Specific phobia is a frequent non-motor feature in stiff man syndrome

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tyf man syndrome is a rare neurological disorder characterised by fluctuating stiffness with superimposed paroxysmal painful spasms of the trunk and legs and absence of neurological signs. Variants of the disorder include progressive encephalomyelitis with rigidity and myoclonus (PERM), with additional neurological signs such as eye movement disturbance, ataxia, or epilepsy not caused by antispastic drug treatment; and the stiff leg syndrome, where stiffness and spasms are confined to the legs. It is controversial whether stiff man syndrome, stiff leg syndrome, and PERM represent different entities or variable manifestations of the same disease process. A high prevalence of autoantibodies against glutamic acid decarboxylase (antiGAD antibodies) in both serum and cerebrospinal fluid, and the frequent association of stiff man syndrome, stiff leg syndrome, and PERM with other autoimmune disorders such as type 1 diabetes or thyroiditis, strongly suggest that stiff man syndrome, stiff leg syndrome, and PERM are closely related manifestations of an autoimmune encephalomyelitis. For the sake of simplicity, we use the term stiff man syndrome for this group of disorders and specifically refer to stiff leg syndrome and PERM only if necessary.

Previous unsystematic or anecdotal communications on small numbers of patients with stiff man syndrome reported phobia or phobic anxiety. “Phobia” in this group of patients means excessive fear and avoidance of circumscribed situations which are assumed by patients to be difficult to master because of an increase in stiffness, paroxysmal spasms, or sudden falls. Such situations comprise crossing a street, climbing downstairs without banisters, or walking unaided. Patients may be incapacitated by this form of phobia at least as much as by the motor symptoms themselves. It has been suggested that the presence of this particular anxiety is one of the reasons for the frequent misdiagnosis of psychogenic movement disorder in these patients. However, case summaries and reviews of the clinical spectrum of stiff man syndrome often do not mention this seemingly common psychological symptom at all.

Here we report the results of a systematic investigation of anxiety in 43 consecutive patients with stiff man syndrome by applying a structured diagnostic interview, the anxiety disorders interview schedule, revised version.

METHODS

Patients

From 1997 to 2002, all patients with a diagnosis of stiff man syndrome seen at the movement disorders clinic of our department of neurology were offered a multidimensional inpatient diagnostic work up that included neurological, neuropsychological, and psychiatric components. All patients who took part in the inpatient evaluation agreed to take part in all three components.

Neurological assessment

Neurological assessment comprised physical status, laboratory tests, clinical neurophysiology testing, and, if necessary, magnetic resonance imaging of the brain, spinal cord, or both. Laboratory tests included oral glucose tolerance and HbA1c, serum levels of vitamin B-12 and folic acid, and screening for autoantibodies against glutamic acid decarboxylase, parietal cells, thyroperoxidase, TSH receptors, and, in cases with a short history, amphiphysin 1 and other antineuronal antibodies. Cerebrospinal fluid was investigated in the majority of patients. Clinical neurophysiology comprised an electromyographic search for continuous motor unit firing in stiff muscles, investigation of exteroceptive spinal (myoclonic reflex spasms) and brain stem reflexes (blink and masseter inhibitory reflexes), and motor, somatosensory, and visual evoked potentials.
Psychiatric assessment

Initially, patients took part in a semistructured clinical psychiatric interview in which they were asked about the onset and course of their disorder, and how they were coping with their neurological symptoms and with the process that led to the current diagnosis. In addition, systematic inquiries were made about a past history of depressive and anxiety symptoms and significant life events preceding the onset of motor symptoms. Basic biographical information was also obtained.

The patients then took part in the anxiety disorders section of the ADIS-R (German version). This structured interview has been developed specifically for a diagnostic assessment of anxiety and related mental disorders. It not only allows one to decide whether diagnostic thresholds for particular disorders have been reached, but it also elicits information that is relevant for planning treatment, such as lists of phobically avoided situations. The ADIS-R and its German version have been shown to yield good or excellent test-retest reliability for all anxiety disorders except generalised anxiety disorder (K values of 0.92 and 0.91, respectively). Both versions have been validated against a whole range of self report questionnaires and other structured interviews.17 18

All interviews were done by the same interviewer (PH) who has had extensive training and experience in undertaking clinical and structured interviews.

In view of the fact that patients with disease related stiffness, spasms, and falls may have realistic fears of certain clinical and structured interviews.

RESULTS

Sample characteristics

Forty three patients with stiff man syndrome were investigated. The sociodemographic details of the patients are listed in table 1. Most patients had already been treated pharmacologically before the time of interview.

In 19 of the 43 patients, task specific phobia developed—that is, phobic anxiety and avoidance of tasks or situations of sufficient severity to warrant the diagnosis. Three additional patients had the same type of phobic anxiety but without behavioural avoidance, so that only a subthreshold diagnosis of phobia is justified in these cases (table 1).

Clinical description of phobia and phobic anxiety

No patient had a history of specific phobia independent of stiff man syndrome. In most cases, the phobia developed after the onset of motor symptoms, with a span ranging from no delay (co-occurrence of motor and phobic symptoms) to a delay of two years. In two patients, however, phobic symptoms clearly started several months before the onset of motor symptoms—for example, one patient developed intense fear when climbing a stair six months before stiffness became evident. Clinically, there was a positive correlation between increasing severity of motor symptoms and the development of phobia in individual cases, although, between individuals, the level of severity of motor symptoms at which phobic symptoms developed varied greatly. In 15 of these 19 patients, specific phobia was still present at the time of interview; in four others phobia had resolved parallel to a reduction in motor symptoms accompanying symptomatic treatment.

| Table 1 Sociodemographic and neurological details of 43 patients included in the study |
|-------------------------------|-------------------|-------------------|-------------------|-------------------|
| Total number | 43 | Male (n [%]) | 14 (32.6) | Female (n [%]) | 29 (67.4) |
| Age (years) | Range 18 to 75 | Mean (SD) | 52.9 (13.8) |
| Duration of disease (years) | Range 0.5 to 35 | Mean (SD) | 7.8 (7.5) |
| Type of disease* (n [%]) | SMS | 22 (51.2) | PERM | 15 (34.9) | SLS | 3 (7.0) | pSMS | 3 (7.0) |
| Antibody status (n [%]) | AntiGAD positive | 30 (69.8) | AntiAMP positive | 2 (4.7) | AntiGAD/AMP negative | 11 (25.6) |
| Present | 19 (44.2) | Phobic anxiety w/o avoidance | 3 (7.0) | Absent | 21 (48.8) |

*No patient was diagnosed with SMS and myoclonus (jerking SMS).

The tasks and situations most commonly inducing fear and avoidance were: walking unaided in open space (17), crossing a street (15), and descending (but not climbing) stairs (12). All patients reported fear of at least one of these three specific tasks; most patients reported two or all three. Further questioning often revealed additional fear of common situations and tasks, such as being among people (8), driving in a car (8), being in a closed room (7), and going into shopping centres (7) or restaurants (5). Usually, these more common situations were only avoided when they involved one of the three specific tasks listed above.

The most common fears were as follows: of growing stiff (16), of falling (14), of losing control (12), of ridiculous behaviour (8), of pain (4), of suffocation (2), of going mad (2), and of a heart attack (2).

Comparison of patients with and without phobia

Twenty one patients with stiff man syndrome did not develop phobia or phobic anxiety. Comparing patients with and without phobia (omitting the three subthreshold cases) did not show significant differences in terms of age, sex, and illness duration (table 2). Most associations between neurological and psychiatric findings were also non-significant. Task specific phobia was evenly distributed among patients with the clinical diagnoses of stiff man syndrome, stiff leg syndrome, and PERM, and among patients with and without antiGAD antibodies (table 2). Moreover, task specific phobia occurred with a similar prevalence among patients with or without a history of falls (63% v 50%) and among those with or without fractures or subluxations (62% v 53%). However, there was a significant association between exaggerated startle reactions and the presence of task specific phobia (table 2).

Psychiatric comorbidity, life events, psychogenic misdiagnosis, and results of psychological treatment

Apart from the presence of specific phobia, the psychiatric interview revealed the following pattern of life time psychiatric comorbidity:

- anxiety and/or adjustment disorders in six patients (two with generalised anxiety disorder)
- depression (major depression or dysthymia) in five patients;
comorbidity. More importantly, there was no consistent phobia in terms of age, sex, disease duration, or psychiatric specific phobia did not differ significantly from those without motor symptoms. Patients with stiff man syndrome and phobia occur independently of stiff man syndrome, but in two patients the phobic symptoms occurred before the onset of motor symptoms (table 2).

In comparison with patients without phobia, in those with phobia there was a significantly higher risk of an initial misdiagnosis of psychogenic movement disorder as a cause for their motor symptoms (table 2).

In 22 patients, no psychiatric comorbidity was detected. Again, comparing the presence or absence of lifetime psychiatric comorbidity in patients with and without specific phobia did not show significant differences, nor did comparison of the rates of important life events before the onset of stiff man syndrome (table 2).

Psychiatric, psychotherapeutic, and behavioural treatment initiated in seven patients following a correct diagnosis of stiff man syndrome, specifically for the phobic symptoms. However, as in other patients with stiff man syndrome and phobias, the phobic symptoms decreased or even disappeared when adequate control of the motor symptoms was obtained by pharmacological means.

**DISCUSSION**

On the basis of a structured diagnostic interview, our study confirms and extends previous unsystematic observations of a high prevalence of task specific phobias in patients with stiff man syndrome. We found that 44% of a consecutive sample of 43 patients with stiff man syndrome developed this particular type of phobia, and another 7% developed subthreshold phobia (anxiety without avoidance). In no patient did a specific phobia occur independently of stiff man syndrome, but in two patients, phobic symptoms developed before the onset of motor symptoms. Patients with stiff man syndrome and specific phobia did not differ significantly from those without phobia in terms of age, sex, disease duration, or psychiatric comorbidity. More importantly, there was no consistent relation between the development of phobia and the clinical manifestation as stiff man syndrome, stiff leg syndrome, or PERM, nor with the presence or absence of anti-GAD antibodies. The study confirms earlier suggestions that the presence of phobia increases the risk of a misdiagnosis of psychogenic movement disorder.

The rate of specific phobia in patients with stiff man syndrome is at least five times greater than in the general population: in the national comorbidity survey, the rate of simple phobia (a synonym for specific phobia) was 11.3% lifetime and 5.5% current. Although psychiatric symptoms and disorders are common in neurological disorders, a rate of about 50% for one specific form of anxiety disorder without particularly raised rates for depression, other anxiety, or other mental disorders has to be considered very high. Dalakas et al mention the same rate of “task specific phobias” in 10 of 20 patients with stiff man syndrome.

Nosologically, the specificity of these phobias points to a classification of “specific phobia, situational subtype” rather than agoraphobia. The term “task specific phobia,” which has already been used to describe the fears of patients with stiff man syndrome, seems to capture well the basic fear underlying this situational type of specificity. In no patient did anxiety and avoidance behaviour become generalised in a way that would justify a diagnosis of agoraphobia (without panic disorder), even later in the course of the illness. On the other hand, the phobia in our patients usually did not involve only the fear of further falling or the fear of free space. Thus, the sub specification of the specific phobia as “space phobia” is not appropriate.

The occurrence of this particular anxiety disorder may be a symptom of, or a reaction to, the neurological disorder, a side effect of drug treatment, or a comorbid condition. Arguing against a merely reactive origin, task specific phobia was distributed fairly evenly among patients with and without gait disturbance or a history of falls, fractures, or subluxations.

Three arguments suggest that the phobic symptoms are a primary manifestation of the disease itself. First, in some patients the phobic symptoms occur before the onset of motor symptoms (this aspect was also described in an earlier case series). Second, there are close temporal correlations between the onset of and decreases in motor and phobic symptoms. Third, phobic cognition remains specific in a

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**Table 2** Comparison of patients with stiff man syndrome with and without task specific phobias

<table>
<thead>
<tr>
<th>Variable</th>
<th>Patients with phobia</th>
<th>Patients without phobia</th>
<th>p Value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total number</td>
<td>19</td>
<td>21</td>
<td></td>
</tr>
<tr>
<td>Male [n (%)]</td>
<td>5 (26.3)</td>
<td>7 (33.3)</td>
<td></td>
</tr>
<tr>
<td>Female [n (%)]</td>
<td>14 (73.7)</td>
<td>14 (66.6)</td>
<td></td>
</tr>
<tr>
<td>Age (years)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>54.4 (14.9)</td>
<td>51.8 (13.6)</td>
<td>NS</td>
</tr>
<tr>
<td>Duration of disease (years)</td>
<td>8.8 (7.1)</td>
<td>5.4 (5.4)</td>
<td>NS</td>
</tr>
<tr>
<td>Type of disease [n (%)]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SMS</td>
<td>11 (57.9)</td>
<td>10 (47.6)</td>
<td></td>
</tr>
<tr>
<td>PERM</td>
<td>7 (36.8)</td>
<td>6 (28.6)</td>
<td></td>
</tr>
<tr>
<td>SLS</td>
<td>1 (5.3)</td>
<td>2 (9.5)</td>
<td></td>
</tr>
<tr>
<td>pSMS</td>
<td>0 (0.0)</td>
<td>3 (14.3)</td>
<td></td>
</tr>
<tr>
<td>Antibody status [n (%)]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AntiGAD positive</td>
<td>16 (84.2)</td>
<td>13 (61.9)</td>
<td></td>
</tr>
<tr>
<td>AntiAMP positive</td>
<td>0 (0.0)</td>
<td>2 (9.5)</td>
<td></td>
</tr>
<tr>
<td>AntiGAD/AMP negative</td>
<td>3 (15.8)</td>
<td>6 (28.6)</td>
<td></td>
</tr>
<tr>
<td>Exaggerated startle response [n (%)]</td>
<td>16 (84.2)</td>
<td>11 (52.4)</td>
<td>&lt; 0.05</td>
</tr>
<tr>
<td>Significant life event before onset [n (%)]</td>
<td>11 (57.9)</td>
<td>8 (44.4)</td>
<td>NS</td>
</tr>
<tr>
<td>Initial psychogenic misdiagnosis [n (%)]</td>
<td>15 (78.9)</td>
<td>8 (38.1)</td>
<td>&lt; 0.02</td>
</tr>
</tbody>
</table>

*Fisher’s exact and t tests, as appropriate.
†Total number for this comparison = 18 (life events could not be ascertained in three patients).

**Note:** AntiGAD, anti-glutamic acid decarboxylase; AntiAMP, anti-ampathysin; GAD, GAD type 1; SMS, stiff man syndrome; SLS, stiff leg syndrome; PERM, nor with the presence or absence of anti-GAD antibodies.
reliable and stereotypical way, closely related to situations that may be difficult to master because of spasms, stiffness, or falls, and without generalisation to other situations or to specific fears that are more common in the general population, such as fear of animals, height, or flying. Nevertheless, the results of our study do not rule out the possibility that the specific phobia is a psychological reaction to the motor symptoms, as even in the few cases with phobic symptoms before the onset of obvious motor symptoms, the latter might have been present in a subtle way unacknowledged consciously by the patient.

The pathogenesis of phobic anxiety in patients with stiff man syndrome remains unknown. Recent hypotheses explain human phobic anxiety through vestibular dysfunction or discuss a possible role of glutamatergic systems. With the vestibular hypothesis, one would expect a predominance of vestibular symptoms over the course of the disease. However, our results do not suggest an association of specific phobia with vestibular symptoms. These findings are consistent with the idea that phobic anxiety in patients with stiff man syndrome is not due to vestibular dysfunction.

Phobic anxiety in stiff man syndrome—was not the case. In patients with stiff man syndrome, contrary to the widely held belief that marked fear reactions are common in stiff man syndrome, spasticity, and without generalisation to other situations or to specific fears that are more common in the general population, such as fear of animals, height, or falling. Nevertheless, the results of our study do not rule out the possibility that the specific phobia is a psychological reaction to the motor symptoms, as even in the few cases with phobic symptoms before the onset of obvious motor symptoms, the latter might have been present in a subtle way unacknowledged consciously by the patient.

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Psychological mechanisms of associative (conditioned) or non-associative learning, or both, will most probably contribute to the development of specific phobia in patients with stiff man syndrome, but the precise nature and interactions of these psychological mechanisms with neurobiological altered thresholds for fear responses remain to be determined.

The significant association of phobia in stiff man syndrome with exaggerated startle responses might be a point of departure for future research on these interactions.

Limitations of our study concern the fact that we did not test the inter-rater reliability of the diagnoses obtained in the structured interview specifically for our sample. However, the good or even excellent reliability for the diagnoses in question obtained with the ADIS-R suggests that inter-rater reliability was not a limiting factor in our findings. A second limitation concerns the lack of a control group of patients with similar movement disorders. Without such a control group it is difficult to rule out the possibility that specific phobia in patients with stiff man syndrome is a psychological reaction to unforeseeable spasms and falls. However, apart from the difficulty in defining and arranging for an adequate control group (though patients with the rare disorder of acquired hyperekplexia might be suitable), the main message of this study—that specific phobia is a frequent non-motor feature of stiff man syndrome—is quite independent of whether it also occurs in other neurological disorders.

Conclusions
Recognition of specific phobia as a common psychological symptom in stiff man syndrome turns it into a truly neuropsychiatric condition. Clinically, the presence of task specific phobia might be a help in the prompt diagnosis of stiff man syndrome, contrary to the widely held belief that marked anxiety is a particular feature of psychogenic movement disorders. Patients with phobias without panic disorder tend not to seek help for their disorder in spite of significant disability, possibly because of embarrassment. Thus early recognition of phobia in patients with stiff man syndrome is particularly important for devising treatment strategies to help them cope with this aspect of their disease.

ACKNOWLEDGEMENT
This study was supported by a grant from the Volkswagen Stiftung (1/71452).

REFERENCES

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Competing interests: none declared