

Original Research Article**Association of Major Depression with Serum Prolidase activity and Oxidative Stress****Abstract**

Aim: Stress is a major causative factor for the progression of major depressive disorder (MDD). The present study was aimed to know the association of serum prolidase activity (SPA) and oxidative stress with the progression of MDD.

Place and Duration of Study: The study was carried out at the Sir Sunder Lal Hospital, Department of Biochemistry, Institute of Medical Sciences (IMS), Banaras Hindu University (BHU), Varanasi, Uttar Pradesh, India. The duration of study was September-2011 to August-2016.

Methodology: 80 patients with MDD and 80 healthy controls of matched age and genders were selected. Serums SPA, total oxidant status (TOS), oxidative stress index (OSI) and total antioxidant status (TAS) were measured spectrophotometrically.

Results: Increased SPA, TOS, and OSI were observed in patients with MDD than healthy controls (all $P < 0.001$). However, TAS was significantly decreased ($P < 0.001$). SPA, TOS and OSI were also increased in patients with > 1 years of MDD than patients with ≤ 1 years of MDD. Positive, linear and significant correlations were observed between duration of MDD and SPA, and TOS, and OSI (all $P < 0.001$). However, negative, linear and significant correlation was observed between duration of MDD and TAS ($P < 0.001$).

Conclusions: The study concluded that SPA and oxidative stress have been significantly increased in the patients with MDD than healthy individuals. Increased SPA and oxidative stress have been significantly correlated to progression of MDD and may be responsible for its pathogenesis.

Key Words: Major Depressive Disorder, Oxidative Stress, Serum Prolidase Activity, Total Oxidant Status, Total Antioxidant Status, Oxidative Stress Index

1. Introduction

Major Depressive disorder (MDD) is a chronic most commonly as well as frequently occurring serious disorder that negatively affects the quality of life. It can alter the morbidity

32 as well as mortality [1]. Including suicide, it is associated with an overall 50% increase in the
33 risk of morbidity [2, 3]. The interaction of both genetic and environmental factor can play a
34 role in the development of depression [4]. Gender, age, socioeconomic status, stressful life
35 events, childhood adversity, and co-morbidities or medical childbirth are counted as risk
36 factor for depression [5].

37 Prolidase is a cytosolic manganese-dependent exopeptidase which cleaves dipeptides with
38 proline or hydroxy-proline at carboxy- terminal end [6]. Proline or hydroxy-proline is an end
39 product of prolidase that participates in collagen metabolism, cell growth and protein
40 synthesis. It involves in deactivations of neuropeptides and can influence the biological as
41 well as conformational properties of neuropeptides [6-8].

42 Oxidative stress is a condition of imbalances between oxidants and antioxidants [9]. Numbers
43 of evidences are supporting the involvement of oxidative and nitrosative stress in the
44 pathophysiology of MDD [10]. Altered status of both oxidants and antioxidants [(which
45 includes reactive oxygen species (peroxide), reactive nitrogen species (NO), glutathione,
46 vitamin E, zinc, coenzyme Q10, manganese superoxide dismutase and catalase)] have
47 reported in the patients with MDD [10, 11]. Several of our previous studies are suggesting
48 that altered status of prolidase activity have correlated to altered status of oxidative stress in
49 different diseases such as non-ulcer dyspepsia, diabetes, diabetic nephropathy, end stage renal
50 disease and Parkinson's disease [6, 8 and 12]. In our previous study, we have been observed
51 altered status of malondialdehyde, nitrite, ceruloplasmin, ascorbic acid and superoxide
52 dismutase in patients with MDD [11]. Thus in the continuation of our previous study, present
53 study aimed with an explorative study on the association of serum prolidase activity, TOS,
54 TAS, and OSI in the patients with MDD.

55

56 **2. Materials and Methods**

57 The study was conducted in the Department of Biochemistry, Institute of Medical Sciences
58 (IMS), Banaras Hindu University (BHU), Varanasi, India from the period of September-2011
59 to August-2016. The study was approved by the ethical committee of the IMS, BHU. Written
60 and signed consent form was taken from every studied subject.

61 A total 80 drug naive and fresh cases of major depressive disorder were included in the study.
62 Cases were diagnosed as per the DSM IV by the consultant psychiatrist. A total 80 healthy
63 subjects of matched age and sex were taken from the general population and considered as
64 control group. Cases and controls, which dependent on tobacco, alcohol or any other
65 substances or aged greater than 60 years were excluded from the study. All the control
66 subjects included for the study were healthy, not addicted to tobacco or alcohol, non-
67 diabetics, normotensive and showed no evidence of any chronic and/or acute infection. The
68 subjects failing to above inclusion criteria were excluded from the study. The subjects who
69 did not agreed to sign informed consent form were also excluded from the study.

70 **2.1 Specimen Collection**

71 From every studied subjects, 5ml of blood was withdrawn by venipuncture (from peripheral
72 vein) method in clean dry glass tube. Serum was separated by centrifugation at 3000 r.p.m.
73 for 10 minutes. All serum samples were stored at -80°C . Care was taken to avoid samples
74 hemolysis. Repetition of thaw of serum samples was also avoided.

75 **2.2 Estimation of serum prolidase activity (SPA)**

76 Reagents such as diluting solution, standard proline solution, 94 mmol/L glycyl-l- proline and
77 Chinard's reagent were prepared. Serum prolidase activity was measured with the use of our

78 previous standardized method [8, 13]. The enzyme activity was expressed in millimolar per
79 minute per liter ($\text{mmol Min}^{-1} \text{L}^{-1}$).

80 **2.3 Estimation of Serum Total Anti-oxidant Status (TAS)**

81 TAS of serum was estimated with using a method developed by Erel (2004) [14]. Following
82 reagents and procedures were used for the estimation of TAS-

83 **Reagent-1:** 75mM Clark and Lubs solution (pH 1.8) was prepared as follows; 5.591 gram of
84 potassium chloride (KCl) was diluted in 1000 mL of deionized water. 6.41 mL of 36.5%
85 hydrochloric acid (HCl) was dissolved in 1000 mL of deionized water. Then finally, 800 mL
86 of above prepared KCl solution was mixed with 200 mL of above prepared HCl solution (pH
87 maintained at 1.8). After this, 3.17 gram of orthodiansidine dihydrochloride (final 10 mM)
88 was mixed in this solution. Mixing was followed by addition of 0.01764 gram of $\text{Fe}(\text{NH}_4)_2$
89 $(\text{SO}_4)_2 \cdot 6\text{H}_2\text{O}$ (final 45 μM). This final reagent was used as reagent-1 for the assay. At 4°C, it
90 is stable for six months. (Note; orthodiansidine is a carcinogenic and very toxic substance.
91 Thus, gloves and face mask were used during the handling of this substance)

92 **Reagent-2:** 7.5 mM of H_2O_2 solution was prepared (0.641mL of 35% H_2O_2 (Sigma Aldrich)
93 was maintained to 1000 mL volume with the Clark and Lubs solution). At 4°C, it is stable for
94 one month.

95 **Procedure:** 2000 μl of Reagent-1 was mixed with 50 μl of serum sample and 100 μl of
96 Reagent-2. Absorbance was taken at 444 nm. The first absorbance was taken before mixing
97 of Reagent-1 and Reagent-2 (this OD was deducted from respective test as sample blank).
98 The last reading of test was taken after 3- 4 minutes of mixing of Reagent-1 and Reagent-2.
99 (Final OD = OD of reagent-1 plus sample plus reagent-2 - OD of reagent-1 plus sample).

100 Finally, TAS was estimated by preparation of standard linear calibration graph of Trolox (a

101 Vitamin-E analogue). For this, 2.0 mM of Trolox was serially diluted as 0.2 mM, 0.4 mM,
102 0.6 mM, 0.8 mM, 1.0 mM, 1.2 mM, 1.4 mM, 1.6 mM, 1.8 mM. Results of TAS were
103 expressed in millimolar Trolox equivalent per liter (mmol Trolox Eq./L).

104 **2.4 Estimation of Serum Total Oxidant Status (TOS)**

105 TOS of serum was estimated with using a method developed by Erel (2005) [15]. Following
106 reagents and procedures were used for the measurement of TOS-

107 **Reagent-1:** 150 μ M xylenol orange (114 mg) and 140 mM NaCl (8.18 gram) were dissolved
108 in 900 mL of 25 mM- H_2SO_4 solution. Then, 100 mL of glycerol was added to this solution
109 (final concentration of glycerol, 1.35 M). The pH of solution was maintained at 1.75. At 4
110 $^{\circ}C$, reagent is stable for six month.

111 **Reagent-2:** 1.96 gram of ferrous ammonium sulfate (5 mM) was mixed with 3.17 gram of o-
112 dianisidine dihydrochloride (10 mM) in 1000 mL of 25 mM- H_2SO_4 solution. At 4 $^{\circ}C$,
113 reagent is stable for six month.

114 **Procedure:** 2250 μ l of Reagent 1 was added to 350 μ l of serum sample and 110 μ l of
115 Reagent 2. The bichromatic absorbance was taken (main wavelength 560 nm, secondary
116 wavelength 800 nm).The first absorbance was taken before mixing of Reagent-1 and
117 Reagent-2 (this OD was deducted from respective test as sample blank). The last reading of
118 test was taken after 3-4 minutes of mixing of Reagent-1 and Reagent-2. For final absorbance,
119 absorbance of a test at 800 nm was deducted from the absorbance at 560 nm of respective test
120 (Final Absorbance = Absorbance at 560 nm – Absorbance at 800 nm).

121 The assay was calibrated with H_2O_2 standard solution. 100 μ mol of standard H_2O_2 solution
122 was serially diluted with deionized water and results were drawn with the help of linear

123 calibrated graph. Thus, results of test was expressed in terms of micromolar hydrogen
124 peroxide equivalent per liter ($\mu\text{mol H}_2\text{O}_2 \text{ Eq./L}$).

125 **2.5 Calculation of Oxidative Stress Index (OSI)**

126 Values of OSI were estimated with the help of following formula [8, 12 and 13]–

$$\begin{aligned} 127 & \qquad \qquad \qquad \text{TOS (mmol H}_2\text{O}_2 \text{ Eq. / L)} \\ 128 \text{ OSI (Arbitrary Unit) } &= \frac{\text{-----}}{\text{TAS (\mu mol Trolox Eq. / L)}} \\ 129 & \qquad \qquad \qquad \text{TAS (\mu mol Trolox Eq. / L)} \end{aligned}$$

130 **2.6 Statistical Analysis**

131 Standard statistical methods were used for the data interpretation. Microsoft office excels
132 worksheet and SPSS (16) software was used for the calculation. Data, which follow normal
133 distributions, were expressed as mean \pm SD (standard deviation). A p-value < 0.05 was
134 considered as significant; student's t-test was used. Pearson's correlation was calculated for
135 the correlative observations.

136 **3. Results**

137 In present study, Non-significant differences of mean age of the cases and controls were
138 observed 39.11 ± 10.64 and 39.70 ± 9.89 years respectively (Table-1).

139 **3.1 Observed status of SPA, TOS, TAS and OSI**

140 The observed SPA, TOS and OSI were significantly increased in the cases than controls ($P <$
141 0.001 , Table-1). However, TAS was significantly decreased in the cases than controls ($P <$
142 0.001 , Table-1).

143

144

145 **3.2 Status of Serum SPA, TOS, TAS and OSI with respect to progression of MDD**

146 Total 80 cases of patients with MDD were categorized on the basis of duration of disease
147 (Group-1; Duration of disease \leq 1 years, range 0.42 – 1 year, n = 50, and group-2; Duration
148 of disease $>$ 1 years, range 1.5 – 6 years, n = 30). In this regards, significantly increased SPA,
149 TOS and OSI were observed among the group of patients with disease duration $>$ 1 year than
150 the patients with disease duration \leq 1 year ($P < 0.001$, Table-2). However, serum TAS was
151 significantly decreased in the group-2 than group-1 ($P < 0.001$, Table-2).

152 **3.3 Correlation of SPA and Serum Oxidative Stress with progression of MDD**

153 On correlative observations, it was observed that positive, linear and significant correlation
154 was observed between duration of disease (MDD), and SPA, and TOS, and OSI ($r = 0.879$,
155 0.646 and 0.695 respectively, all $P < 0.001$) (Figure-1A, 1B and 1D). However, negative
156 linear and significant correlation was observed between duration of disease (MDD) and
157 serum TAS ($r = - 0.619$, $P < 0.001$) (Figure-1C).

158 **4. Discussion**

159 In our previous study we have observed that increased oxidative stress in term of different
160 individual oxidative stress markers in the patients with MDD [11]. Thus in present study we
161 planned to observe oxidative stress in term of TOS, TAS and OSI in the patients with MDD.
162 Along with this, serum prolidase activity (SPA) was also assessed. Present explorative study
163 included 80 subjects of both cases of MDD and healthy individuals of matched age and sex.
164 All the studied subjects have age below 60 years old.

165 Prolidase is an enzyme which cleaves the glycyl-l-proline and provides proline as end
166 product [6]. Proline is usually circulated in the central nervous system (CNS), and may be act
167 as neuromodulator in synaptic transmission [16, 17]. It is believed that glutamate is involved

168 in the etiology of depression [18]. Several present literatures represented that proline and
169 glutamate receptor interacts with each other [19]. It was shown that proline inhibited the
170 glutamate release in cerebrospinal fluid (CSF), which induced the glutamatergic signaling, in
171 the hippocampus [20, 21]. It has also reported in literature that increased proline may be
172 neurotoxic and damage brain by the decrease in glutamate uptake [22]. In present study, it is
173 observed that serum prolidase activity was increased in the patients with major depressive
174 disorder than the healthy individuals (Table 1). Same results are reported by Kokacya et al.,
175 2014, in Turkey population [23]. In addition to this, we also evaluated the serum prolidase
176 activity with respect to progression of major depression in term of duration of depression and
177 correlation. It is observed that serum prolidase activity was significantly increased in the
178 patients with > 1 years of MDD than the patients with ≤ 1 years of MDD (Table 2). Increased
179 serum prolidase activity was also significantly, positively and linearly correlated to duration
180 of MDD ($r = 0.879$, $P < 0.001$) (Figure 1A). Thus it seems that increased prolidase activity
181 can leads to increase in proline concentration in circulation. Increased proline concentration
182 may interfere with glutamate signaling of depressive patients and may be responsible for the
183 progression and pathogenesis of major depressive disorder. But further explorative study is
184 needed for better clarification.

185 An imbalance in the control of oxidants and antioxidants in human system leads to oxidative
186 stress. Previously, we have reported that the status of different oxidants (malondialdehyde
187 and nitrite) and antioxidants (superoxide dismutase, ascorbic acid and ceruloplasmin) were
188 not in balance manner in the patients with MDD. It's observed value have been altered as
189 compared to healthy individuals [11]. Delwing et al., 2003, reported that proline itself is able
190 to increase the oxidative stress in the brain [24]. In present study, we have been observed
191 significantly increased TOS and OSI in the patients with MDD than healthy individuals.
192 However, significantly decreased TAS has been observed (all $P < 0.001$, Table 1). Same

193 observations are done by Kokacya et al., 2014, in Turkey population [23]. Along with this,
194 we have been also observed association of oxidative stress with the progression of MDD.
195 TOS and OSI were significantly increased in the patients with > 1 years of MDD than the
196 patients with ≤ 1 years of MDD ($P < 0.001$, Table 2). While, TAS was significantly decreased
197 ($P < 0.001$, Table 2).

198 On correlative observations, it has been observed that duration of MDD was significantly,
199 positively and linearly correlated to TOS, and OSI ($r = 0.646$, $P < 0.001$ and $r = 0.695$, $P <$
200 0.001 respectively) (Figure 1B and 1D). While, significant and negative correlation was
201 observed between duration of MDD and TAS ($r = - 0.619$, $P < 0.001$) (Figure 1C). Thus, it
202 seems that altered different oxidative stress markers as well as increased proline (as product
203 of prolidase) may lead to increase in oxidative stress in patients with MDD. This increase in
204 oxidative stress may be responsible for the pathogenesis and progression of MDD.

205 **5. Conclusions**

206 The study concluded that serum prolidase activity, total oxidant status and oxidative stress
207 index have been significantly increased in the patients with major depressive disorder than
208 healthy controls. However total antioxidant status has been significantly decreased. It is also
209 concluded that serum prolidase activity and oxidative stress have been progressively
210 increased with the progression of major depressive disorder and may be associated with its
211 pathogenesis.

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276 **Table-1: Status of serum prolidase activity (SPA) and oxidative stress (TOS, TAS, and**
 277 **OSI) in cases and controls.**

Variables	Cases (n= 80)	Controls (n= 80)	P- value
Age (Years)	39.11 ± 10.64	39.70 ± 9.89	NS
Gender	M- 48, F- 32	M- 45, F- 35	NS
SPA (mmol Min ⁻¹ L ⁻¹)	112.92 ± 24.19	91.19 ± 20.91	< 0.001
TOS (µmol H ₂ O ₂ Eq./ L)	16.86 ± 2.59	13.17 ± 2.78	< 0.001
TAS (mmol Trolox Eq./L)	0.95 ± 0.33	1.56 ± 0.54	< 0.001
OSI (Arbitrary Unit)	21.22 ± 10.88	10.01 ± 5.16	< 0.001

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279 SPA – Serum Prolidase Activity, TOS – Total Oxidant Status, TAS – Total Antioxidant
 280 Status, OSI – Oxidative Stress Index, n – Numbers of Subjects, NS – Non Significant, M-
 281 Male, F-Female

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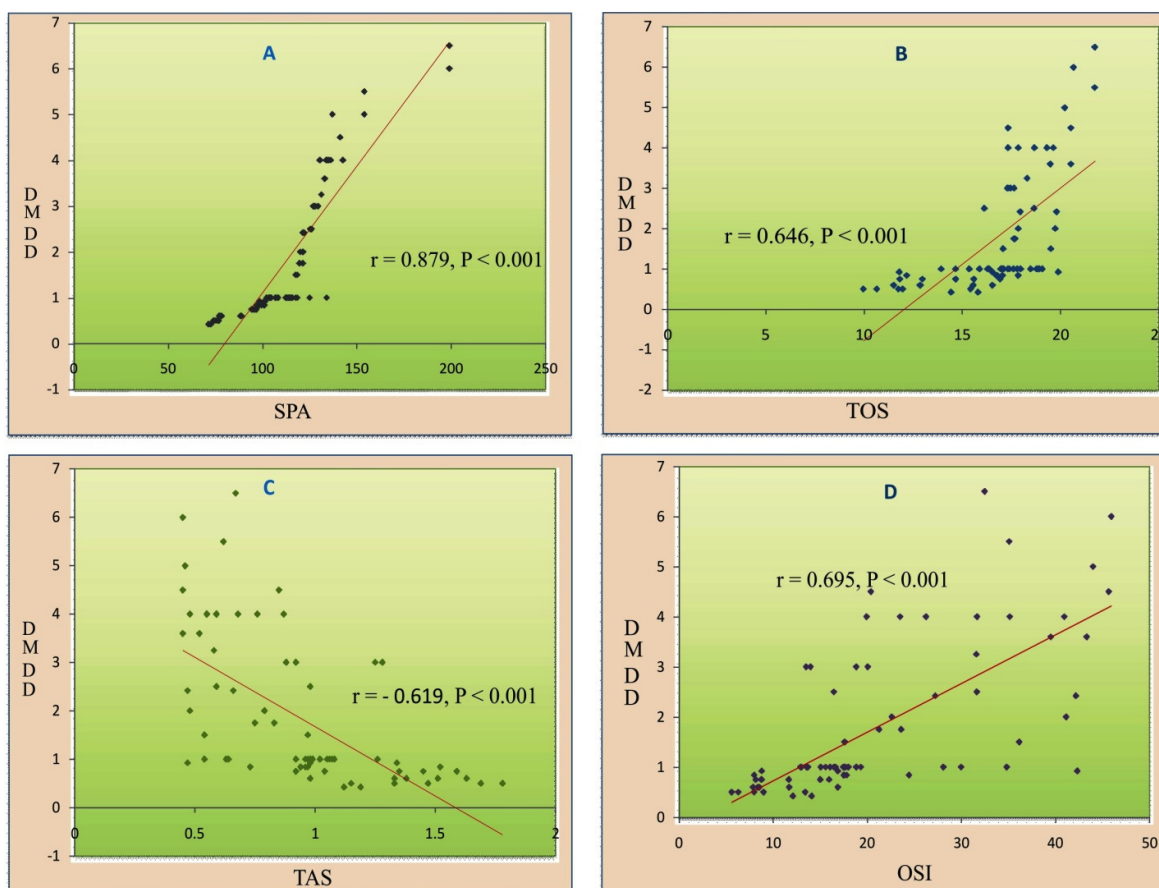
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287 **Table-2: Status of serum prolidase activity (SPA) and serum oxidative stress (TOS, TAS**
 288 **and OSI) with respect to disease duration (duration of major depression).**

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Variables	Duration of disease \leq 1 years Range 0.42 – 1 year, n = 50	Duration of disease $>$ 1 years Range 1.5 – 6 years, n = 30	P- value
SPA ($\text{mmol Min}^{-1} \text{L}^{-1}$)	99.50 ± 14.98	135.31 ± 19.67	< 0.001
TOS ($\mu\text{mol H}_2\text{O}_2 \text{Eq./L}$)	15.70 ± 2.43	18.78 ± 1.47	< 0.001
TAS (mmol Trolox Eq./L)	1.10 ± 0.28	0.69 ± 0.23	< 0.001
OSI (Arbitrary Unit)	15.85 ± 6.90	30.18 ± 10.45	< 0.001

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292

293 **Figure-1:** Scattered diagram showing Pearson's correlation between duration of depression
294 and SPA (Fig A), and TOS (Fig B), and TAS (Fig C), and OSI (Fig D). r = Pearson's
295 correlation coefficient, D-MDD = Duration of major depressive disorder in years.