participants with FM were as follows: 1) aged ≥18 years; 2) female; 3) clinical diagnosis of FM per the American College of Rheumatology 2011 diagnostic criteria;\textsuperscript{44} 4) pain duration of at least 6 months; 5) moderate pain level defined by at least 30 on a 0–10 pain VAS or > on a 0–10 pain VAS and medical record review during initial screening. Healthy controls were matched to the FM participants on age and body mass index (BMI). Potential control participants were excluded if they reported pain of >0 on a 0–10 pain VAS or had any history of chronic pain conditions.

**Current stage of menstrual cycle**

During study screening, day of menstrual cycle was recorded by asking for the last day of the previous menstrual period. Research has demonstrated that autonomic regulation of the heart fluctuates during the menstrual cycle with HRV being lower in the luteal phase (days 15–28) than in other phases.\textsuperscript{23,36} The results of these studies suggest that sympathetic nervous system activity is dominant during the luteal phase, possibly due to higher concentrations of progesterone.

All participants were studied in the early follicular phase of the menstrual cycle or in the low hormone phase of oral contraceptive use.

**Procedures**

**Study conditions and physiologic assessment**

The laboratory assessment was completed in one visit (~90 minutes). Study participants completed the self-report measures (described below) and were then instrumented and tested in the supine position. Instrumentation was followed by a 5-minute acclimation period for the participant to adjust to the laboratory environment and to ensure equipment was functioning correctly. Following the acclimation period, the participants were given instructions for three breathing conditions, each lasting 15 minutes with a 5-minute rest period between each condition. The first condition was normal breathing with the following instruction: “For the next 15 minutes, I would like you to breathe normally and rest quietly.” The second breathing condition was slow-paced breathing with the following instruction: “For the next 15 minutes, I would like you to breathe with the visual metronome, inhaling as the bar goes up, and exhaling as the bar goes down.” The slow-paced breathing visual metronome was set for an inhalation period of 5 seconds and an exhalation period of 5 seconds and was running on a laptop set up for easy viewing by the participant. The final condition was assisted breathing using a C-PAP machine with the following instruction: “For the next 15 minutes, I would like you to breathe with the mechanical ventilator.” The C-PAP machine...
was set for a respiration rate of six breaths per minute to match the slow-paced breathing condition.

The physiologic assessment configuration for this study included the measurement of electrocardiogram (ECG), continuous blood pressure, and RR. ECG was recorded using 3-leads (Cardiocap) placed in the Lead II configuration. Continuous blood pressure was recorded on a beat-to-beat basis by finger photoplethysmography (Finometer) regularly verified by automated sphygmomanometry on the contralateral arm. RR was recorded using thoracic and diaphragmatic belted strain-gauges (Respiratrace). All physiologic measures were recorded using the Windaq data acquisition system at a sampling rate of 500 Hz.

Standardized psychologic assessment

Pain VAS
The VAS is a 100-mm line with the endpoints anchored by “0: no pain” and “100: worst pain imaginable.” The participants were instructed to place a mark on the line indicating the present pain level.

Center for Epidemiologic Studies Depression Scale (CES-D)
The CES-D is a 20-item measure of the presence and severity of depressive symptoms. A cutoff score of ≥16 is associated with depressed outpatients. Test–retest reliability, internal consistency, and convergent validity are adequate.

State-Trait Anxiety Inventory (STAI)
The STAI is a 40-item measure used to assess individual differences related to state and trait anxiety constructs. Scores on this scale range from 20 to 80, and higher scores indicate greater anxiety. The STAI has consistently demonstrated good psychometric properties in patients with chronic pain.

Pittsburgh Sleep Quality Index (PSQI)
The PSQI is a 12-item comprehensive measure of sleep quality. It has exhibited test–retest stability (full scale r=0.85) and good overall internal consistency (α=0.83) and provides a valid and reliable assessment of overall sleep quality and disturbance.

Multidimensional Fatigue Symptom Inventory (MFSI)
The MFSI is a 30-item measure designed to identify five facets of fatigue: global experience, somatic symptoms, cognitive symptoms, affective symptoms, and behavioral symptoms. Participants rate each statement according to how true it has been for them over the past 7 days.

Autonomic activity
HRV was used as a measure of autonomic activity. To calculate HRV, the ECG signal was filtered and transformed into normal-to-normal (NN) intervals using the Windaq Waveform Browser software. These data were imported into the Nevrokard Advanced HRV analysis software, version 10.1.0, for time domain analyses (Nevrokard Kiauta, k.d., Izola, Slovenia). For this study, the time domain measure of root mean square of successive differences of NN intervals (RMSSD) is reported.

sBRS index was also calculated and reported. The time domain sBRS index is based on the quantification of sequences of three heart beats in which SBP consecutively increases or decreases by at least 0.5 mmHg, accompanied by changes in the same direction of the NN intervals by at least 5 ms of the subsequent beats. To complete the time domain sBRS, ECG and continuous blood pressure data were imported into the Nevrokard Baroreflex Sensitivity Analysis software, version 5.7.0.

Data analysis
Data were analyzed using the SPSS statistical package (version 21.0; IBM Corporation, Armonk, NY, USA). Descriptive statistics included mean and standard deviations for all demographic, questionnaire, and physiological data. Any outlying scores were compared to the original data to ensure there were no data entry errors. The alpha level for all analyses was set at p<0.05. The study hypotheses were evaluated by using repeated measures analysis of variance (ANOVA) for within group comparisons. Study groups were compared on self-report and normal breathing physiological measures using ANOVA. Means and standard deviations of the physiological data for each full 15-minute recording period are reported for HR, mean arterial pressure (MAP), and RR. The HRV and sBRS values were calculated using the Nevrokard software using data from each 15-minute recording period.

Results

Demographics
Study participants were patients with a diagnosis of FM (n=20) and healthy controls (n=14) matched on age (mean age FM group =46.2 [SD =9.0] vs control group =47.0 [SD =9.0], p=0.801) and BMI (mean BMI FM group =28.3 [SD =8.5] vs control group =26.8 [SD =4.7], p=0.535). The FM group patients reported a mean present VAS pain score of 31.8 mm (SD =16.9) and a mean average VAS score over the past week of 50.3 mm (SD =18.9). Control group participants did not report any pain at the start of laboratory assessment.

The effects of slow-paced versus mechanically assisted breathing
Affect and physical functioning
The FM group reported significantly higher rates of depression (CES-D; *F*=35.32, *p*<0.001), trait and state anxiety (STAI; trait *F*=18.97, *p*<0.001; state *F*=9.91, *p*=0.004), sleep dysfunction (PSQI; *F*=28.42, *p*<0.001), and fatigue (MFSI total score; *F*=56.76, *p*<0.001) compared to the control group. Table 1 summarizes the details on affect and physical functioning measures.

Normal breathing physiologic variables
During normal breathing, the FM group had significantly higher HR (*F*=6.93, *p*<0.014), but significantly lower RMSSD (*F*=16.03, *p*<0.001) and sBRS (*F*=5.671, *p*=0.010) compared to the control group. No significant differences between the two groups were found on MAP and RR. Table 2 provides details regarding physiologic data during normal breathing.

Change in physiologic variables
When compared across study conditions, the FM group had significant change in HR (*F*=3.35, *p*=0.048), RMSSD (*F*=6.05, *p*<0.014) and sBRS (*F*=7.28, *p*=0.012) compared to the control group. No significant changes that occur with controlled breathing techniques in patients with FM. Slow-paced breathing, the practice of slow

### Table 1 Psychological and functional measures at baseline

<table>
<thead>
<tr>
<th>Variables</th>
<th>FM, mean (SD)</th>
<th>Control, mean (SD)</th>
<th><em>F</em></th>
<th><em>p</em>-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>CES-D</td>
<td>19.7 (8.7)</td>
<td>3.4 (5.9)</td>
<td>35.3</td>
<td>0.000</td>
</tr>
<tr>
<td>STAI-Trait</td>
<td>43.5 (12.6)</td>
<td>26.0 (5.5)</td>
<td>19.0</td>
<td>0.000</td>
</tr>
<tr>
<td>STAI-State</td>
<td>39.4 (13.3)</td>
<td>25.8 (7.0)</td>
<td>9.91</td>
<td>0.004</td>
</tr>
<tr>
<td>PSQI</td>
<td>11.0 (3.8)</td>
<td>4.2 (2.9)</td>
<td>28.4</td>
<td>0.000</td>
</tr>
<tr>
<td>MFSI (total score)</td>
<td>39.4 (19.4)</td>
<td>-9.9 (16.4)</td>
<td>56.8</td>
<td>0.000</td>
</tr>
<tr>
<td>General</td>
<td>16.7 (5.4)</td>
<td>3.5 (6.4)</td>
<td>39.9</td>
<td>0.000</td>
</tr>
<tr>
<td>Emotional</td>
<td>7.8 (5.3)</td>
<td>1.0 (1.9)</td>
<td>19.5</td>
<td>0.000</td>
</tr>
<tr>
<td>Physical</td>
<td>12.5 (5.8)</td>
<td>1.3 (2.7)</td>
<td>42.2</td>
<td>0.000</td>
</tr>
<tr>
<td>Mental</td>
<td>11.5 (5.3)</td>
<td>1.8 (2.3)</td>
<td>38.5</td>
<td>0.000</td>
</tr>
<tr>
<td>Vigor</td>
<td>9.0 (4.2)</td>
<td>17.5 (4.5)</td>
<td>29.8</td>
<td>0.000</td>
</tr>
</tbody>
</table>

**Note:** *p*<0.05.

**Abbreviations:** FM, fibromyalgia; SD, standard deviation; CES-D, Center for Epidemiological Studies Depression measure; STAI, State Trait Anxiety Inventory; PSQI, Pittsburgh Sleep Quality Index; MFSI, Multidimensional Fatigue Symptom Inventory.

### Table 2 Physiologic measures between study groups during normal breathing

<table>
<thead>
<tr>
<th>Variables</th>
<th>FM, mean (SD)</th>
<th>Control, mean (SD)</th>
<th><em>F</em></th>
<th><em>p</em>-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart rate (beats per minute)</td>
<td>73.2 (10.4)</td>
<td>64.0 (7.1)</td>
<td>6.93</td>
<td>0.014*</td>
</tr>
<tr>
<td>MAP (mmHg)</td>
<td>79.2 (10.8)</td>
<td>82.5 (10.5)</td>
<td>0.65</td>
<td>0.427</td>
</tr>
<tr>
<td>sBRS (ms/mmHg)</td>
<td>8.9 (4.5)</td>
<td>14.8 (7.5)</td>
<td>7.28</td>
<td>0.012*</td>
</tr>
<tr>
<td>RMSSD (ms)</td>
<td>25.2 (19.7)</td>
<td>42.9 (19.7)</td>
<td>7.43</td>
<td>0.011*</td>
</tr>
<tr>
<td>Respiratory rate (breaths per minute)</td>
<td>14.7 (3.7)</td>
<td>12.8 (2.0)</td>
<td>2.69</td>
<td>0.112</td>
</tr>
</tbody>
</table>

**Note:** *p*<0.05.

**Abbreviations:** FM, fibromyalgia; SD, standard deviation; MAP, mean arterial pressure; sBRS, spontaneous baroreceptor sensitivity; RMSSD, root mean square of the SD of normal-to-normal intervals.

### Table 3 Repeated measures between experimental breathing conditions for the FM group

<table>
<thead>
<tr>
<th>Variables</th>
<th>Breathing condition, mean (SD)</th>
<th><em>F</em></th>
<th><em>p</em>-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart rate (beats per minute)</td>
<td>Normal: 72.92 (10.7)</td>
<td>Slow-paced: 72.18 (12.7)</td>
<td>3.35</td>
</tr>
<tr>
<td>MAP (mmHg)</td>
<td>Normal: 79.06 (11.3)</td>
<td>Slow-paced: 79.74 (8.6)</td>
<td>2.86</td>
</tr>
<tr>
<td>sBRS (ms/mmHg)</td>
<td>Normal: 9.32 (4.8)</td>
<td>Slow-paced: 9.87 (3.4)</td>
<td>5.67</td>
</tr>
<tr>
<td>RMSSD (ms)</td>
<td>Normal: 21.48 (8.6)</td>
<td>Slow-paced: 23.04 (7.7)</td>
<td>16.03</td>
</tr>
<tr>
<td>Respiratory rate (breaths per minute)</td>
<td>Normal: 14.89 (3.9)</td>
<td>Slow-paced: 6.05 (1.7)</td>
<td>44.19</td>
</tr>
</tbody>
</table>

**Notes:** *p*<0.05.

**Abbreviations:** FM, fibromyalgia; SD, standard deviation; MAP, mean arterial pressure; sBRS, spontaneous baroreceptor sensitivity; RMSSD, root mean square of the SD of normal-to-normal intervals.

Discussion
This physiologic study provides insight into the autonomic changes that occur with controlled breathing techniques in patients with FM. Slow-paced breathing, the practice of slow
timed breaths in response to anxiety-provoking stimuli, has been shown to alter autonomic nervous system (ANS) activity in multiple ways. In particular, previous studies have shown that sympathetic nervous system (SNS) activity decreases, as evidenced by changes in HR and blood pressure, while parasympathetic nervous system (PSNS) activity increases, as revealed by changes in HRV indices and sBRS. 

These physiological changes have been associated with a reduction in symptoms of chronic pain syndromes, including fear, anxiety, and catastrophizing. By understanding the mechanisms behind the therapeutic effects of slow-paced breathing, treatments can be refined and applied to other chronic pain conditions. Our work challenges the hypothesis that changing RR alone can drive the ANS changes responsible for the improvement in symptoms that occurs with slow-paced breathing.

In the present study during normal breathing, FM patients had measures of increased SNS activity and decreased PSNS activity compared to healthy pain-free controls. These results were anticipated, based on numerous prior studies characterizing the altered ANS activity in FM patients. The variation in time interval between heart beats, is generally determined by the balance of afferent input into the sinoatrial node. SNS activity leads to increases in HR but decreases in HRV. PSNS activity is responsible for most determinants of HRV, such as RSA, the change in NN interval between inspiration and expiration. PSNS input to baroreceptors in the carotid sinus and aortic arch increases their sensitivity to small changes in blood pressure, which increases HRV. Therefore, indices of HRV and sBRS activity have been used as reliable surrogates for PSNS activity in FM patients.

We know that controlled breathing techniques can alter the maladaptive changes in ANS activity that exacerbate symptoms of chronic pain, but have little understanding of how this occurs. To clarify the role of RR on autonomic changes, we measured markers of ANS activity in FM and control patients under both slow-paced and mechanically assisted breathing conditions. When comparing FM patients between slow-paced and mechanically assisted breathing, HR decreased and measures of both HRV and sBRS increased in the mechanically assisted condition. These changes occurred in response to anxiety-provoking stimuli, has been shown to alter autonomic nervous system (ANS) activity in multiple ways. In particular, previous studies have shown that sympathetic nervous system (SNS) activity decreases, as evidenced by changes in HR and blood pressure, while parasympathetic nervous system (PSNS) activity increases, as revealed by changes in HRV indices and sBRS. These physiological changes have been associated with a reduction in symptoms of chronic pain syndromes, including fear, anxiety, and catastrophizing. By understanding the mechanisms behind the therapeutic effects of slow-paced breathing, treatments can be refined and applied to other chronic pain conditions. Our work challenges the hypothesis that changing RR alone can drive the ANS changes responsible for the improvement in symptoms that occurs with slow-paced breathing.

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### Table 4 Repeated measures between experimental breathing conditions for the control group

<table>
<thead>
<tr>
<th>Breathing condition, mean (SD)</th>
<th>Normal</th>
<th>Slow-paced</th>
<th>Mechanically assisted</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart rate (beats per minute)</td>
<td>64.25 (8.1)</td>
<td>64.72 (9.9)</td>
<td>63.50 (9.4)</td>
</tr>
<tr>
<td>MAP (mmHg)</td>
<td>82.53 (7.6)</td>
<td>79.97 (8.6)</td>
<td>89.24 (10.2)</td>
</tr>
<tr>
<td>sBRS (ms/mmHg)</td>
<td>15.46 (9.1)</td>
<td>12.19 (4.8)</td>
<td>13.62 (7.1)</td>
</tr>
<tr>
<td>RMSSD (ms)</td>
<td>49.97 (19.6)</td>
<td>39.50 (9.3)</td>
<td>48.69 (13.9)</td>
</tr>
<tr>
<td>Respiratory rate (breaths per minute)</td>
<td>12.54 (1.0)</td>
<td>4.41 (0.7)</td>
<td>5.45 (1.0)</td>
</tr>
</tbody>
</table>

**Notes:**
- When superscripts are the same between two conditions on a measure, post hoc comparison indicates no significant difference between condition means.
- When superscripts are different, post hoc comparison indicates significant difference between group means at p < 0.01.

**Abbreviations:** SD, standard deviation; MAP, mean arterial pressure; sBRS, spontaneous baroreceptor sensitivity; RMSSD, root mean square of the SD of normal-to-normal intervals.

### Figure 1 Baroreceptor function across conditions by group.

**Note:** *p < 0.05.

**Abbreviations:** FM, fibromyalgia; sBRS, spontaneous baroreceptor sensitivity.
even though RR was not significantly different between the two conditions. In contrast, these measures were not significantly changed in the control group from one breathing condition to the next. These results indicate a unique increase in PSNS activity and potentially a decrease in SNS activity among FM patients undergoing mechanically assisted breathing compared to slow-paced breathing.

These data raise several interesting questions. Despite similar RRs between the slow-paced and mechanically assisted breathing conditions, measures of PSNS activity among FM patients were clearly improved during mechanically assisted breathing. This suggests that simply decreasing RR alone does not alter autonomic activity or explain the efficacy of slow-paced breathing for FM patients. Mechanically assisted breathing may have triggered alterations in higher level autonomic centers that change the balance between SNS and PSNS activity, which were not replicated with slow-paced breathing. Although the spinal cord and peripheral autonomic ganglia play an important role in short-term ANS activity, supraspinal structures such as the hypothalamus and limbic system are crucial for long-term and anticipatory ANS regulation.35 There is overlap between the central respiratory and autonomic centers related to the connection between the hypothalamus and medullary respiratory nuclei.16

One potential explanation for our findings is that voluntary slow-paced breathing is mediated via cortical pathways, whereas mechanically assisted breathing activates hypothalamic pathways that are integrated with autonomic regulation. It is important to note that, unlike FM patients undergoing a multidisciplinary treatment program, our subjects had no prior experience with slow-paced breathing. It is reasonable to believe that, with training, slow-paced breathing could activate the involuntary hypothalamic pathways that cause long-term adaptations in autonomic regulation. More research will be needed to define these supraspinal pathways and how they are altered by controlled breathing techniques.

Given the fact that sympathetic activity and HRV indices are correlated with pain intensity in FM patients, it is important to develop treatments that improve autonomic regulation. However, prior research disagrees on exactly how altered ANS activity causes exacerbation of pain symptoms. Findings of both sympathetic hyperactivity and hypo-reactivity suggest that the physiologic relationship between SNS activity and chronic pain may be more complex than we once thought.21,27 In addition, a few studies have actually shown normal muscle sympathetic nerve activity and normal autonomic reactivity tests in patients with FM, challenging the idea that SNS overactivity is a primary physiologic mechanism in FM.15,22 An alternative theory is that deconditioning results in peripheral ANS alterations (high SNS tone and low PSNS tone) that cause regional ischemia, which leads to widespread pain.21 This could explain why exercise is such an effective therapy for FM patients and others with chronic pain syndrome. In reality, there is likely a complex interplay between deconditioning, peripheral ANS activity, and central autonomic drive that is responsible for the varied symptoms seen in FM patients.

There are several limitations of this study that narrow the conclusions that can be drawn. This is a relatively small
study with a total of 34 participants; hence, the results may not apply to all patients with FM, especially given the varied clinical presentations seen with this disease and the lack of understanding regarding its pathophysiology. The study design did not allow for randomization of breathing conditions, thus there is a potential for an order effect that may have influenced the data. In addition, it is unclear whether some healthy controls could actually have abnormalities in ANS activity amenable to controlled breathing techniques, depending on their comorbidities and risk factors.\(^\text{40}\) We did not directly assess sympathetic activity. Our assessments using sBRS and RMSSD are generally considered measures of vagal tone or the lack thereof when low. We did not collect data on CO\(_2\) levels or operational lung volume (stretch), both of which may have contributed to our findings.\(^\text{31,42}\)

The questions raised by these data will require further study to elucidate the complex interactions between ANS activity and chronic pain, in addition to the role that controlled breathing techniques can play as part of a multidisciplinary treatment approach. Future directions should include larger studies with patients experienced at slow-paced breathing to test the hypothesis that there is a learning and/ or entrainment component responsible for the differences in ANS activity seen between slow-paced and mechanically assisted breathing. Incorporating functional MRI could also shed light on how central autonomic processes are altered with these techniques. By further understanding the physiology behind the symptoms of FM and the treatments shown to be efficacious, we will be able to better refine and individualize these therapies in the future.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**


