Affective Behavior Dysregulation Was Induced by Chronic Administration of Copper in Wistar Rats

Mouloud Lamtai*, Sihame Ouakki, Oussama Zghari ©, Abdelhalem Mesfioui, Aboubaker El Hessni, Ali Ouichou

Unit of Nervous and Endocrine Physiology, Laboratory of Genetics, Neuroendocrinology and Biotechnology, Faculty of Science, University Ibn Tofail, Kenitra, Morocco
Email: *mouloud-lamtai@hotmail.fr, ouichou@hotmail.com

Abstract

As both deficiency and excess of copper (Cu) can be harmful, dysregulation in its homeostasis has been connected with various neurological disorders. The present study was undertaken to examine whether Cu chronic administration can induce alterations of affective behavior especially anxiety and depression levels in male and female rats. Twenty-four rats, for each gender, divided in control and three test groups (n = 6), were injected intraperitoneally with saline (0.9% NaCl) or CuCl₂ (0.25 mg/kg, 0.5 mg/kg and 1 mg/kg) for 8 weeks. After treatment period, animals were tested in the open-field, elevated plus maze tests for anxiety-like behavior, and forced swimming test for depression-like behavior. Results demonstrated that Cu administered chronically, exerts an anxiogenic effect in rats. In the OFT, Cu decreases the TCA and NRC parameters without modifying the locomotor activity represented by the NTS parameter. With regard to EPM, Cu decreases TOA and EOA parameters without modifying the TAE parameter. A significant increase in depression-like symptoms was also exhibited by Cu treated rats (p < 0.001). A dose of 1 mg/kg CuCl₂ showed maximum anxiety-like and depression-like symptoms as compared to controls as well as from the other two doses indicating dose-dependent effects of chronic Cu administration. Overall, these results suggest that intoxication with Cu has potentially deleterious effects on brain as reflected in behavioral dysfunctions such as depression and anxiety.

Keywords

Copper, Depression-Like, Anxiety-Like, Rat
1. Introduction

Mental disorders are increasing in number worldwide. The number of people suffering from depression and/or anxiety increased by years lived with disability globally [1]. Depressive and anxiety disorders are common at all ages. In addition, their effects on well-being and daily functioning are enormous and comparable to those of major chronic physical illnesses [2] [3].

Generally, patients with depression often have features of anxiety disorders, and those with anxiety disorders commonly also have depression. Both disorders may occur together, it can be difficult to discriminate between them, 85% of patients with depression also experience significant symptoms of anxiety, while comorbid depression occurs in up to 90% of patients with anxiety disorders [4] [5].

Several studies have been done in the past decades. However, the etiology of depression and anxiety has not been fully explained. In an effort of searching the risk factors related to these pathologies, a number of epidemiological studies and meta-analyses have highlighted risk factors related to lifestyle, medical history and exposure to occupational and environmental pollutants, such as pesticides, polluted air and heavy metals [6].

Exposure to metals from environment, which are highly neurotoxic and have no other biological functions, such as aluminum (Al), lead (Pb) and cadmium (Cd), has long been debated as a potential environmental risk factor for mood disorders [7] [8]. However, nowadays, people are mainly focusing on biologically important metals such as copper (Cu), because their imbalance is related to a profound physiological alteration including the central nervous system (CNS) [9] [10].

Cu is an essential trace element and its toxicity is far lower than that of other heavy metals, it can be toxic once the safe dose is exceeded. The optimal levels need to be carefully determined [11]. The widespread potential exposure of human to Cu occurs via consumption of contaminated food and water with copper and also by inhalation of industrial or cosmetics contained fumes or dust with it [12].

It is known that Cu can ensure the well-functioning of different biological systems including CNS [13], It has a crucial role in erythropoiesis, myelin formation, synthesis of hormones, antioxidant protection and immune system modulation [14]. Serving as a structural component and cofactor for many proteins and enzymes comprising the free radical scavenger superoxide dismutase (Cu, Zn-SOD), dopamine monooxygenase, cytochrome oxidase, lysyl oxidase and ceruloplasmin, Cu regulates a large number of bodily biochemical processes [15] [16].

Among the body organs, the brain is the one of the most Cu-rich (next to the heart and liver) [17]. This metal is capable of crossing the blood-brain barrier and is distributed throughout most regions of the brain, especially, in hippocampus, basal ganglia, cerebellum and synaptic membranes [17] [18]. Cu is called the
“Emotional mineral”. The reason for this is that Cu and imbalances related to it have such a profound impact on the CNS. Low Cu level may result in incomplete development, while excess concentration maybe injurious [19].

Epidemiological studies have demonstrated that high concentrations of Cu are closely associated with reduced cognitive function [20] and symptoms of Alzheimer's disease (AD) [21], Menkes disease (MD), Wilson's disease (WD), Parkinson's disease (PD) and Huntington disease (HD) [19] [22] [23]. Also, an inverse relationship has been confirmed between serum concentrations of Cu and behavioral changes [24] [25] [26]. A meta-analysis demonstrated that serum Cu was slightly but significantly increased in patients with anxiety and depression relative to healthy controls, suggesting a possible role for Cu in these pathologies [27] [28] and its influence on the neurotransmission implicated in emotional processes, such as serotonergic, dopaminergic, glutamatergic, noradrenergic and GABAergic systems [29] [30].

Limited are studies which have addressed the potential importance of Cu in the development of neuropsychiatric disorders, including anxiety and depression [31] [32] [33]. There are only few studies on this topic, and future researches are needed to explore the underlying mechanisms. Thus, in this study, we decided to evaluate the effects of chronic administration of Cu on animal affective behavior in male and female rats.

2. Material & Methods

2.1. Animals and Experimental Conditions

Male and female Wistar rats aged 8 to 9 weeks and weighing about (120 ± 20) g were used in this study. Animals were raised in the Animals House Unit in Faculty of life sciences, University Ibn Tofail. They were maintained in plastic cages (36 cm long, 20 cm wide and 15 cm high) with stainless steel wire lids (bedded with wood shavings); The animals were maintained at a constant temperature (22˚C degrees ± 1), 12 hours light/12 hours dark with free access to food and water. All animal procedures were approved by the Animal Ethics.

A total of 48 rats were divided into 4 groups of 6 animals (for each sex): The first group served as the control and the animals daily received an intraperitoneal injection of 0.9% NaCl. The second group rats were treated with Cu at dose of 0.25 mg/kg, the third group rats were treated with Cu at dose of 0.5 mg/kg, while the last group rats were treated with Cu at dose of 1 mg/kg.

Saline solution or CuCl₂ (obtained from SIGMA-ALDRICH) used in the present work were injected intraperitoneally and chronically at the rate of one injection per day and this during 8 weeks according. All injections are carried out between 16:00 and 16:30.

2.2. Neurobehavioral Tests

Animals of the four groups were examined with three different behavioral tests twenty four hours after end of the 8 weeks corresponding to the treatment pe-
period (at the rate of one test per day). Anxiety behavior in all animals was measured in the following order: in Open field test (OFT) and the Elevated plus maze (EPM). The depression behavior was measured in Forced swimming test (FST) [34] [35] [36]. The tests were performed between 8 am and 12 am.

**Anxiety-Like Measurement**

**Open Field Test:** The OFT has been widely used to assess the anxiety-like behavior in rodents [37] [38]. The behaviour of rats was assessed in a wooden box measuring 100 × 100 × 40 cm high. The floor was divided by black lines into 25 small squares (20 cm × 20 cm), defined as 9 central and 16 peripheral squares. Immediately after a rat was placed gently in the centre of the open field, the movement of the animal was scored for 10 min. During this time, the time spent in the center of the area (TCA) and the number of returns to the center (NRC) were measured. Central perimeter residence time and the number of returns to the central area were used as a measure of anxiety. The central area of a novel environment is anxiogenic and aversive and the behavioral inhibition appears therefore as an avoidance behavior towards the central zone of the OFT. The number of total squares (NTS) was taken as an indicator of locomotor activity. The apparatus was cleaned with 70% ethyl alcohol and water solution prior to behavioral testing to remove residues left by previously tested rats.

**Test of the Elevated Plus Maze (EPM):** EPM is frequently used as a measure for evaluating the risk assessment and anxiety behavior of an ethologically derived animal model [39] [40]. This test is based on the creation of a conflict between the exploratory drive of the rat and its innate fear of open and exposed areas. EPM is prepared seems like a plus shape and 40 - 70 cm height from floor. The Apparatus was consisting of 2 open and 2 closed arms (50 cm × 10 cm). Two of the opposing arms are closed by 40 cm high side and end walls, having an open roof. In order to avoid fall, the other two arms (open arms) were surrounded by 0.5 cm high edge and at the intersection of the four arms there is a central platform (10 cm × 10 cm). Exactly over this central platform, a 100-W lamp was placed. The rats were individually placed onto the central platform facing one of the open arms and were observed their behaviour for 5 min while freely exploring the maze. During this session, the following parameters of anxiety-related behavior were measured: 1) entries into open arms (EOA), 2) time spent on the open arms (TOA), 3) and number of full entries into the arms (the total number of the introduction to open and closed arms) (TAE). The animal was considered to have entered an arm when all four limbs were inside the arm. Decreased anxiety-like behavior is indicated by a significant statistical increase of parameters in open arms (time and/or entries). Treatments that reduced open arm exploration were considered to be anxiogenic. The total number of the entries into all arms provides general hyperactivity. To eliminate any lingering olfactory cues, the maze was cleaned with a 70% ethyl alcohol after each test.

**Depression-Like Measurement in Forced Swimming Test (FST):** To assess the depressive-like behavior, the FST is used [41]. It is an apparatus comprised
of a glass tank having height of 50 cm and width of 30 cm, which contained wa-
ter at the height of 30 cm and temperature of 25˚C. In this glass tank animals
were individually forced to swim for 5 min. The height of water was adjusted to
prevent the escape of the rat from the cylinder, also in order that its tail could
not touch the bottom of the glass tank. When the rats are placed in an inescapa-
ble cylinder which is filled with water then the duration of immobility was
measured. The latency to the first bout of immobility was also recorded. The
animal is considered immobile when it makes no further attempts to escape and
only tries to keep its head above the water by ceasing all active behaviors (i.e.
struggling, swimming and jumping). In the present test, high percent time float-
ing is interpreted as an increased depressive-like response [41] [42].

2.3. Statistical Analysis

Statistical analyses were performed by using SPSS statistical software package
version 22. Behavioral Data are presented as the means ± standard error of the
means (S.E.M). Normality and homogeneity of the data were confirmed before
ANOVA, differences among the experimental groups were assessed by two-way
ANOVA followed by the Tukey test. In all experiments, differences were consi-
dered significant when p < 0.05, very significant when p < 0.01 and highly sig-
nificant when p < 0.001.

3. Results

3.1. Effect of Copper on the Levels of Anxiety-Like Measured in
the OFT

- **Time Spent in the Central Area (TCA) (Figure 1(a))**:

  The treatment factor significantly affected the TCA (F(3.32) = 11.94, p < 0.001).
The results summarized in Figure 1 show that: In males and females, Cu at dose
of 1 mg/kg was associated with a significant decrease of TCA compared with the
control animals (p < 0.001 and p < 0.01 respectively). The metal induced mean
average decrease of 9%, 14% and 34% in males; of 30%, 30% and 57% in females
respectively at doses of 0.25, 0.5 and 1 mg/kg respectively. In addition, no stati-
sically significant difference was observed with comparing different treated Cu
groups (p > 0.05).

  The sex effect was visible when considering the relative comparison (TCA% BL)
between respective treated Cu groups in males and females. Indeed, the fe-
males of the groups Cu-0.25, Cu-0.5 and Cu-1 showed a TCA significantly lower
compared to males of similar groups (p < 0.01, p < 0.05 and p < 0.01 respectively).

- **Number of Returns to the Center (NRC) (Figure 1(b))**:

  The treatment factor significantly affected the NRC (F(3,32) = 7.93, p < 0.001).
In males, at doses of 0.5 and 1 mg/kg, it decreases the NRC compared with the
control group (p < 0.01), whereas at 0.25 mg/kg Cu was not effective (p > 0.05).
Cu induced mean average decrease of 3%, 31% and 30% at doses of 0.25, 0.5 and
1 mg/kg respectively. Indeed, there is a difference statistically significant between
Figure 1. (a) Total amount time spent in the center (TCA); (b) Number of return into center area of the arena in the open-field behavior apparatus (NRC) and (c) Number of total squares (NTS) in the open field by female and male rats after 2 month of treatment with 0.9% of NaCl (Control), 0.25 mg/kg (Cu-0.25) 0.5 mg/kg (Cu-0.5) and 1 mg/Kg (Cu-1) of Cu. Results are expressed as mean ± SEM. The significance level is 0.05. *p < 0.05, **p < 0.01, ***p < 0.001.

Cu-0.25/Ni-0.5 and Cu-0.25/Ni-1 groups (p < 0.01). No difference was noted between Cu-0.5/Cu-1 groups (p > 0.05).

In females, Cu significantly reduced the NRC compared with the control group (p < 0.01). It induced mean average decrease of 12%, 8% and 65% at doses of 0.25, 0.5 and 1 mg/kg respectively. No statistically significant difference was observed with comparing different treated Cu groups (p > 0.05).

The sex effect was clear when considering the relative comparison (NRC% BL) between respective treated Cu groups in males and females. Indeed, the females of the group Cu-1 showed a NRC significantly lower compared to males of similar groups (p < 0.01).

- **Number of Total Squares (NTS) (Figure 1(c)):**
  
In contrast to TCA and NRC parameters, Cu was no significant effect on lo-
comotors activity (NTS) represented whatever the dose considered (p > 0.05).

3.2. Effect of Cu on Anxiety Levels Measured in Elevated Plus Maze Test (EPM)

- **Time Spent in Open Arms (TOA) (Figure 2(a))**: Statistic analysis showed that TOA was significantly affected by the Cu treatment (F(3,32) = 25.65, p < 0.001).

  In males, Cu affects TOA in dose-dependent manner in comparison with the control group (Cont/Cu-0.5: p < 0.01 and Cont/Cu-1: p < 0.001), whereas at 0.25 mg/kg Cu was not effective (p > 0.05). Cu induced mean average decrease of 12%, 20% and 40% at doses of 0.25, 0.5 and 1 mg/kg respectively. In addition, there is a statistically significant difference between the groups Cu-0.25/Cu-1 (p < 0.05). In contrast, no difference was noted between Cu-0.25/Cu-0.5 and Cu-0.5/Cu-1 groups (p > 0.05). In females also, the metal affects TOA in dose-dependent

![Figure 2](image)

**Figure 2.** (a) Number of entries in exposed arms (EOA); (b) Total amount of time spent in exposed arms (TOA) and (c) Total number of arms entries (TEA) in elevated plus maze by female and male rats after 2 month of treatment with 0.9% of NaCl (Control), 0.25 mg/kg (Cu-0.25) 0.5 mg/kg (Cu-0.5) and 1 mg/Kg (Cu-1) of Cu. Results are expressed as mean ± SEM. The significance level is 0.05. *p < 0.05, **p < 0.01, ***p < 0.001.
manner between 0.5 and 1 mg/kg in comparison with the control group (Cont/Cu-0.5: p < 0.05 and Cont/Cu-1: p < 0.001), whereas at 0.25 mg/kg Cu was not effective (p > 0.05). It induced mean average decrease of 11%, 18% and 43% at doses of 0.25, 0.5 and 1 mg/kg respectively. In addition, there is a statistically significant difference between the groups Cu-0.25/Cu-1 (p < 0.001) and Cu-0.5/Cu-1 (p < 0.01). In contrast, no difference was noted between Cu-0.25/Cu-0.5 groups (p > 0.05).

- **Entry to Open Arms (EOA) (Figure 2(b))**: The results summarized in Figure 2(b) show that: In males and females, Cu at dose of 0.25, 0.5 and 1 mg/kg was associated with a non-significant decrease of TCA compared with the control animals (p > 0.05).

  Similar results were observed when considering the relative comparison (EOA % BL) between treated Cu and control groups. Thus, Cu induced mean average decrease of 19%, 29% and 35% in males at doses of 0.25, 0.5 and 1 mg/kg respectively; and a mean average decrease of 18% and 44% in females, at doses of 0.5 and 1 mg/kg respectively.

- **Total Entries in Arms (TEA) (Figure 2(c))**: Locomotors activity was unaffected by any treatment ($F_{(3,32)} = 1.72$, p > 0.05), and no effect of sex ($F_{(1,32)} = 1.06$, p > 0.05) was noted. The values of all groups were comparable.

### 3.3. Effect of Copper on Depressive-Like Performances Measured by Forced Swimming Test (FST)

- **Immobility Time (TIM) (Figure 3(a))**: Statistical analysis showed that TIM was significantly affected by the Cu treatment ($F_{(3,32)} = 31.82$, p < 0.001). In males and females, Cu increases the TIM compared with the control group (Cont/Cu-0.25: p < 0.01, Cont/Cu-0.5: p < 0.001 and Cont/Cu-1: p < 0.001). Indeed, no statistically significant difference was observed with comparing different treated Cu groups (p > 0.05).

  Similar results were observed when considering the relative comparison (TIM % BL) between treated Cu and control groups. Thus, Cu induced mean average increase of 74%, 91% and 92% in males; of 94%, 130% and 132% in females at doses of 0.25, 0.5 and 1 mg/kg respectively.

- **Struggling Time (TST) (Figure 3(b))**: This parameter was affected by Cu treatment ($F_{(3,32)} = 28.77$, p < 0.001). In males, at doses of 0.25, 0.5 and 1 mg/kg, it decreases the TST compared with the control group (p < 0.01). The metal induced mean average decrease of 19%, 22% and 19% at doses of 0.25, 0.5 and 1 mg/kg respectively. No statistically significant difference was observed with comparing different treated Cu groups (p > 0.05).

  In females, Cu affects TST in dose-dependent manner between 0.25 and 1 mg/kg in comparison with the control group (Cont/Cu-0.25: p < 0.01, Cont/Cu-0.5: p < 0.001 and Cont/Cu-1: p < 0.001). It induced mean average decrease of 15%, 23% and 26% at doses of 0.25, 0.5 and 1 mg/kg respectively. In addition, there is a difference statistically significant between Cu-0.25/Cu-1 groups (p < 0.05). No
Figure 3. (a) Immobility time expressed in seconds (s) (TIM); (b) Struggling time in Forced swimming test expressed in seconds (s) by female and male rats after 2 month of treatment with 0.9% of NaCl (Control), 0.25 mg/kg (Cu-0.25) 0.5 mg/kg (Cu-0.5) and 1 mg/Kg (Cu-1) of Cu. Results are represented as mean ± SEM. The significance level is 0.05. *p < 0.05, **p < 0.01, ***p < 0.001.

difference was noted between Cu-0.25/Cu-0.5 and Cu-0.5/Cu-1 groups (p > 0.05).

4. Discussion

The main objective of this study was to determine the effects of chronic exposure to Cu on animal behavior, in particular on affective disorders. The assessment of anxiety-like and depression-like behaviors is based on the use of validated OFT, EPM, and FST behavioral tests.

The present study showed that Cu, administered chronically, exerts an anxiogenic effect in rats. Cu decreases the TCA and NRC parameters in the OFT, and TOA and EOA parameters in EPM without modifying the locomotor activity. The results are in agreement with animal data that showed anxiety behavior in Cu intoxication [43]. Using the dark-light box task, adult rats injected intraperitoneally with Cu at a dose of 10 mg/kg bw for 3 consecutive days, have a strong preference for the dark compartment compared with controls, suggesting the anxiogenic effect of this metal [13] [43]. No further studies have addressed such neurobehavioral outcomes of Cu exposure in mammals.

Our work also showed that in FST, Cu caused an increase in TIM and a decrease in TST in males and females, highlighting the depressant effect of this metal. A dose of 1 mg/kg CuCl₂ showed maximum anxiety-like and depression-like symptoms as compared to controls. The Cu effects observed in our study are similar
to numerous heavy metals (Cd, Al and Ni) [7] [8] [10].

It has been known that abnormal levels of several trace metals such Cu are associated with psychiatric disorders [44]. Several studies investigated serum Cu concentrations in psychiatric patients. Generally, serum Cu has been suggested as a marker of anxiety and depression [14]. For examples, Islam et al. demonstrated that generalized anxiety disorder patients display elevated Cu [28]. Patients with high Cu associated with Wilson’s disease often suffer from depression, anxiety and psychosis [14]. Manser et al. Schlegel-Zawadzka et al. as well as Schlegel-Zawadzka and Nowak showed that serum Cu levels were significantly higher in depressed patients as compared to controls, by 14%, 22% and 21%, respectively [16] [45] [46]. Similar observations were also made later by Narang et al. [27]. Also, it was proven that antidepressant treatment significantly reduced serum Cu levels [27] [47].

It is speculated that increased brain Cu plays a significant role in the progression of abnormal emotional behavior [44]. The question then arises, what role does Cu play in the course of affective disorders?

These findings may be explained by the fact that, a high levels, Cu has been reported to be a neurotoxic metal that have a profound impact on the CNS [19], especially on structures implicated in emotional processes. This metal is capable of crossing the blood-brain barrier and is distributed throughout most regions of the brain, especially, in hippocampus, basal ganglia, cerebellum and synaptic membranes [17] [18]. Several studies have provided evidence concerning the effect of Cu on the neurotransmission implicated in normal functioning and emotional processes, such as serotonergic, dopaminergic, glutamatergic, noradrenergic, and GABAergic systems [29] [30].

Serotonin (5-hydroxytryptamine, 5HT) is a major neurotransmitter that controls many functions, ranging from mood and behaviour through to sleep and motor functions [48] [49] [50]. This neurotransmitter involve projections from the dorsal raphe nucleus to limbic structures implicated in anxiety and depression related behaviors [51] [52]. The principal mechanisms whereby Cu modulates 5HT are still far from being fully understood. However, some hypotheses were evolved. The disturbed anxiety and depression states resulting from Cu exposure, may involve a direct neuromodulation of the serotonergic system by decreasing its tone [13]. As known, the loss of 5 HT innervation in the brain is one of the most recognized neuronal basics of anxiety and depression disorders [53]. It is reported that Cu can modulate the 5 HT biosynthesis [13]. Generally, 5HT arises from tryptophan by reaction involving tryptophan hydroxylase, an enzyme which requires for its activity the presence of metal such as Cu [13], but at high concentration, Cu may induce conformational changes leading to its inhibition and an alteration in the amount of monoamine stored and released from the nerve terminals [54]. In addition, an interaction between Cu and 5 HT has been suggested [50]. It is showed that oxidation of 5-HT occurred in the presence of Cu [55]. Cu effectively converts functional 5-HT into a dimeric compound [50]. A studies have indicated that the products of 5-HT oxidation...
such 5,5′-dihydroxy-4,4′-bitryptamine (DHBT), have the potential to be, neurotoxic [55] [56] [57]. The administration of DHBT to mice resulted in death within 24 h [58].

The loss of monomeric 5-HT may help account for behaviour disturbances observed, but the central control of the animal behavior is very complex and involves several other neuronal systems, especially DA [13]. The dopaminergic system seems to be another target of Cu poisoning [13]. An excess of this metal may be associated with DA dysregulation [59]. In the study of Przybyłkowski et al., an increase in Cu content in different brain regions and slight decrease in DA levels in the striatum was demonstrated [60], a DAergic structure involved in the control of anxiety and depression [13]. Rats show a reduction of DA and its metabolites contents in striatum with an inhibition of Tyrosine Hydroxylase mRNA expression in substantia nigra pars compacta [61]. It has been demonstrated that Cu reduces brain levels of DA in common carp (Cyprinus carpio) after exposure to Cu sublethal levels [62]. Cu also provokes damage of the DAergic innervations in the nigrostriatal system of rats [13]. In addition, Cu has a selective toxicity for neocortical neurons provoking neuronal apoptosis [63], a process initiated through pathways involving the production of oxidative stress and free radicals-induced neurotoxicity [64]. This process is widely believed to be one of the causes of neuronal death [65].

In our study, sex dependent effect was established; the effects of Cu on depression-like and anxiety-like being slightly pronounced in females than in males. A plausible explanation of the gender-related differences suggests the involvement of sexual hormones in the modulation of affective disorders. It has been demonstrated that estrogen promotes retention of Cu [66] [67]. As a result, females may be particularly susceptible to the neurotoxic effects of Cu.

5. Conclusion

In conclusion, brain is greatly targeted to damage by toxic agents. Along with evidence derived from our study where exposure to Cu constitutes a great threat being associated with injurious effects on mental health, the chronic exposure of Cu even at low doses might result in behavioral aberrations and also emphasizes the risk of environmental exposure of this toxic metal to humans.

Acknowledgements

Thanks are due to F. ABOUBAKR, TY. SMAMRI, and Y. CHAHIROU from the Unit of Nervous and Endocrine Physiology, Laboratory of Genetics, Neuroendocrinology and Biotechnology, University Ibn Tofail, Kenitra, Morocco, for their help and assistance.

Conflicts of Interest

The authors declare no conflict of interest.
References


DOI: 10.4236/nm.2019.102009 147 Neuroscience & Medicine


