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Thiol-disulfide homeostasis: an integrated approach with biochemical and clinical aspects

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Abstract: Dynamic thiol-disulfide homeostasis (TDH) is a new area that has begun to attract more scrutiny. Dynamic TDH is reversal of thiol oxidation in proteins and represents the status of thiols (-SH) and disulfides (-S-S-). Organic compounds containing the sulfhydryl group is called thiol, composed of sulfur and hydrogen atoms. Disulfides are the most important class of dynamic, redox responsive covalent bonds built between two thiol groups. For many years, thiol levels were analyzed by several methods. During last years, measurements of disulfide levels have been analyzed by a novel automated method, developed by Erel and Neselioglu. In this method, addition to thiol (termed as native thiol) levels, disulfide levels were also measured and sum of native thiol and disulfide levels were termed as total thiol. Therefore, TDH was begun to be understood in organism. In healthy humans, TDH is maintained within a certain range. Dysregulated dynamic TDH has been implicated several disorders with unknown etiology. A growing body of evidence has demonstrated that the thiol-disulfide homeostasis is involved in variety diseases, such as diabetes mellitus, hypertension, nonsmall cell lung cancer, familial Mediterranean fever (FMF), inflammatory bowel diseases, occupational diseases, gestational diabetes mellitus and preeclampsia. These results may elucidate some pathogenic mechanism or may be a predictor indicating diagnostic clue, prognostic marker or therapeutic sign. In conclusion, protection of the thiol-disulfide homeostasis is of great importance for the human being. Evidence achieved so far has proposed that thiol-disulfide homeostasis is an important issue needs to elucidate wholly.

Key words: Thiol, disulfide, oxidant stress, thiol-disulfide homeostasis

1. Introduction

Human life is maintained by oxygen and aerobic processes, and reactive oxygen species (ROS) are harmful by-products of human organism metabolism [1]. Reactive oxygen species, composed of several molecules such as superoxide anion radicals (O2−), hydrogen peroxide (H2O2), and hydroxyl radicals (•OH), are normally produced by cells during metabolic processes such as protein synthesis and mitochondrial metabolism [2].

The oxidative effects of ROS are neutralized by the antioxidant capacity of cells and this battle with oxidant stress maintains homeostasis [1]. Within the cell, the redox couples are controlled in a location-specific manner, especially in mitochondria, endoplasmic reticulum (ER), and nuclei [3]. Additionally, extracellular compartments supply defensive barriers against external oxidants. Cysteine (Cys) and its disulfide, cystine (CySS) compose the major low-molecular weight thiol/disulfide couple in human plasma. The Cys/CySS pool is central redox control point in the biological signaling [4,5].

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2. Dynamic thiol-disulfide homeostasis

Organic compounds containing the sulfhydryl group is called thiol (-SH), composed of sulfur and hydrogen atoms. The thiols have high vulnerability to the oxidation due to their –SH group. Disulfides (-S-S-) are the most important class of dynamic, redox responsive covalent bonds built in between two thiol groups. Dynamic thiol-disulfide homeostasis (TDH) is reversal of thiol oxidation in proteins and represents the levels of thiols and disulfides. It is an important parameter associated with several biochemical processes, including regulation of protein function, stabilization of protein structure, protection of proteins against irreversible oxidation of cysteine residues, chaperon function, regulation of enzyme functions and transcription [6–9].

2.1. Thiol and disulfide detection

Up to date, thiol, disulfide, sulfur-containing amino acid, reduced glutathione and oxidized glutathione levels were analyzed by several methods.

Ellman [10] and Hu [11] have developed a technique using 5,5′-dithiobis-(2-nitrobenzoic) acid (DTNB) in other words Ellman’s reagent as a chromogen to measure the thiol levels.
Recently, Erel and Neselioglu [12] modified Ellman’s method. In this new method, it was aimed both thiol and disulfide levels in blood. Firstly, available thiol in serum/plasma was detected without pretreatment. This first result was accepted as native thiol (NT). Then, a pretreatment step was carried out to reduce dynamic disulfide bonds to free sulphydryl groups by using sodium borohydrate (NaBH₄). Second measurement was performed and second result was accepted as total thiol (TT). In this step, extra reduction of DTNB and further reduction of formed disulfide bonds may occur. To prevent this positive interference, remaining NaBH₄ remnants were completely removed using formaldehyde. Lastly, half of the difference between TT and NT was accepted disulfide. During measurement procedure, 2-mercaptoethanol was used to achieve a linear calibration curve. Also, they used two wavelengths, main wavelength 415 nm, secondary wavelength 700 nm (optionally bichromatic). In this endpoint assay, the first absorbance was taken before the mixing of Reagent 2 and Reagent 3 and the last absorbance was taken when the reaction trace draws a plateau. Reagent ingredients and volumes were presented in Table. In this method, thiol (-SH) was represented with NT, while oxidized thiol (-S-S) was represented by disulfide. Total thiol is composed of sum of the NT and disulfide levels. This method has analyzed NT, total thiol (TT), disulfide levels and ratios of disulfide to native thiol, disulfide to total thiol and native to total thiol to elucidate dynamic TDH in organism.

Beside to the colorimetric methods, several measurement methods were developed such as fluorometric methods [13], bioluminescence analysis [14], chromatographic methods [15].

2.2. Researches on thiol-disulfide homeostasis
During last years, TDH has started to be measured more widely. Therefore, TDH has started to be understood more exactly. Dysregulated TDH has been implicated several disorders with unknown etiology. A growing body of evidence has demonstrated that TDH is involved in the several diseases. Researchers have established that changing in NT, TT, disulfide levels and/or ratios. These results may elucidate some pathogenic mechanism or may be a predictor indicating diagnostic clue, prognostic marker or therapeutic sign.

2.2.1. Cardiac pathologies
In a study investigating association between TDH and severity of coronary atherosclerosis, high syntax score was found to be associated with decreased NT level and NT to disulfide ratio [16]. Also, it has been demonstrated that NT, TT and disulfide levels of stable angina pectoris and acute myocardial infarction patients were significantly lower than control patients [17,18]. Protecting power of extracellular cysteine pairs in disulfides protect for the reactive thiol groups and supporting effect on protein stability and function may be reason for lower disulfide level [19]. Lower disulfide levels may result from mentioned mechanism.

It has proposed that total thiol is independently diagnostic predictor of coronary syndrome X [20]. Also, decreased NT and increased disulfide levels are associated with coronary artery ectasia [21] and decreased NT to disulfide ratio was independently associated with slow coronary flow [22]. Additionally in a study investigating association between childhood obesity, reduced thiol parameters were decreased whereas oxidized thiol parameters were increased [23].

In both primary hypertension and masked hypertension patients, disulfide and ratios in favor of disulfide were found to be increased. Furthermore disulfide to NT ratio to be an independent indicator of both systolic and diastolic blood pressure [24–26].

In an interesting study, researchers have investigated whether TDH can predict the occurrence of anthracycline-induced cardiac toxicity. They were concluded that NT may elicit more objective data for physicians to detect which patients have high risk for cardiac toxicity [27].

2.2.2. Endocrine disorders
Authors have demonstrated that NT and TT levels of prediabetes, type 1 and type 2 diabetes mellitus were lower while disulfide levels and disulfide to NT and disulfide

| Insert | Volume (μL) | Ingredient |
|--------|-------------|------------|
| Reagent 1* | 10 | 10 mM NaCl in methanol–water solution, 50 v/v |
| Reagent 1** | 10 | 10 mM NaBH₄ in methanol–water solution, 50 v/v |
| Reagent 2 | 110 | 6.715 mM formaldehyde and 10.0 mM EDTA in tris buffer, 100 mM, pH: 8.2 |
| Reagent 3 | 10 | 10 mM DTNB in methanol |
| Sample | 10 | Serum or plasma |

*for native thiol, **for total thiol.
to TT ratios were higher in patients compared to healthy volunteers [28–32].

Authors have demonstrated that TDH shifted toward disulfide formation in patients with autoimmune subclinical hypothyroidism and thyroid autoantibodies are positively correlated with disulfide to NT ratio [33].

2.2.3. Neurological diseases
It has been established that NT and TT levels of Alzheimer's disease, diabetic axonal polyneuropathy patients and children with febrile seizures were found to be statistically lower than controls. On the contrary, disulfide/TT ratio was found to be statistically higher than controls [34–36].

Authors have demonstrated that in epileptic patients, TDH was not affected. This unchanged homeostasis may result in both selecting patients taking mono- or poly- antiepileptic medication and preferring seizure-free period in collection of patient samples [37].

It has been demonstrated increased NT and TT levels in the attention deficit hyperactivity disorder and migraine patients [38,39].

It has been demonstrated that disulfide levels and ratios of disulfide to NT and disulfide to TT were significantly higher in multiple sclerosis patients in the relapse than in patients in the remission [40]. Similarly, it has been demonstrated that NT and TT levels in Parkinson's disease, panic disorder and acute ischemic stroke patients were detected to be significantly lower than healthy control subjects. Furthermore, it has been demonstrated that lower NT levels were associated with higher infarct volume and higher National Institutes of Health Stroke Scale (NIHSS) score [41–43].

Headache including migraine- and tension-type headache in childhood was investigated in a study. It has been established that there was no significant difference in patient group compared to control group in terms of TDH. However, it has been established that disulfide to NT and disulfide to TT ratios were higher in the migraine group. In tension-type headache group, a negative correlation was found between thiol levels and Pediatric Migraine Disability Assessment (PedMIDAS) [44].

2.2.4. Psychiatric diseases
It has been demonstrated that NT, TT levels were significantly lower in schizophrenia and heroin addiction patients. Furthermore, authors have demonstrated a negative correlation between NT, TT levels and Positive and Negative Syndrome Scale (PANSS) [45,46].

Disruptive episodes of mania/hypomania and depression are called as bipolar affective disorder. In patients group, NT and TT levels were lower than remission and control group whereas no significant difference is detected between groups [47].

Repetitive transcranial magnetic stimulation (rTMS), a treatment modality based on magnetic pulses administered to cerebral cortex, may result in decreased serum NT and TT levels. It has been proposed that decreased thiol levels were associated with decreased inflammatory processes and decreased antioxidant activity [48].

2.2.5. Respiratory system diseases
In acute pulmonary embolism and childhood asthma patients, it has been established that disulfide level and disulfide to NT ratio of patient group were found to be higher compared to control group [49,50]. Contrarily, in chronic obstructive pulmonary disease, asthma and asthma-chronic obstructive pulmonary disease overlap syndrome patients, TDH parameters were found similar among three groups. Furthermore, TDH parameters were not changed among smokers, nonsmokers and ex-smokers [51].

Authors have shown that NT, TT and disulfide levels in nonsmall cell lung cancer patients were significantly decreased compared with control subjects. It has been suggested that decrease of TDH parameters might be the prognostic marker of the tumor aggression and malignant disease [52].

2.2.6. Head and neck pathologies
Authors have demonstrated that NT levels were significantly lower in patients with benign paroxysmal positional vertigo, seasonal allergic rhinitis, Bell's palsy, obstructive sleep apnea and nasal polyposis [53–57].

2.2.7. Gastrointestinal system disorders
It has been demonstrated that in patients with celiac disease, acute pancreatitis and inflammatory bowel diseases NT and TT levels were found to be lower, whereas disulfide level was found to be higher compared to healthy subjects [58–60].

2.2.8. Infectious diseases
In acute brucellosis, Crimean-Congo hemorrhagic fever, viral and bacterial tonsillopharyngitis NT and TT levels were found to be lower in patients than healthy control subjects. [61–63].

2.2.9. Rheumatologic diseases
Familial Mediterranean fever (FMF) is found to be associated with lower NT and TT levels than healthy control subjects. During attack periods, it has been demonstrated that NT and TT levels are found to be lower than attack free periods. In fact it has been established that NT and TT levels were associated with colchicine dosage [64]. Additionally, authors have investigated FMF in terms of effect of different mutations in the MEFV gene on TDH. Authors have proposed that TDH was not sufficient indicator in different mutations in MEFV gene. Nevertheless, it has been demonstrated lower NT, TT and disulfide levels, disulfide to NT and disulfide to TT ratios in FMF patients. Authors have explained these decreased levels transformation form disulfide to S-nitrosothiol, sulfenic acid, sufinic acid and sufonic acid [65].
It has been demonstrated that TT levels were significantly lower in ankylosing spondylitis and juvenile idiopathic arthritis patients compared to control group. Additionally, authors have proposed that thiol levels might be a useful marker of disease activation and there was a significant negative correlation TT levels and visual analog scale (VAS); TT levels and Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) [66, 67]. Additionally, authors have demonstrated that decreased native thiol and total thiol levels and increased WOMAC score and disulphide levels were independently associated with increased risk of late-stage osteoarthritis [68].

Fibromyalgia syndrome is related to thiol-disulfide imbalance shifted to the reductive state according to a recent paper. In this study, NT levels were found to be significantly higher and disulfide levels were found to be significantly lower in patients with fibromyalgia syndrome than control group. This situation was connected with proliferative alteration explaining with higher NT and lower disulfide levels [69].

2.2.10. Dermatologic disorders
Thiol-disulfide homeostasis in patients with basal cell carcinoma, psoriasis and seborrheic dermatitis has changed as the increased NT. It has been suggested that increased NT levels cause proliferation, whereas reduced NT along with the increased disulfide results in cell apoptosis [70–72].

It has been demonstrated that TDH was shifted towards to disulfide formation in patients with chronic spontaneous urticaria whereas TDH was no changed in patients with acute urticarial [73].

In atopic dermatitis patients, serum disulfide levels and disulfide to NT and disulfide to TT ratios were found to be lower compared with healthy control children. Moreover, disulfide to NT ratio was positively correlated and NT to TT ratio was negatively correlated with higher scores assessing atopic dermatitis severity. Authors have proposed that decreased disulfide level might result from rebound phenomenon to continue protective influence by facilitating the neutralization of oxidative state [74].

It has been demonstrated that TDH was no changed in patients with alopecia areata, an autoimmune disease characterized scar-free hair loss. This may result from patients which has mild alopecia areata disease [75].

2.2.11. Ophthalmic disorders
Authors have established that NT and TT levels were lower whereas disulfide level was higher in keratoconus and pseudoexfoliation syndrome patients compared with control group [76–78].

Native thiol and TT levels were found to be lower in central serous chorioretinopathy and macular degeneration patients compared to control subject [79–81].

In another study, authors have investigated age-related macular degeneration and they have established that NT levels were significantly decreased and disulfide levels were significantly increased in patients compared to control subjects [82].

2.2.12. Obstetric and gynecologic disorders
As one most common endocrine disease, polycystic ovary syndrome has been shown to be related to higher thiol and lower disulfide levels. Authors have proposed that antioxidants hinder the atresia of antral follicles and the thiol play an essential role in cell proliferation, division and apoptosis. Elevated thiol levels may result from anti-oxidant response to oxidative load due to obesity or a consequence of physiological changes in polycystic ovary syndrome such as anovulation, multiple follicular development and apoptosis [83]. In another study investigating polycystic ovary syndrome, NT and TT levels were found to be significantly lower in patients with overweight polycystic ovary syndrome than normal weight polycystic ovary syndrome and oxidative stress was more marked in overweight polycystic ovary syndrome. Furthermore, it has been demonstrated that lipid accumulation index was statistically significantly higher than in patients with overweight polycystic ovary syndrome than those in other three groups. Authors have established that lipid accumulation index and TDH may play an important role in the pathogenesis of cardiovascular disease in polycystic ovary syndrome [84].

In a study investigating postmenopausal osteoporosis it has been demonstrated that disulfide to NT ratio was higher in patients after adjustment for age and menopause duration and there was a negative correlation between the disulfide to NT ratio and bone mineral density of the lumbar vertebrae [85].

Authors investigated whether there is an association between TDH and premature ovarian failure. It has been demonstrated that TDH was shifted in favor of disulfide formation [86]. In patients with uterine myoma it has been demonstrated that NT, TT and disulfide levels were found to be lower than healthy control subjects. These results have been commented that proliferative phase of uterine myoma might be a possible cause and disulfide levels were found to be lower than control group [87]. It has been demonstrated that NT and TT levels in patients with endometriosis were lower than control group [88].

It has been demonstrated that TDH was shifted in favor of oxidative status in preeclampsia, pregnant women diagnosed with idiopathic recurrent pregnancy loss, pregnant women diagnosed with FMF, pregnant women with fetal neural tube defects and vaginitis patients [89–93].

There are several studies investigating association between TDH and gestational diabetes mellitus [94–97].
It has been demonstrated that NT and TT levels were found to be lower than control subjects and decreased NT levels have an increased risk of possible adverse perinatal outcomes [95]. It has been demonstrated that thiol levels were significantly lower whereas disulfide levels were significantly higher in patients with gestational diabetes mellitus compared with both pregnant women with impaired glucose tolerance and uncomplicated pregnant subjects [96]. Authors have investigated pregnant women with gestational diabetes mellitus, obese pregnant women and healthy pregnant women in terms of TDH of cord blood. Native thiol levels were found to be lower in patients with gestational diabetes mellitus compared to other two groups. Total thiol levels were found to be lower in patients with gestational diabetes mellitus compared to obese pregnant women. Disulfide levels were found to be higher in patients with gestational diabetes mellitus compared to healthy pregnant women and in obese pregnant women compared to healthy pregnant women. Authors have suggested that infants of obese or diabetic mothers are exposed to increased oxidative stress [94]. In another study, authors have demonstrated that 50 grams glucose challenge test-positive pregnant women have increased disulfide levels after glucose loading compared to baseline [97].

Thiol-disulfide homeostasis has been investigated in patients with abortus imminens. It has been demonstrated that NT and TT levels were found to be lower than control group. Authors have proposed that during pregnancy, insufficient antioxidant defense may be an etiologic factor in the pathogenesis of abortus imminens [98].

It has been demonstrated that NT ad TT levels were significantly lower and disulfide level was significantly higher in pregnant women with hyperemesis gravidarum and intrahepatic cholestasis and in fetus with nuchal cord during labor than control group [99–101].

In another study, authors have demonstrated that maternal NT to TT ratio, NT and TT levels were significantly lower while ratios of disulfide to NT and disulfide to TT were significantly higher in pregnancies complicated by intra uterine growth restriction compared to control subjects. Authors have proposed that decreased NT and TT levels might diminish maternal serum $H_2S$ level and inhibit its vasodilatation effect through nitric oxide synthesis [102].

It has been demonstrated that NT and TT levels were lower and disulfide to NT and disulfide to TT ratios were higher in pregnancies complicated by obstructive sleep apnea syndrome compared to nonpregnant control subjects [103].

At birth, the timing of umbilical cord clamping has important for cardiovascular system, pulmonary system and blood volume of neonate. In a study investigating effect of three different cord clamping procedures (i.e., early clamping, delayed clamping, and cord milking) on TDH, it has been demonstrated that NT and TT levels were statistically significantly lower in the early clamping group compared with the delayed clamping and cord milking group as an indicator of oxidative stress. In conclusion, they have recommended delayed cord clamping and cord milking due to beneficial effects on the neonates [104].

In a study investigating TDH in preterm infant, researchers have collect patient blood at baseline, first week and third week after birth. It has been demonstrated that NT and TT levels were increased in each analysis and disulfide levels and disulfide to NT and disulfide to TT ratios were increased at first week and decreased at third week. The ratio of NT to TT was decreased at first week and increased at third week. Authors have proposed that increased thiol levels might result from cysteine containing amino acid administration or breast milk that enhances antioxidative defense compared to enteral feeding. Moreover, increased disulfide level at first week may due to oxidative damaged-treatment including phototherapy and antibiotics. Furthermore, it is speculated that healing from respiratory diseases might contribute to diminish in disulfide levels at third week [105].

2.2.13. Urological disorders
In a study executed on prostate cancer patients, authors have compared prostate cancer patients before and six month after radical prostatectomy operation and control group in term of TDH. In the study, NT, TT and disulfide levels were found to be lower in patients before operation compared to control subjects. There was a significant negative correlation between levels of prostate specific antigen and levels of NT and TT in patient group before operation and control group. Similarly, NT, TT and disulfide levels were found to be lower in patients after operation compared to control subjects. Furthermore, authors have stated that TDH begins to shift towards thiol following operation [106]. Similarly, authors have demonstrated that NT and TT levels decreased in patients underwent transrectal ultrasound guided prostate biopsy [107].

It has been demonstrated that TDH has shifted towards disulfide in patients with varicocele as one of the most common causes of male infertility [108].

2.2.14. Occupational diseases
In two study investigating TDH in operating theater personnel and asphalt workers, authors have demonstrated that TDH has shifted to disulfide formation resulting from increased disulfide level, increased disulfide to NT ratio [109, 110].

In a study investigating occupationally arsenic-exposed workers in terms of TDH, disulfide level, disulfide to NT ratio and disulfide to TT ratio were found to be increased in workers exposed to arsenic, lead and asphalt fume than
control group. Furthermore, it has been demonstrated that there are positive correlations between urinary arsenic level and disulfide level and urinary arsenic level and disulfide to NT ratio. Also, a positive correlation was detected between lead and disulfide levels [111–113].

It has been demonstrated that NT levels were found to be significantly lower in professionals working in radiation environments than control subjects. However, there was no difference between two groups in terms of TT and disulfide levels. The results have been explained that NT would only have been affected by initial stages of these processes [114].

When TDH parameters were investigated silica and trichloroethylene exposure group, it has been established higher disulfide levels in exposed workers than healthy volunteers. Furthermore, there was a negative correlation between NT, TT, NT to TT ratio and urinary trichloroethylene levels and a positive correlation between disulfide to NT and disulfide to TT ratios and urinary trichloroethylene levels [115,116].

2.2.15. Other disorders
In an interesting study, authors have investigated TDH parameters in gunshot injuries. It has been demonstrated that NT, TT and disulfide levels were decreased whereas disulfide to NT and disulfide to TT ratios were increased in patient with gunshot injury. Furthermore, NT levels were found to be an independent indicator of Revised Trauma Scale (RTS) and Glasgow Coma Scale (GCS). Authors have speculated that low NT levels might result from attempt to detoxify oxidative stress and ROS generated due to trauma [117].

The authors have demonstrated that NT and TT levels are found to be lower and disulfide level was found to be higher in patients with acute appendicitis compared to control group. In fact this shift towards disulfide formation was valid in perforated appendicitis patients compared to nonperforated appendicitis patients [118]. Similarly, in breast cancer patients, NT and TT levels are found to be lower and disulfide level was found to be higher than control subjects [119].

In a study investigating effect of elective laparoscopic cholecystectomy and open inguinal-femoral hernia repair operation on TDH, the authors have demonstrated that during operation, NT, TT and disulfide levels were found to be decreased. Additionally, 24 h after laparoscopic surgery, NT, TT and disulfide levels were found to be increased but not return to preoperative levels [120].

In operated aortic aneurysm and acute aortic syndrome patients, NT and TT levels were increased and disulfide level and disulfide to NT ratio were decreased after 6 months of operation. Authors have concluded that thiol and disulfide levels are distinctive tests between healthy subjects and aneurysm or acute aortic syndrome [121].

It has been demonstrated that in patients with carbon monoxide poisoning and smokers NT and TT levels were decrease whereas disulfide level was increased [122, 123].

Authors have demonstrated that NT, TT and disulfide levels decreased in patient with multiple myeloma, a malignancy of bone morrow. Because the ratios were similar in all groups, authors have asserted that there is a balance between thiol and disulfide levels and the balance determine the systemic effects [124].

It has been demonstrated that TDH parameters were no changed in maple syrup urine disease patients. Authors have decided that good metabolic control and proper dietary compliance in maple syrup urine disease patients may prevent oxidative stress [125].

Recently, it has been provided that thiol-disulfide homeostasis might be a notable key for evaluating the severity of burns and predicting the survival [126].

3. Conclusion
Thiol-disulfide homeostasis is reversal of thiol oxidation in proteins and represents the levels of thiols and disulfides. Thiol-disulfide homeostasis is an important parameter associated with several biochemical processes. Dysregulated thiol-disulfide homeostasis has been implicated several disorders with unknown etiology. The evidence so far has proposed that thiol-disulfide homeostasis is an important issue and needs to be elucidated wholly.

References
1. Yi MC, Khosla C. Thiol-disulfide exchange reactions in the mammalian extracellular environment. Annual Review of Chemical Biomolecular Engineering 2016, 7: 197-222.
2. Halliwell B, Gutteridge JM. Lipid peroxidation, oxygen radicals, cell damage, and antioxidant therapy. Lancet 1984, 1: 1396-1397.
3. D'Autreaux B, Toledano MB. ROS as signalling molecules: mechanisms that generate specificity in ROS homeostasis. Nature Reviews Molecular Cell Biology 2007; 8: 813-824.
4. Jones DP, Go YM, Anderson CL, Ziegler TR, Kinkade JM Jr et al. Cysteine/cystine couple is a newly recognized node in the circuitry for biologic redox signaling and control. The FASEB Journal 2004; 18: 1246-1248.
5. Jones DP, Mody VC Jr, Carlson JL, Lynn MJ, Sternberg P Jr. Redox analysis of human plasma allows separation of pro-oxidant events of aging from decline in antioxidant defenses. Free Radical Biology Medicine 2002; 33: 1290-1300.
17. Altiparmak IH, Erkuş ME, Sezen H, Demirbag R, Gunebakmaz
16. Kundi H, Erel Ö, Balun A, Çiçekçıoğlu H, Cetin M et al.
15. Reeve J, Kuhlenkamp J, Kaplowitz N. Estimation of glutathione
14. Hinze WL, Riehl TE, Singh HN, Baba Y . Micelle-enhanced
13. Cohn VH, Lyle J. A fluorometric assay for glutathione
12. Erel O, Neselioglu S. A novel and automated assay for thiol/
11. Hu ML. Measurement of protein thiol groups and glutathione
10. Ellman GL. Tissue sulfhydryl groups
9. Sanchez-Rodriguez MA, Mendoza-Nunez VM. Oxidative stress
8. Brulsauer L, Gauthier MA, Leroux JC. Disulphide-containing
7. Chen W , Zhao Y , Seefeldt T, Guan X. Determination of thiols
6. Ellgaard L, Sevier CS, Bulleid NJ. How are proteins reduced in
5. Altiparmak IH, Erkuş ME, Sezen H, Demirbag R, Gunebakmaz
4. Gumusyayla S, Vural G, Bektas H, Deniz O, Neselioglu S et al.
3. Ates I, Altay M, Yilmaz FM, Topcuoğlu C et al.
2. Y asar Durmus S, Sahin NM, Ergin M, Neselioglu S, Aycan Z
1. Altiparmak IH, Erkus ME, Sezen H, Demirbag R, Gunebakmaz

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65. Balta B, Erdogan M, Alisik M, Kiraz A, Akalin T et al. Does thiol-disulphide balance show oxidative stress in different MEFV mutations? Rheumatology International 2018; 38: 97-104.

66. Dogru A, Balkarli A, Cetin GY, Neselioglu S, Erel O et al. Thiol/disulfide homeostasis in patients with ankylosing spondylitis. Bosnian Journal of Basic Medical Sciences 2016; 16: 187.

67. Altinmel Acoglu E, Erel O, Yazilislas F, Bulbul M, Oguz MM et al. Changes in thiol/disulfide homeostasis in juvenile idiopathic arthritis. Pediatrics International 2018; 60 (6): 593-596.

68. Ozler K, Erel O, Gokalp O, Avcioglu G, Neselioglu S. Is there a relationship between dynamic thiol/disulfide homeostasis and osteoarthritis progression? Archives of Physiology and Biochemistry 2019: 1-7.

69. Fidan F, Alkan BM, Uğurlu FG, Bozkurt S, Sezer N et al. Dynamic thiol/disulphide homeostasis in patients with fibromyalgia. Archives of Rheumatology 2017; 32: 112-117.

70. Demirseren DD, Cicek C, Alisik M, Demirseren ME, Aktaş A et al. Dynamic thiol/disulfide homeostasis in patients with basal cell carcinoma. Cutaneous and Ocular Toxicology 2017; 36: 278-282.

71. Emre S, Demirseren DD, Alisik M, Aktaş A, Neselioglu S et al. Dynamic thiol/disulfide homeostasis and effects of smoking on homeostasis parameters in patients with psoriasis. Cutaneous and Ocular Toxicology 2017; 36: 393-396.

72. Emre S, Kalkan G, Erdogan S, Akta A, Ergin M. Dynamic thiol/disulfide balance in patients with seborrheic dermatitis: a case-control study. Saudi Journal of Medicine and Medical Sciences 2020; 8: 12.

73. Akbas A, Kilinc F, Sener S, Aktaş A, Baran P et al. Investigation of thiol-disulphide balance in patients with acute urticaria and chronic spontaneous urticaria. Cutaneous and Ocular Toxicology 2017; 36: 205-210.

74. Uysal P, Avcil S, Neselioglu S, Bicer C, Catal F. Association of oxidative stress and dynamic thiol-disulfide homeostasis with atopic dermatitis severity and chronicity in children: a prospective study. Clinical and Experimental Dermatology 2018; 43: 124-130.

75. Kilinc F, Sener S, Akbas A, Neselioglu S, Erel O et al. Investigation of dynamic thiol-disulfide homeostasis in alopecia areata patients. Journal of Advances in Medicine and Medical Research 2017; 21 (6): 1-7.

76. Sagdic HM, Ucar F, Tetikoglu M, Aktaş S, Ozcura F et al. Investigation of dynamic thiol-disulfide homeostasis in age-related cataract patients with a novel and automated assay. International Ophthalmology 2017; 38: 655-661.

77. Gulpamuk B, Koc M, Karatepe MS, Yildiz A, Erel O et al. Novel assay assessment of oxidative stress biomarkers in patients with keratoconus: thiol-disulfide homeostasis. Current Eye Research 2017; 42: 1215-1219.

78. Tetikoglu M, Aktaş S, Sagdic HM, Ozcura F, Ucar F et al. Thiol disulfide homeostasis in pseudoxefoliation syndrome. Current Eye Research 2017; 42: 876-879.

79. Turkoglu EB, Dikci S, Celik E, Erel O, Neselioglu S et al. Thiol/disulfide homeostasis in patients with central serous chorioretinopathy. Current Eye Research 2016; 41: 1489-1491.

80. Altinkaynak H, Kurkuoglu PZ, Caglayan M, Yorgun MA, Yuksel N et al. A novel marker in acute central serous chorioretinopathy: thiol/disulfide homeostasis. International Ophthalmology 2018; 38: 175-181.

81. Arikian Yorgun M, Toklu Y, Altinkaynak H, Tanriverdi B, Ergin M et al. A novel tool for the assessment oxidative stress in age-related macular degeneration: thiol/disulfide homeostasis revisited. Current Eye Research 2016; 41: 1584-1589.

82. Aktaş S, Sağdıç HM, Tetikoglu M, Aktaş H, Özcura F et al. Dynamic thiol/disulfide homeostasis in patients with age-related macular degeneration. Arquivos Brasileiros de Oftalmologia 2017; 80: 234-237.

83. Yıldırım M, Turkyilmaz E, Neselioglu S, Alisik M, Avsar AF. Dynamic thiol-disulphide status in polycystic ovary syndrome and its association with the pathogenesis of the disease. Gynecologic and Obstetric Investigation 2017; 82: 54-59.

84. Özler S, Oztas E, Tokmak A, Ergin M, Isci E et al. The association of thiol/disulfide homeostasis and lipid accumulation index with cardiovascular risk factors in overweight adolescents with polycystic ovary syndrome. Clinical Endocrinology 2016; 84: 516-523.

85. Korkmaz V, Kurdoğlu Z, Alisik M, Turgut E, Sezgin OO et al. Thiol/disulfide homeostasis in postmenopausal osteoporosis. European Journal of Obstetric & Gynecology Reproductive Biology 2017; 216: 24-26.

86. Isik H, Sahbaz A, Timur H, Aynioglu O, Atalay Mert S et al. The use of thiol/disulfide as a novel marker in premature ovarian failure. Gynecologic and Obstetric Investigation 2017; 82: 113-118.

87. Eroğlu S, Haskul I, Aziz V, Yurtçu E, Karatas F et al. Dynamic thiol/disulfide homeostasis in patients with Uterine Myoma. European Journal of Obstetric & Gynecology Reproductive Biology 2017; 216: 24-26.

88. Turkyilmaz E, Yıldırım M, Cendeş BD, Baran P, Alisik M et al. Evaluation of oxidative stress markers and intra-extracellular antioxidant activities in patients with endometriosis. European Journal of Obstetric & Gynecology Reproductive Biology 2016; 199: 164-168.

89. Desdiçioglu R, Yıldırım M, Özbek B, Khic G, Erel O et al. Dynamic thiol/disulfide homeostasis in patients with vaginitis. Journal of Gynecology and Obstetrics 2018; 6: 1.

90. Özler S, Erel O, Oztas E, Ersoy AO, Ergin M et al. Serum thiol/disulfide homeostasis in pre eclampsia. Hypertension in Pregnancy 2015; 34: 474-485.

91. Yucel A, Sanhal CY, Daglar K, Kara O, Uygun D et al. Thiol/disulfide homeostasis in pregnant women with familial Mediterranean fever. Redox Report 2016; 21: 287-291.

92. Erkenekli K, Sanhal CY, Yucel A, Bicer CK, Erel O et al. Thiol/disulfide homeostasis in patients with idiopathic recurrent pregnancy loss assessed by a novel assay: Report of a preliminary study. Journal of Obstetrics & Gynaecology Research 2016; 42: 136-141.
105. Unal S, Ozel S, Karabulut E, Kahyaoglu S, Neselioglu S et al. Oxidative-Anti-oxidative Markers in Pregnant Women with Fetal Neural Tube Defects. Fetal and Pediatric Pathology 2019: 1-10.

106. Hanikoglu F, Hanikoglu A, Kucuksayan E, Alisik M, Gocener AA et al. Dynamic thiol/disulphide homeostasis before and after radical prostatectomy in patients with prostate cancer. Free Radical Research 2016; 50: S79-S84.

107. Tokgoz H, Tas S, Giray O, Yalcinkaya S, Tokgoz O et al. The change in serum Thiol/Disulphide homeostasis after transrectal ultrasound guided prostate biopsy. International Brazilian Journal of Urology 2017; 43: 455-461.

108. Gul M, Bugday MS, Erel O. Thiol-disulphide homeostasis as an oxidative stress marker in men with varicocele. Andrologia 2018.

109. Kozanhan B, Inanli I, Deniz CD, Iyisoy MS, Neselioglu S et al. Dynamic thiol disulphide homeostasis in operating theater personnel exposed to anesthetic gases. American Journal of Industrial Medicine 2017; 60: 1003-1009.

110. Yilmaz OH, Bal C, Neselioglu S, Buyuksekerici M, Gunduzoz M et al. Thiol/disulphide homeostasis in asphalt workers. Archives of Environmental and Occupational Health 2016; 71: 268-272.

111. Buyuksekerici M, Bal C, Sercant U, Gunduzoz M, Alisik M et al. Alteration of thiol-disulphide homeostasis in workers occupationally exposed to arsenic. Archives of Environmental and Occupational Health 2018; 73: 90-95.

112. Bal C, Agis ER, Gunduzoz M, Buyuksekerici M, Alisik M et al. Dynamic disulphide/thiol homeostasis in lead exposure denoted by a novel method. Toxicology and Industrial Health 2017; 33: 426-430.

113. Bal C, Agis ER, Buyuksekerici M, Gunduzoz M, Tutkun L et al. Occupational exposure to asphalt fume can cause oxidative DNA damage among road paving workers. American Journal of Industrial Medicine 2018; 61: 471-476.

114. Koc U, Tan S, Ertem AG, Gumus M, Ozbek B et al. Evaluation of thiol-disulphide homeostasis in radiation workers. International Journal of Radiation Biology 2017; 93: 705-710.

115. Gunduzoz M, Bal C, Buyuksekerici M, Neselioglu S, Nadir Ozis T et al. Evaluation of dynamic disulphide/thiol homeostasis in silica exposed workers. Balkan Medical Journal 2017; 34: 102-107.

116. Bal C, Buyuksekerici M, Koca C, Agis ER, Erdogan S et al. The compromise of dynamic disulphide/thiol homeostasis as a biomarker of oxidative stress in trichloroethylene exposure. Human & Experimental Toxicology 2016; 35: 915-920.

117. Buyukaslan H, Gulacti U, Goldemir MT, Giden R, Celik H et al. Serum thiol levels and thiol/disulphide homeostasis in gunshot injuries. European Journal of Trauma and Emergency Surgery 2019; 45: 167-174.

118. Ozayci S, Karateke F, Turan U, Kuvvetli A, Kilavuz H et al. A novel oxidative stress mediator in acute appendicitis: thiol/disulphide homeostasis. Mediators of Inflammation 2016; 2016: 6761050.

119. Eryilmaz MA, Kozanhan B, Solak I, Cetinkaya CD, Neselioglu S et al. Thiol-disulphide homeostasis in breast cancer patients. Journal of Cancer Research & Therapeutics 2019; 15: 1062-1066.
120. Polat M, Ozcan O, Sahan L, Ustundag-Budak Y, Alisik M et al. Changes in thiol-disulfide homeostasis of the body to surgical trauma in laparoscopic cholecystectomy patients. Journal of Laparoendoscopic & Advanced Surgical Techniques 2016; 26: 992-996.

121. Akkus O, Kaypakli O, Koca H, Topuz M, Kaplan M et al. Thiol/disulphide homeostasis in thoracic aortic aneurysm and acute aortic syndrome. Biomarkers in Medicine 2018; 12 (4).

122. Ergin M, Caliskanturk M, Senat A, Akturk O, Erel O. Disulfide stress in carbon monoxide poisoning. Clinical Biochemistry 2016; 49: 1243-1247.

123. Solak I, Cetinkaya CD, Gederet YT, Kozanhan B, Erel O et al. Effects of smoking on thiol/disulfide homeostasis. European Review for Medical and Pharmacological Sciences 2018; 22: 2477-2482.

124. Guney T, Kanat IF, Alkan A, Alisik M, Akinci S et al. Assessment of serum thiol/disulfide homeostasis in multiple myeloma patients by a new method. Redox Report 2017; 22: 246-251.

125. Zubarioglu T, Kiykim E, Cansever MS, Neselioglu S, Aktugu-Zeybek C et al. Evaluation of dynamic thiol/disulphide homeostasis as a novel indicator of oxidative stress in maple syrup urine disease patients under treatment. Metabolic Brain Disease 2017; 32: 179-184.

126. Ergin Tuncay M, Erkilic A, Gunes A, Nural C, Erel O. A remarkable point for evaluating the severity of burns: thiol-disulfide profile. Burns 2019.