Cognitive–behavioural group treatment for a range of functional somatic syndromes: randomised trial

Andreas Schröder, Emma Rehfeld, Eva Ørnbøl, Michael Sharpe, Rasmus W. Licht and Per Fink

Background
Many specialty-specific functional somatic syndrome diagnoses exist to describe people who are experiencing so-called medically unexplained symptoms. Although cognitive–behavioural therapy can be effective in the management of such syndromes, it is rarely available. A cognitive–behavioural therapy suitable for group treatment of people with different functional somatic syndromes could address this problem.

Aims
To test the efficacy of a cognitive–behavioural therapy (Specialised Treatment for Severe Bodily Distress Syndromes, STreSS) designed for patients with a range of severe functional somatic syndromes.

Method
A randomised controlled trial (clinicaltrials.gov, NCT00132197) compared STreSS (nine 3.5 h sessions over 4 months, n=54) with enhanced usual care (management by primary care physician or medical specialist, n=66). The primary outcome was improvement in aggregate score on subscales of the 36-item Short Form Health Survey (physical functioning, bodily pain and vitality) at 16 months.

Results
Participants receiving STreSS had a greater improvement on the primary outcome (adjusted mean difference 4.0, 95% CI 1.4–6.6, P=0.002) and on most secondary outcomes.

Conclusions
In the management of functional somatic syndromes, a cognitive–behavioural group treatment was more effective than enhanced usual care.

Declaration of interest
None.

Functional somatic syndromes such as fibromyalgia, irritable bowel and chronic fatigue syndrome are a major public health issue. These syndromes are prevalent worldwide and in all medical settings, and when severe pose a major burden on those with the syndrome, on health services and on society as a whole.1–4 Although psychiatry has promising treatment strategies at its disposal, few people with these conditions are seen by mental health specialists.5–8 Instead, their management is typically shared between primary care and secondary care medical services. It consists largely of repeated investigations by a number of specialist medical services, each of which focuses on diagnosing and managing disease in specific bodily systems, and on the provision of symptomatic treatment when no disease is found.9–11 Better approaches to treatment are available: cognitive–behavioural therapy can be effective in the management of functional somatic syndromes.12–16 Although, the fragmented nature of care imposed by different specialist medical clinics and the unacceptability to patients of referral to mental health services pose obstacles to its efficient delivery and uptake.7,10,17

To address these barriers to treatment we developed a complex intervention based on CBT, entitled Specialised Treatment for Severe Bodily Distress Syndromes (STreSS). This was designed as a common treatment for patients with a variety of severe functional somatic syndromes (grouped under the unifying diagnostic category of bodily distress syndrome, multi-organ type18,19) to be suitable for delivery as a group treatment in a general hospital setting. In this study we aimed to compare the efficacy of STreSS for patients with severe functional somatic syndromes with that of enhanced usual care on self-reported physical health 16 months after randomisation.

Method
The study took place at a university general hospital in Aarhus, Denmark. In advance of the project, information about a novel model for the assessment and treatment of functional somatic syndromes was sent to all primary care physicians and hospital wards in the western part of Denmark (Jutland), which covers a population of approximately 2 million persons living in both urban and rural areas. We requested the referral of patients with such syndromes and diagnostic analogues, described in our recruitment letter as ‘patients with a long-term illness course due to medically unexplained or functional somatic symptoms who may have received diagnoses like fibromyalgia, chronic fatigue syndrome, whiplash associated disorder, or somatoform disorder’. Between March 2005 and December 2006 the case notes of all patients referred were screened for eligibility. People who were considered likely to meet inclusion criteria were invited to undergo a clinical assessment to determine eligibility. To be eligible for participation individuals had to have a chronic (i.e. of at least 2 years duration) bodily distress syndrome of the severe multi-organ type, which requires functional somatic symptoms from at least three of four bodily systems, and moderate to severe impairment in daily living.18,19 This unifying definition captures most patients with severe functional somatic syndrome diagnoses,19 and has a prevalence of 3–4% in both primary and secondary care.18 We included only adults aged 20–45 years, as we regarded the possibility of improvement to be lower in older people and the exclusion of disease to be more difficult. Patients with severe psychiatric morbidity (psychotic and bipolar disorders, alcohol or drug misuse) were also excluded, although we did include...
people with other psychiatric disorders (e.g. anxiety, depression) and medical disorders (e.g. diabetes, asthma) if these conditions did not explain the somatic symptoms. For practical reasons we excluded people involved in litigation, those who were pregnant and those who were not fluent in the Danish language (operationalised as non-Scandinavian origin).

In a block randomisation protocol (six blocks with 20 individuals per block) participants were randomised by means of a computer algorithm that used predefined concealed random numbers and stratified for gender and psychiatric lifetime comorbidity status. In each block 9 participants were randomised to receive STreSS and 11 to receive enhanced usual care. We used the ratio 9:11 because we expected a higher attrition rate in those allocated to enhanced usual care. Patients and their referring doctors were informed of the assignment through standard letters. The study was approved by the Danish Data Protection Agency and the local research ethics committee. All participating patients gave written informed consent before enrolment.

Treatments

Enhanced usual care

Figure 1 depicts the treatment elements provided to each group. All participants underwent a thorough biopsychosocial assessment (‘a’ and ‘b’ in Fig. 1) prior to randomisation to ensure that their symptoms were not due to an undiagnosed medical condition. The assessment included a review of all clinical records; the semi-structured psychiatric interview Schedule for Clinical Assessment in Neuropsychiatry (SCAN), which includes an extensive section about physical symptoms; a physical and neurological examination; and a laboratory screening battery. In cases in which the clinical assessment suggested an undiagnosed medical condition we consulted a relevant specialist, or requested further diagnostic tests before the person was considered eligible for participation. All patients were given individualised information about the nature, course and treatment options for their symptoms (‘b’). We informed each patient’s primary care and hospital doctors of the patient’s diagnosis together with a summary of the medical history (‘c’). In cases of comorbid anxiety and depression, written individualised advice on treatment was given to the patient’s primary care physician. The assessments were made by the same psychiatrists who delivered the STreSS treatment.

Alongside the information on treatment allocation, participants allocated to enhanced usual care were informed that their primary care physician, assisted by the referring specialist where appropriate, would take responsibility for their further treatment. In Denmark 98% of the population have a primary care doctor who acts as gatekeeper for the specialised healthcare system. The aim of conveying the patients’ diagnosis and medical history to the primary care doctor was to achieve a shift from diagnostic procedures to the management of somatic symptoms and comorbid mental illness. Primary care physicians in Denmark are generally well trained in psychiatric care, and many are part of psychiatric supervision groups. There was no restriction on the psychological or pharmacological interventions that could be given to these patients, or on new referrals to secondary care services. Since it is likely that the clinical assessment (‘a’, ‘b’) and the feedback from it (‘c’) enhanced the ‘usual care’ provided to these patients (‘d’), we refer to it as ‘enhanced usual care’.

Intervention

The STreSS intervention consisted of the aforementioned clinical assessment, ‘usual care’ provided by patients’ primary care physician, and several specific additional elements (‘e’ to ‘i’ in Fig. 1). Patients allocated to STreSS received nine modules of manualised psychotherapy, based on a cognitive–behavioural approach, each of 3.5 h duration and delivered to groups of nine patients by two psychiatrists. Sessions were given at weeks 1, 2, 3, 4, 6, 8, 10, 12 and 16 after randomisation. Details about the STreSS
treatment modules are given in the Appendix. Treatment modules of one of the six groups were videotaped and evaluated by two independent psychologists for adherence to the treatment manual. The psychiatrists were consultants (E.R., P.F.) or senior residents (A.S. and one other colleague) in psychiatry with at least 2 years of training in cognitive–behavioural treatment, experience with group treatment and expertise in the field of functional somatic syndromes. The senior residents were supervised by E.R.

The STreSS treatment also included a letter to the patient’s primary care doctor (available from the authors on request) which included management recommendations and offered a telephone consultancy service and close cooperation with social authorities and the patients’ employer, when needed. The STreSS psychiatrists did not prescribe drug treatment or make referrals to other specialists themselves, but could give advice to the patient’s primary care physician to do so. This consultancy service may have further influenced the usual care provided to those allocated to the STreSS intervention, and usual care may therefore have differed between treatment groups.

Usual care and patients’ diagnoses

Details regarding usual care for the period from randomisation to the trial end-point at 16 months were obtained from Danish health registries. We analysed daytime consultations in primary care, contacts with psychiatrists or psychologists in private practice, and use of out-patient and in-patient mental health services. We also recorded use of antidepressants and prescription pain medication from patients’ reports at 4 months, 10 months and 16 months. Using the clinical assessment and records we made diagnoses of the most common functional somatic syndromes according to previously reported diagnostic algorithms.19

Outcome measures

Participants completed self-rated measures at baseline (a few days prior to randomisation) and again at 4 months, 10 months and 16 months after randomisation (Fig. 1). Questionnaires were sent by post and administered by independent research assistants who were unaware of the allocation of participants. The primary outcome was the mean change in aggregate score on the 36-item Short Form Health Survey (SF-36) subscales ‘physical functioning’, ‘bodily pain’ and ‘vitality’ from baseline to 16 months, i.e. 1 year after the STreSS intervention was completed.23 These three widely used subscales were chosen because they cover key aspects of physical health that are commonly impaired in patients with functional somatic syndromes.24 The aggregate score was calculated as the mean of the z-scores of the three scales, and transformed into a t-score (mean 50, s.d. = 10, in the 1990 general US population) following the procedure for the calculation of norm-based SF-36 scores.25,26 The aggregate score ranges from 15 to 65 and has a mean of 52.5 in the general Danish population. A 4-point increase on the aggregate score equals a change of 0.5 s.d. unit in the study sample and may be regarded as a clinically significant difference.26 To facilitate comparison with other trials, we also report the more widely used SF-36 Physical Component Summary (PCS), which in the study protocol was used for power calculation (see below). On this scale, a 4–7 point improvement is regarded as a clinically relevant change.26

Secondary outcomes were treatment response, defined as an improvement in the SF-36 aggregate score of at least 4 points; improvement in social functioning (SF-36 social functioning scale) and emotional wellbeing (SF-36 mental health scale); and reduction of illness worry (7-item subscale of the Whiteley Index), physical symptoms (90-item Symptom Checklist – Revised somatisation subscale) and severity of depression and anxiety (8-item Symptom Checklist scale).28–30

Statistical analysis

The power calculation was based on the SF-36 PCS scores.25 A sample size of 120 participants was estimated to provide 83% power, at the 5% significance level, to reject the null hypothesis of no difference in improvement between the two groups, when a difference of 5 points on the PCS was assumed and losses to follow-up of 15% in the STreSS group and 30% in the comparison group were allowed. However, a validation of the SF-36 questionnaire in the study sample, based on patients’ reports at referral, demonstrated serious shortcomings with the SF-36 PCS.31 We found an unexpected moderate negative correlation of the physical and mental component summary measures, which are constructed as independent measures. According to the SF-36 manual, a low or zero correlation of the physical and mental components is a prerequisite of their use.32 Moreover, three SF-36 scales that contribute considerably to the PCS did not fulfil basic scaling assumptions.31 These findings, together with a recent report of problems with the PCS in patients with physical and mental comorbidity,32 made us concerned that the PCS would not reliably measure patients’ physical health in the study sample. We therefore decided before conducting the analysis not to use the PCS, but to use instead the aggregate score as outlined above as our primary outcome measure. This decision was made on 26 February 2009 and registered as a protocol change at clinicaltrials.gov on 11 March 2009. Only baseline data had been analysed when we made our decision and the follow-up data were still concealed. A post hoc power analysis revealed that power was slightly reduced (80% instead of 83%) by the change of primary outcome definition.

All analyses were done on an intention-to-treat basis. We fitted a mixed model with random intercept for the SF-36 aggregate score and all secondary outcomes at baseline, 4 months, 10 months and 16 months, with separate treatment effects calculated for each time point. All analyses were adjusted for gender, age, work status, lifetime psychiatric comorbidity and clinician-rated impairment, and corrected with a cluster effect for treatment group.27 Adjustment variables were defined before commencing analyses, and chosen since we regarded them to be potential moderators of change. Using this mixed model, we first tested whether the two groups differed with regard to changes over time on the primary outcome. Next, adjusted change scores from baseline to 4 months, 10 months and 16 months were calculated for each group for the primary and all secondary outcomes. We then calculated comparison effect sizes (adjusted Cohen’s d) for each time point. Finally, we calculated the relative risk for treatment response at 16 months and the number needed to treat (NNT) to achieve one additional treatment response. Further details regarding statistical analysis are available as an online supplement; this supplement also includes a sensitivity analysis in which missing values were replaced by means of multiple imputations. All analyses were done using Stata version 11 for Windows. Since the intervention group was smaller than the comparison group, it was not possible to mask the statistician to group assignment. The trial was registered with clinicaltrials.gov, number NCT00132197.

Results

Of 278 consecutively referred patients screened for eligibility, 114 were not eligible for the trial and 17 did not agree to participate (Fig. 2). Thus, 147 patients underwent the clinical assessment. Of these, 5 patients declined participation and 22 were not
eligible. The remaining 120 patients were enrolled and randomly assigned: 66 to enhanced usual care and 54 to STreSS. Outcome data were obtained for 111 patients (93%) at one or more time points after baseline and for 94 patients (78%) at 16 months (the completers group). Contrary to our expectation, the proportions completing the study were similar in both intervention and usual care groups: 44 (81%) v. 50 (76%), P=0.45. An attrition analysis found no significant difference between the completer and non-completer groups with respect to baseline characteristics.

**Baseline characteristics of participants and treatment received**

Demographic and clinical characteristics of the study participants are presented in Table 1. Patients’ conditions were long-lasting, and a majority were receiving benefits (unemployment, sick leave, disability pension). Most patients qualified for several functional somatic syndrome diagnoses and about a third had a current comorbid mental disorder. All patients fulfilled the diagnostic criteria for at least one somatoform disorder.

Of the 54 patients allocated to STreSS, 45 (83%) received STreSS, delivered in 6 treatment groups consisting of median 7.5 patients (range 6–9) by a total of 4 psychiatrists, involved in median 2.5 treatment groups (range 1–6). 6 (11%) did not receive STreSS: 2 felt too ill to participate, 2 did not want to work in a group, 2 were not able to spend the time. 3 (6%) discontinued STreSS: 1 moved to an undisclosed address, 1 relapsed with alcohol abuse, 1 could not tolerate the group.

Of the 66 allocated to enhanced usual care, 66 (100%) received enhanced usual care, provided by their primary care physician, occasionally assisted by a mental health specialist (cf. text). No restrictions on psychological or pharmacological interventions. No restrictions on referrals to secondary care or to mental health services.

**Fig. 2** Trial profile.

ITT, intention to treat; STreSS, Specialised Treatment for Severe Bodily Distress Syndromes.
We found no difference in the ‘usual care’ (element ‘d’ in Fig. 1) delivered during the 16 months after randomisation between the two groups. Patients had median 7 (IQR 3–13) v. median 9 (IQR 5–15) daytime consultations in primary care in the intervention and usual care groups respectively. A similar proportion of participants in the two study groups had contact with a mental health professional: STreSS group n = 11 (20%), median number of consultations 7 (IQR 3–10); usual care group n = 22 (33%), median number of consultations 7 (IQR 3–11). Three participants in the enhanced usual care group and one in the STreSS group received in-patient psychiatric care. At 4 months a slightly higher proportion of patients in the enhanced usual care group (40%, n = 22) than in the STreSS group (22%, n = 11) reported daily use of prescription pain medication (two-sample test of proportion z = 1.98, P = 0.047), whereas there was no difference at 10 months and 16 months (40–44% in the usual care group v. 27% in the STreSS group). We found no difference in the proportion of patients taking antidepressant medication at any time point (32–36% in the usual care group v. 35–40% in the STreSS group).

Main results

Raw means and standard deviations for the primary outcome and the more widely used SF-36 PCS at baseline and at 4 months, 10 months and 16 months after randomisation are given in Table 2; Fig. 3 shows mean scores with 95% confidence intervals derived from the regression model for the primary outcome (a) and the secondary outcomes (b) to (f). The treatment groups differed significantly on the primary outcome with regard to changes over time (Wald $\chi^2(3) = 18.0$, $P = 0.0004$), with an adjusted difference in mean SF-36 aggregate score change from baseline to 16 months of 4.0 points (95% CI 1.4–6.6, $P = 0.002$). Participants in the STreSS group showed a mean improvement in their physical health during that time of 4.0 points (95% CI 2.0–6.0, $P < 0.001$) or 0.5 s.d. unit, which was the predefined boundary for a clinically relevant change, but we observed no change in the enhanced usual care group (~0.02 point, 95% CI 1–1.6, $P = 0.98$). Adjusted comparison effect sizes (Cohen’s $d$) for all time points are also given in Fig. 3. At 16 months a moderate effect size was found for the primary outcome measure (0.50, 95% CI 0.18–0.83).

The outcome measures by the more widely used SF-36 PCS were similar; these are provided here for comparison and were not part of the primary analysis. The adjusted difference in mean change from

### Table 1: Patient characteristics

<table>
<thead>
<tr>
<th></th>
<th>Intervention group (n = 54)</th>
<th>Usual care group (n = 66)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years: mean (s.d.)</td>
<td>35.4 (6.3)</td>
<td>36.2 (6.5)</td>
</tr>
<tr>
<td>Gender: female, n (%)</td>
<td>40 (74)</td>
<td>55 (83)</td>
</tr>
<tr>
<td>Referred from secondary care, n (%)</td>
<td>8 (15)</td>
<td>12 (18)</td>
</tr>
<tr>
<td>Education, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basic school (years 7–10)</td>
<td>31 (57)</td>
<td>36 (55)</td>
</tr>
<tr>
<td>Further education</td>
<td>23 (43)</td>
<td>30 (45)</td>
</tr>
<tr>
<td>Work status, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Employed or student</td>
<td>26 (48)</td>
<td>21 (32)</td>
</tr>
<tr>
<td>Of these on sick leave</td>
<td>7 (13)</td>
<td>8 (12)</td>
</tr>
<tr>
<td>Unemployed</td>
<td>21 (39)</td>
<td>26 (39)</td>
</tr>
<tr>
<td>Disability pension or flexible work</td>
<td>7 (13)</td>
<td>19 (29)</td>
</tr>
<tr>
<td>Severe multi-organ bodily distress syndrome, n (%)</td>
<td>54 (100)</td>
<td>66 (100)</td>
</tr>
<tr>
<td>Number of sub-syndromes: mean (s.d.)</td>
<td>3.5 (0.5)</td>
<td>3.5 (0.5)</td>
</tr>
<tr>
<td>Prevalence, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Musculoskeletal tension and pain syndrome</td>
<td>50 (93)</td>
<td>62 (94)</td>
</tr>
<tr>
<td>Gastrointestinal distress syndrome</td>
<td>44 (81)</td>
<td>53 (80)</td>
</tr>
<tr>
<td>Cardiopulmonary distress syndrome</td>
<td>44 (81)</td>
<td>52 (79)</td>
</tr>
<tr>
<td>General distress syndrome</td>
<td>53 (98)</td>
<td>63 (95)</td>
</tr>
<tr>
<td>Number of functional somatic symptoms: mean (s.d.)</td>
<td>32.3 (7.5)</td>
<td>32.6 (10.0)</td>
</tr>
<tr>
<td>Illness duration, median (IQR)</td>
<td>6.7 (3–14)</td>
<td>9.5 (4–15)</td>
</tr>
<tr>
<td>Clinician-rated impairment in daily living, n (%)</td>
<td>17 (31)</td>
<td>15 (23)</td>
</tr>
<tr>
<td>Moderate</td>
<td>37 (69)</td>
<td>51 (77)</td>
</tr>
<tr>
<td>Severe</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Functional somatic syndromes, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chronic fatigue syndrome</td>
<td>30 (56)</td>
<td>41 (62)</td>
</tr>
<tr>
<td>Fibromyalgia</td>
<td>38 (70)</td>
<td>40 (61)</td>
</tr>
<tr>
<td>Irritable bowel syndrome</td>
<td>19 (33)</td>
<td>24 (36)</td>
</tr>
<tr>
<td>Non-cardiac chest pain</td>
<td>28 (52)</td>
<td>34 (52)</td>
</tr>
<tr>
<td>Hyperventilation syndrome</td>
<td>10 (19)</td>
<td>12 (18)</td>
</tr>
<tr>
<td>Hyperventilation syndrome</td>
<td>10 (19)</td>
<td>12 (18)</td>
</tr>
<tr>
<td>Tension-type headache</td>
<td>41 (76)</td>
<td>48 (73)</td>
</tr>
<tr>
<td>At least one of the above diagnoses</td>
<td>53 (98)</td>
<td>60 (91)</td>
</tr>
<tr>
<td>Current psychiatric comorbidity (DSM-IV codes), n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Major depressive disorder (296.x–296.3x)</td>
<td>9 (17)</td>
<td>14 (21)</td>
</tr>
<tr>
<td>Dysthymia (300.4)</td>
<td>2 (4)</td>
<td>3 (5)</td>
</tr>
<tr>
<td>Anxiety disorder (300.1, 300.21–300.23)</td>
<td>10 (19)</td>
<td>12 (18)</td>
</tr>
<tr>
<td>At least one of the above diagnoses</td>
<td>16 (30)</td>
<td>24 (36)</td>
</tr>
<tr>
<td>Lifetime psychiatric comorbidity, n (%)</td>
<td>31 (57)</td>
<td>40 (61)</td>
</tr>
<tr>
<td>Somatoform disorders (DSM-IV codes), n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Somatisation disorder (300.81)</td>
<td>21 (39)</td>
<td>33 (50)</td>
</tr>
<tr>
<td>Undifferentiated somatoform disorder (300.82)</td>
<td>33 (61)</td>
<td>33 (50)</td>
</tr>
<tr>
<td>Pain disorder (307.80)</td>
<td>22 (41)</td>
<td>19 (29)</td>
</tr>
<tr>
<td>Hypochondriasis (300.7)</td>
<td>6 (11)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>At least one of the above diagnoses</td>
<td>54 (100)</td>
<td>64 (100)</td>
</tr>
<tr>
<td>Health-related quality of life: mean (s.d.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Physical health aggregate score (15–65)</td>
<td>36.4 (8.8)</td>
<td>34.6 (7.5)</td>
</tr>
<tr>
<td>Social functioning score (0–100)</td>
<td>57.6 (24.6)</td>
<td>54.2 (29.7)</td>
</tr>
<tr>
<td>Mental health score (0–100)</td>
<td>61.5 (15.9)</td>
<td>59.3 (20.1)</td>
</tr>
<tr>
<td>Illness severitya</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Physical symptoms score (0–4; mean (s.d.))</td>
<td>1.65 (0.67)</td>
<td>1.66 (0.71)</td>
</tr>
<tr>
<td>Illness worry score (0–4; median (IQR))</td>
<td>1.21</td>
<td>(0.57–1.86)</td>
</tr>
<tr>
<td>Depression/anxiety score (0–4; median (IQR))</td>
<td>0.81</td>
<td>(0.50–2.13)</td>
</tr>
</tbody>
</table>

IQR, interquartile range.

a. Diagnosis based on functional somatic symptoms in the past 2 years, according to diagnostic interview and review of clinical records.

b. Allowing more than one diagnosis per patient.

c. Only patients who currently met full criteria for a major depressive episode.

d. Without specific phobias (300.29).

e. Short Form Health Survey (SF-36) questionnaire at baseline; higher scores indicate better health.

f. Scored on the 9-item Symptom Checklist – Revised Somatisation subscale.

i. Measured using the 7-item Worry-scale.

j. Measured using the 7-item Symptom Checklist.

### Table 2: Comparison of 36-item Short Form Health Survey aggregate scores (primary outcome) and the Physical Component Summary (provided to facilitate comparison with other trials) at baseline and follow-up

<table>
<thead>
<tr>
<th></th>
<th>STreSS group Mean (s.d.)a</th>
<th>Usual care group Mean (s.d.)a</th>
<th>Unadjusted difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>SF-36 aggregate score (physical functioning, bodily pain, vitality)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>36.4 (8.8)</td>
<td>34.6 (7.5)</td>
<td>1.8</td>
</tr>
<tr>
<td>4 months</td>
<td>39.9 (9.6)</td>
<td>33.7 (6.6)</td>
<td>6.2</td>
</tr>
<tr>
<td>10 months</td>
<td>40.7 (9.3)</td>
<td>33.8 (8.3)</td>
<td>7.4</td>
</tr>
<tr>
<td>16 months</td>
<td>40.8 (10.9)</td>
<td>33.8 (8.1)</td>
<td>7.0</td>
</tr>
<tr>
<td>SF-36 Physical Component Summary</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>32.5 (11.0)</td>
<td>31.7 (9.6)</td>
<td>0.7</td>
</tr>
<tr>
<td>4 months</td>
<td>36.7 (12.1)</td>
<td>30.2 (9.3)</td>
<td>6.4</td>
</tr>
<tr>
<td>10 months</td>
<td>36.3 (12.3)</td>
<td>29.7 (10.0)</td>
<td>6.6</td>
</tr>
<tr>
<td>16 months</td>
<td>38.7 (13.0)</td>
<td>29.8 (11.0)</td>
<td>8.8</td>
</tr>
</tbody>
</table>

SF-36, 36-item Short Form Health Survey; STreSS, Specialised Treatment for Severe Bodily Distress Syndromes.

a. Means are raw values.
Although patients allocated to usual care did not improve at baseline to 16 months on the SF-36 PCS was 6.2 points (95% CI 2.5–9.9, P = 0.001). Participants allocated to STreSS improved by 5.6 points (95% CI 2.5–8.7, P < 0.001), whereas participants allocated to usual care remained substantially unchanged (–0.6 points, 95% CI –2.7 to 1.4; P = 0.54). Comparison effect sizes for the SF-36 PCS were 0.37 (95% CI 0.18–0.58), 0.43 (95% CI 0.24–0.62) and 0.61 (95% CI 0.26–0.98) at 4 months, 10 months and 16 months respectively (data not shown in Fig. 3).

**Secondary outcomes**

Figure 4 gives the proportions of patients in each treatment group who had improved by at least 4 points (treatment response) or 8 points (large improvement) on the primary outcome from baseline to 16 months. The relative risk for treatment response was 1.8 (95% CI 1.1–3.2, P = 0.03) in favour of STreSS. The NNT to achieve one additional treatment response with STreSS compared with enhanced usual care was 5 (95% CI 2–38, P = 0.03). Over half (56%) of patients in the usual care group reported their physical health to be worse than before randomisation, which was the case for only a quarter (25%) of the STreSS group.

The difference between the treatment groups increased over time, with statistical significance for all secondary outcomes (see Fig. 3) except for the anxiety and depression severity score. Although patients allocated to usual care did not improve at
any time point on any of the secondary outcomes, patients allocated to StReSS experienced an immediate reduction in physical symptoms and illness worry, which was sustained or even increased from 4 months to 16 months. However, differences in social functioning and mental health between the two groups were first observable at 10 months. This sequence suggests that improvement in social functioning and mental health might be a consequence of symptom relief, rather than a mechanism of change.

**Discussion**

For patients with severe functional somatic syndromes combined under the unifying category of multi-organ bodily distress syndrome, the StReSS intervention produced a greater improvement in self-reported physical health than that achieved by usual care enhanced by a thorough initial clinical assessment. The treatment effect was of clinical significance and was sustained at follow-up. We also noted evidence that the intervention led to greater improvements than usual care in patients’ social functioning and mental health, and to marked reductions in physical symptoms and illness worry.

To the best of our knowledge, this is the first report of a single CBT approach applicable to the management of a group of patients with different severe functional somatic syndromes conducted in secondary care. Most of the trials reported to date have included patients with a single functional somatic syndrome diagnosis, or comprising a special group such as Gulf War veterans. On the other hand, studies of somatiform disorders in mental health settings are often conducted in highly selected veterans. Our study suggests that patients with the unifying diagnosis of bodily distress syndrome can feasibly be treated together regardless of the functional somatic syndrome diagnosis they have been previously given.

The current evidence for CBT is largely limited to short-term follow-up, and recent meta-analyses have called for measurement of longer-term outcomes in trials of treatment for functional somatic syndromes. Our study provides that evidence; the positive effect induced by the StReSS intervention was sustained during the 12-month follow-up period. Moreover, since StReSS is a group treatment, the psychiatrists delivering it spent fewer hours per patient than required for individually delivered treatments. The group format was well accepted by the majority of patients, and the rates of withdrawal and effect sizes observed in this trial are comparable to those reported in previous studies. The StReSS intervention may therefore be more cost-efficient than individually delivered CBT, although this needs further investigation.

Compared with psychological interventions delivered in mental health settings, an important strength of the StReSS intervention is its integrated approach. This treatment provides patients with a positive and evidence-based understanding of their illness that aims to transcend the mind–body dualism inherent in the current diagnostic classifications, and that contrasts with the usual approach to management, which tends to suggest either physical or mental disease. Although the behavioural and cognitive treatment elements of StReSS are targeted at maintaining psychosocial and lifestyle factors that can be influenced by behavioural interventions, psychoeducation about functional somatic syndromes specifically acknowledges known biological factors in their aetiology.

Patients learn that although their somatic symptoms are best understood as pathophysiological responses to prolonged or severe mental or physical stress on the basis of genetic susceptibility, both psychological and behavioural factors are known to be involved in the initiation and maintenance of those symptoms, and that these factors therefore are the target of the treatment. This balanced conceptualisation may not only increase patients’ motivation to engage in psychological treatment, but also help them to accept their vulnerability to developing new somatic symptoms, and to adapt their lifestyle accordingly in order to prevent relapse.

**Limitations**

The findings of this trial have to be discussed against the background of its potential limitations: we included only patients with multi-organ bodily distress syndrome, and hence with multiple functional somatic symptoms. Multiple symptoms are a marker of illness severity, and therefore the efficacy of StReSS for patients with less severe functional somatic syndromes remains unclear. Furthermore, although we included consecutively referred patients from both primary and secondary care, the study sample is not fully representative of people with severe functional somatic syndromes, since only young and middle-aged patients were included. Therefore, we do not know whether StReSS would be effective in older people. A final point regarding generalisability is the fact that all patients in the study underwent a thorough clinical assessment prior to treatment allocation. We therefore cannot be sure that StReSS would be as effective in people who are not offered this kind of assessment, which might have increased the participants’ motivation to engage in the intervention.

The efficacy of StReSS in this trial may have been influenced by the fact that treatment was carried out by a few highly skilled psychiatrists, and our findings therefore require replication. However, the fact that patients in the comparison group also received a specific intervention (the clinical assessment) is likely to have improved their outcome, and potentially diminished the treatment effect observed. The interpretation of the effect sizes reported in this study is limited by the fact that the primary...
outcome was an aggregate measure of three SF-36 scales that has not yet been used in other clinical trials. However, we found similar effect sizes on the standard SF-36 PCS. Finally, although we were able to demonstrate a clear treatment effect, with the point estimate reaching the pre-specified clinically important difference of 4 points, the wide confidence intervals do not definitely establish a clinically significant treatment effect. A much larger sample would be needed to determine this.

The STreSS intervention is composed of several components, but the design of the study does not allow us to determine whether one component is more active than another in achieving change. Accordingly, our aim was simply to assess whether the whole complex intervention as delivered was acceptable and effective in improving functioning and health-related quality of life. However, it is unlikely that the observed effect was driven by systematic differences in the usual care delivered in both treatment groups, since we found no such difference in use of primary care and mental health services, or in use of prescription pain medication and antidepressants during the 16-month study period.

We are not yet able to provide cost comparison data, and the cost-effectiveness of the STreSS intervention therefore remains currently unclear. A thorough cost-effectiveness analysis that also includes societal costs such as sick leave and social benefits is in progress.

Implications
This study suggests that it is feasible and acceptable to patients and physicians to implement a single cognitive–behavioural group treatment for people with different severe functional somatic syndromes, thereby offering a pragmatic and effective treatment approach for a large group of people who are usually regarded as difficult to treat and often described as unwilling to accept psychological treatment. This novel unified treatment approach may be preferable to the implementation of different specialised treatments for single functional somatic syndromes in each secondary medical care service.

Further research
This trial is only a first step to evaluating an effective management strategy for patients with severe functional somatic syndromes. The STreSS intervention needs to be compared with other active treatments, and the possible additional effect of pharmacotherapy also requires investigation. Two further trials will address these questions. The results of our study need to be confirmed in large, multicentre trials designed to explore the effectiveness and cost-effectiveness of STreSS or similar complex interventions in people with functional somatic syndromes before widespread implementation is justified. These trials should be powered to allow subgroup analyses of the effect of STreSS on different types of functional somatic syndromes and on patients with and without psychiatric comorbidity.

In summary, STreSS provides a promising unified approach to the management of people with severe functional somatic syndromes that overcomes existing shortcomings in organisation of care.

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Andreas Schröder, MD, PhD, Emma Rehfeld, MD, Eva Ömøbel, MSc, Research Clinic for Functional Disorders and Psychosomatics, Aarhus University Hospital, Aarhus, Denmark; Michael Sharpe, MD, Psychological Medicine Research Department of Psychiatry, University of Oxford, UK; Rasmus W. Licht, MD, PhD, Mood Disorders Research Unit, Aarhus University Hospital, Risølev; Per Fink, MD, PhD, DfMedInc, Research Clinic for Functional Disorders and Psychosomatics, Aarhus University Hospital, Aarhus, Denmark

Correspondence: Dr Andreas Schröder, Research Clinic for Functional Disorders and Psychosomatics, Aarhus University Hospital, DK-8000 Aarhus C, Denmark. E-mail: andreas.schroeder@aaahus.m.dk

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Appendix
Specialised Treatment for Severe Bodily Distress Syndromes (STreSS)
Module 1: Introduction (week 1)
Enhancing motivation to deal with painful and disabling bodily symptoms. Full acknowledgement of patients’ suffering. Introduction to cognitive–behavioural therapy. Introduction of group members.

Module 2: Bodily symptoms and their interpretation (week 2)
Registering and differentiating bodily symptoms. Challenging inflexible symptom attributions.

Module 3: Illness perceptions, stress response and treatment goals (week 3)
Diagnostic labels for and subtypes of bodily distress syndromes. Biological, psychological and social factors contributing to the development and maintenance of bodily distress. Impact of negative illness perceptions. Defining individual treatment goals for each patient.

Module 4: Negative automatic thoughts and dysfunctional behaviours (week 4)
(Re-)connecting bodily symptoms with emotions, thoughts and behaviours. For each patient, identification of perpetuating factors (thoughts and behaviours) that contribute to disability.

Module 5: Cognitive distortions and emotional awareness (week 6)
Ongoing work with the connection of bodily symptoms, emotions, thoughts and behaviours. Identification of cognitive distortions. Construction of alternative responses. Enhancing emotional awareness.

Module 6: From illness behaviour to health behaviour I (week 8)
Ongoing work with the connection of bodily symptoms, emotions, thoughts and behaviours. Looking back: connecting life events and bodily distress. Looking forward: boosting pleasurable activities.

Module 7: From illness behaviour to health behaviour II (week 10)
Restoring sleep, balanced diet and physical exercise. Evaluating social network and interpersonal relationships. Evaluating work status. Revision and adjustment of individual treatment goals.

Module 8: Becoming your own therapist; relapse prevention (week 12)
Adapting lifestyle to improved functioning. Recapitulation of dysfunctional thoughts and behaviours, and construction of alternative beliefs. Providing problem-solving skills. Drawing up individual treatment manual for possible relapse.
Module 9: How to maintain learned skills and coping strategies (week 16)
Review of concepts taught and skills learned in the STreSS programme.

Treatment delivery
Each module consists of 3.5 h of therapy. Treatment is delivered by two psychiatrists to groups of seven to nine patients. Each patient is allocated to a contact psychiatrist who is primarily responsible for his or her treatment. An English version of the treatment manual is available from the authors upon request (and will be available at www.functionaldisorders.dk).

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Statistical analysis

Primary analysis

All analyses were done in Stata version 11 on an intention-to-treat basis, i.e. all patients and all available data were included in the analysis. We fitted a mixed model with random intercept for the 36-item Short Form Health Survey (SF-36) aggregate score (primary outcome) at baseline, 4 months, 10 months and 16 months, with separate treatment effects calculated for each time point (i.e. the model treated time as a fixed categorical effect). The model adjusted for gender, age, work status, lifetime psychiatric comorbidity and clinician-rated impairment, and was corrected with a cluster effect for treatment group. Adjustment variables were defined before commencing analyses, and chosen since we regarded them to be potential moderators of change. The model was checked by diagnostic plots of the residuals.

Using this mixed model we first tested whether the two groups differed with regard to changes over time on the primary outcome, i.e. tested the hypothesis that the interaction estimates of time group for all three follow-up time points were equal to zero. This was done by means of the Wald $\chi^2$ test (see Fig. 3). In a next step, adjusted change scores from baseline to 4 months, 10 months and 16 months were calculated for each group for the primary outcome. We then calculated comparison effect sizes (adjusted Cohen's $d$) for each time point by dividing the interaction estimate of group time (which represents the adjusted between-group difference in mean change from baseline to this time point) by the pooled standard deviation at baseline (see Fig. 3(a)). The same statistical model was used for the analysis of the standard Physical Component Summary (PCS) of the SF-36, which is provided for comparison only, and for the secondary outcome measures (see Fig. 3(b–f)).

Calculation of probability of treatment response and number needed to treat

Probabilities of treatment response (i.e. improvement of at least 4 points on the SF-36 aggregate score from baseline to 16 months) in both groups were calculated from log odds derived from a simple logistic regression model (i.e. a model with intervention as the only covariate). The calculation of probabilities on group level does not allow adjustment for baseline variables. We calculated relative risks (RRs) from a simple generalised linear model with a binomial family and a log link function. Number needed to treat (NNT) was estimated through $1/\text{risk difference}$ between the two groups. Both RR and NNT were estimated using unadjusted data and stated with 95% confidence intervals.

Sensitivity analysis

A sensitivity analysis was carried out for the primary outcome, using the same mixed model with random intercept as described above, but based on multiple imputation of missing outcomes. We did this analysis since we found considerable drop-out at 16 months, to ensure that the results were stable and that the reported group differences could not be explained by attrition. The multiple imputations were made by means of a multivariate normal data augmentation method (50 unique data-sets), applied to the intervention and comparison group separately, and where each used an additional measurement of the respective scale (obtained at referral, i.e. before the clinical assessment) together with the adjustment variables as covariates.

Results based on multiple imputations (sensitivity analysis)

Results based on multiple imputations differed only marginally from the main analysis, and did not change the interpretation of the trial results. Based on multiple imputation, the adjusted difference in mean SF-36 aggregate score change from baseline to 16 months was 4.1 points (95% CI 1.5–6.7, $P=0.002$). Patients in the Specialised Treatment for Severe Bodily Distress Syndromes (STreSS) intervention group improved 4.0 points (95% CI 2.0–6.0, $P<0.001$), whereas patients in the enhanced usual care group showed no improvement (−0.1 points, 95% CI −1.7 to 1.6, $P=0.94$).

Based on multiple imputation, probability of treatment response (i.e. improvement of at least 4 points on the SF-36 aggregate score) was 0.47 (95% CI 0.32–0.61) in the STreSS group and 0.26 (95% CI 0.14–0.38) in the enhanced usual care group. The relative risk for treatment response was 1.84 (95% CI 1.02–3.21, $P=0.04$) in favour of STreSS. The number needed to treat (NNT) to achieve an additional treatment response with STreSS compared with enhanced usual care was 5 (95% CI 3–53, $P=0.03$).

Conclusions

The sensitivity analysis where all missing values were replaced by means of multiple imputations yielded very similar results to the intention-to-treat-analysis based on observed data. Attrition did not introduce a bias in the trial results, and the results reported in our paper seem to be statistically robust.
Cognitive–behavioural group treatment for a range of functional somatic syndromes: randomised trial
Andreas Schröder, Emma Rehfeld, Eva Ørnbøl, Michael Sharpe, Rasmus W. Licht and Per Fink
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