EVALUATION OF SERUM-FREE IRON AND GLYCATED HEMOGLOBIN IN UNCONTROLLED TYPE-II DIABETIC PATIENTS

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ABSTRACT

Objective: Diabetes mellitus (DM) is a common health problem in the world. Free iron may contribute to the pathogenesis and progress of this disease and its complications. Iron causes hyperinsulinemia by decreasing the insulin uptake and metabolism by hepatocytes. Elevated iron stores are commonly found in insulin resistance. In its free form, it is known to induce oxidation of biomolecules by producing harmful hydroxyl radicals. In this study, we aimed to estimate and compare the serum levels of free iron in diabetes and healthy individuals.

Methods: This study included 244 subjects in two groups. Group-I comprised 204 subjects with DM and Group-II comprised 40 healthy subjects. Blood sugar, free iron, and glycated hemoglobin were analyzed in blood samples using standard kits. The results of all the parameters were expressed as mean±standard deviation. Student t-test was done to assess the statistical significance between two groups. The association between the parameters was studied by Pearson correlation.

Results: In this study, we found a significant increase in serum free iron in Group-I (p<0.01) when compared with Group-II. A significant correlation between the serum free iron and glycated hemoglobin (r=0.59; p<0.001) and fasting blood sugar (r=0.43; p<0.001) was found.

Conclusion: The elevated serum free iron in uncontrolled diabetes may contribute to oxidative stress, which may be associated with complications of diabetes.

Keywords: Free iron, Glycated hemoglobin, Hyperinsulinemia, Diabetes mellitus.

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INTRODUCTION

Iron is a very important transition metal for the cells in the body, and its abnormal homeostasis is associated with the pathogenesis of various chronic diseases, including diabetes [1,2]. Iron plays a key role in vital biochemical activities [3]. Iron is potentially hazardous because of its ability to participate in the generation of powerful oxidant species such as hydroxyl radical [4,5]. Dietary iron is a critical determinant of body's iron status since once absorbed it is not actively excreted [6,7]. However, the ability of iron to cycle between its two stable oxidation states is also potentially to generate reactive oxygen or nitrogen species (ROS or RNS) such as hydroxyl radical via Fenton's and Haber-Weiss reactions [3,6,8]. To control and balance the production of ROSs and RNSs, the cell builds up a set of antioxidants and detoxifying enzymes such as superoxide dismutase, catalase, and glutathione peroxidase that can scavenge excessive ROSs or RNSs beyond the antioxidant capacity of the organism called oxidative stress is encountered in many pathological conditions such as diabetes and diabetic complications [3,6,9,10]. Iron can be released from ferritin by the action of reducing agents that convert Fe³⁺ into Fe²⁺ [1,11,12]. Apoferritin, the protein fraction of ferritin that holds oxidized iron molecules. When the concentrations of antioxidants are low, the reducing potential and anaerobiosis progressively increase, facilitating a rapid release of iron from ferritin. The overall result of oxidative reactions is an increase in the availability of free iron from the ferritin [1,11,13]. Iron influences insulin action and it interferes with insulin inhibition of glucose production by the liver [11,14-16]. Hepatic extraction and metabolism of insulin are reduced with increasing iron stores, leading to peripheral hyperinsulinemia. In fact, the initial and most common abnormality seen in iron overload conditions is liver insulin resistance [11,15].

The exact mechanism of iron-induced diabetes is mediated by 3 key mechanisms: “Insulin deficiency, insulin resistance, and hepatic dysfunction” [4,17]. The mechanisms for insulin resistance include the possibility of iron overload causing resistance directly or through hepatic dysfunction. Patients with unexplained hepatic iron overload mostly found to be insulin resistant, which suggests a common etiologic link among hepatic iron, hepatic dysfunction, and insulin resistance [18]. This study was aimed to determine serum free iron in person with diabetes and healthy individuals.

METHODS

This study was done at the Department of Biochemistry, Sri Ramachandra Medical College and Research Institute, Chennai - 116, and it was approved by our Institutional Ethical Committee (REF: CSP/12/JUL/24/111).

Selection of participants

The total number of subjects was 244. Study subjects were categorized into 2 groups.

Diabetes mellitus (DM) group consists of 204 subjects. The exclusion criteria are Type-I DM, hyperthyroidism, cancer, and pancreatitis. Inclusion criteria are Type-2 DM >30 years of age.

The healthy group consists of 40 subjects. Exclusion criteria are smokers, alcoholics, pregnant and lactating women. The inclusion criteria are >30 years of healthy persons without any disease.

Sample type

Serum, plasma, and whole blood.
Sample collection
The blood sample from 244 individuals was collected in fasting state using vacutainers with sodium fluoride and EDTA and without anticoagulants sterile gel tube. These tubes were centrifuged at 3500 rpm for 10 minutes, and serum was separated.

Blood sample analysis
Fasting blood sugar (FBS) by hexokinase method and free iron by direct iron assay using chromophore ferene were analyzed in blood sample using standard kits in Dimension RxL autoanalyzer, USA, and glycated hemoglobin (HbA1c) by high-performance liquid chromatography method was analyzed in blood sample using Bio-rad autoanalyzer, USA. Glucose present in plasma in expressed as mg/dl. Iron present in serum is expressed as µg%. HbA1c present in whole blood is expressed as g%.

Statistical analysis
The results of all the parameters were expressed as mean±standard deviation. Student t-test was done to assess the statistical significance between two groups. The association between the parameters was done by the correlation coefficient.

RESULTS
Our present study was done on 204 Type-II diabetes and 40 healthy subjects. A direct relationship was found between serum free iron and Type-II DM. We found a significant increase in serum free iron in Group-I 98.33±51.27 (p<0.01) when compared with Group-II 78.07±33.13 (Table 1).

In this study, we found a significant positive correlation between the serum free iron and FBS (r=0.43; p<0.001) (Fig. 1) and HbA1c (r=0.59; p<0.001) (Fig. 2), respectively.

DISCUSSION
In this present study, we have observed a significant increase in serum free iron in Group-I cases, compared to Group-II cases who were in good glycemic control [19]. Hyperglycemia observed in Group-I causes increased glycation of protein and releases the iron in its free state. Several studies have shown that serum free iron is significantly increased in Type-II DM [20,21]. This observation is contradicted by the studies of EI-Nabarawy et al. [1]. Glycemic control reflects the interaction between hyperinsulinemia with iron. This is further supported by evidence suggesting that hyperglycemia precedes the elevation of iron in diabetes [22,23].

In this present study, we found a positive correlation between FBS and free iron as well as free iron and HbA1c in Group-I. Increased glycation of proteins as a result of poor glycemic control triggers the increased release of free iron from glycated proteins like hemoglobin which generates a vicious cycle of hyperglycemia, glycation of hemoglobin, and increase in levels of free iron [18]. The relationship between free iron and Type-2 diabetes is bidirectional [24]. Insulin influences the iron uptake and storage by increasing the cell surface transferring receptors [25], reciprocally iron influences the insulin activity by interfering with glucose uptake and utilization [26]. This increased presence of free iron pool will enhance oxidant generation leading damage to biomolecules [8]. Free iron is known to induce oxidation of biomolecules through Fenton’s and Haber–Weiss reaction by producing harmful hydroxyl radicals [8,24]. Free radical formation may play a role in the pathogenesis of diabetes by disrupting insulin action and total body glucose disposal. The iron act as a powerful pro-oxidant and the oxidative stress is increased in glucose-intolerant states [27-29]. The presence of poor glycemic control, hyperglycemia, iron overload, and protein glycation will all lead to oxidative stress causing the early appearance of microvascular complications (retinopathy and nephropathy) [8,24,30]. The limitations of the present study, we have not assessed the markers of oxidative stress and insulin resistance.

Table 1: Demographical data and biochemical parameters in the study groups

| Variables         | Group I (n=204) | Group II (n=40) |
|-------------------|----------------|----------------|
| Age (in years)    | 55.74±10.72    | 40.35±8.83     |
| Sex (female: male)| 78:126         | 27:13          |
| FBS (mg%)         | 155.78±58.20***| 91.72±8.82     |
| HbA1c (g%)        | 8.33±2.12***   | -              |
| Serum free iron (µg%) | 98.33±51.27** | 78.07±33.13    |

**p<0.01, ***p<0.001. FBS: Fasting blood sugar, HbA1c: Glycated hemoglobin.

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Fig. 1: Correlation between serum free iron and fasting blood sugar in Group-I

Fig. 2: Correlation between serum free iron and glycated hemoglobin in Group-I

CONCLUSION
The significantly increased serum free iron and HbA1c in DM reveals the fact that serum free iron contributes indirectly to oxidative stress in DM.
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