A comprehensive study of oxidative stress in patients with somatic symptom disorder

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Abstract

Objective: To investigate oxidative stress parameters [total oxidant status (TOS), total antioxidant status (TAS), oxidative stress index (OSI), paraoxonase (PON), arylesterase (ARE) and thiol-disulphide homeostasis] in patients who were diagnosed as having somatic symptom disorder in accordance with Diagnostic and Statistical Manual of Mental Disorders-5. Methods: The study included 41 medication-free patients with somatic symptom disorder and 47 age, sex, and sociodemographic-matched healthy individuals. The patients were administered the Patient Health Questionnaire-15, Somatic Symptom Amplification Scale, Beck Depression Inventory, and Beck Anxiety Inventory. TOS, TAS, OSI, PON, ARE, thiol, disulphide levels, and routine biochemical parameters were compared between the two groups. Results: TOS, OSI, disulphide levels, disulphide/native thiol, and disulphide/total thiol ratios were found significantly higher in the patient group compared with the control group (p < 0.001). There was no significant difference in PON, ARE, and TAS parameters between the two groups (p > 0.05). Conclusion: This study showed that the level of oxidants increased and oxidative balance was impaired in somatic symptom disorder. Oxidative stress may play a role in the aetiopathogenesis of this disorder. This is the first study to report an association between oxidative stress and somatic symptom disorder.

Significant outcomes

- Our study revealed that oxidative balance is disrupted in favour of oxidants in patients with somatic symptom disorder.
- Lipid peroxidation seems not to have a role in aetiology of somatic symptom disorder.
- Approaches to reduce oxidative stress may be an alternative option for the prevention and treatment of this disorder.

Limitations

- This research was a cross-sectional study conducted at a single centre with a relatively small group of subjects.
- The oxidative stress parameters were not studied in cerebrospinal fluid (CSF) samples; therefore, serum levels of these oxidative stress parameters may not be fully correlated with their CSF levels.

Introduction

According to fifth edition of the Diagnostic and Statistical Manual of Mental Disorders (DSM-5), somatic symptom disorder is characterised by excessive thoughts, feelings of fear or worry, and anxious behaviours of an individual lasting for 6 months or longer, thinking that the individual has a serious disease as a result of misinterpretations of somatic symptoms (1,2). Although there are some biopsychosocial theories about its aetiopathogenesis, the aetiological basis of somatic symptom disorder is not yet known (3).

Oxidative stress is defined as the disruption of the balance between cellular reactive oxygen radicals and antioxidant mechanisms that serve to protect the body against oxidative stress. It is thought that the brain is prone to oxidative damage for various reasons (4). The main reason is that the brain has a high level of oxygen consumption for its metabolism. In addition, the presence of a large amount of redox-active metals such as iron and copper in the brain catalyzes the formation of reactive oxygen radicals. Furthermore, multiple polyunsaturated fatty acids, which exist in large amounts in neuronal cell membranes, are used as substrate in lipid peroxidation (5). The presence of a relatively modest level of important endogenous...
antioxidant molecules, along with the liability to oxidative reactions, causes the brain’s defense mechanisms against oxidative stress to become vulnerable (6). Hence, there is a great number of studies in the literature evaluating the relationship between psychiatric disorders and oxidative stress. It has been demonstrated that there is a relationship between oxidative stress and various psychiatric disorders such as depression, anxiety disorders, autism, schizophrenia, and bipolar disorder (7,8).

Few studies have examined the association between somatization and pathways related with oxidative stress, including immune-inflammatory pathways and the tryptophan metabolite (TRYCAT) pathway (9,10). These studies showed that there was an increase in inflammatory processes in patients with somatoform disorders (11). It was reported that activation of inflammatory processes triggered oxidative stress, and that oxidative damage might be related with somatic symptoms (12). However, no studies have shown the status of specific oxidative stress indicators in patients with somatic symptom disorder. Furthermore, it is thought that new studies are needed on this topic due to the manifestation of new probable oxidative stress mechanisms, which may have a role on the aetiology of mental disorders (12).

**Aims of the study**

In this study, we aimed to investigate oxidative stress parameters including total oxidant status (TOS), total antioxidant status (TAS), paraoxonase (PON), arylesterase (ARE), and thiol-disulphide homeostasis in patients with somatic symptom disorder. Consequently, it was aimed to establish the possible relationship between these oxidative stress parameters and total illness duration and severity of somatic symptoms to contribute to new hypotheses regarding the aetiology of this disorder.

**Material and methods**

**Subjects**

The study participants comprised 46 patients who were admitted to the psychiatry outpatient clinic of Ankara Ataturk Research and Training Hospital between January 2016 and September 2017. Patients who were diagnosed as having somatic symptom disorder in accordance with the diagnostic criteria of DSM-5 were screened. Three patients who were suspected of having a rheumatologic disorder and two patients who did not agree to participate in the study were excluded. Accordingly, the study population comprised 41 patients (24 females, 17 males) aged between 18 and 65 years who were medication free for at least 8 weeks. The patients were diagnosed as having somatic symptom disorder by a psychiatrist using a semi-structured psychiatric interview based on the DSM-5.

The exclusion criteria were the presence of any comorbid chronic disease (including fibromyalgia and chronic fatigue syndrome, and any neurologic, endocrinologic or metabolic disorder) or psychiatric disorder (e.g., mood disorders, anxiety disorders, or psychotic disorders), use of any medication, vitamin, or herbal supplement in the last 2 months, use of cigarettes or alcohol, and a history of acute trauma or infection in the last 2 months. To exclude the diagnosis of fibromyalgia, patients with pain symptoms were evaluated according to the ‘ACR 2010 fibromyalgia diagnostic criteria’ (13).

The control group was constituted by 47 healthy individuals (34 females, 13 males) who were matched for age, sex, and other sociodemographic features, and had no medical diseases or psychiatric disorders according to the Symptom Check List 90-Revised test and in the evaluation of the psychiatric interview. Alcohol users, smokers, and those who used any herbal supplements or medication, or had a history of acute infection or trauma in the last 2 months were excluded from the study.

**Psychiatric measurements**

In the clinical assessment, the Patient Health Questionnaire (PHQ-15) was administered to all patients to evaluate the severity of the somatic symptoms. The PHQ-15 is a 15-item Likert-type self-report scale, which was developed by Spitzer et al (14). The scale asks questions about somatic symptoms and their severity (15). Each item is assessed using scores between 0 and 2.

The Somatic Symptom Amplification Scale (SSAS), which was developed by Barsky et al. (16) to measure the amplification/exaggeration of normal somatic senses, was also administered to the patient group. The Turkish validity and reliability study of the scale was performed by Gülec et al. (17). The SSAS is a Likert-type scale comprising 10 questions, which are graded from 1 to 5.

The Beck Depression Inventory (BDI) and Beck Anxiety Inventory (BAI) were conducted to the patient group to assess the severity of their symptoms of depression and anxiety. The BDI is a 17-item Likert-type self-report scale developed by Beck et al. (18) that indicates the emotional, somatic, cognitive, and motivational symptoms of depression. The validity and reliability studies of Turkish version of the scale were conducted by Hisli et al. (19). The BAI is a 21-item scale that was developed by Beck et al. (20). The Turkish validity and reliability study of the scale was performed by Ulusoy (21). Each item of both tests is evaluated with a scores between 0 and 3.

**Blood samples**

After 8 h of fasting, 5 cc venous blood was drawn from all participants into ethylene diamine tetra acetic acid tubes in the morning. Specimens were centrifuged at 1800 g for 10 min. After separation, the serum samples were frozen and stored at −80°C until required for biochemical analysis. Biochemical analysis was conducted when the samples were dissolved after being kept at +4°C for 12 h.

From the samples collected from both groups, routine haemogram; liver, kidney, and thyroid function tests; vitamin B12; folate; ferritin levels; and lipid profiles were studied. Measurements of oxidative stress parameters (TOS, TAS, PON, ARE, and total thiol, native thiol, and disulphide) were performed using commercial kits (RelAssay, Gaziantep, Turkey) on a Roche Hitachi Cobas c501 (Indianapolis, IN, USA) automatic analyzer.

**Measurement of oxidative stress parameters**

The total antioxidant response and TOS were determined using the automated methods described by Erel (22,23). The ratio of TOS to TAS gives the oxidative stress index (OSI) value. The OSI value was calculated using the following formula:

\[
\text{OSI (arbitrary unit)} = \frac{\text{TOS} (\mu\text{mol H}_2\text{O}_2 \text{ Eqv/l})}{\text{TAS} (\text{mmol Trolox Eqv/l})} \times 100
\]

PON and ARE are enzymes that protect cells against lipid peroxidation. The serum activity of PON and ARE was measured using commercially available kits (Rel Assay Diagnostics). PON activity was measured using paraoxon substrate by recording the 37°C absorbance of the colour formed as a result of paraoxon hydrolysis at 412 nm, and results are expressed as U/l. To measure ARE activity, a newly non-salt-stimulated method is used, with
phenyl acetate used as substrate. One unit of ARE activity is determined as μmol phenol formed in 1 min, and the results are expressed as kU/l (24).

Serum thiol-disulphide homeostasis was studied with the new automated measurement method developed by Erel and Neselioglu (25). Dynamic disulphide bonds (–S–S–) are reduced to functional thiol groups (–SH) by sodium borohydride (NaBH₄). The remaining unused NaBH₄ is completely removed by formaldehyde. The native thiol (–SH) and total thiol (–SH + –S–S–) content of the sample is measured using the modified Ellman reagent. The amount of disulphide bond is obtained by half of the difference between total thiol and native thiol. In addition, disulphide/native thiol, disulphide/total thiol, native thiol/total thiol ratios were calculated as percentage (%).

**Statistical analysis**

The IBM SPSS Statistics 21 software program was used to analyse the study data. The normality of continuous variables was tested using the Shapiro–Wilk test. Descriptive statistics [mean ± standard deviation (SD)] are given for continuous variables. Sex variance is expressed as percentage. For the results of the oxidative stress parameters, mean ± SD is given.

The continuous variables of both groups were compared using the independent samples t-test. Univariate analysis of covariance was performed to investigate the effects of potential confounders [age, sex, body mass index (BMI)]. The relationship between the levels of the oxidative stress parameters of the patient group, and total illness duration, and the scores of the PHQ-15 and SSAS scales were examined using Spearman correlation analysis. The statistical significance level was determined as p < 0.05.

**Results**

The study consisted of 88 subjects, including 41 patients and 47 healthy individuals. The demographic features of the patient and control groups are shown in Table 1. There was no statistically significant difference between the groups in terms of age, sex, BMI, marital status, and educational status (p > 0.05). The average total illness duration of the patient group was 53.54 ± 53.7 months. The scores of the psychiatric evaluation scales of the patient group are also given in Table 1.

When the biochemical findings of the patient and control groups were examined, there was no statistically significant difference between the two groups in terms of liver, kidney, and thyroid function tests, lipid profiles, ferritin, folate, vitamin B₁₂, white blood cell counts, haemoglobin, and platelet levels (p > 0.05).

Both groups were compared in terms of oxidative stress parameters. It was found that parameters indicating oxidant molecules increased. The native thiol level of the patient group was significantly lower than that of the control group, but the disulphide level was significantly higher (p < 0.001). The TOS, OSI, disulphide/native thiol [(–S–S–)/(–SH)], and disulphide/total thiol [(–S–S–)/(–SH + –S–S–)] levels of the patient group were higher than those of the control group (p < 0.001). In addition, the native thiol/total thiol [(–SH)/(–SH + –S–S–)] level was significantly lower for the patient group than in the control group.

<table>
<thead>
<tr>
<th>Table 1. Demographic characteristics of the two groups</th>
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<td><strong>Age</strong></td>
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<td>Sex (n, %)</td>
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<td>Educational status (n, %)</td>
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<td>Marital status (n, %)</td>
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<tr>
<td>Other</td>
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<tr>
<td>BMI (kg/m²)</td>
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<tr>
<td>Duration of illness (month)</td>
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<td>PHQ-15 score</td>
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BAI, Beck Anxiety Inventory; BDI, Beck Depression Inventory; BMI, body mass index; PHQ-15, Patient Health Questionnaire-15; SSAS, Somatic Symptom Amplification Scale.

Values are presented as mean ± SD.
and that inflammation triggered oxidative stress (26). Reported that there was an increase in proinflammatory processes, conducted with patients with somatoform disorders, it was order as compared with the control group. In some studies TOS, TAS, and OSI levels is the first to investigate the relationship between oxidative stress and somatic symptom disorder.

Discussion

In this study, it was found that oxidative balance was disrupted in favour of oxidants in patients with somatic symptom disorder. We found that TOS, OSI, disulphide, disulphide/native thiol and disulphide/total thiol parameters were higher in patients with somatic symptom disorder than in healthy individuals. This study is the first to investigate the relationship between oxidative stress and somatic symptom disorder.

TOS, TAS, and OSI

The prominent finding of this study was that TOS and OSI levels are significantly higher in patients with somatic symptom disorder as compared with the control group. In some studies conducted with patients with somatoform disorders, it was reported that there was an increase in proinflammatory processes, and that inflammation triggered oxidative stress (26–29). The increase in oxidant load causes a decrease in antioxidant molecules, and thus oxidative damage in many molecules such as DNA and protein (30).

The damaged molecules become the new target epitopes of autoimmunity. Among neurotransmitters, serotonin is also one of the targets of this autoimmunity (31). It is known that the serotonergic system inhibits the response to painful stimuli (32). In addition to causing serotonergic neuronal damage, oxidative stress can also decrease the production of serotonin from tryptophan by increasing the level of indoleamine 2,3-dioxygenase enzyme activity (33). Moreover, it was demonstrated that a decrease in tryptophan levels and an increase in its degradation products might cause psychological stress, fatigue, and increased pain and autonomic symptoms (34–36). The fact that TOS levels were found to be high for somatic symptom disorder in our study can be associated with the aforementioned proinflammatory processes in this patient group.

Although some studies expressed how the immune system and proinflammatory cytokines could cause somatisation symptoms (37,38), no significant relationship has been detected between TOS levels and somatosensory amplification scores or the severity of somatic symptoms. The use of self-reporting scales to assess symptom severity could be a confounding factor concealing the relationship between them.

In this study, the TAS level of the patients with somatic symptom disorder was not different from those of the control group. Similarly, in another study, it was revealed that there was no difference in some anti-inflammatory cytokines between patients with somatisation disorder and the control group (9). The TAS levels of our study group can be interpreted as a ‘rebound increase’ to compensate for the increase in oxidant molecules. However, the increased OSI ratio indicating general oxidative stress balance, demonstrates that this probable reactive increase is inadequate to balance the oxidative stress, and the oxidative balance is disrupted in favour of oxidants. The fact that no significant difference was detected in our study between the patient and control groups for PON and ARE antioxidant enzymes further shows that the status of oxidative stress in somatic symptom disorder might not be associated with the disruption in the antioxidant system, but with the increase in oxidant molecules.

Lipid peroxidation

It is known that PON and ARE enzymes inhibit lipid peroxidation by protecting high-density lipoprotein (HDL) and low-density lipoprotein cholesterol against oxidative reactions, and
have antioxidant and anti-inflammatory features. It was found that the peroxidation of HDL, which is seen in some diseases characterised by oxidative damage, is related with decreased PON-1 activity (28). No studies in the literature have evaluated the role of lipid peroxidation in the pathogenesis of somatic symptom disorder. However, we found that there was non-significant difference between the two groups for PON and ARE enzyme levels. This result shows that lipid peroxidation might not have a role in the aetiopathogenesis of this disorder.

**Thiol-disulphide homeostasis**

In this study, we also found that there was a decrease in native thiol levels and an increase in disulphide levels for the patient group. Consequently, the thiol-disulphide balance changed in favour of the reduced side. Thiols can undergo oxidation reactions with ROS, and they lose electrons from sulphydryl (–SH) groups to reduce them. Therefore, they protect cells against oxidative damage and disulphide bonds (–S–S) are formed as a result of these reactions (23). Glutathione is the intracellular molecule that contains the most thiol groups and is one of the important antioxidant defense systems in the brain (39). It was demonstrated that there was a decrease in antioxidant molecules such as glutathione, zinc, and coenzyme Q10 in patients with depression and somatisation symptoms such as chronic fatigue syndrome (29,40).

In some studies, it was reported that the thiol-disulphide balance was disrupted in favour of disulphide, as in this study, in a variety of neurologic and psychiatric diseases such as depression, schizophrenia (41), drug abuse (42), attention-deficit/hyperactivity disorder (43), and Alzheimer’s disease (44). Despite the different groups of disorders with differing aetiological and clinical features, there are overlapping findings regarding the disruption in thiol-disulphide balance among them, which might be due to the role of psychological and biologic stress in the aetiologies and clinical courses of the diseases.

There are certain limitations to our study. One of which is that the study was conducted in a single centre and with a relatively limited number of subjects. Another is that our research was a cross-sectional study. In addition, our findings may be inadequate to fully reflect the oxidative balance of the brain because oxidative stress parameters were only studied for serum samples. Studies with CSF or changes in receptors with regard to oxidative parameters might yield clearer results.

There are many unknown factors in terms of the aetiology, diagnosis, and treatment of somatic symptom disorder. Our study contributes to the understanding of the aetiology of somatic symptom disorder by showing that oxidative balance is disrupted in favour of oxidants in the disorder. Different therapeutic interventions that reduce oxidant load could be considered in the treatment of somatic symptom disorder. This result needs to be supported by new studies with larger sample groups. Further studies are also needed to investigate the potential effects of antioxidant treatment on oxidative stress parameters and symptoms of somatic symptom disorder.

**Acknowledgements.** This article is the result of a resident thesis and the authors would like to thank all subjects who participated in the study and the staff of the biochemistry laboratory of Ankara Ataturk Research and Training Hospital for their assistance. Authors’ contribution: All authors contributed equally to all steps of this manuscript such as the design of the study, data collection, data interpretation, literature review, content revision, and final approval.

**Financial Disclosure.** This study was supported by a grant given by Ankara Yıldırım Beyazıt University Scientific Research Project Unit with the project number 2670.

**Conflicts of Interest.** None.

**Ethical Standards.** The authors assert that all procedures that contributed to this work complied with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, as revised in 2008. Approval for the study was granted by the Ankara Yıldırım Beyazıt University Institutional Ethics Committee, and written informed consent was obtained from all participants.

**References**


