Simple Modified Colorimetric Methods for Assay of Total Oxidative Stress and Antioxidant Defense in Plasma: Study in Diabetic Patients

Abstract

Background: It is well established that oxidative stress and antioxidant defense of our body is associated with diabetes mellitus. Researchers have used different methods for estimation of oxidative stress and antioxidant defense, all of which have been criticized because of their limitations. Highly sophisticated methods are also not feasible in most of the clinical laboratories. The current study was aimed to estimate Total Oxidative Stress (TOS) and Total Antioxidant Defense (TAD) levels in plasma in patients with diabetes mellitus by two simple colorimetric tests developed and standardized in our laboratory and to find out their relationship with the fasting blood glucose (FBG) levels.

Methods and Findings: Plasma TOS and TAD levels and fasting blood glucose (FBG) were measured, in sixty two recently diagnosed diabetic patients and compared with similar number of healthy volunteers as controls. The TOS values in patients (28.35 ± 11.21 mM of H₂O₂/l) are significantly higher (P<0.001) than that in the control group (11.01 ± 4.55 mM of H₂O₂/l) whereas the TAD values of patients (113.71 ± 100.98 mM/l equivalent of trolox) are significantly (Z<0.001) less than that of the controls (338.87 ± 167.34 mM/l equivalent of trolox). There is a significant positive correlation of FBG levels with TOS (r=0.269, P=0.034, Figure 3) and a significant negative correlation with TAD values(r=-0.267, P=0.036, Figure 4).

Conclusions: The present study further reveals that diabetes mellitus is associated with oxidative stress. Plasma total oxidative stress and antioxidant defense is well correlated with FBG levels. Future study containing measurement of TOS and TAD values in plasma or in other body fluids can be conducted using these simple colorimetric and cost-effective methods even in a small clinical laboratory.

Keywords: Type 2 diabetes mellitus, Total oxidative stress, Total antioxidant defense, Fasting blood glucose

Introduction

In the recent years the measurement of oxidative stress and antioxidant status in plasma, other body fluids and extracts has been considered to be included in the methodology by many researchers in their studies. Monitoring oxidative stress in humans is achieved by assaying products of oxidative damage or by investigating the potential of an organism, tissue or body fluids to withstand further oxidation. Unfortunately, there is little consensus about the selection of parameters of oxidative stress or antioxidant state in defined patients or diseases. This is not only due to the uncertainty whether or not a certain parameter is playing a causative role. Either distinct antioxidant components are assayed or the total antioxidant potency is estimated by the combined reducing activities of a given body fluid, especially plasma [1]. The plasma total antioxidant capacity is modulated either by radical overload or by intake of dietary antioxidants and can therefore be regarded as more representative of the in vivo balance between oxidizing species and antioxidant compounds, known and unknown, measurable and not measurable, than the concentration of single, selected antioxidant. Moreover, the methods of determination described in literature represent very different levels of analytical practicability, costs, and quality. Generally accepted reference ranges and interpretations of pathological situations are lacking as well as control materials. So the availability of rapid and simple assays which measures the combined effect of the overall oxidative stress and antioxidant defenses in blood is of fundamental importance to provide an index of ability to resist oxidative damage.

Many analytical methods have been developed at present to determine the antioxidant activity such as the total radical trapping antioxidant parameter (TRAP) assay [2], the oxygen radical absorbance capacity (ORAC) assay [3] as well as dichlorofluorescein-diacetate (DCFH-DA) [4], phycocerythrin (R-Pe) [5] and crocin bleaching assays [6] are included in the first category of methods. The second approach to measure antioxidant capacity in plasma is to quench a stable and pre-formed radical that does not act as pro-oxidant. Trolox Equivalent Antioxidant Capacity (TEAC) assay [7] Ferric Reducing Ability of Plasma (FRAP) assay [8] cyclic voltammetry [9] procedure are included in this second category of methods.

However, the test kits for photometric determinations applicable to small laboratories or Points of Care are increasingly available. In the screening of such simplified procedures, we found the FORT (Free Oxygen Radical Test) [10,11] and FORD (Free Oxygen Radical Defense) method developed by Callegari, Parma, Italy, Catellani Group, has been used in some studies [10].

We found the non-lyophilized chromogen in solution procured and used in our laboratory (lyophilized form used in FORT assay-4-amino-N-isopropyl-aniline hydrochloride) is very unstable and even when stored at -20°C, the reagent degraded within 3-4days. Other researchers also observed the same and criticized the method [12]. The chromogen (N,N-dimethyl-p-phenylenediamine sulphate) used for FORD test was found to be more stable even at room temperature. The current study was aimed to estimate Total Oxidative Stress (TOS) and Total Antioxidant Defense (TAD) levels in plasma in patients with diabetes mellitus by two simple colorimetric tests developed and standardized in our laboratory and to find out their relationship with the fasting blood glucose (FBG) levels.

Methods

Human subjects, controls, selection criteria: Sixty-two diabetic patients (30 males and 32 females) were enrolled for the present study, with their ages ranging from 20 to 50 years, and similar number of healthy voluntary controls (37 males and 25 females) of the same age group was recruited for the study. The study was preapproved by the Institutional Ethics Committee of N.R.S. Medical College, Kolkata, India. Patients having other endocrine disorders like thyroid disorders, pregnant mothers, patients suffering from polycystic ovarian disease, renal failure, any malignant disease and patients receiving antioxidant agents are excluded from the study.

Sample Collection: 5ml of whole blood (after 8 to 10 hours calorie deprivation) was drawn aseptically from superficial vein, in heparin containing vials, centrifuged for 5 minutes at 2500 r.p.m., plasma was used for estimation of different biochemical parameters.

Assay of Total Oxidative Stress (TOS) And Total Antioxidant Defense (TAD) in plasma:

Reagent Preparation:

1. Chromogen Reagent solution: Freshly prepared 100 milimolar working chromogen solution (N,N-dimethyl- p-phenylenediamine sulphate, Aldrich, Sigma), by dissolving 23.5 mg in 10 ml of 20 milimolar PBS (pH 7.4) and kept in 4-8°C for use during assay.

2. Acetate Buffer (pH 5.2): prepared by mixing 0.1N acetic acid (570 µl to 500 ml deionized water) and 0.1N sodium acetate (2.72g in 200ml) in 42: 158 proportions.

3. 20 Molar PBS (pH 7.4): Prepared by mixing 80.4ml of Solution A (Solution A prepared by dissolving 11.47 gm of disodium hydrogen phosphate dissolved in 1000 ml of deionized water) to 19.6 ml of Solution B(Solution B prepared by dissolving 9.08 gm of potassium dihydrogen phosphate to 1000ml of deionized water), and pH adjusted to 7.4

4. 7.6 Molar (0.26 mg/l) H₂O₂ prepared by adding 88 microliter of 30% H₂O₂ to 100 ml of deionised water

5. 100 m Molar FeCl₃ ( MW 162.21) solution; prepared by adding 1.622 gm FeCl₃ in 100ml of deionized water

6. 2.5 milimolar Trolox solution; prepared by dissolving 15.6 mg Trolox (MW 250.29) in 25 ml of 20 m Molar PBS and stored in 0-4°C in aliquots.

Estimation of Total Oxidative Stress (TOS) in plasma: [10, 11]

Principle: This test is based on iron catalysed breakdown
of hydroperoxides into alkoxyl (RO•-) and peroxy (ROO•-) radicals which reacts with the chromogen (N,N-dimethyl- p-phenylenediamine sulphate) towards formation of a colored compound the absorbance of which is photometrically detectable. The intensity of the color correlates directly with the quantity of radical compounds, according to the Lambert-Beer’s law, and it can be related to the oxidative status of the sample.

**Assay Procedure:** 100 microliters of plasma diluted to 20 times in PBS, was dissolved in one ml of acetate buffer. 25microliters of working chromogen solution added, absorbance taken at 505nm by 6 min time-scan in UV-VIS spectrophotometer. The absorbance values obtained at 4 to 6 minutes for each sample against blank, were compared to the curve obtained using \( \text{H}_2\text{O}_2 \).

**Standardization:** Standard curve of TOS by modified FORT was prepared using different dilutions of hydrogen peroxide \( \text{H}_2\text{O}_2 \) in millimolar concentrations per liter and difference in absorbance values taken at 505nm in a six-minute time-scan between 6th and 4th minute (increase in absorbance noted maximum in this period) as depicted in **Figure 1**. Each data is the mean of three different values by the same procedure performed in four different occasions. The assay was completed in 6 minutes.

Intra assay and inter assay coefficients of variation: The maximum intra assay variation at 4 min was 4.762 and that at 6 min was 4.145 and inter assay variations at 4 and 6 minutes were 2.717 and 2.105 respectively for TOS. The maximum sensitivity of the assay was 1.22 milimol/l \( \text{H}_2\text{O}_2 \) and the linearity was up to 120milimol/l.

**Estimation of Total Antioxidant Defense (TAD) in Plasma:** [10]

**Principle:** In an acidic medium (pH=5.2) and a suitable oxidant (FeCl₃), the chromogen (N, N-dimethyl- p- phenylenediamine sulphate) develops a stable and colored radical cation that is photometrically detectable at 505 nm at 37° C. Antioxidant compounds in the sample reduce the radical cation of the chromogen, quenching the color and producing a discoloration of the solution, which is proportional to their concentration. The absorbance values obtained for the samples are compared with a standard curve obtained using Trolox.

**Assay procedure:** One ml of acetate buffer (pH=5.2) taken in a test tube. 25 microliter chromogen reagent that contains N, N-dimethyl- p- phenylenediamine sulphate and 10 microliter \( \text{FeCl}_3 \) solutions are added. 10 microliter of 20 times diluted plasma added to the mixture, the antioxidant compounds in the sample reduce the chromogen, quenching the color and producing a discoloration of the solution, which is proportional to their concentration.

**Standardization:** Standard curve of assay of TAD was prepared by using different dilutions of trolox, a potent antioxidant and it quenches the color of the chromogen maximally between 4th and 6th minute. With different concentrations of trolox, (6-hydroxy-2, 5,7,8-tetramethylchroman-2-carboxylic acid), a water soluble form of vitamin E, the difference in absorbance between the absorbance values at 505nm at 4th and 6th minute of the 6 minutes time scan was plot and a calibration curve was constructed (**Figure 2**).

Intra-assay and inter-assay coefficients of variation: The maximum intra-assay variations at 4 and 6 minutes were 7.914 and 9.009 respectively and inter-assay variation at 4minutes and 6minutes were 2.173 and 4.717 respectively, whereas the linearity ranged from 0.25 to 10.0 mmol/l equivalent of trolox.

**Estimation of Fasting Blood Glucose:** Estimation of fasting blood glucose (FBG) was done by GOD-POD method using standardized reagent kit.

**Storage:** All the above reagents in our method should be stored for future use in 6 to 8 degree for two weeks except the chromogen reagent (N, N-dimethyl- p- phenylenediamine sulphate) and \( \text{FeCl}_3 \) which were required to be prepared freshly before the tests. Plasma samples were stored in -40˚C deep freezer. Tests were performed within a week after sample collection.

**Statistical Analysis:** Data were expressed as mean ± standard deviation (SD), comparison of data was done using unpaired two-tailed Students’ t-test and Pearson’s correlation, \( P<0.05 \) was considered as significant. Statistical analysis was done using Microsoft Office Excel-2007 and SPSS Statistics version 2020.
Results

The clinical and biochemical findings of the current study based on the diabetic patients are summarized in Table 1.

In our study, the fasting blood glucose level in patients (118.07 ± 25.75 mg/dl) are significantly (P<0.001) higher than that of the healthy controls (79.06 ± 12.64 mg/dl). We found that TOS values in patients (28.35 ± 11.21 milimol of H\textsubscript{2}O\textsubscript{2}/l) are also significantly higher (P<0.001) than that in the control group (11.01 ± 4.55 milimol of H\textsubscript{2}O\textsubscript{2}/l). In contrast, the TAD values of patients (113.71 ± 100.98 milimol/l equivalent of trolox) are significantly (Z<0.001) less than that of the controls (338.87 ± 167.34 milimol/l equivalent of trolox). There is a significant positive correlation of FBG levels with TOS (r=0.269, P=0.034, Figure 3) and a significant negative correlation with TAD values (r=-0.267, P=0.036, Figure 4).

Discussion

Diabetes mellitus has already being established as a disease associated with oxidative stress [13-15]. Several studies have reported that the disease causation and progression is due to increased oxidative stress along with deficient antioxidant defense mechanisms [16]. Free radicals play a major role in the etiopathogenesis of this multifactorial disease [17-19]. Glucose toxicity and generation of different advanced glycated end products (AGE), is associated with the generation of free radicals. It has been shown that elevated extra and intra-cellular glucose concentrations result in an oxidative stress, which is defined as an imbalance between prooxidants and antioxidants. Several mechanisms seem to be involved in the genesis of this oxidative stress, which has been reported both in experimental diabetes in animals and in type 1 and type 2 diabetic patients: glucose auto-oxidation, protein glycation and formation of advanced glycation end-products and the polyol pathway. Reciprocally, oxidative stress is involved in the origin of type 1 diabetes, especially via the apoptosis of pancreatic beta-cells, as well as insulin resistance in type 2 diabetes. Macromolecules such as molecules of extracellular matrix, lipoproteins and deoxyribonucleic acid are also damaged by free radicals in diabetes mellitus [20].

Comprehensive studies with total oxidative stress and total antioxidant defense status in this condition are limited. Direct measurement of free radical levels in biological fluids is extremely difficult, since their half life is short. Electron Paramagnetic Resonance (EPR) spectroscopy is the only direct method for detecting radicals as such with the help of ‘spin-trap’ reagent, but is still far from being a ‘routine procedure’ [21,22]. However, most of the studies have been conducted with different methods consisting of some of the oxidative stress parameters in the tissues or in plasma separately, but the total effects of both the oxidative stress as well as the antioxidant defense have been reported in very few literatures. The three most widely used colorimetric methods for measuring oxidative stress are i) TMB peroxidase assay (3,5,3’,5’-tetramethyl benzene) ii) d-ROMs test (ROM-Reactive Oxygen Metabolites), sometimes referred to as the FORT test. iii) FOX (Ferrous Oxidation in Xylenol orange) assay. [23]. Some authors have reported that dROMs or FORT test does not measure the true oxidative stress, but only the ceruloplasmin oxidase activity [21,24,25]. Other authors have claimed that inspire the fact that dROMs test mainly measures ceruloplasmin oxidase activity but it is related to the oxidative stress [26]. Assesment of solitary effect of a deietary antioxidant say flavenoids may be measured in vitro, but inside the body its effects are influenced by other substances. Various assays have been described in which the putative antioxidant is added to a reaction mixture in which free radicals are generated such as 2,2'-azinobis3-ethylbenzthiazoline-6-sulfonic acid (ABTS) assay, ferric ion-reducing ability (FRA) assay etc. [27]. Commerically marketed kits to measure TAD are very expensive and time consuming [28].

Because of the limitations of original FORT test (the nonlyophilized form of original chromogen in solution turned out unstable), we have used a different chromogen based on the above principle.
Table 1. Clinico-biochemical parameters of the diabetic patients.

<table>
<thead>
<tr>
<th>Variables</th>
<th>Patient mean ± SD</th>
<th>Control mean ± SD</th>
<th>P value</th>
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</thead>
<tbody>
<tr>
<td>Age(years)</td>
<td>44.03 ± 5.27</td>
<td>43.63 ± 5.56</td>
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<tr>
<td>Sex(M/F)</td>
<td>30/32</td>
<td>37/25</td>
<td></td>
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<tr>
<td>Body mass index(BMI)</td>
<td>24.15 ± 4.09</td>
<td>24.81 ± 2.63</td>
<td>NS</td>
</tr>
<tr>
<td>Fasting Blood Glucose(mg/dl)</td>
<td>118.07 ± 25.75</td>
<td>79.06 ± 12.64</td>
<td>&lt;0.001*</td>
</tr>
<tr>
<td>Total Oxidative Stress:TOS (milimol of H_{2}O_{2}/l)</td>
<td>28.35 ± 11.21</td>
<td>11.01 ± 4.55</td>
<td>&lt;0.001*</td>
</tr>
<tr>
<td>Total Antioxidant Defence: TAD(milimol/l equivalent of trolox)</td>
<td>113.71 ± 100.98</td>
<td>338.87 ± 167.34</td>
<td>&lt;0.001*</td>
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The same chromogen was also useful for measurement of antioxidant defense. We have further modified and standardised to develop these two modified simple colorimetric methods in our laboratory. As both these tests do not require any sophisticated instruments and are cost effective, may be used by the researchers intending to study oxidative stress parameters working even in a small sized clinical laboratory and in a rural set-up.

**Conclusions**

We have developed two simple modified colorimetric tests for study of Total Oxidative Stress (TOS) and Total Antioxidant Defense (TAD) in our laboratory using a single chromogen, N,N-dimethyl-p-phenylenediamine sulphate. The present study further reveals that plasma total oxidative stress and antioxidant defense is well correlated with FBG levels in diabetes. Future study containing measurement of TOS and TAD values in plasma or on other body fluids can be conducted using these simple colorimetric and cost-effective methods even in a small clinical laboratory.

**Strength and weakness of the study:** As both these tests do not
require any sophisticated instruments and are cost effective, may be used by the researchers intending to study oxidative stress parameters working even in a small sized clinical laboratory and in a rural set-up which may not have a spectrophotometer but a simple colorimeter which can measure FBG, TOS and TAD. But the nonlyophilized form of chromogen is not very stable in the solution and practically everyday the chromogen and FeCl₃ have to be prepared before the two tests were done. The sample size for the current study was small as it was a pilot study. Further studies are required to validate the results of the current study with large sample size.
References