Factors associated with severity of symptoms in patients with chronic unexplained muscular aching

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SUMMARY Subjects with chronic, diffuse, unexplained muscular aching were recruited—21 from a primary care practice, nine from a rheumatology practice, and two from a pain clinic. No additional criteria were used to select subjects. Subjects with mild or moderate symptoms differed from those with severe symptoms with respect to the following characteristics: the presence of fatigue on awakening, the number of tender points, difficulty in sleeping, and the degree of tenderness in typical fibromyalgic areas as measured by a dolorimeter. These findings suggest that muscular aching is likely to be of greater severity if other symptoms or signs of fibromyalgia are also present.

Key words: fibromyalgia, soft tissue rheumatism, tender points, sleep disorder.

Fibromyalgia is a syndrome whose primary symptom is chronic diffuse unexplained muscular aching. Other features of the syndrome include sleep disturbance, predictable sites of localised tenderness, and a high sensitivity to exogenous modulating factors. As there is no pathognomonic feature for fibromyalgia there has been considerable debate as to what criteria are essential for diagnosis. In particular, interest has focused on the diagnostic value of tender points: which points should be assessed and how many are necessary to establish the diagnosis. Some investigators believe that the presence of tender points is an indispensable condition.

In this study we examine the way in which the criteria for fibromyalgia relate to the severity of symptoms in the general group of patients who have chronic, diffuse, unexplained muscular aching. Many of the patients in this study did not meet other criteria for fibromyalgia. The results indicate which features of the fibromyalgia syndrome are most important in identifying patients with severe disease.

Patients and methods

RECRUITMENT OF SUBJECTS

The 32 patients for this study were recruited from three sources: 21 from a hospital based family medicine clinic, nine from a private rheumatology practice, and two from a hospital based pain clinic. All patient contact and evaluation was conducted by the same person (EK). Patients from the rheumatology practice had been treated within the past six months for soft tissue rheumatism with no identifiable cause. Patients from the family medicine and pain clinics were initially identified in an epidemiology study of chronic, diffuse, unexplained muscular aching. They had been screened by questionnaire, personal interview, physical examination, and review of the medical records. No laboratory tests were performed for the purposes of this study, but all available medical records were reviewed, and subjects were excluded if laboratory tests or x rays provided a possible explanation for the subjects' symptoms. The presence of tender points was not used to identify subjects for this study.

All subjects agreeing to participate in the study were required to meet the following criteria: (a) age between 21 and 70 years; (b) muscular aching or stiffness, or both in at least three distinct locations; (c) symptoms of greater than three months' duration; (d) symptoms not secondary to osteoarthritis or other diagnoses, such as trauma, tendinitis, bursitis, inflammatory disease, or hypothyroidism.

INFORMATION OBTAINED

Subjects completed an extensive questionnaire that
provided the following information: demographic data, severity of muscle aching, problems with sleeping, presence of fatigue on awakening, frequency of aching and stiffness on awakening, presence of symptoms related to somatisation, and effects of various modulating factors on the degree of their muscle aching.

Muscle aching was assessed by the response to the question: ‘Please record the degree of muscle aching’. The subjects were asked to circle a response on a five point scale from absent to severe. Responses were divided into two categories on the basis of severity: (1) mild or moderate muscle aching and (2) moderately severe or severe aching. Although there is likely to be some overlap between the moderate and moderately severe responses, subjects in group 2 should in general have more severe symptoms than those in group 1.

Subjects were asked if they had any of the following problems with sleeping: falling asleep, waking frequently, or waking early. For each of these questions there were three possible responses: rarely or never, sometimes, or frequently (a quarter of the time or more). Subjects were considered to have problems sleeping if they responded that they had any of these problems frequently.

The symptom of non-restorative sleep was assessed by asking the subjects: ‘In general, how do you feel after you wake up in the morning?’ There were four possible responses to this question: refreshed, mildly fatigued, moderately fatigued, or severely fatigued. Subjects who chose either of the last two responses were considered to have non-restorative sleep.

To evaluate the presence of somatisation disorder we asked the subjects about the presence of symptoms that have previously been shown to be associated with this disorder: faintness, abdominal fullness, dizziness, pain or fullness in the chest, shortness of breath, pounding heart, numbness, headaches, and abdominal pain relieved by bowel movement. Each of these symptoms was scored as a ‘0’ (rarely or never), ‘1’ (occasionally), or ‘2’ (frequently). A subject was considered to have one of the above symptoms if the symptom was recorded as occurring frequently. An average score for all the somatisation symptoms was also calculated for each subject.

Factors that have been reported to modulate the symptoms of fibrositis were assessed for their ability to improve or worsen muscle symptoms. The factors studied included cold, damp, hot, and dry weather, physical activity, emotional tension, fatigue, rest, recreation, menstruation, pregnancy, and heat application.

ASSESSMENT OF TENDER POINTS

The physical examination consisted of measurement of height and weight, inspection for obvious physical abnormality, and assessment for tender points. Table 1 lists the areas assessed for tender points. These areas were defined by Campbell et al.

Eighteen of these areas are considered typical locations for tender points in patients with fibromyalgia, and the remaining seven locations were used as control sites.

Each of the 25 defined areas was assessed for tender points both by palpation and with the aid of the dolorimeter. An area was considered to be a discrete tender point by palpation only when it was reported as significantly more tender than the surrounding areas.

The value of a dolorimeter for measuring pressure sensitivity has been described elsewhere. The device we used was a simple spring loaded scale (0-00-4-50 kg) with a 0-95 cm flat round metal tip. It was used to apply a specified amount of pressure to any given point. If discrete tender points were identified the dolorimeter was then applied specifically to them, otherwise it was applied to the centre of the tender point area. The dolorimeter was then depressed and the patient asked to identify when the pressure began to hurt (tenderness level A) and when the pain became unbearable (tenderness level B). It was stressed that this was an evaluation of the patient’s sensitivity to pressure rather than of the patient’s ability to tolerate or endure the pressure. The pressure was discontinued when the patient acknowledged unbearable discomfort verbally or by flinching, grimacing, or withdrawal. We then recorded the absence or presence of discrete points of tenderness, the pressure levels associated with pain, and the nature of the patient’s response.

Subjects with severe muscle aching were compared

Table 1 Description of the principal areas examined for tender points

<table>
<thead>
<tr>
<th>Fibrositic areas</th>
<th>Control areas</th>
</tr>
</thead>
<tbody>
<tr>
<td>Occiput: 2 cm below occipital crest, 1 cm lateral to midline</td>
<td>Forehead: midline just below scalp line</td>
</tr>
<tr>
<td>Intertransverse ligaments: posterior to transverse processes C4–6</td>
<td>Forearm: volar aspect mid.forearm</td>
</tr>
<tr>
<td>Trapezius: midpoint of upper border</td>
<td>Thumb: over thumbnail with thumb placed on table</td>
</tr>
<tr>
<td>Paraspinous: 3 cm lateral to midline at level of mid-scapula</td>
<td>Shin: over bony prominence of mid-shin</td>
</tr>
<tr>
<td>Second costochondral junction: upper border of second rib just lateral to costochondral junction</td>
<td></td>
</tr>
</tbody>
</table>
with those with mild or moderate aching on the basis of
each of the following tender point counts: (a) the number of discrete tender points found in all 25
areas tested (see Table 1); (b) the number of discrete tender points in the seven control areas (see Table 1); (c) the number of discrete tender points in the 18 fibrositis areas (see Table 1); (d) the
number of tender points in the same 18 areas using the
definition of tender point as a maximum
tolerable dolorimeter reading below a threshold of
2-60 kg/cm². This threshold is the same pressure per
square centimetre as that used by Campbell et al.,
though the tips of the dolorimeters used in the two
studies differed.

In addition to the number of tender points we also
constructed four indices of tenderness based on
dolorimeter readings: (a) the averages of pressures
recorded at all locations when the patient first noted
pain or discomfort (tenderness level A); (b) the
average pressures for all locations when the degree
of applied pressure became unbearable (tenderness
level B); (c) the average tenderness level B for
traditional fibromyalgia locations; (d) the average
tenderness level B for control locations.

**Statistical Methods**

Each categorical symptom was tested for an association
with severity of muscle aching using the \( \chi^2 \)
contingency table analysis without correcting for
continuity. Ordinal variables were tested for an
association with the severity of muscle aching using
the Wilcoxon rank sum test. Logistic regression with
SAS statistical programs was used to determine
whether a combination of patient characteristics was
more predictive of degree of muscle aching than any
one characteristic alone.

**Results**

Twenty eight of the 32 subjects were women. Four
subjects had less than a high school education, and
four subjects completed college. The age of the
subjects ranged from 21 to 66 years (mean 41·2).
Duration of symptoms ranged from one to 27 years
(mean 9·1). Age of onset of the symptoms was less
than age 20 for 26% of the subjects and more than
age 40 for 24% of the subjects.

Tables 2 and 3 present the association between
the degree of muscle aching and a number of patient
characteristics. There were statistically significant
associations between muscle aching and the following
patient characteristics: number of areas with discrete
tender points, number of tender points as defined by
sensitivity to dolorimeter pressure below a given
threshold, degree of tenderness as determined by
the dolorimeter in all 25 sites or for the fibromyalgia

| Table 2 Mean levels of characteristics associated with the fibrositis syndrome |
|-----------------------------|-----------------|------------------|
| Characteristic              | Degree of muscle aching | p Value* |
|                            | Mild to moderate (n=17) | Moderately severe to severe (n=15) |
| Number of discrete tender points (all points) | 12.3 (n=16) | 17.3 | <0·03 |
| Number of tender points based on dolorimeter? | 6·7 (n=14) | 11·0 | <0·03 |
| Dolorimeter pressure all areas (TLA): (kg/cm²) | 2·5 (n=16) | 1·9 | <0·04 |
| Dolorimeter pressure all areas (TLB): (kg/cm²) | 3·3 (n=14) | 2·8 | <0·04 |
| Dolorimeter pressure fibrositis areas (TLB): (kg/cm²) | 3·2 (n=16) | 2·5 | <0·04 |
| Dolorimeter pressure control areas (TLB): (kg/cm²) | 3·8 (n=14) | 3·4 | NS |
| Somatisation score | 0·75 (n=14) | 0·69 | NS |

*The Wilcoxon rank sum test was used to determine p values. NS indicates p>0·1.

+An area was considered a tender point if the maximum tolerable
pressure at the point was below a threshold of 2·6 kg/cm².

\( \chi^2 \) test was used to determine significance level. NS indicates p>0·1.
sites alone, sleep problems, and non-restorative sleep.

We found no association between the degree of muscle aching and awakening with aching and stiffness, the overall somatisation score or any individual symptom of somatisation, or feeling tired during the day.

Although not shown in these tables, we found that the presence of modulating factors, the age of the subjects, duration of symptoms, or age of onset of symptoms were not associated with severity. We also found that most subjects were affected by modulating factors. A hot shower improved the symptoms of 89% of the subjects and cold weather, damp weather, fatigue, and overactivity each made the symptoms worse for more than 75% of the subjects.

The variable most strongly associated with increased muscle aching was non-restorative sleep. We used logistic regression to test whether other factors offered significant additional predictive value when used with the presence of non-restorative sleep. Only the number of discrete tender areas made an additional significant contribution to predicting the severity of muscle aching.

Table 4 gives a distribution of the number of discrete tender points in the 18 areas commonly used to assess tenderness. All of the subjects with severe symptoms had at least 10 tender points but so did most subjects with milder symptoms. To maximise the difference between the two groups in the percentage of subjects with severe aching, we divided the subjects into two groups: those with 13 or more tender points and those with fewer tender points.

Table 5 illustrates the additive effect of tender points and non-restorative sleep on predicting severe symptoms. Subjects who had more than 12 tender points and who did not have restorative sleep were very likely to have severe symptoms, subjects with neither condition did not have severe symptoms, and those with only one of the conditions had an intermediate chance of severe symptoms. Although subjects who had non-restorative sleep were more likely to have 13 or more tender points, the symptoms often occurred independently of each other, and both had statistically significant associations with severity of muscle aching in the logistic regression.

**Discussion**

Our results suggest that awakening with fatigue and the number of tender points are significantly and independently associated with the degree of muscle aching in patients presenting with diffuse chronic muscle aching of unknown aetiology. Although in other studies the number of tender points has been considered more important than non-restorative sleep, the latter was more significantly associated with severity of symptoms in our study. In addition, the presence of non-restorative sleep may be more important clinically than the tender point count because it is information which is more easily obtained and does not require the setting of an arbitrary threshold value. It was not possible to determine with the information available in our study whether non-restorative sleep was a cause or consequence of the musculoskeletal symptoms.

Symptoms of somatisation disorder were not

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**Table 4  Distribution of tender point counts**

<table>
<thead>
<tr>
<th>Number of discrete areas of tenderness*</th>
<th>Degree of muscle aching</th>
<th>Moderate severe to severe</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mild to moderate</td>
<td>Cumulative %</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Number</td>
<td>%</td>
</tr>
<tr>
<td></td>
<td>0-4</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>5-9</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>10-14</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>15-19</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>20-23</td>
<td>3</td>
</tr>
</tbody>
</table>

*The number of discrete areas of tenderness was not recorded for one subject with mild to moderate muscle aching.

**Table 5  Factors independently associated with severity of muscle aching**

<table>
<thead>
<tr>
<th>Non-restorative sleep**</th>
<th>13 or more tender points*</th>
<th>Proportion with severe symptoms</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>%</td>
<td>Number</td>
</tr>
<tr>
<td>No</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Yes</td>
<td>25</td>
<td>2</td>
</tr>
<tr>
<td>Yes</td>
<td>40</td>
<td>2</td>
</tr>
<tr>
<td>Yes</td>
<td>85</td>
<td>11/13</td>
</tr>
</tbody>
</table>

*p<0.05 with logistic regression; **p<0.01 with logistic regression.
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significantly related to the degree of muscular aching in this study, though the relation may become more apparent with larger samples.

Several factors were found to modulate symptoms for most subjects in this study. There was no association between the degree of the musculoskeletal symptoms and the ability of the modulating factors to affect these symptoms.

The major difference between the design of this study and that of most others evaluating the individual symptoms of the fibromyalgia syndrome is that we selected subjects only on the basis of the musculoskeletal complaint. They were achy patients without identifiable organic musculoskeletal disease. Previous studies compared subjects with strictly defined fibromyalgia with subjects without this condition.1-7 These studies found that certain symptoms are much more common in those with fibromyalgia. A limitation of these studies, however, is that the symptoms of fibromyalgia, particularly the presence of a critical number of tender points, were used to identify those with fibromyalgia. Therefore subjects with fibromyalgia necessarily had a greater prevalence of these symptoms than the controls. The circular reasoning involved in these studies has been noted previously.13 14

A second difference between our study and most previous studies is that we recruited most patients from a general medical clinic. Therefore, it is likely that many of the subjects in this study had less severe muscular aching and fewer associated symptoms than the subjects in the other studies. The heterogeneity of the patients in this study facilitated a study of factors associated with severity of musculoskeletal symptoms.

Unfortunately, without a gold standard to identify fibromyalgia it is not possible to evaluate the use of certain symptoms in making the diagnosis. Although our design reduces the problem of circular reasoning, the results of our investigation only indirectly relate to the problem of defining fibromyalgia. The study indicates, however, which features of the fibromyalgia syndrome are most associated with severe symptoms.

Our results support the model for fibromyalgia proposed by Masi and Yunus.15 In this model the fibromyalgia syndrome is not a discrete entity but a continuum from no tender points and no muscle aching to many tender points and severe muscle aching. Our results suggest that this model should be expanded to include non-restorative sleep and perhaps other factors in the fibromyalgia syndrome. The model suggests that the syndrome of fibrosis has been recognised as a clinical entity because those patients who have many of the features of the syndrome are most likely to have severe symptoms and therefore are most likely to present to the physician. The model does not suggest why the features of the syndrome occur together or whether the musculoskeletal symptoms or the other features of the syndrome represent the primary abnormality. More research is required to answer these questions.

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References
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