Patients with severe and disabling medically unexplained symptoms are common in the healthcare system. Many of these patients are heavy users of healthcare and incur large costs to health services and society. Moreover, hospital clinicians, primary care physicians and mental health specialists often find these patients difficult to treat.\(^1,2\) Medically unexplained symptoms are classified as somatoform disorders, a DSM-IV mental health category, and in parallel as functional somatic syndromes such as irritable bowel syndrome or fibromyalgia that are typically within the realms of general and specialist medicine.\(^3,4\) All of these disorders are characterised by distressing somatic symptoms that significantly disrupt daily life. Pain is one of the most frequent and clinically relevant complaints alongside functional complaints such as diarrhoea, dizziness and fatigue.\(^5–7\) Evidence has accumulated showing that psychotherapy is a promising option (e.g. depression, health anxiety or healthcare use) would not necessarily be affected.

**Background**
Multisomatoform disorder is characterised by severe and disabling bodily symptoms, and pain is one of the most common and impairing of these. Furthermore, these bodily symptoms cannot be explained by an underlying organic disorder. Patients with multisomatoform disorder are commonly found at all levels of healthcare and are typically difficult to treat for physicians as well as for mental health specialists.

**Aims**
To test whether brief psychodynamic interpersonal therapy (PIT) effectively improves the physical quality of life in patients who have had multisomatoform disorder for at least 2 years.

**Method**
We recruited 211 patients (from six German academic outpatient centres) who met the criteria for multisomatoform disorder for a randomised, controlled, 12-week, parallel-group trial from 1 July 2006 to 1 January 2009 (International Standard Randomised Controlled Trial Number ISRCTN23215121). We randomly assigned the patients to receive either 12 weekly sessions of PIT (n = 107) or three sessions of enhanced medical care (EMC, n = 104). The physical component summary of the Short Form Health Survey (SF-36) was the pre-specified primary outcome at a 9-month follow-up.

**Results**
Psychodynamic interpersonal therapy improved patients’ physical quality of life at follow-up better than EMC (mean improvement in SF-36 score: PIT 5.3, EMC 2.2), with a small to medium between-group effect size (d = 0.42, 95% CI 0.15–0.69, P = 0.001). We also observed a significant improvement in somatisation but not in depression, health anxiety or healthcare utilisation.

**Conclusions**
This trial documents the long-term efficacy of brief PIT for improving the physical quality of life in patients with multiple, difficult-to-treat, medically unexplained symptoms.

**Declaration of interest**
None.

**Method**
We conducted this study at six university departments of psychosomatic medicine in Germany (Munich, Düsseldorf, Hannover, Heidelberg, Münster and Regensburg). We recruited patients from the out-patient departments of neurology and orthopaedics private practice. The independent clinical trials unit at the University of Düsseldorf stored all the data, regularly monitored all project sites and analysed the primary and secondary outcome data. The ethics committees at all sites approved the study.
Participants

Between June 2006 and December 2007 we used the 15-item Patient Health Questionnaire (PHQ-15) and the 36-item Short Form Health Survey (SF-36) to screen consecutive patients, aged 18–77 years, who presented at somatic out-patient departments.13,14 Trained psychologists and physicians from the respective departments of psychosomatic medicine interviewed the patients with possible multisomatoform disorder (PHQ-15 scores >10) and markedly impaired physical quality of life (SF-36 scores <40) using the somatoform disorders and hypochondria sections of the Structured Clinical Interview for DSM-IV (SCID).15 We modified the interview to check for the presence of multisomatoform disorder according to the published criteria.16 A multisomatoform disorder diagnosis requires a minimum of three current somatoform symptoms (pain, dizziness, bowel dysfunction, fatigue, etc.) that are functionally disabling and that an organic disease or another mental disorder cannot sufficiently explain, and a history of somatoform symptoms on at least half of the days over at least 2 years, resulting in healthcare use.12 We reviewed the medical charts and contacted the treating physicians to rule out possible or unclear organic explanations for the symptoms.

Patients were eligible for trial inclusion if they fulfilled the criteria for multisomatoform disorder and had at least one pain-related symptom as well as an SF-36 physical component summary (PCS) score of 40 or less. We selected only those patients with significant impairments of physical quality of life (according to the proposed DSM-5 criteria)8 to ensure that the results of these interventions could be compared regarding their differential effect on this central clinical feature. We excluded patients with insufficient cognitive abilities (a Mini-Mental State Examination score <27),16 severe, chronic and disabling somatic disease or severe comorbid mental disorder that caused major impairment in social functioning (e.g. schizophrenia, severe forms of bipolar disorder or substance misuse), as well as those who were at risk of suicide, were undergoing psychotherapy at the time of the screening or could not speak German. Additionally, we excluded a small number of patients with a DSM-IV diagnosis of hypochondriasis (n = 5 in the screened sample) because its main feature consists of excessive worry about illness and the intractable belief that one has an undiagnosed physical disease; patients with somatoform disorders, on the other hand, are impaired primarily by their physical symptoms. Patients with hypochondriasis require a focus on health anxiety, for which effective treatments are available, such as CBT.17 We did not exclude patients with major depression or an anxiety disorder if there was no major clinically detectable impairment in social functioning.

Randomisation and assessment

We obtained written informed consent after fully explaining the study to the participants. A computer program generated a blocked randomisation list and the Coordination Centre for Clinical Trials covertly applied this list to our sample. Specifically, the centre applied stratified random blocks of four, six or eight patients. After receiving informed consent, we submitted a randomisation request and the centre returned the result for the patient in question within 24 h. The patients completed the baseline assessment independently immediately after providing informed consent and delivered the assessment to the study therapist in person or by post. Thereafter, the therapist disclosed the group assignment to the patient and treatment began. When asked, 28% of the patients assigned to the EMC group disapproved of their assignment, but this opinion was not associated with study attrition.

The assessment questionnaires at the end of the therapy and for the 9-month follow-up period were handed out by the therapist after the last session or sent by post shortly afterwards. The patients completed the questionnaires independently and returned them, usually by post.

Interventions

The experimental intervention consisted of 12 weekly sessions of PIT specifically adapted to the needs of patients in bodily distress. The first session lasted up to 90 min; all other sessions were approximately 45 min long. Conceptually, this interpersonal variant of psychodynamic therapy is similar to modern variants such as mentalisation-based therapy, in that it assumes that developmentally based deregulations of (bodily) self-experience and relationships rather than unconscious conflicts are the primary basis for the symptoms.18,19 The PIT treatment manual was developed jointly and pilot-tested over the course of 3 years.20 It describes three phases. In the first phase the emphasis lies on building a therapeutic relationship by underscoring the legitimacy of bodily complaints. This relationship is achieved by assessing bodily complaints, clarifying the associated disappointments and other negative experiences with the healthcare system and introducing the possibility that there are links between emotional states and bodily complaints. In addition, bodily relaxation training is introduced and preferably maintained during the three phases of therapy. Finally, the patients are educated about the emergence of their symptoms in an appropriate and non-catastrophising manner. These basic measures are comparable to those applied in currently approved CBT approaches.21

The second phase of PIT (consisting of at least six sessions) focuses on clarifying the patient’s emotions as they relate to symptoms within the person’s own life. To this end, the links between bodily complaints and emotions that were introduced in the first phase are explored in greater detail. Furthermore, these links are discussed in the context of actual relationships. Associations are also established between the experience of severe bodily distress and earlier relationships. This goal is based on the assumption that early dysfunctional or traumatic bodily relationship experiences shape the tendency to experience disabling bodily symptoms that the patient cannot explain or control later in life.20 Identifying these dysfunctional patterns is assumed to be essential for improving the regulation of bodily self-experience and symptoms. The third phase of the therapy concentrates on termination issues, such as taking stock and planning additional psychological, social or pharmacological therapeutic measures for well-defined psychological symptoms or bodily symptoms without medical explanation. Finally, the patients move to the post-treatment phase.

After recruitment we treated the participants in the outpatient departments of psychosomatic medicine. The therapists, four psychologists and four physicians with at least 3 years of training in psychotherapy, who were employed at these departments, were trained centrally and jointly in Heidelberg and Munich in the use of the manual by the group who developed the manual (P.H., H.G., E.G., J.K., C.L., N.S. and G.S.). The local principal investigators supervised all the psychotherapies. We recorded all the sessions using video- or audiotape. Based on these recordings, we used a modified version of the Sheffield Psychotherapy Rating Scale to rate adherence to the experimental intervention.22 We rated the non-specific therapeutic and PIT interventions according to their frequency and intensity. We also included an additional 13 items to cover disorder-specific conditions relevant to treating multisomatoform disorder. The interrater reliability (Cronbach’s α) for this modified scale was...
clinically small, moderate and large. In a previous PIT trial more directly than the mental component summary. Patients addresses the major concerns of the patients included in our trial both pain and pain-related disability.

The median number of days of restricted activity per month was in six areas of physical functioning or a moderate reduction in change. This increase could result from a moderate improvement mean increase of 5 points (corresponding to an effect size of 0.5) also reduced from 5 to 0. As a consequence, we determined that a model of the development and maintenance of their symptoms. This model included advice on symptom management, specific pharmacological or psychotherapeutic treatments for either well-defined psychological symptoms or pain and healthcare use (e.g. avoiding unnecessary somatic diagnostic and/ or therapeutic interventions). The patients had three approximately half-hour sessions at 6-week intervals with physicians specifically trained in EMC (also based on a manual) at their respective departments of psychosomatic medicine. Nine specialists at the referring out-patient department provided EMC. The therapists controlled treatment adherence using checklists and covered 94% of all the assessed elements in at least one session. We considered this type of medical care to be ‘enhanced’, as (unlike typical clinical practice) it strictly observed recommended national guidelines.

During the treatment phase we told the participants not to initiate additional treatments and to report primary or specialist care contacts. At the end of therapy the clinicians recommended additional psychotherapeutic or somatic treatments and medication for the patients in a comparable manner to the PIT intervention.

Measures

We assessed the patients at baseline (t1), at the end of treatment (3 months after baseline, t2) and – to evaluate the long-term therapeutic effects – at a follow-up assessment 9 months after the end of treatment (t3). As the outcome assessment consisted of self-report measures, masking procedures were not applicable. All the measures were delivered in their validated German versions. In keeping with the current debates regarding treatment success in functional somatic syndromes, a lesser degree of symptom impairment or improvements in functional bodily health (as opposed to complete relief of symptoms) often already indicate an effective treatment. Therefore, we felt that it was important to use a health-related quality of life measure as the primary outcome, and selected the improvement in the physical component summary (range 0–100) of the SF-36,14 as measured at t3. The PCS has been shown to be a valid and change-sensitive indicator of bodily functioning and quality of life; moreover, it addresses the major concerns of the patients included in our trial more directly than the mental component summary. Patients regarded PCS changes of 2.5, 4.5 and 6.5 points respectively as clinically small, moderate and large. In a previous PIT trial investigating patients with a functional somatic syndrome,20 those who received PIT improved by a mean of 5.5 points on the PCS. The median number of days of restricted activity per month was also reduced from 5 to 0. As a consequence, we determined that a mean increase of 5 points (corresponding to an effect size of 0.5) in an individual patient would indicate a clinically significant change. This increase could result from a moderate improvement in six areas of physical functioning or a moderate reduction in both pain and pain-related disability.

As the sustainability of potential treatment effects is particularly important in a chronic condition such as multisomatoform disorder, we chose improvement at t3 rather than t2 as the primary end-point. This decision was in keeping with the observation that the effects of psychotherapy often increase after the end of the treatment.27 The secondary outcome measures assessed aspects of psychopathology or potential mediators of change. The measures reported here included the MCS (range 0–100) of the SF-36; the Patient Health Questionnaire, which contains a somatisation module (PHQ-15, range 0–30) as well as a depression module (PHQ-9, range 0–27);21 and the Whiteley Index Short Form (WI-7) to assess health anxiety.28,29 The questions on the WI-7 (range 0–7) refer to the preceding 4 weeks, and a cut-off score of 4 or more points indicates clinically relevant health anxiety. We recorded the number of times a patient visited either a primary care physician or specialist, the use of antidepressants and the use of psychotherapy in the 3 months preceding each time point (except for other psychotherapy at t2, as this was an exclusion criteria) to assess aspects of healthcare use.

Statistical analysis

Analyses, under the direction of the independent clinical trials unit at the University of Düsseldorf, were based on the intent-to-treat method for all participants and confirmed that the missing data for the primary and secondary outcomes were missing at random. This assumption holds when the missing data can be traced back to the observed data and no study-specific reason can account for their absence. Multiple imputations were used to replace missing data, which consisted mainly of patients who were lost to follow-up (n = 26). We performed sensitivity analyses to estimate the potential differences in the imputed outcomes vs. those derived from incomplete data. A sample size of 110 patients per group combined with an expected attrition rate of 20% provided 80% power to detect a moderate treatment effect size of 0.5. We compared the t1 clinical and demographic variables of the two treatment groups using chi-squared tests for dichotomous or nominally distributed variables and t-tests for continuous variables.

We tested the treatment effects in both groups across t1, t2 and t3 using linear mixed models (LMM) by analysing the differences between baseline and post-treatment or 9-month follow-up respectively and controlling for study site, age, gender and education. We report the confidence intervals for the differences between the groups. We analysed the secondary outcomes and clinical characteristics such as depression and anxiety in a similar manner. To control for differences at t1 we included the initial number of primary care and specialist visits in the LMM. We compared changes in antidepressant and psychotherapy use between the groups using Mann–Whitney U-tests. The Bonferroni correction for multiple tests was used and resulted in adjusting the alpha level for significance to 0.01 (0.05/5). We performed the statistical analyses using SPSS for Windows version 18.0. The reporting of this trial follows the Consort criteria.

Results

In total 211 participants gave informed consent and were randomised to PIT (n = 107) or EMC (n = 104); three withdrew consent to participate immediately after randomisation (Fig. 1). The withdrawal rates were 19% for PIT and 27% for EMC. Respectively, 10% and 15% of the patients in the PIT and EMC groups did not provide data at t3. The attrition rates...
Discussion
This study suggests that clinically significant improvements can be achieved using a short-term intervention consisting of 12 PIT sessions in patients who experience chronic and disabling bodily symptoms that have no recognised treatment.8 In particular, this randomised controlled trial demonstrates that, in comparison between the sites proved to be negligible. At t1, 44% and 39% of the patients in the PIT and EMC groups respectively fulfilled the DSM-IV criteria for major depression according to the PHQ-9 (i.e. having at least five depressive symptoms, at least one of which was depressed mood or loss of interest or pleasure, over a 2-week period and having these symptoms represent a change from previous functioning). Furthermore, 49% and 45% of the patients in the PIT and EMC groups respectively received antidepressants, which were stopped for both groups when their depressive symptoms improved. Using the cut-off of 4 points on the WI-7 scale the majority of patients in both groups were significantly impaired by clinically relevant health anxiety (55% in the PIT group and 56% in the EMC group).

Clinical outcomes
Table 2 shows the primary and secondary outcomes for PIT and EMC groups at the three assessment points. In terms of the primary outcome – physical quality of life as measured by the SF-36 PCS at t3 – patients in both groups improved from t1. However, the improvement of the patients in the PIT group was significantly greater than that of the EMC group, according to the LMM analysis and in tests of actual group differences at t3. The mean increase for the PIT group was 5.3 points, which passed the level of clinical significance that was set before we collected the data. Specifically, 43% of the PIT group who received psychotherapy reported 5 or more points of improvement at t3 (26% for the EMC patients; odds ratio 2.13, 95% CI 1.19–3.80). The between-group effect size (d = 0.42) was moderate. The difference between the groups became apparent in the period between t2 and t3, in which the PCS ratings of the PIT group increased further and those of the EMC group decreased slightly.

We also observed treatment effects from t1 to t3 for both groups in the patients’ secondary outcomes. Health anxiety improved in both groups; in addition, the PIT group improved their PHQ-15 somatisation and PHQ-9 depression scores, and the EMC group improved their MCS scores. We observed differential treatment effects between the groups for the PHQ-15 somatisation score, which was significantly improved in the PIT group 9 months after the end of the treatment. Despite the rate of major depressive syndrome significantly decreasing from 44% at t1 to 26% at t3 in the PIT group and remaining essentially constant in the EMC group (39% at both assessment points), there was no significant difference between the groups in the corresponding change in their PHQ-9 depression scores.

Healthcare use
Table 3 presents data on the participants’ use of healthcare. Although the number of healthcare visits for primary care and specialists remained stable for both groups, we observed a reduction in antidepressant use immediately at t2. At t3 antidepressant use remained stable for the PIT group, whereas it increased for the EMC group. We also observed a significant difference in psychotherapy use; however, neither group frequently used psychotherapy at t1 (PIT n = 3; EMC n = 12).

Characteristics at t1
At baseline we found no significant difference in age, gender, marital status, education or employment status between the two study groups (Table 1). Patients in both groups were chronically ill with multiple symptoms; on average they had more than ten somatoform symptoms (according to the SCID). The mean duration of these symptoms was 10 years. The only characteristic difference between the participating sites was participant age; in particular, differences in the treatment effect did not significantly differ between the groups. Nine patients who received PIT and twelve EMC group patients who did not complete the treatment sent back the questionnaires at the 9-month follow-up. These patients demonstrated comparable improvements in PCS score of 3.2 (PIT) and 3.8 points (EMC). Two patients were admitted to a mental hospital for a short period owing to suicidal thoughts during the treatment phase (one in PIT and one in EMC). However, both patients eventually continued with their treatments. 

Fig. 1 Study profile (t2, end of therapy; t3, 9-month follow-up).
Psychodynamic interpersonal therapy in multisomatoform disorder

with EMC, brief PIT provides greater improvement in physical quality of life at follow-up for those with multiple chronic and disabling medically unexplained symptoms and a high rate of comorbid depression and anxiety.27

Although a few controlled trials have evaluated the merits of CBT in patients with multiple somatoform symptoms,8,27 to our knowledge this is the first randomised controlled trial to demonstrate the efficacy of PIT in the same population. With a between-group effect size of 0.42, PIT provided a moderate benefit to patients,31 which is consistent with the reported treatment effects of CBT in patients with bodily distress.32 We could demonstrate this effect even though interpersonal and in particular psychodynamic techniques usually employ more therapeutic sessions than used in our PIT intervention. Furthermore, the efficacy of PIT might be underestimated in an unselected group of patients following a randomisation procedure.33

Kleinstaeuber et al recently meta-analysed 27 trials mainly applying cognitive–behavioural approaches, behavioural medical interventions or reattribution for multiple medically unexplained symptoms.27 The effects of these studies also proved to be small to modest for the majority of approaches and observable primarily for changes in physical symptoms ($d = 0.40$ at 1-year follow-up) and – with reservations – for somatoform symptoms and

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**Table 1** Characteristics of the participants at baseline

<table>
<thead>
<tr>
<th></th>
<th>Psychodynamic interpersonal therapy (n = 107)</th>
<th>Enhanced medical care (n = 104)</th>
<th>t or $\chi^2$</th>
<th>p$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at entry, years: mean (s.d.)</td>
<td>47.9 (10.8)</td>
<td>48.0 (12.4)</td>
<td>t = −0.05</td>
<td>0.96</td>
</tr>
<tr>
<td>Female, n (%)</td>
<td>67 (63)</td>
<td>72 (69)</td>
<td>$\chi^2 = 1.45$</td>
<td>0.23</td>
</tr>
<tr>
<td>Education, n (%)</td>
<td></td>
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<td></td>
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<tr>
<td>&lt; 10 years</td>
<td>48 (45)</td>
<td>37 (36)</td>
<td>$\chi^2 = 1.85$</td>
<td>0.40</td>
</tr>
<tr>
<td>10–12 years</td>
<td>39 (36)</td>
<td>40 (39)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13+ years</td>
<td>21 (20)</td>
<td>26 (25)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Married, n (%)</td>
<td>71 (67)</td>
<td>64 (62)</td>
<td>$\chi^2 = 0.41$</td>
<td>0.52</td>
</tr>
<tr>
<td>Employment status, n (%)</td>
<td></td>
<td></td>
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<tr>
<td>Employed</td>
<td>44 (41)</td>
<td>44 (42)</td>
<td>$\chi^2 = 0.09$</td>
<td>0.77</td>
</tr>
<tr>
<td>Retired</td>
<td>27 (25)</td>
<td>18 (17)</td>
<td>$\chi^2 = 1.78$</td>
<td>0.18</td>
</tr>
<tr>
<td>Other</td>
<td>37 (35)</td>
<td>41 (39)</td>
<td>$\chi^2 = 0.70$</td>
<td>0.40</td>
</tr>
<tr>
<td>Symptoms</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Number of SCID somatoform symptoms (actual), mean (s.d.)</td>
<td>10.0 (3.9)</td>
<td>10.6 (4.0)</td>
<td>t = −1.01</td>
<td>0.28</td>
</tr>
<tr>
<td>Symptom duration, years: mean (s.d.)</td>
<td>10.4 (5.5)</td>
<td>10.8 (5.5)</td>
<td>t = −0.45</td>
<td>0.66</td>
</tr>
</tbody>
</table>

SCID, Structured Clinical Interview for DSM-IV.

a. Two-sided tests.

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**Table 2** Outcome measure scores at baseline, end of therapy and 9-month follow-up

<table>
<thead>
<tr>
<th></th>
<th>Psychodynamic interpersonal psychotherapy (n = 107)</th>
<th>Enhanced medical care (n = 104)</th>
<th>Group difference</th>
<th>Mean (95% CI)$^a$</th>
<th>p$^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Primary outcome</strong></td>
<td></td>
<td></td>
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<tr>
<td>SF-36 Health Survey PCS</td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>29.0 (6.4)</td>
<td>29.6 (7.0)</td>
<td>−0.6 (−2.43 to 1.21)</td>
<td>0.51</td>
<td></td>
</tr>
<tr>
<td>End of therapy</td>
<td>31.3 (8.1)</td>
<td>32.6 (11.0)</td>
<td>−1.3 (−3.90 to 1.37)</td>
<td>0.70</td>
<td></td>
</tr>
<tr>
<td>Follow-up</td>
<td>34.3 (9.3)</td>
<td>31.8 (9.5)</td>
<td>2.5 (0.16 to 5.09)</td>
<td>0.001</td>
<td></td>
</tr>
<tr>
<td><strong>Secondary outcomes</strong></td>
<td></td>
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<tr>
<td>SF-36 Health Survey MCS</td>
<td></td>
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<tr>
<td>Baseline</td>
<td>36.5 (12.6)</td>
<td>36.0 (12.4)</td>
<td>0.6 (−2.85 to 3.94)</td>
<td>0.75</td>
<td></td>
</tr>
<tr>
<td>End of therapy</td>
<td>41.9 (12.0)</td>
<td>38.3 (13.2)</td>
<td>3.6 (0.20 to 7.06)</td>
<td>0.08</td>
<td></td>
</tr>
<tr>
<td>Follow-up</td>
<td>40.1 (13.3)</td>
<td>38.9 (12.7)</td>
<td>1.1 (−2.25 to 4.55)</td>
<td>0.73</td>
<td></td>
</tr>
<tr>
<td>PHQ Somatisation score</td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>Baseline</td>
<td>15.2 (5.2)</td>
<td>14.8 (4.7)</td>
<td>0.4 (−0.94 to 1.77)</td>
<td>0.48</td>
<td></td>
</tr>
<tr>
<td>End of therapy</td>
<td>13.8 (5.3)</td>
<td>14.0 (5.3)</td>
<td>−0.2 (−1.72 to 1.14)</td>
<td>0.29</td>
<td></td>
</tr>
<tr>
<td>Follow-up</td>
<td>12.7 (5.8)</td>
<td>13.8 (5.3)</td>
<td>−1.12 (−2.65 to 0.31)</td>
<td>0.31</td>
<td></td>
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<tr>
<td>PHQ Depression score</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>12.9 (5.7)</td>
<td>12.4 (5.7)</td>
<td>0.5 (−1.03 to 2.06)</td>
<td>0.47</td>
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<tr>
<td>End of therapy</td>
<td>10.3 (6.0)</td>
<td>11.1 (6.2)</td>
<td>−0.8 (−2.57 to 0.84)</td>
<td>0.10</td>
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<tr>
<td>Follow-up</td>
<td>10.5 (6.0)</td>
<td>11.2 (6.5)</td>
<td>−0.8 (−2.46 to 0.93)</td>
<td>0.08</td>
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</tr>
<tr>
<td>Health Anxiety/Whiteley Index (WI-7)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>3.9 (2.1)</td>
<td>4.1 (2.0)</td>
<td>−0.2 (−0.79 to 0.34)</td>
<td>0.42</td>
<td></td>
</tr>
<tr>
<td>End of therapy</td>
<td>3.5 (2.3)</td>
<td>3.5 (2.2)</td>
<td>0.0 (−0.62 to 0.63)</td>
<td>0.52</td>
<td></td>
</tr>
<tr>
<td>Follow-up</td>
<td>3.1 (2.3)</td>
<td>3.4 (2.3)</td>
<td>−0.3 (−0.88 to 0.31)</td>
<td>0.62</td>
<td></td>
</tr>
</tbody>
</table>

MCS, Mental Component Summary; PCS, Physical Component Summary; PHQ, Patient Health Questionnaire; SF-36, 36-item Short Form Health Questionnaire.

a. Confidence interval of the mean difference between groups (therapy group minus control group) at each assessment.
b. Mixed linear models, between-group effects for therapy controlling for study site, age, gender and education.
healthcare use ($d = 0.33$ for each dimension). Another meta-analysis of psychodynamic psychotherapy for somatic symptom disorders demonstrated medium to high short-term and long-term effects on depression, anxiety and somatic symptoms.$^8$

Somatic symptom disorders refer to chronic pain (nearly half of the studies), irritable bowel syndrome and a number of organically explained diseases. The authors stated that – despite the magnitude of the effects – the results were limited by the variable methodological quality of the identified studies and the heterogeneity in the analyses.$^8$

Participants in our study who received EMC reported small improvements in health-related quality of life, with a within-group effect size of $0.30$. This improvement is considerably greater than the control effect in similar psychotherapy trials (a mean control group effect size of $0.13$ was observed in data collected from Abbass et al)$^9$ and thus contributes to the between-group effect size being moderate as reported here. We could not demonstrate a significant improvement in depression for PIT compared with EMC. This finding suggests that the assumption that an improvement in bodily function is solely or primarily related to an improvement in depression is incorrect. Kroenke et al described the same independence between bodily distress and depression: the majority of CBT studies improved patients’ symptom severity but not their psychological distress.$^7,9$

With the Bonferroni correction the reductions in the use of psychotherapy and antidepressants in the PIT group were not significant. This finding is not surprising, as we did not design this study to demonstrate such an effect. Moreover, the data regarding use of such therapies could potentially have been biased by the fact that neither the study personnel nor the patients had been masked to randomisation, and recommendations regarding future therapeutic procedures had been a part of the protocol. However, these data indicate that the observed improvement in physical quality of life for the PIT group cannot be explained by an increase in healthcare use. This conclusion recognises that the differential effects between the PIT and EMC groups became apparent only after the end of treatment. This result may signify that one important effect of the experimental intervention was an increase in specific interpersonal and health-related self-efficacy (i.e. the ability to care for one’s own health in the context of others, both in and outside of the healthcare system). Previous studies have described comparable trajectories of change, including post-treatment improvements in psychotherapy, especially in PIT.$^7$  

We designed and carried out our study as an explanatory trial (with regard to the control of related conditions, the selection of the sample by using a widely used structured clinical interview, the delivery of standardised therapy and the strict allocation to the respective treatment condition). However, some features could allow it to be considered also as a pragmatic trial:34 we intended to examine the effect of PIT closely linked to routine clinical practice, with a group of patients characterised by some degree of heterogeneity, and we selected outcomes that represented a broad range of health gains.

**Strengths and limitations**

Strengths of the study include the rigorously controlled and monitored trial, its multisite setting, the relatively large sample size and moderate withdrawal rate, and the thorough joint pre-trial development of its manual among all principal investigators. Our intention was to address a population of at least moderately impaired patients with multiple medically unexplained bodily symptoms without confinement to a particular functional syndrome, presenting at somatic out-patient departments. For this purpose we chose the diagnostic category of multisomatoform disorder, which has been used occasionally in other trials, such as that by Dickinson et al.35 The advantage of using this diagnostic construct as a common point of reference across single functional somatic symptoms and syndromes lies in its clinical relevance and its advanced integrative approach.$^8$

Several limitations of our study, which are mainly related to the setting of the trial, the selection of patients and the comparability of the control intervention, should be discussed. The selection of the participating centres was not random, and the willingness to cooperate may have biased the selection. Accordingly the sample of out-patient departments cannot – in a strict sense – be deemed to be representative of multisomatoform disorders as a whole, as certain forms of medically unexplained bodily symptoms are likely to be treated in disciplines not represented in our study. The inclusion of patients with pain as

### Table 3 Use of healthcare and antidepressant medication at baseline, end of therapy and 9-month follow-up

<table>
<thead>
<tr>
<th></th>
<th>Psychodynamic interpersonal psychotherapy ($n = 107$)</th>
<th>Enhanced medical care ($n = 104$)</th>
<th>Comparison</th>
<th>$P$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of healthcare visits, mean (s.d.)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primary care</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>3.2 (3.3)</td>
<td>3.9 (3.6)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>End of therapy</td>
<td>3.9 (5.1)</td>
<td>3.4 (3.8)</td>
<td>0.22$a$</td>
<td>0.68</td>
</tr>
<tr>
<td>Follow-up</td>
<td>2.5 (2.6)</td>
<td>3.5 (3.6)</td>
<td>0.75$a$</td>
<td>0.39</td>
</tr>
<tr>
<td>Specialist consultation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>7.1 (6.0)</td>
<td>6.9 (7.6)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>End of therapy</td>
<td>7.4 (7.4)</td>
<td>5.3 (6.6)</td>
<td>3.25$a$</td>
<td>0.11</td>
</tr>
<tr>
<td>Follow-up</td>
<td>7.4 (8.8)</td>
<td>7.5 (7.2)</td>
<td>0.15$a$</td>
<td>0.73</td>
</tr>
<tr>
<td>Concurrent antidepressant medication, n (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>52 (49)</td>
<td>46 (44)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>End of therapy</td>
<td>41 (38)</td>
<td>44 (42)</td>
<td>3.57$b$</td>
<td>0.17</td>
</tr>
<tr>
<td>Follow-up</td>
<td>39 (36)</td>
<td>48 (46)</td>
<td>6.24$b$</td>
<td>0.04</td>
</tr>
<tr>
<td>Psychotherapy at follow-up, n (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>End of therapy</td>
<td>6 (6)</td>
<td>6 (6)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Follow-up$^c$</td>
<td>3 (3)</td>
<td>12 (12)</td>
<td>$−1.98^c$</td>
<td>0.05</td>
</tr>
</tbody>
</table>

*a. Mixed models, between-group comparison of differences between baseline ($t_1$) and end of therapy ($t_2$), and between $t_1$ and 9-month follow-up ($t_3$) respectively, controlling for initial frequency of visits and study site.

$b. Chi$-squared test.

$c. No other psychotherapy allowed during treatment.

$d. Mann–Whitney U$-test.
the presenting symptom limits our ability to generalise these results to other somatoform disorders – although pain is by far the most frequent type of symptom in these disorders.

Finally, another limitation concerns the comparability of the two interventions: we are unable to determine whether the observed treatment effects in the experimental and control groups were due to the treatment itself or to non-specific effects related to the different amount of time spent with patients. Sumathipala et al demonstrated that structured care is as effective as CBT in patients with less severe somatic symptoms, both therapies consisting of six sessions of 30 min each. However, two arguments speak against this possibility: first, our differential treatment effect became apparent only after $t_2$ and second, in the PIT group the treatment effect was positively associated with the extent and intensity of interpersonal elements as measured with the adherence rating scale. We considered using a waiting-list condition to represent the natural course of the disease as an alternative to our control group. The decision to offer EMC was made for reasons that were both ethical (patients should receive the best available care, and EMC follows valid evidence-based guidelines) and practical (to ensure study participation).

Future research

Three immediate questions arise from the results of this study. First, does PIT produce improvements in the quality of life of patients with multisomatoform disorder comparable with those provided by CBT? Second – and perhaps of greater concern to health service planners – how does PIT perform compared with the care that patients are likely to receive in a real-world setting? Third, given that two studies have already shown that PIT – unlike other approaches such as reattribution – is one of few psychological interventions to demonstrate real cost savings, can we demonstrate such an effect for our study as well? So cost-effectiveness and cost-utility analyses for PIT compared with EMC in patients with multisomatoform disorder should be considered.

Implications of the study

Patients with severely disabling and chronic bodily distress who seek help from different somatic out-patient departments can be helped using PIT. This therapy also allowed patients to increase their physical functioning after the intervention. Enhanced medical care also provides relief for severely ill patients, but its effects are not of comparable sustainability. We need more information about this intervention’s long-term sustainability and differential effects (i.e. those related to gender, individual psychological disease models and physiological data). Furthermore, we need to evaluate the benefits of this therapy in relationship to its potential cost savings. Knowing this information would enable us to better tailor our intervention and to increase its impact.

References


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Psychodynamic interpersonal therapy in multisomatoform disorder


Brief psychodynamic interpersonal psychotherapy for patients with multisomatoform disorder: randomised controlled trial
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