Acute Fluoxetine Treatment Produce Anxiolytic Effects without Modulating Behaviour Response on Exposure to Forced Swimming Test
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Abstract

Stress is a condition which disturbs physiological and psychological homeostasis mechanism. Depression is a severe psychiatric disorder. Biogenic amine theory of depression illustrate that the low level of brain 5 hydroxy tryptamine (5-HT) and catecholamine leads to depressive symptoms. Immunological challenges can alter the tryptophan (TRP) metabolism, it is clinical indication of depression but stress also shift TRP metabolism. Fluoxetine is more effective anxiolytic drug as compared to others antidepressants. The aim of the current investigation is to examine the effects of fluoxetine administration on tryptophan metabolism and disposition in forced swimming test (FST) in rats. Albino Wistar rats were separated into three groups. Each group had 5 rats. Control animals received vehicle (DMF:Saline, 1:3 v/v) while test group treated with vehicle or fluoxetine (20 mg/kg) (i.p) 3.5 hr prior to FST . Present study shows that holo, total enzyme activity was inhibited when rats subjected to FST and serum total TRP concentration was decreased while liver, brain TRP, 5-HT and 5-HIAA levels were increased in vehicle treated FST rats. Pretreatment with fluoxetine did not reduce immobility however it inhibited holo, total and apo enzyme activity. Pretreatment of fluoxetine also decreased serum total TRP and brain 5HT concentration while increased liver TRP, brain TRP, and 5-HIAA in FST rats when compared with their respective controls. Acute treatment with fluoxetine did not reduced immobility but it decrease brain 5-HT concentration by converting into 5-HIAA in FST rats which shows anxiolytic effect of fluoxetine.

Key words: Forced swimming Test, Fluoxetine, Tryptophan, 5-hydroxy tryptamine, 5-hydroxy indole acetic acid, anxiolytic.

INTRODUCTION

Stress was initially defined by Hans Selye in 1936; He states that it is a condition which disturbs the physiological and psychological homeostasis mechanism of an organism [1]. Moreover chronic stress could be responsible to the progression of mental disorders. He proposed that the main factor which control stress response is hypothalamic-pituitary-adrenal (HPA) axis. Chronic stress can lead over secretion of glucocorticoids, increased concentration of cortisol and corticosterone can damage the hippocampal neurons. Depression is psychiatric disorders which affect 20% of an adult people overall the world. There are many factors which involve in pathophysiology of depression one of them is dysfunction of monoaminergic system.

This system is responsible for the behavioral and visceral characteristic of mood disorders [2]. The biogenic amine theory of depression illustrate that the low amount of brain 5 hydroxy tryptamine (5-HT) and noradrenaline or catecholamine lead to depressive symptoms[3-4].One of the essential factors in etiology of mental disorders like depression and schizophrenia is disturbance of serotonergic system[5].

To evaluate the mode of action of antidepressant and etiology, the serotonin, neurotransmitter system is extensively investigated. Long term administration of different class of antidepressant (AD) could improve the serotonergic function and proof the involvement of serotonin in depression [6]. Serotonergic activity is improved by ADs drugs. The deficiency of serotonergic system
might be possible due to substrate unavailability, reduced tryptophan hydroxylase activity, any abnormalities in 5-HT up take or release, defect in 5-HT receptor or contact with other neurotransmitters. Any changes of tryptophan (TRP) metabolism are a sign of clinical depression caused by immunological challenges [7] but stress also alter TRP metabolism.

Among all AD selective serotonin reuptake inhibitors (SSRIs) are commonly prescribed AD for last era whereas earliest groups of most widely used AD were tricyclic antidepressant (TCA) and (mono amine oxidase inhibitor) MAO [8]. Fluoxetine, a type of SSRI, is supposed to be more effective anxiolytic drug as compared with other SSRIs [9, 10]. Several researches show that fluoxetine is effective against mood disorders like depression, panic and anxiety [11, 12]. Acute fluoxetine treatment enhances synthesis of serotonin in brain[13]. The increase in the rate of synthesis of serotonin after fluoxetine administration is secondary to its primary inhibitory action of liver tryptophan 2,3-dioxygenase (TDO) enzyme [14], which also enhances plasma free TRP concentration in animal. Chronic AD treatment has been found to diminish the sensitivity of corticotrophin releasing factor in neurons rather than their basal activity due to stress[15], which may contribute to normalization of HPA activity in patients undergoing psychopharmacological treatment. Earlier it was reported that chronic treatment with fluoxetine shows behavior anxiolytic effects devoid of reducing neuroendocrine responses to conditioned stress in rats[16]. Currently it was investigated that chronic treatment with imipramine reverse depressive-anxiety-like behaviors by maintaining adrenocorticotropic hormone, and reducing interleukin-1β in the brain of rats when subjected to experimental periaical lesion[17].

Forced swimming test (FST) is an animal model of depression. The FST is used to check AD action, drugs administered between pretest and test session reduce the interval of behavioral immobility. There is a link between dysfunction in monoamine transmission (5-HT, or nor-epinephrine NE) and depression [18]. Aim of current research is to find out the effects of fluoxetine administration on tryptophan metabolism and disposition in forced swimming rats.

MATERIALS AND METHODS

Animal and treatment

All protocols were strictly accompanied agreement with the national research council of laboratory animals. Ethical permission was taken from institutional animal Ethics committee, University of Karachi, Pakistan. Animals subjected to any treatment or procedures were made painless or minimize pain. Locally bred fifteen male Albino Wistar rats (150-200 gm body weight) were separated into three groups in plastic cage. Each group had five rats. They were kept at room temperature provide water and food for adjustment of animal to their environment before any treatment of experiment. Control group treated with saline (0.95%NaCl) (2ml/kg) injected intraperitoneally (LP) according to body weight. Test group also treated with saline followed subjected to forced swimming test. Drug treated test group injected fluoxetine HCl (20 mg/kg) (LP) dissolved in vehicle (DMF: Saline) (1:3 v/v) after 3.5 hr subjected to FST. All animals were killed after specific treatment and time by decapitation using guillotine. Whole sample (serum, brain and perfused liver) were kept immediately at -70 ºC till investigation.

Biochemical Determinations

TDO enzyme activity was estimated in rat liver homogenate. We took 2 gm of frozen liver tissue homogenized in 13 ml of 0.14 M KCl (pH 7.0) at 0ºC using polytron homogenizer spinning at 13000 rpm for 2-3 minutes either in the presence (total enzyme activity) of added haematin 2µM (haematin dissolved in 0.1M NaOH) or in absence (holo enzyme activity) as mention in detail[14]. The apo enzyme activity was estimated as the difference between total and holo enzyme activities. Serum total TRP, Brain TRP, 5-HT and 5-HIAA were measured by spectrofluorimetric procedures [19].

Drug preparation

Fluoxetine (Merck) was dissolved in DMF:Saline (1:3 v/v) and administered i.P at a dose of 20 mg/kg according to the body weight of rats and control groups treated with saline (0.95% NaCl).

Forced Swimming Test (FST)

For behavioral study, animals were exposed to FST to make a model of depression. We put the animal in glass tank which height was 46 cm long and width was 20 cm, tank was filled with water, the depth of water was 30 cm and temperature was 25 ±2ºC. The depth of water allowed the rats to easily swim and float in the tank without touching the bottom of tank. All test group rats were placed separately in tank for 15 minutes pretest, after 15 minutes rats were removed from water dried with towel and returned back to their cage. Twenty four hour later rats were treated with saline/drug, 3.5 hour after injection rats were again exposed to FST for 5-minutes. Control animals (untreated) were not exposed to force swimming test [20-21].

Behavioral analysis

Behavior was noted during the test swimming session using time-sampling method [9], one of three behaviors (climbing, swimming, immobility) was recorded at every five second. When animal was showing lowest movement required to stay afloat scored as immobility. When the animal was showing
actively swimming and greater movement required to stay afloat in tank scored as swimming and when the animals was showing vigorous thrashing movements with its forepaws scored as climbing.

RESULTS

Effects of FST and acute fluoxetine-HCl administration on TDO activity in rats

Table 1 shows the effects of FST and acute fluoxetine on hepatic TDO activity in rats. Data analyzed by student t-test shows holo and total enzyme activities significantly inhibited, (30.8%; P<0.001) and (20.4%; P<0.001) respectively in saline injected force swim rats when compared with untreated rats. While total and apo enzyme activities were inhibited (18.7%; P<0.001, 33.3%; P<0.01) in fluoxetine pretreated force swim rats, when compared with saline injected force swim rats. Holo enzyme activity was inhibited by 31.7% (P<0.001), (35.4%; P<0.001) and (40%; P<0.001) respectively in fluoxetine treated force swim rats when compared with untreated control rats.

Effects of FST and acute fluoxetine on serum total and liver TRP concentration in rats

Table 2 shows the effects of FST on serum total and liver TRP concentration in fluoxetine-pretreated rats. Data analyzed by student t-test show that serum total TRP concentration was significantly decreased by 52.5% (P<0.01) in saline injected FST rats when compared with (16.8%; P<0.01) in fluoxetine pretreated FST rats. Serum TRP concentration was decreased (60.5% P<0.001) in fluoxetine treated FST rats when compared with saline injected control rats. Hepatic TRP concentrations were significantly increased (50.3%; P<0.001) in saline injected FST rats when compared with untreated control rats and (70.5%; P<0.001) in fluoxetine treated FST rats when compared with untreated control rats.

Effects of FST and acute fluoxetine –HCl administration on brain indole concentration in rats

Table 3 shows effects of FST on brain indoles in fluoxetine-pretreated rats. Data analyzed by student t-test show that brain TRP, and 5-HIAA concentrations were significantly increased by 58.3% (P<0.001), 66.6% (P<0.001), and 80 % (P<0.001), in saline injected FST rats when compared with untreated control rats. While, brain 5-HT concentration was decreased (14%; P<0.05) and 5-HIAA concentrations was increased (33.3%; P<0.001) in fluoxetine-pretreated FST rats when compared with saline treated FST rats. Brain TRP, 5-HT and 5-HIAA concentrations were increased (67.5%; P<0.001) (43.3%; P<0.001) and (140%; P< 0.001) respectively in fluoxetine-pretreated rats when compared with untreated control rats.

Effects of acute fluoxetine-HCl on forced swimming test

Table 4 shows that acute fluoxetine-HCl did not produce any significant effect on immobility.

Table 1: Effects of Forced Swimming Test and acute Fluoxetine-HCl (20mg/kg) (I.P) on hepatic tryptophan 2,3-dioxygenase (TDO) activity in rats

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Tryptophan 2,3-dioxygenase activity (µM of Kynurenine formed /h/g wet.wt.of liver)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Untreated control</td>
</tr>
<tr>
<td>Holo enzyme</td>
<td>2.17±0.1</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Total enzyme</td>
<td>4.15±0.06</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Apo enzyme</td>
<td>2.0± 0.07</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Experimental details are given in method’s section. All values are mean ± SEM for each group (n=5) animals. Statistical analysis performed using student t-test. Saline treated FST group when compared with untreated control is indicated by †P<0.001. Fluoxetine-HCl treated FST group when compared with saline treated control is indicated by *P<0.01 and when compared with untreated control is indicated by δP<0.001.

Table 2: Effects of forced swimming test and acute Fluoxetine (20mg/kg) on serum total and liver TRP concentrations in rats

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Untreated control</th>
<th>Saline+ Forced swim</th>
<th>Fluoxetine + Forced swim</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serum Total TRP µg/ml</td>
<td>15.1±1.65</td>
<td>7.17±0.3†</td>
<td>5.96±0.5*δ</td>
</tr>
<tr>
<td></td>
<td></td>
<td>-52.5%</td>
<td>-16.8%</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>-60.5%</td>
</tr>
<tr>
<td>Liver TRP µg/g</td>
<td>7.8±0.3</td>
<td>11.73±0.37††</td>
<td>13.3±1.078</td>
</tr>
<tr>
<td></td>
<td></td>
<td>50.3%</td>
<td>-70.5%</td>
</tr>
</tbody>
</table>
Experimental details are given in method’s section. All values are mean ± SEM for each group (n=5) animals. Statistical analysis performed using student t-test. Saline treated FST group when compared with untreated control is indicated by †P<0.01, ††P<0.001. Fluoxetine-HCl treated FST group when compared with saline treated control is indicated by *P<0.05 and when compared with untreated control is indicated by δP<0.001.

Table-3: Effects of forced swimming test and acute fluoxetine (20mg/kg) on serum total and liver TRP concentrations in rats

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Untreated control</th>
<th>Saline+ Forced swim</th>
<th>Fluoxetine + Forced swim</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brain TRP</td>
<td>1.2±0.07</td>
<td>1.9±0.07† 58.3%</td>
<td>2.01±0.0488 67.5%</td>
</tr>
<tr>
<td>Brain 5-HT</td>
<td>0.6±0.04</td>
<td>1.0±0.05† 66.6%</td>
<td>0.86±0.05*§ 43.3% -14%</td>
</tr>
<tr>
<td>Brain 5-HIAA</td>
<td>0.25±0.05</td>
<td>0.45±0.04† 80%</td>
<td>0.6±0.04**δδ 140% 33.3%</td>
</tr>
</tbody>
</table>

Experimental details are given in method’s section. All values are mean ± SEM for each group (n=5) animals. Statistical analysis performed using student t-test. Saline treated FST group when compared with untreated control is indicated by †P<0.001. Fluoxetine-HCl treated FST group when compared with saline treated control is indicated by *P<0.05,**P<0.001 and when compared with untreated control is indicated by δP<0.01, δδP<0.001.

Table-4: Effects of acute administration of fluoxetine (20 mg/kg) on immobility in FST rats

<table>
<thead>
<tr>
<th>Parameter</th>
<th>FST</th>
<th>Fluoxetine + FST</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immobility time (S)</td>
<td>242±0.09</td>
<td>238±0.07</td>
</tr>
</tbody>
</table>

Experimental details are given in material and method’s section. Rats subjected to FST received an acute administration of Fluoxetine (20 mg/kg) and control rats received an equal volume of saline. All values are mean ± SEM of five rats.

DISCUSSION

Currently SSRI commonly used AD and produce their AD response by enhancing synaptic concentration of serotonin. The FST is the most extensively used animal test for monitoring action of AD. The aim of the current study is to examine whether acute administration of pretreated fluoxetine in FST rats produces anxiolytic effect.

The result shows that single administration of fluoxetine is given I.P at a dose of 20mg/kg, do not reduce the immobility time or depressive behavior of rats subjected to FST may be chronic treatment would produce any significant effect on behavioral activity. These results are similar with the other findings, which demonstrate that SSRI are lacking of any activity in this test in rats, whereas in mice they reduce the immobility time [22-24]. The mechanism of action of acutely administered SSRI leading to increased cerebral 5-HT synthesis by inhibiting the activity of liver TDO enzyme. In the same way we also found decreased holo, total enzyme activity, serum total TRP concentration and increases liver TRP concentration followed subjected to FST. The increase level of TRP in liver, provide increase availability for 5-HT synthesis. So, in the present study we also found increased brain TRP, serotonin and 5-HIAA levels in FST rats as compared to the saline treated control however fluoxetine treatment did not reverse the depressive symptoms but it decreases the 5-HT level in brain by converting into 5-HIAA. Pretreatment with fluoxetine also elevate brain TRP level in FST rats which shows fluoxetine inhibit the TDO enzyme activity. In contrast kreiss and Lucki [25] (1995) have also found transiently elevated levels of 5-HT in the striatum as compared to hippocampus. Similarly Kirby and Lucki [26] have also found acute administration of fluoxetine (15 mg/kg) slightly decreased lateral septum 5-HT; chronic treatments with SSRI are commonly used for several anxiety disorders; at the start of the treatment showed anxiety in patients. Similarly anxiety associated behaviors have seen in animal studies after a single injection with SSRIs. Fluoxetine (20mg/kg) totally blocked exploration area induced anxiogenic effects in mice [27]. Repeated injection of fluoxetine (1.0 mg/kg and 5.0 mg/kg) has also produced anxiolytic effect in chronic mild stress rats [28].

CONCLUSION

In conclusion, our results showed depressive likes behaviors following exposure to FST. Moreover we also found increased level of neurotransmitter serotonin (5-HT) due to inhibition of TDO enzyme activity. Acute treatment with fluoxetine did not reduced immobility but it reduce brain 5-HT concentration by converting into 5-HIAA in FST rats which shows anxiolytic effect of fluoxetine. However more investigations needs to explain the chronic
treatment of fluoxetine in FST rats and level of other neurotransmitter like catecholamine should be monitored.

REFERENCES

ACKNOWLEDGMENT
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