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Rask, Mette Trøllund; Ørnbøl, Eva; Rosendal, Marianne; Fink, Per Klausen

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Long-Term Outcome of Bodily Distress Syndrome in Primary Care: A Follow-Up Study on Health Care Costs, Work Disability, and Self-Rated Health

Mette Trollund Rask, PhD, Eva Ørnbøl, MSc, Marianne Rosendal, PhD, and Per Fink, DMSc, MD

ABSTRACT

Objective: The upcoming International Classification of Diseases, 11th Revision for primary care use suggests inclusion of a new diagnostic construct, bodily (di)stress syndrome (BDS), for individuals with medically unexplained symptoms. We aimed to explore the long-term outcome of BDS in health care costs, work disability, and self-rated health.

Methods: Consecutive patients consulting their family physician for a new health problem were screened for physical and mental symptoms by questionnaires (n = 1785). A stratified subsample was examined with a standardized diagnostic interview (n = 701). Patients with single-organ BDS (n = 124) and multiorgan BDS (n = 35), and a reference group with a family physician–verified medical condition (n = 880) were included. All included patients completed a questionnaire at 3, 12, and 24 months of follow-up. Register data on health care costs and work disability were obtained after 2 and 10 years of follow-up, respectively.

Results: Patients with BDS displayed poorer self-rated health and higher illness worry at index consultation and throughout follow-up than the reference group (p ≤ .001). The annual health care costs were higher in the BDS groups (2270 USD and 4066 USD) than in the reference group (1392 USD) (achieved significance level (ASL) ≤ .001). Both BDS groups had higher risk of sick leave during the first 2 years of follow-up (RRsingle-organ BDS = 3.0; 95% confidence interval [CI] = 1.8–5.0; RRmultiorgan BDS = 3.4; 95% CI = 1.5–7.5) and substantially higher risk of newly awarded disability pension than the reference group (HRsingle-organ BDS = 4.9; 95% CI = 2.8–8.4; HRmultiorgan BDS = 8.7; 95% CI = 3.7–20.7).

Conclusions: Patients with BDS have poor long-term outcome of health care costs, work disability, and subjective suffering. These findings stress the need for adequate recognition and management of BDS.

Key words: somatoform disorders, functional somatic syndromes (not MeSH), sick leave, Health care costs, primary health care.

INTRODUCTION

Symptoms that are not attributable to any conventionally defined disease are highly prevalent across all medical settings. Such medically unexplained or functional somatic symptoms represent a spectrum ranging from normal or self-limiting symptoms to severe and persistent conditions (1). Patients with persistent functional somatic symptoms constitute a vast challenge to both the health care system and the social security system; they are characterized by high rates of anxiety and depression (2,3), they are frequent attenders to health care services and contribute to high costs for Primary Health Care, MCS = mental component summary, MCS = mental component summary, PCS = physical component summary, SCL = symptom check list, SCAN = Schedules for Clinical Assessment in Neuropsychiatry, SCID = Structured Clinical Interview for DSM disorders, SF-36 = Short Form Health Survey, SSD = Somatic symptom disorder, WHO = World Health Organization, WHO-CIDI = WHO Composite International Diagnostic Interview

ASL = achieved significance level, BCa = bias-corrected and accelerated, BDS = bodily distress syndrome, CAGE = Cutting down, Annoyance by criticism, Guilty feeling, Eye openers Questionnaire, DREAM = Danish register for evaluation of marginalization, DRG = diagnosis-related group, DSM = Diagnostic and Statistical Manual of Mental Disorders, ES = effect sizes, FIP = family physician, GLM = generalized linear model, ICD-11-PHC = the International Classification of Diseases for Primary Health Care, SF-36 = Short Form Health Survey, SSD = Somatic symptom disorder, WHO = World Health Organization, WHO-CIDI = WHO Composite International Diagnostic Interview

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health care costs (4,5), and they may have an increased risk of sickness absence and work disability (6–9).

Previous studies support the evidence for an unfavorable outcome of conditions involving persistent functional somatic symptoms, but these studies are largely cross-sectional or based on self-reported questionnaires and/or less well-defined diagnostic constructs. Thus, we lack evidence on the long-term consequences of persistent functional somatic symptoms based on well-defined diagnostic criteria.

The field of persistent functional somatic symptoms is characterized by a lack of consistent terminology and overlapping diagnoses. Recently, a new concept of bodily distress syndrome (BDS) was introduced. This concept derives from empirical studies and seems to capture most patients with somatoform disorders and functional somatic syndromes (e.g., fibromyalgia, chronic fatigue syndrome, and irritable bowel syndrome) (10,11). The BDS builds on specific physical symptom patterns representing 4 symptom groups: cardiopulmonary, gastrointestinal, musculoskeletal, and general symptoms group. Bodily distress syndrome can be further divided into 2 severity groups: severe multiorgan type and less severe single-organ type (Fig. 1) (11). This new unifying concept has been suggested to replace diagnoses of somatoform disorders and has been incorporated into the current draft of the International Classification of Diseases for Primary Health Care (ICD-11-PHC) (12,13).

As BDS is a newly introduced diagnostic concept, only little knowledge exists on the prognosis of patients with BDS. Therefore, we performed a longitudinal study aiming to determine the long-term outcome of BDS in health care costs, work disability, and self-rated physical and mental health (including illness worry). The original patient sample in which the diagnostic criteria for BDS were developed was used in this study (11).

METHODS

The study was a questionnaire- and register-based follow-up study of a cohort established in 2000 for the Functional Illness in Primary Care (FIP) study comprising data on adult patients who consulted their family physician (FP) for a new health problem. The FIP study was an intervention study focusing on the effect of an educational program for training FPs in recognition and management of patients with functional symptoms (14). The FIP study was carried out in a 2-phase design consisting of a patient screening questionnaire and a standardized psychiatric interview. The FIP study design and the recruitment of patients have been described extensively elsewhere (15). The study was approved by the Science Ethics Committee in the former County of Aarhus, the Scientific Research Evaluation Committee of the Danish College of General Practitioners, and the Danish Data Protection Agency.

Participants and Setting

In the FIP study, 1785 consecutive patients aged 18 to 65 years attending their family physician (n = 38) for a new health problem during a 3-week period in March and April 2000 gave informed consent to participate and were included. Patients who were of non-Scandinavian origin, who were severely ill or demented, or who were not enlisted with the participating FPs were excluded from participation (15).

1. The patient is moderately to severely impaired by the stated number of symptoms from at least one of the following symptom groups:

   ≥3 Cardiopulmonary symptoms
   - Palpitations/heart pounding, precordial discomfort, breathlessness without exertion,
   - Hyperventilation, hot or cold sweats, dry mouth, trembling or shaking, churning in stomach/”butterflies,” flushing or blushing

   ≥3 Gastrointestinal symptoms
   - Abdominal pains, frequent loose bowel movements, feeling bloated/full of gas/distended,
   - Regurgitations, diarrhea, nausea, burning sensation in chest or epigastrium, constipation,
   - Vomiting (other than during pregnancy)

   ≥3 Musculoskeletal symptoms
   - Pains in arms or legs, muscular aches or pains, pains in the joints, feelings of pain or localized weakness, backache, pain moving from one place to another, unpleasant numbness or tingling sensations

   ≥3 General symptoms
   - Concentration difficulties, impairment of memory, excessive fatigue, headache, dizziness

   ≥4 Symptoms from any of the above symptom groups

2. Illness duration ≥ 6 months

3. Relevant differential diagnoses have been ruled out

Bodily Distress Syndrome (BDS)

Single-organ type
- Fulfilling criteria for 1-3 symptom groups

Multi-organ type
- Fulfilling criteria for at least 4 symptom groups

FIGURE 1. Diagnostic criteria for BDS.
Study Design and Procedure

The patients completed a screening questionnaire in the FP's waiting room just before entering the consultation. This questionnaire included the Symptom Check List (SCL-8), which assesses anxiety and depression; the somatization subscale of the SCL-90, which screens for 12 common physical symptoms, the Whitely Index (Whitely-7), which assesses illness worry, and the 4-item Cutting down, Annoyance by criticism, Guilty feeling, Eye openers (CAGE) screening questionnaire for alcohol abuse (16). Included patients also completed the Medical Outcome Study's Short Form Health Survey (SF-36), which assesses physical and mental health (17), and sociodemographic data were obtained. A detailed description of the screening questionnaire is provided elsewhere (16).

A stratified sample consisting of a random selection of one ninth of the patients and all patients with a high score on the screening questionnaires ($n = 894$) was invited for a psychiatric research interview (Fig. 2). The stratified sampling procedure was used solely for the initial identification of patients with high likelihood of being cases to help reduce the number of noncase interviews. We did not use the information on distress from the questionnaires for categorization of patients or diagnostic purposes; categorization was based on the results of the psychiatric interview. A total of 701 patients (78.4%) accepted to participate in the psychiatric research interview. Patients who had a low score on the screening questionnaire, who were younger, or who were males were more likely to decline participation than other patients (15). For most patients, the interview was performed within a week after the initial contact.

Psychiatric Research Interview

The interview was based on the Schedules for Clinical Assessment in Neuropsychiatry (SCAN), version 2.1. (18). The SCAN is endorsed by the World Health Organization (WHO) and is a standardized semistructured interview performed by trained medical doctors and covering all types of mental disorders, including a separate section that screens for a wide range of physical symptoms. This section allows the rater to assess whether present symptoms are explained by a medical condition or rather should be seen as functional symptoms, and whether these symptoms are considered to be disturbing for the patient or have been an issue receiving medical attention. Based on the physical symptom screening, diagnoses for a variety of functional somatic syndromes (e.g., chronic fatigue syndrome and fibromyalgia) were made.

![Flow chart](image-url)

**FIGURE 2.** Flow chart. *FPs stated a well-defined medical condition as the main problem in 1009 patients; 296 of these were SCAN interviewed of which 35 fulfilled the criteria for single-organ BDS and 7 fulfilled the criteria for multiorgan BDS and were classified as such.
can be generated. The interviews were performed by 6 physicians who had been certified at the WHO SCAN training center in Aarhus and who had psychiatric, medical, and surgical residency. The interrater reliability between the 6 interviewers were found to be high (kappa = 0.88) for the ICD-10 somatoform disorders and other psychiatric diagnoses (15).

FP Assessment
Immediately after the index consultation, the FP completed a questionnaire inquiring on the presence of chronic physical disease/psychiatric disorders, including categorization of the patient's main health problem as either “Well-defined medical condition” (n = 1009), “Probably well-defined medical condition” (n = 395), “Medically unexplained symptoms” (n = 229), “Psychiatric disorder with physical manifestations” (n = 95), or “No physical health complaints” (n = 39). Family physician's rating of main problem was missing for 18 patients.

Patient Grouping
The SCAN interviews were used to generate a BDS single-organ group and a BDS multiorgan group according to the BDS criteria (11,12). We compared these 2 groups to a reference group of patients with a well-defined medical condition (as rated by their FP). In total, 29% of the 1009 patients who attended due to a well-defined medical condition were SCAN interviewed; 35 patients were found to meet the criteria for single-organ BDS, whereas 7 patients were found to meet the criteria for multiorgan BDS. These 42 patients were included only in one of the BDS groups and excluded from the medical condition group. Ultimately, 880 patients were included in the medical condition group, 124 in the BDS single-organ group, and 35 in the BDS multiorgan group (Fig. 2).

Follow-Up
The patients were asked to complete a mailed questionnaire at 3, 12, and 24 months after the index consultation, including scales measuring self-rated health and illness worry. Register data on health care costs was retrieved for a period of 3 years before index consultation and throughout 2 years after, whereas register data on sick leave and disability were obtained for the 10 years following the index consultation.

Self-Rated Health and Illness Worry
The Physical Component Summary (PCS) and the Mental Component Summary (MCS) from the SF-36 were used as measures of self-rated health (19), and the Whiteley-7 scale as a measure of illness worry (20,21).

Healthcare Costs
The Danish health care system is almost entirely tax financed, and Danish residents are generally offered medical care free of charge. In this study, the costs of primary care, secondary care, and prescribed medication were obtained from the National Health Service Register in the former County of Aarhus. We obtained data from 3 years before index consultation through 2 years after, but the data on prescribed medicine were limited to include only the 6 months immediately before the index consultation due to legal restrictions on registration of medicine use in Denmark. The analyses did not include costs of laboratory tests and x-rays requested in primary care. All costs for inpatient and daytime admissions and outpatient and emergency ward contacts were extracted from the Danish National Patient Register and the Danish Psychiatric Central Register. Nonpsychiatric costs were calculated as diagnosis-related group case-mix prices by the Diagnosis-Related Group pricing office of the Danish Health and Medicines Authority (2004 fixed prices). Psychiatric hospital care costs were calculated from the average 2004 fixed prices for hospital bed days and outpatient contacts with the finance administration team of the Psychiatric Hospital in Aarhus. All costs were adjusted for time at risk, and the object for analysis was cost per year. Index consultation was included in the 2 years of follow-up.

Work Disability
Danish citizens who have received social benefits or any other welfare payments are registered in the Danish Register for Evaluation of Marginalization (DREAM) (22). Social transfers recorded in DREAM represent 5 categories: benefits to otherwise self-supporting individuals (e.g., statutory maternity pay), benefits related to the labor market (e.g., unemployment benefit or social assistance), temporary health-related benefits (sickness benefit and vocational rehabilitation benefit), permanent health-related benefits (full and partial disability pension), and public (old-age or early) retirement pension. Furthermore, death and emigration of registered individuals are recorded in DREAM.

Danish law stipulates that sickness benefit can be granted for a maximum of 12 months, after which the individual must either return to work, will be eligible for social assistance, or may be awarded disability pension by the municipal authorities. Until a recent restriction, individuals aged 18 to 65 years with permanently reduced work ability were eligible for disability pension. Partial disability pension is granted to individuals with partial loss of working capacity on a permanent basis, and “flexible working” may be arranged, whereas full disability pension is a permanent departure from the labor market.

As a measure of work disability, data on temporary (sickness benefit and vocational rehabilitation) and permanent (full and partial disability pension) health-related benefits were retrieved from DREAM for the 3 patient groups. Due to the registration procedure in DREAM, health-related benefits were recorded in weeks.

Data Analysis
Descriptive statistics were used to summarize patients’ characteristics at index consultation. The SF-36 PCS and MCS scores were calculated according to the validated Danish version of the SF-36, with higher scores expressing better health (23). The Whiteley-7 was transformed into a scale ranging from zero to 100 by the following expression: (patient raw score – lowest possible score) / (highest possible score – lowest possible score) × 100. Group comparisons were performed by χ² test for categorical data, Student t test for normally distributed data and Mann-Whitney U test for non-normally distributed data.

To further analyze the development in PCS, MCS, and Whiteley-7 scores over time within and between groups, we estimated mixed models with random intercept for each of these outcomes. The general shape of the models is a group-specific level at index consultation and a group-specific level, that is, one level, at the remaining time points. This was modeled through 2 variables (time and group) with potential interaction effects. The models were graphically depicted, and mean differences with corresponding 95% confidence intervals (CIs) and effect sizes (ESs) were calculated as measures of within-group changes.

We accounted for skewed health care costs with an excess of zeros by estimating sample means of health care costs with bias-corrected and accelerated 95% CIs. Tests of equality of health care costs for patients with a medical condition and for patients with BDS were done by computing the bootstrap test statistic achieved significance level (ASL) based on 1000 replications (24).

Weekly sick leave status was recorded for each patient throughout the follow-up period, excluding weeks of permanent disability pension, age-related retirement, emigration, or death (missing values). To compare the risk of sick leave between patient groups, we applied a generalized linear model from the binomial family using log-link. Risk ratios (RRs) were used as a measure of association. Corresponding 95% CI were assessed by performing cluster-robust variance estimation to account for the expected dependency of awarded temporary health-related benefits in different weeks for the same patient (25).

The Cox proportional hazard model was used to compare the risk of incident award of full or partial disability pension during 10 years of follow-up among the 3 patient groups. Patient age was used as time scale and was hence appropriately adjusted for. Patients entered the study at the time of
the index consultation (delayed entry), that is, they were not observed until the index consultation, and they were censored at the time of death, emigration, or public retirement. The proportional-hazards assumption was graphically assessed using log-log plots. Hazard ratios (HRs) with corresponding 95% CI were used as a measure of association.

In all analyses, patients with a medical condition constituted the reference group. Crude estimates were presented as were estimates adjusted for patient age, sex, chronic illness, comorbid major depressive episode, anxiety disorder, and intervention. Two-sided \( p \) values < .05 were considered to be statistically significant, except for analyses of health care costs in which a Bonferroni correction was applied, and the level of significance was set at \( p < .0025 \) (2 group comparisons on 5 different measures of costs at 2 different time points). Analyses were conducted by Stata statistical software, version 11.

RESULTS

Patients' Characteristics

Patients in the 2 BDS groups were slightly older, they were more often females, and they were less prone to have finished

<table>
<thead>
<tr>
<th>TABLE 1. Patients' Baseline Characteristics(^a)</th>
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<tbody>
<tr>
<td>Bodily Distress Syndrome</td>
</tr>
<tr>
<td>---------------------------</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Age, M (SD)</td>
</tr>
<tr>
<td>Female, (n(%))</td>
</tr>
<tr>
<td>Married/with a partner, (n(%))</td>
</tr>
<tr>
<td>Vocational training, (n(%))</td>
</tr>
<tr>
<td>Unskilled</td>
</tr>
<tr>
<td>Skilled</td>
</tr>
<tr>
<td>Higher education (\leq 4)</td>
</tr>
<tr>
<td>Higher education &gt; 4 years</td>
</tr>
<tr>
<td>Other education</td>
</tr>
<tr>
<td>Labor market dropout, (n(%))</td>
</tr>
<tr>
<td>Available for labor market</td>
</tr>
<tr>
<td>Partial or full disability pension</td>
</tr>
<tr>
<td>Public retirement pension</td>
</tr>
<tr>
<td>Medical condition according to FP, (n(%))</td>
</tr>
<tr>
<td>Definitely</td>
</tr>
<tr>
<td>Probably</td>
</tr>
<tr>
<td>Chronic illness according to FP</td>
</tr>
<tr>
<td>Psychiatric comorbidity, (n(%))(^b)</td>
</tr>
<tr>
<td>Major depressive episode</td>
</tr>
<tr>
<td>Anxiety disorder</td>
</tr>
<tr>
<td>Somatoform disorder (DSM-IV), (n(%))(^b)</td>
</tr>
<tr>
<td>Functional somatic syndromes, (n(%))(^b,c)</td>
</tr>
<tr>
<td>Chronic fatigue syndrome</td>
</tr>
<tr>
<td>Fibromyalgia</td>
</tr>
<tr>
<td>Pain syndrome</td>
</tr>
<tr>
<td>Irritable bowel syndrome</td>
</tr>
<tr>
<td>Noncardiac chest pain</td>
</tr>
<tr>
<td>Hyperventilation syndrome</td>
</tr>
<tr>
<td>At least one of the above syndromes</td>
</tr>
</tbody>
</table>

\(^a\)Missing data: marital status, 133; vocational training, 186; labor market dropout, 1; chronic illness according to FP, 4; medical condition according to FP, 1.

\(^b\)Non–SCAN-interviewed patients in the medical condition group were considered not to have psychiatric comorbidity, somatoform disorder, or any functional somatic syndrome, as they did not have a high score on the screening questionnaire.

\(^c\)Diagnostic criteria and algorithms for the 6 functional somatic syndromes followed Fink and Schröder (10).
a higher educational program compared to patients in the medical condition group. Furthermore, patients with BDS had more frequently been granted partial or full disability pension (20.2% and 34.3% vs 3.6%) and thus were less likely to be available for the labor market at index consultation than patients in the medical condition group (Table 1).

The FPs found more patients in the BDS groups to have a chronic illness compared to the medical condition group. Furthermore, major depressive episode and anxiety disorder (as based on the SCAN interview) were more frequent in the BDS groups. Most of the patients with BDS had at least one functional somatic syndrome: 82.3% of single-organ BDS and 94.3% of multiorgan BDS (Table 1).

Self-Rated Health and Illness Worry
Self-rated physical health, mental health, and illness worry are displayed in Table 2. For all 3 scales and at all time points (index and 3-, 12-, and 24-month follow-ups), completers and noncompleters were equally distributed across the 3 patient groups. We did not find any differences in the baseline scores on PCS, MCS, or Whiteley-7 when comparing patients who completed the questionnaires with noncompleters or patients lost to follow-up at any follow-up time points (data not shown).

At baseline and all follow-up time points, patients with BDS had lower self-rated physical and mental health and higher illness worry than patients with a well-defined medical condition (Table 2). Within each group, baseline scores were compared with follow-up scores adjusted for age, sex, chronic illness according to the FP, anxiety disorder, major depressive episode, and intervention. From baseline to 24 months follow-up, physical health improved slightly in the multiorgan BDS group (mean difference = 3.9; 95% CI = 0.3–7.5) and in the medical condition group (mean difference = 3.5; 95% CI = 2.2–4.8). Mental health improved only in the single-organ BDS group (mean difference = 4.0; 95% CI = 1.1–7.0), whereas illness worry decreased in both the single-organ BDS group (mean difference = −9.6; 95% CI = −13.6 to −5.7), and the medical condition group (mean difference = −8.0; 95% CI = −9.8 to −6.2). The improvements in self-rated health were rather small (ES = 0.29–0.39), although statistically significant, and only reduced illness worry may be of clinical relevance (ES = 0.70–0.79) (Fig. S1, Supplemental Digital Content 1, http://links.lww.com/PSYMED/A329).

Health Care Costs
The total annual healthcare costs incurred by the 2 BDS groups were higher than the costs incurred by the medical condition group (follow-up estimates: single-organ BDS: mean = 2270 USD, ASL = 0.001; multiorgan BDS: mean = 4066 USD, ASL < 0.001) (Table 3). Patients with multiorgan BDS generally displayed higher health care costs across all medical settings and types of services (except for psychiatric care) during follow-up, whereas patients with single-organ BDS incurred higher costs only in connection with primary care consultations and prescribed medications. The follow-up costs incurred by general hospital care accounted for 57%, 51%, and 73% of the total costs in the groups with single-organ BDS, multiorgan BDS and medical condition, respectively. Psychiatric care costs accounted for 22% of the total health care costs in the group with multiorgan BDS (service use: n = 8 (23%)), only 3% in patients with single-organ BDS (service use: n = 8 (6%)), and 4% in patients with a well-defined medical condition (service use: n = 7 (<1%)).

Work Disability
The average risk of sick leave in the first 3 months after the index consultation was higher in patients with single-organ BDS (RR = 4.9; 95% CI = 2.7–8.9) and multiorgan BDS (RR = 6.7; 95% CI = 3.1–14.4) than in patients with a well-defined medical condition (Table 4). The risks decreased over time; after 10 years of follow-up, no statistically significant differences were observed between the groups.

Marginalization in terms of permanently reduced or loss of work ability seemed relatively stable over time in the 3 groups (Fig. 3). A higher proportion of patients with BDS received disability pension, most pronounced in the multiorgan BDS group, compared to patients with a well-defined medical condition. It should be noted that migration between categories was evident, for example, patients receiving disability pension (full or partial) were transferred to public retirement no later than the age of 65 years, whereas others who were available for the labor market at the index consultation were awarded disability pension during the follow-up period. During the 10 years of follow-up, patients with BDS were more likely to be excluded from the labor force due to ill health; the multiorgan BDS group was 8 times as likely (HR = 8.7; 95% CI = 3.1–20.7) and the single-organ BDS group 4 times as likely (HR = 4.9; 95% CI = 2.8–8.4) to receive a new award of disability pension than patients with a well-defined medical condition (Table 5).

DISCUSSION
This study suggests that BDS (both single-organ and multiorgan) is a disabling condition with substantial long-term impact on both the individual and the society. Bodily distress syndrome was associated with lower self-rated health, higher illness worry, higher health care costs, and higher risk of work disability throughout the follow-up period.

Physical component summary scores of approximately 41 and MCS scores of approximately 47 have been demonstrated in samples of primary care patients who meet the criteria for either somatization disorder, abridged somatization disorder, or multisomatoform disorder (26). These
<table>
<thead>
<tr>
<th></th>
<th>Physical component summary</th>
<th>Mental component summary</th>
<th>Whiteley-7</th>
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<tbody>
<tr>
<td></td>
<td>SF-36</td>
<td>BDS Multiorgan Type (n = 35)</td>
<td>Well-Defined Medical Condition (n = 880)</td>
</tr>
<tr>
<td></td>
<td>n</td>
<td>M</td>
<td>SD</td>
</tr>
<tr>
<td>Baseline</td>
<td>109</td>
<td>40.4</td>
<td>10.9</td>
</tr>
<tr>
<td>3 months</td>
<td>91</td>
<td>41.3</td>
<td>11.9</td>
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<tr>
<td>12 months</td>
<td>88</td>
<td>42.9</td>
<td>11.8</td>
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<tr>
<td>24 months</td>
<td>80</td>
<td>40.5</td>
<td>12.1</td>
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<tr>
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<td>SD, standard deviation; Q1–Q3, interquartile range.</td>
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*Pairwise comparisons (2-sample t test or Mann-Whitney U test) of single-organ BDS vs well-defined medical condition and multiorgan BDS vs well-defined medical condition showed statistically significant differences (p = <.0001) on both SF-36 and Whiteley-7 at all time points (at index consultation and at 3, 12, and 24 months of follow-up).
scores are comparable to the scores found for patients with single-organ BDS, whereas patients with multiorgan BDS seemed to report considerably more impaired physical and mental health status. The degree of impairment in both BDS groups was remarkably higher than previously found for chronic physical conditions such as arthritis, chronic lung disease, diabetes, and chronic heart disease in the general population (27).

Illness worry and the related interaction with somatic symptom burden have been found to predict health care use (28). Our data may indicate that patients with a medical condition and, to some degree, patients with single-organ BDS, feel reassured after consulting their FP as we observed a decrease in illness worry right after the index consultation in both groups. We found patients with multiorgan BDS to remain highly worried over time. On the one hand, changes in illness worry may reflect changes in symptom status and self-evaluated health. On the other hand, our results may reflect previous findings that indicate that FPs tend to feel more comfortable and more successful with the task of explaining and reassuring patients with medical conditions and less severe functional symptoms, whereas patients with severe and persistent conditions are often found to be burdensome and difficult to manage (29–31).

We found patients with multiorgan BDS to incur higher health care costs across all medical settings (except for psychiatric care) during follow-up, whereas higher costs in patients with single-organ BDS were primarily due to use of primary care and prescribed medications. There may be several explanations for these findings. First, our findings may reflect the gradient in severity between single-organ BDS and multiorgan BDS and different needs of care. Second, our findings may indicate that FPs are better capable of handling patients with single-organ BDS, but need specialist support to manage patients with multiorgan BDS in line with the clinical recommendations (32). The results on psychiatric care costs need cautious interpretation, as very few individuals requested psychiatric services. In general, our findings are consistent with the existing literature, as both former population-based studies and clinical studies demonstrate high health care use and increased costs for patients with functional somatic symptoms or somatoform disorders (4,33,34).

Our study results indicate that BDS is strongly related to work disability. Several studies have found an increased risk of sick leave and/or disability in patients with a high functional symptom burden or functional somatic syndromes (7,35–37). We have previously demonstrated a

| TABLE 3. Annual Health Care Costs Before and After Index Consultation in USD | Bodily Distress Syndrome | Test for Equality of Means |
| Health Service Costs | Single-Organ Type (n = 124) (a) | Multiorgan Type (n = 35) (b) | Well-Defined Medical Condition (n = 880) (c) | a vs c | b vs c |
| Health Service Costs | Mean (Bca 95% CI) | Mean (Bca 95% CI) | Mean (Bca 95% CI) | ASL | ASL |
| Overall Before | 2668 (2043–3700) | 3542 (2390–5793) | 1245 (1095–1436) | <0.001 | 0.001 |
| After | 2270 (1771–3200) | 4066 (2927–7139) | 1392 (1160–1849) | 0.001 | <0.001 |
| Primary care Before | 475 (380–753) | 436 (350–587) | 168 (153–187) | <0.001 | <0.001 |
| After | 498 (402–654) | 529 (414–689) | 212 (196–233) | <0.001 | <0.001 |
| Medicine reimbursement b | Before | 504 (361–716) | 753 (434–1494) | 184 (153–224) | <0.001 | 0.002 |
| After | 415 (289–619) | 581 (354–1043) | 110 (89–137) | <0.001 | <0.001 |
| General hospital care Before | 1425 (1059–1942) | 1711 (1082–2674) | 865 (738–1022) | 0.001 | 0.006 |
| After | 1291 (921–2028) | 2065 (1467–2741) | 1012 (822–1527) | 0.15 | 0.001 |
| Psychiatric care Before | 263 (26–1429) | 642 (98–2033) | 28 (8–95) | 0.10 | 0.019 |
| After | 67 (26–174) | 891 (152–3757) | 59 (3–278) | 0.51 | 0.038 |

BCa, Bias-corrected and accelerated; ASL, achieved significance level.

a Except for medicine reimbursement, estimates are based on the 3 years before index consultation through 2 years after; “after” period includes the month of the index consultation: 100 USD = 133.603 EUR.

b Only 6 months before the index consultation (converted to 1-year annual costs) through 2 years after.
A strong relationship between somatoform disorders and disability pension, a relationship that has also been found in prospective studies of fibromyalgia-associated symptoms and the number of pain sites (38–40). After 10 years of follow-up, we could no longer demonstrate any differences in the risk of sick leave between our patient groups. This may be explained by the higher proportion of patients with BDS who had been granted disability pension. As disability pension is a permanent departure from the labor market, patients who already receive disability pension are not at risk of receiving sick leave benefits. Patients with disabilities and restricted work ability may not necessarily receive health-related benefits, but they may instead be granted unemployment benefit or social assistance. A Dutch study found that sick-listed patients with a high somatic symptom burden and decreased functioning were at higher risk of redundancy (37). As a consequence, our analyses of work disability may only partly describe the seriousness of the problem. Thus, the demonstrated group differences may be of an even greater magnitude.

Somatic symptom disorder (SSD) has been introduced in the Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition (41). The main criterion of the diagnosis is that the patient must present at least one distressing somatic symptom (criteria A). The symptom can be of any origin, that is, even a symptom caused by cancer. In addition, the patient must have excessive thoughts, feelings, or behaviors (e.g., health anxiety) related to the symptom (criteria B) (42). As a consequence, the SSD seems to be a new permutation of the DSM-IV hypochondria diagnosis rather than a replacement of the DSM-IV somatization disorder and related diagnoses. The SSD does not use somatic symptom characteristics (e.g., pattern, number, character, or type) for diagnostic purposes. Consequently, the SSD is a very different construct than the BDS; the BDS is solely defined by somatic symptom patterns, and no emotional or behavioral symptoms are needed for the diagnosis (but they may be important for treatment). The data material used in this study formed part of the study in which the BDS diagnostic construct was developed. Back then, we included only medically unexplained symptoms for diagnostic purposes. However, our 10-year clinical experience with BDS and the results from a new study in primary care indicate that it is obsolete to seek to define each symptom as either medically unexplained or not (43). Rather, the FP should ask for symptoms of BDS to identify the unique symptom patterns or illness picture described in the diagnostic criteria (Fig. 1). As always in medical practice, the FP must also exclude differential diagnoses, that is, other conditions that may present with a similar symptom pattern.

**Strengths and Limitations**

A major strength of our study is that we had complete follow-up of outcomes related to health care costs, sick
leave, and work disability (except for emigrated patients), as our data were obtained from Danish national registers. Danish Register for Evaluation of Marginalization has formerly been found to provide valid and high-quality data on disability pension and sickness benefits (22,44), and the completeness of the registers used for the calculations of health care costs are generally considered to be high because the data are continuously updated by state authorities and are used for reimbursement purposes (45,46). Moreover, our use of register data reduced the risk of recall bias, which may be present in studies relying on self-reported measures of health care use and sick leave. Due to the longitudinal design, a rather high proportion of patients were lost to follow-up on the self-reported outcomes. However, as noncompleters were equally distributed across the 3 patient groups at all follow-up points and that no differences between completers and noncompleters were detected in baseline scores, we expect nonresponse to have had no major influence on results.

Another major strength of the study was that the BDS groups were generated from acknowledged standardized diagnostic interviews. Instead of relying on subjective reports by the FPs or the patients on functional symptoms (47), trained physicians performed a systematic screening for a high number of symptoms and rated these as functional symptoms according to predefined criteria. Compared to other standardized psychiatric research interviews, for example, the Structured Clinical Interview for DSM disorders (SCID) or the WHO Composite International Diagnostic Interview (WHO-CIDI), the SCAN interview is better suited for the research purposes. The SCID and WHO-CIDI are both diagnosis focused and include only symptoms that are relevant for DSM-IV or ICD-10 psychiatric diagnoses (48,49), whereas the SCAN interview is symptom driven with a bottom-up approach focusing on psychopathology rather than on specific diagnoses (18). The SCAN is thus much more comprehensive. In addition, because the SCAN is not bound to any diagnostic system, it is better suited for somatic symptoms and complaints, areas in which the SCID and CIDI are both weak.

As the BDS groups were based on the SCAN interview, these groups are expected to represent highly valid classifications. In the medical condition group, a minority of patients (either randomly selected or with a high score on the screening questionnaire) were SCAN interviewed. Therefore, some of the noninterview patients assigned to the medical condition group may have been undetected cases of BDS. Such misclassification of patients would bias the observed differences toward the null. Therefore, the study results may be conservative estimates, and we have no reason to believe that misclassification poses a particular problem to the validity of our study conclusions. Our choice of control group implied that also patients with a high score on the screening questionnaire were included in the control group. However, these high-scoring control group patients were all SCAN interviewed and distress could not be attributed to BDS;

<table>
<thead>
<tr>
<th>Reference group (n=879)</th>
<th>BDS single-organ type (n=124)</th>
<th>BDS multi-organ type (n=35)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emigration</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Age retirement</td>
<td>40</td>
<td>40</td>
</tr>
<tr>
<td>Disability pension</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Death</td>
<td>20</td>
<td>20</td>
</tr>
</tbody>
</table>

**FIGURE 3.** Percentage of patients who left the labor market during 10 years of follow-up according to the 3 patient groups. BDS, bodily distress syndrome.

**TABLE 5.** Risk of New Awards of Permanent Health-Related Benefits According to Patient Groups During 10 Years of Follow-Up

<table>
<thead>
<tr>
<th></th>
<th>Hazard Ratios (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Crude</td>
</tr>
<tr>
<td>Well-defined medical condition (reference)</td>
<td>1</td>
</tr>
<tr>
<td>BDS single-organ type</td>
<td>5.8 (3.6–9.3)</td>
</tr>
<tr>
<td>BDS multi-organ type</td>
<td>8.0 (3.8–16.9)</td>
</tr>
</tbody>
</table>

BDS, bodily distress syndrome.

\(^a\) Adjusted for age, sex, chronic illness, major depressive episode, anxiety disorder, and intervention.
the high scores were probably rather related to their underlying medical condition or mental distress. A general population sample could have constituted an alternative control group, but we would have expected this to have caused even more pronounced findings.

One study limitation was that the applied methods of health care cost analyses did not allow us to adjust for FP-rated chronic illness. The FP rated whether the patient had a chronic illness, but the FP did not make any specification of this illness. Therefore, we could not differentiate between chronic illness due to BDS and chronic illness in the form of a chronic comorbid medical condition. Consequently, we do not know how much of the increased health care costs may be attributed to a comorbid medical condition. However, in the analyses that were adjusted for chronic illness, we saw only minor differences between the crude and the adjusted estimates.

Finally, although we followed the included patients for an extensive period of time, we did not measure patient status of BDS at follow-up. Schedules for Clinical Assessment in Neuropsychiatry interviews were performed at baseline, and no reinterview was made during follow-up. Functional disorders have previously been shown to represent unstable conditions, depending on the diagnostic criteria applied (8,50,51). Based on the present study, we cannot conclude whether patients with BDS continuously fulfilled the criteria for either single-organ or multiorgan BDS or whether their baseline condition worsened or improved over time.

CONCLUSIONS

Patients with single-organ or multiorgan BDS were found to have unfavorable long-term outcomes and to be costly for society. Our study strongly supports the clinical use and the prognostic value of the new concept of BDS. Somatoform disorders and functional somatic syndromes may be treated effectively with combinations of cognitive behavioral therapy, antidepressants, graded exercise, and relaxation techniques (52,53). Correspondingly, promising results have been shown for treatment of severe BDS (54,55). These findings stress the need for improved recognition and implementation of the BDS diagnosis in primary care followed by research on effective treatment of patients with BDS.

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