Acupuncture Promoted the Recovery of the Neural Plasticity Functions and the Repair of Neural Plasticity-Related Proteins in the Prefrontal Cortex of CUMS-induced Depressed Rats

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Research

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Abstract

Background: Acupuncture therapy in Traditional Chinese Medicine has been widely believed as a complementary treatment for depression with numerous clinic and research reports. However, mechanisms of neural plasticity about the positive effects of acupuncture still rarely reported. The purpose of our paper aim to investigate the efficiency of acupuncture in curing depressive-like behaviors and discuss the mechanism underlying neural plasticity and its related proteins BDNF/PSD95/SYN/PKMZ in prefrontal cortex.

Methods: 32 rats with the same baseline were divided into 4 groups at random: control group(8rats), model group(8rats), acupuncture group(8rats) and flouoxetine group(8rats). Depression rat models were induced by chronic unpredictable mild stress combined with single cage isolation for 6 weeks. Acupuncture group and flouoxetine group then give 3-weeks treatment during the later 3 weeks of modeling procedure. The open field test (OFT), elevated-plus-maze (EPM) and sucrose preference test (SPT) were executed to estimate the depressive behaviors. The number of nerve cells, the length of dendrites, and the spines density in prefrontal cortex were observed by Golgi staining. The expression of prefrontal cortex BDNF, PSD95, SYN and PKMZ protein were detected by western blot and their related mRNA were detected by RT-PCR.

Results: Acupuncture could alleviate depressive-like behaviors and promote the recovery of the neural plasticity functions in prefrontal cortex, which exerted similar antidepressant effect as flouoxetine, increasing cells number, prolonging the dendrites length, enhancing the spine density.

Conclusion: Acupuncture therapy could alleviate depressive-like behaviors which may contribute to the recovery of the neural plasticity functions and the repair of neural plasticity-related proteins BDNF/PSD95/SYN/PKMZ in the prefrontal cortex of CUMS-induced depressed rats.

1. Introduction

Depression, which is considered to be one of the chronic neurological diseases with continuous symptoms including low self-esteem, helplessness and anhedonia. Patients with severe depression have the highest suicide rates among human being[1]. According to the latest analysis of the World Health Organization's survey, more than 300 million people have suffered from severe depression in the world and with a growth rate of about 18% in the past decade, thus, depression has become an major cause of human emotional disorders in the world[2, 3]. Studies demonstrated that depression could induce long-term neural plasticity changes in some specific brain regions and the repair of neurogenesis and synaptic plasticity are strongly related to periodontal health and the prognosis of most depressive patients[4–6]. Most antidepressant treatment produces therapeutic action by enhancing neuroplasticity[7]. Acupuncture has been demonstrated to be one of the effective therapies for depression and the pathologies of which are related to neural plasticity[8, 9].
Acupuncture is a vital component of Oriental medicine which widely believed as an effective complementary treatment for depression, with fewer by-effects and lower financial burden. The study about acupuncture promoting neural plasticity has become a research hotspot in the medical community recently [10, 11]. Researches showed that acupuncture plays an essential role in protecting neurons, which may relieve the symptoms of patients via mediating on neural plasticity[8, 9, 12]. In addition, doctors have achieved significant results in the clinical treatment of depression patients by using acupuncture[13–15]. It is believed that acupuncture is associated with the regulation of cortical striatum reward / motivation circuit in some specific brains of depression patients[16]. Previous study of our research found that acupuncture can reverse the changes of behavior on depression model rats. It can also play a role in treating depression-like behavior by regulating β-CaMKII in the lateral habenular nucleus of depression model rats in the signaling pathway[17]. But the deep-seated mechanism needs to be further explored.

Prefrontal cortex (PFC) is part of the limbic cortex, which participate in many stress-sensitive psychological disorders because of its rich neural interconnectivity[18, 19]. Researches has demonstrated that neuroplasticity in limbic areas is disrupted in depression, particularly the neural plasticity-related proteins in prefrontal cortex which plays a vital role in pathologies of depression[20–22]. Brain derived neurotrophic factor (BDNF) which is a plasticity-related protein widely distributed in the brain and central nervous system[23, 24]. BDNF plays a critical role in the survival and differentiation of neurons during the development of the brain central nervous system, thus, involving in neurogenesis and neuroplasticity in the brain central nervous system[25, 26]. Postsynaptic density 95 (PSD-95) is a kind of protein connecting structurally to synaptic signaling complexes and receptors [27]. Synaptophysin (SYN) is a marker protein located in the overall glycoprotein of presynaptic nerve development, which is the member of presynaptic vesicle boutons[28, 29]. PSD-95 and SYP are two synapse marker proteins which involved in synaptic signal transmission[30].

PKMZ is short for Protein kinase Mzeta, which is an atypical protein kinase C subtype in some specific part of the brain. Storing memory and maintaining long-term potentiation are the functions of PKMZ. Studies have shown that Protein kinase Mzeta in prefrontal cortex mediates rats depressive-like behaviors and associated with the morphology of neural dendritic spines[31, 32]. It is currently believed that the damage of neural plasticity-related proteins in some specific part of the brain might associated with the pathologies of depression. The efficacy of acupuncture in the intervention of depression has been widely reported. However, few studies demonstrated the relationship among acupuncture, prefrontal cortex and the neural plasticity-related proteins.

The object of current research was to explore the antidepressant mechanism underlying acupuncture based on the neuroplasticity and its neural plasticity-related proteins BDNF/PSD95/SYN/PKMZ in the prefrontal cortex of CUMS-induced depressed rats. To our best knowledge, it was an unprecedented research to demonstrated the antidepressant effects of acupuncture underling its neural plasticity-related proteins BDNF/PSD95/SYN/PKMZ in prefrontal cortex.
2. Materials And Methods

2.1. Animals and experimental design

48 SPF SD male rats weighing 120–150 g, which purchased from the company of Shanghai Slake Laboratory Animal Co., Ltd. with the license number SCXK (Shanghai) 2017-0005. After one week of adaptive feeding, the rats body weight, open field test and sucrose preference test were investigated before the experiment, which ensure that all experimental rats characteristics were in the same baseline. We choose 32 rats with the same baseline of body weight and behavior which met the requirements.

The 32 rats were divided into 4 groups at random: control group (N = 8), CUMS group (N = 8), acupuncture group(N = 8) and fluoxetine group (N = 8). Rats in the control group reared together, while rats in the other three groups were socially isolated by placing each rat in a single cage and underwent CUMS procedure for 6 weeks. Acupuncture group and fluoxetine group then give 3-weeks treatment during the later 3weeks of modeling procedure (Fig. 1).

2.2. Establishment of animal model

According to previous study [33, 34], the depression rat model was induced by chronic unpredictable mild stress combined with single cage isolation for 42 days. We chose one stimulus every day and same stimulus could not occur continuously, the modeling procedure and schedule were shown in Table 1. Weight monitoring and behavioral test were conducted to estimate the state of rats. The success criteria of depressive model establishment showed the significant reduction of behavioral score and body weight.

| stress                                      | day     |
|---------------------------------------------|---------|
| Fasting (24 h)                              | 1,13,19,32,42 |
| 45°C hot water stimulation(5min)           | 2,10,24,34,41 |
| restraint (3 h)                             | 3,17,20,30,40 |
| water deprivation (24 h)                    | 4,14,25,28,39 |
| day and night inversion (24 h)              | 5,11,21,36,38 |
| 4°C cold water stimulation(5min)           | 6,18,26,35,37 |
| squirrel cage tilt (45°, 24 h)              | 7,15,22,33,36 |
| noise stimulation (3 h, 80 dB)              | 8,12,27,31  |
| tail clamping (3 min)                       | 9,16,23,29  |

2.3. Acupuncture and fluoxetine intervention
As shown in Fig. 2A, rats in acupuncture group were given acupuncture treatment at Shangxing (DU23) and Daling (PC7) before daily CUMS stimulation. Acupuncture operation: Flat needling Shangxing (DU23) toward the direction of the nose, the needle was inserted as deep as 3-5mm; Obliquely stab Daling (PC7) toward the direction of Neiguan (PC6), the needle was inserted as deep as 2-3mm. The needles used were sterilized disposable stainless steel needles (0.18 mm diameter; Hanyi, Beijing, China). As shown in Fig. 2B, rats in fluoxetine group were given fluoxetine (PHR1394-1G, Sigma-aldrich) solution which (0.21mg/ml) by gavage (1ml/100g) before daily stimulation.

2.4. Body weights and general status.

Rats body weights were monitored once a week, observe the rat's mental state, eating status, activity level and hair gloss every day.

2.5. Behavioral tests

Behavioral tests including open field test, elevated-plus-maze and sucrose preference test were executed at the end of experiment.

2.5.1. Open field test

OPT was executed to test the locomotor activity and depressive-like behaviors of rats, the operational steps is conducted as previous study[35]. We put the rats in the test environment for adaption prior to the experiment. The open field region consists of the black acrylic device (100cm in length and width, 40cm in height). When the test started, rats were first placed in the center of area, then allowed to travel freely during the whole area. We observed the autonomic activity of rats during the 5 minutes test (recorded by video camera and analyzed by SMART software). The change of total crossing distance and central residence time were assessed as the criteria of depressive symptom. After every test, 75% ethanol was used to disinfect the experimental site.

2.5.2. Elevated-plus-maze test

Elevated-plus-maze test was used to estimate the anxiety-like behavior of depressive rats as reported in previous studies[36, 37]. The elevated plus-maze devices have one central zone (10 × 10 cm) together with two open arms (50 × 10 × 40 cm) and two closed arms (50 × 10 × 40 cm), the arm specifications is 50cm in length, 10cm in width and 40cm in height. When the test started, rats were first placed in the center of area and their heads facing the open arms. We observed the autonomic activity of rats for crossing the open arms and the closed arms during the 5 minutes test (recorded by video camera and analyzed by SMART software). After every test, 75% ethanol was used to disinfect the experimental site.

2.5.3. Sucrose preference test

Sucrose preference test was executed to examine the anhedonia of rats, the operational steps is conducted as previous study[38]. We trained rats to adapt to sucrose-water 2 days before the real test begun: each cage was equipped with one 1% sucrose bottle and one pure water bottle, we exchanged the
position of two bottles every 12 hours. After the adaptive phase, water and food prohibition in rats for 24 hours. Then start the formal test: each rat was given two bottles (one 1% sucrose and one pure water) which had been quantified beforehand, we removed bottles after 12 hours and calculated the liquid consumption. Sucrose preference (%) = \(\frac{\text{sucrose solution consumption}}{\text{sucrose solution consumption} + \text{pure water consumption}}\) × 100.

2.6. Extraction and detection

Fasting and water deprivation for 12 hours before the extraction procedure. After animals were anesthetized with 25% pentobarbital sodium (50mg/kg, intraperitoneal injection), we used cervical dislocation to prevent pre-synaptic and post-synaptic effects of anesthesia. Abdominal aorta blood collection and brain removal after euthanasia, then the prefrontal cortex harvested rapidly under ice. The number of nerve cells, the length of dendrites, and the density of dendritic spines were observed by Golgi staining. The expression of prefrontal cortex BDNF, PSD95, SYN and PKMZ protein were detected by western blot and their related mRNA were detected by RT-PCR.

2.6.1. Golgi Staining and Evaluation of Dendritic Spine Density

After the rat euthanasia, immediately take the prefrontal cortex and fix it in the fixative for more than 48h. Cut the prefrontal cortex tissue into tissue blocks with thickness of 2-3mm according to the tissue site to be observed, gently rinse the brain tissue with normal saline for several times, then place in a 45ml round bottom EP tube, add Golgi-cox staining solution to completely submerge the brain tissue, and place in a cool and ventilated place and avoid light to treat for 14 days (after 48h soak, change the new staining solution, and then change the new staining solution every 3 days, total of 14 days). Immerse in distilled water for 3 times, pour in 80% glacial acetic acid to immerse the tissue overnight, wait for the tissue to become soft, then wash with distilled water, and place into 30% sucrose. Cut the tissue into 100 microns with oscillating microtome, paste it on a gelatin slide, and dry in the dark overnight. Treat dried tissue slides with concentrated ammonia water for 15 minutes, wash with distilled water for 1 minute, treat with acid hardening fixing solution for 15 minutes, wash with distilled water for 3 minutes, dry and seal the section with glycerin gelatin. Microscope inspection, image acquisition and analysis, obtain panoramic images of brain tissue by panoramic multi-layer scanning with digital slice scanner. Analyses were performed according to standard protocols of our laboratory [39]. The number of spinous cells was observed in the pyramidal neuron dendrites of the prefrontal cortex. We used Sholl analysis method for dendritic tree examination[40]. The density of the spine was quantified and the number of spines along the length was calculated[41].

2.6.2. Western blot

The contents of BDNF, PSD95, SYN and PKMZ in prefrontal cortex were determined by Western-blot. The methods was described previously[42]: We took 30µg samples from brain tissue and added with 4×loading buffer, put them in boiling water for 10 min, 10000 rpm centrifugation for 10 min; SDS-PAGE
gel with 10% or 12% according to protein molecular mass was prepared for electrophoretic separation of protein samples; SDS-PAGE gel protein was transferred to PVDF membrane by electrophoresis; PVDF was immersed in sealed solution containing 5% skimmed milk powder after electrophoresis, incubated for 60 minutes during room temperature condition; PBST washed membrane was incubated at 3×5min. Add the corresponding antibody, stay at 4 Celsius condition overnight, then wash PBST for 3 times, 20 min each time, add the corresponding HRP labeled two anti-solution, incubated for 60 minutes during room temperature condition, PBST wash 3 times, 10 min each time. The PVDF film was reacted with the freshly prepared enhanced chemiluminescence agent (ECL) solution to 2min, and was rapidly exposed and developed in the darkroom. The gray value of the target protein was analyzed by the gel image system.

2.6.3. Reverse transcription-polymerase chain reaction (RT-PCR).

We searched the sequences of BDNF, PSD95, SYN and PKMZ mRNA codes from the consensus coding sequence database of the National Biotechnology Information Center ((NCBI)). The detailed primer sequences for BDNF, PSD95, SYN, PKMZ and the beta-actin control are shown in Table 2. PCR primers were developed before the RT-PCR operation.

| mRNA Primers | Sequence(5'-3') | Product leng(bp) |
|--------------|----------------|------------------|
| Rat-β-actin-F | CTGGCTCCTAGCACCATGAA AAAACGCAGCTCAGTAACAGTC | 180 |
| Rat-β-actin-R | GCCTCCTCTGCTCTTTTCT | 141 |
| Rat-β-BDNF-F | GCGGTACCCACTCACTA | 142 |
| Rat-β-BDNF-R | ACACCCATTGCCCAGAAC | 209 |
| Rat-β-PSD95-F | TCTCCACGCAGTCTAAAG | 166 |
| Rat-β-PSD95-R | ACAGCAGTGTTCGCTTTCA |  |
Add 500ul deproteinized solution 12000rpm to centrifuge for 45s, add 500ul rinse solution to rinse twice, 12000rpm centrifuge for 45s, elute RNA and determine the concentration to run glue identification. 1ug was extracted from each sample for reverse transcription, and each sample was detected by BDNF, PSD95, SYN, PKMZ and the beta-actin primers respectively. Finally, 2µL PCR products were detected by agarose gel. The electrophoresis conditions were 140V, 15min, gel imaging. We analyzed the gel using ImageJ software (National Institutes of Health, Bethesda, Maryland).

2.7. Statistical analysis

All data were processed by SPSS 21.0 software. The statistical data of each test index are expressed by means of mean and standard deviation (x ± s). Firstly, the normality and square deviation are tested. If the variance is homogeneous, LSD method is selected; if the variance is uneven, games Howell method is selected for variance analysis; if not, rank sum test is selected. The difference was statistically significant (p < 0.05).

3. Results

3.1. Effects of CUMS and intervention treatments on body weight.

As shown in Fig. 3B, before the experiment, there is no differences in rats body weight baseline among all groups (p > 0.05); while with the progress of the experiment, the weight gain trend of the CUMS- model rats all showed lower levels comparing with the control rats (p < 0.001). As we can see in Fig. 3C, after the experiment, CUMS rats showed a significant reduction in body weight (p < 0.001) comparing with the control group; Rats in acupuncture group showed a significant growth in body weight (p < 0.001) comparing with the CUMS group; while there is no significant differences in rats body weight between CUMS and Fluoxetine group (p > 0.05).

3.2. Effects of CUMS and intervention treatments on behavior tests.

As shown in Fig. 4A, the results of open-field test showed a significant reduction in total crossing distance of model rats (p < 0.01) comparing to the control rats; After the intervention treatments, acupuncture group and the fluoxetine group all showed an increase trend of total distance comparing to the model group, but there are not significant differences (p > 0.05). As shown in Fig. 4B, the results of the elevated plus maze showed a significant reduced time in open arms in the model group comparing to the control group (p < 0.001); the reduced time in open arms was significantly increased in the acupuncture group (p < 0.05) comparing to the model group, but the fluoxetine group showed no statistical significance (p > 0.05). As we can see from the results of sucrose preference test (Fig. 4C), the model group showed a significant decreased in the 1% sucrose intake comparing to the control group (p < 0.01); After the intervention treatments, the intake of 1% sucrose was significantly increased in the acupuncture group and the fluoxetine group (p < 0.01, p < 0.01) comparing to the model group.
3.3. Effects of CUMS and intervention treatments on Golgi staining.

As shown in Fig. 5, Golgi staining of rat prefrontal cortex, the arrangement of cells distribute in the control rats were more neatly, and showed a clearer outline than CUMS group. The cells in CUMS group were disorderly arranged with unclear outline, and the number of dendrites is decreased, the length of dendrites is shorter. After the intervention treatments, the acupuncture group and the fluoxetine group all increased cells number, prolonged the dendrites length, enhanced the spine density comparing with the CUMS model group.

3.4. Effects of CUMS and intervention treatments on morphological character of dendritic arborization and spine density of prefrontal cortex.

As shown in Fig. 6, the dendritic arborization were rich and well developed in control group A; After the CUMS procedure, the development dendritic spine impaired in the CUMS group (Fig. 6B); After the intervention treatments, acupuncture and fluoxetine are able to restore the impaired dendritic spines, with enlarged spiny dendritic branches in the acupuncture group(Fig. 6C) and the fluoxetine group(Fig. 6D) comparing with the CUMS model group. As shown in Fig. 6E, the CUMS model group showed lower level of intersection per shell of pyramidal neuron arborization comparing with control group; After the intervention treatments, acupuncture group and the fluoxetine group all increase the level of intersection per shell. As shown in Fig. 6F, the sum of intersections of dendrites in the CUMS group showed a significant decrease(p < 0.01) comparing to the control group; After the intervention treatments, the sum of intersections of dendrites in acupuncture group and the fluoxetine group all showed significant increases (p < 0.01, p < 0.05). As shown in Fig. 6G, the CUMS model group showed lower level of density of dendritic spines comparing with control group, but no statistical significance (p > 0.05); After the intervention treatments, the fluoxetine group showed a significant increase of the dendritic spine density comparing with the CUMS model group(p < 0.05), while the acupuncture group showed no statistical increase(p > 0.05).

3.5. Effects of CUMS and intervention treatments on the protein expression of BDNF, PSD95, SYN and PKMZ in prefrontal cortex measured by western-blot.

As shown in Fig. 7, the protein expression of BDNF, PSD95, SYN and PKMZ in prefrontal cortex of the CUMS model group decreased(p < 0.05, p < 0.01, p < 0.01, p < 0.01) comparing to the control group; After the intervention treatments, the protein expression of BDNF, PSD95, SYN and PKMZ in prefrontal cortex of
the acupuncture group and the fluoxetine group all increased compared with the CUMS model group (p < 0.05).

3.6. Effects of CUMS and intervention treatments on the mRNA level of BDNF, PSD95, SYN and PKMZ in prefrontal cortex measured by RT-PCR.

As shown in Fig. 8, the mRNA level of BDNF, PSD95, SYN and PKMZ in prefrontal cortex of the CUMS model group all decreased (p < 0.05, p < 0.01, p < 0.01, p < 0.001) comparing to the control group; After the intervention treatments, the mRNA level of PSD95, SYN and PKMZ in prefrontal cortex of the acupuncture group and the fluoxetine group all increased compared with the CUMS model group, showing significant differences (p < 0.05 or p < 0.01).

4. Discussion

In our current research, the CUMS model was established successfully according to the results of behavior test. Under the 6-weeks chronic unpredictable mild stress, rats showed less sucrose consumption in SPT, less total distances crossing in OPT as well as less open arms during EPM than rats in control groups. And their body weight grew slowly that the control rats. It is gratifying to find that those depressive-like behavior can be ameliorated by prolonged acupuncture treatment and fluoxetine intervention. These consequences indicated that the acupuncture and fluoxetine are effective to treat the CUMS induced depressive rats, thus showing the antidepressant-effect as proved in previous study[43, 44].

At present, antidepressant drugs are still in the leading position in clinic[45], fluoxetine in particularly; however, most antidepressant drugs have slow onset and low efficacy with severe side effects[46]. Acupuncture, as a safe and effective alternative therapy for depression with fewer side effects. The choice of acupoints is a key factor to ensure the curative effects of acupuncture. Daling (PC7) is considered to be the original acupoint of pericardium meridian. According to the meridian-collateral theory in Traditional Chinese Medicine, Daling (PC7) can not only relieve the fire evil enveloped by the heart, but also has great efficacy in treating mental diseases which can calm the mind and benefit the brain nerves. Shangxing (DU23) located in the governor meridian, and is good at supervising and awakening the mind. These two points are especially effective for mental illness. Therefore, the acupoints of Daling and Shangxing were elected in the acupuncture treatment group, and fluoxetine was used as intervention control group because it is one of the most commonly used antidepressants in clinic. To further explore the deep-seated mechanism of antidepressant-effect of acupuncture and fluoxetine, the Golgi staining and evaluation of dendritic spine density were used. Then, Western blot (WB) and Reverse transcription-polymerase chain reaction (RT-PCR) were be used to detect the neural plasticity-related proteins BDNF/PSD95/SYN/PKMZ and their mRNA level.
Neural plasticity, also known as brain plasticity, containing structural plasticity, functional plasticity and the cellular and molecular mechanisms associated with them[47]. Literature research indicated that the biological mechanism of depression associate with the neural plasticity in specific brain areas, especially the prefrontal cortex[48]. The number and function of synapses in central nervous system of depression were obviously disordered[49]. As a basic mechanism of neuronal adaptation, neural plasticity is destroyed in animal models of emotional disorder and stress. Chronic stress can promote or aggravate depression and destroy neural plasticity, while antidepressant treatment has the opposite effect and can enhance neural plasticity[50]. Our current results of Golgi staining showed that the arrangement of cells distributed in the CUMS rat prefrontal cortex was disorderly arranged with unclear outline, and the number of dendrites is decreased, the length of dendrites is shorter. Morphological analyses of dendritic arborization and spine density of CUMS rat prefrontal cortex also impaired. After the intervention treatments, acupuncture and fluoxetine all increased cells number, prolonged the dendrites length, enhanced the spine density comparing with the CUMS model group. Moreover, acupuncture and fluoxetine are able to restore the impaired dendritic spines, with enlarged spiny dendritic branches, increasing level of intersection per shell and intersections of dendrites comparing with the CUMS model group. Our results are consistent with the report: CUMS can promote or aggravate depression and destroy neural plasticity, while acupuncture and fluoxetine treatment have the opposite effect and can enhance neural plasticity.

Synaptic formation and neural plasticity are regulated by complex molecular protein interactions. The destruction of many key proteins is related to the susceptibility to depression, including brain derived neurotrophic factor (BDNF), post synaptic density 95(PSD-95), synaptophysin (SYN) and protein kinase Mzeta (PKMZ). We choose those proteins as they were proved to be the neural plasticity-related proteins. BDNF can regulate the neural plasticity and synaptic transmission by increasing Ca2+ levels and mitochondrial movement in neurons. The transportation and distribution of mitochondria play a vital role in BDNF-mediated synaptic transmission. Research show that BDNF enhances the release of neurotransmitter to protect neuroplasticity[51]. Endogenous BDNF is effective for cannabinoid-mediated neurogenesis[52]. As a scaffold of synaptic components required for synapse development, PSD-95 is the main organizer of signal complexes on the postsynaptic membrane. Impaired synaptic function can lead to neuropsychiatric diseases, collectively referred to as synaptic diseases. The genetic decrease in PSD-95 protein levels may affect postsynaptic function and plasticity[53]. The synaptophysin mutant synapse is a weak donor at the presynaptic terminal. Previous research indicated that synaptophysin can regulate the formation of activity-dependent synapses. Previous studies have shown that the decreased levels of neuronal synapses can change neuronal circuits and the process might be associated with the mechanism of mental diseases[54]. Protein kinase Mzeta (PKMZ) is a constitutively active N-terminal truncated form of PKC-$\zeta$, has long been implicated in a cellular correlate of learning, long-term potentiation (LTP). The regulation of PKM-$\zeta$ in the brain can affect cerebellar-dependent learning and memory[55]. Plasticity-related protein kinase Mzeta can promote neurite shaft protrusion which is closely related to mental illness.
In our current results, the protein expression and mRNA level of BDNF, PSD95, SYN and PKMZ in prefrontal cortex of the CUMS model group decreased. After the intervention treatments, the protein expression and mRNA level of BDNF, PSD95, SYN and PKMZ in prefrontal cortex of the acupuncture group and the fluoxetine group all increased comparing to the CUMS model group. Those results are consistent with previous assumption that acupuncture promote the repair of neural plasticity-related proteins BDNF/PSD95/SYN/PKMZ in the prefrontal cortex of CUMS-induced depressed rats.

5. Conclusions

In conclusion, the current research suggests that acupuncture is effective to treat the CUMS induced depressive rats, which could ameliorate the depression-like behavior, increase cells number, prolong the dendrites length and enhance the spine density. The recovery of the neural plasticity functions by enlarging spiny dendritic branches, increasing the level of intersection per shell and intersections of dendrites. These alterations may contribute to the repair of neural plasticity-related proteins BDNF/PSD95/SYN/PKMZ in prefrontal cortex. The present study provides new insights into the antidepressant effects of acupuncture underling neural plasticity and its related proteins. However, there are some limitations need to be improved: more brain region should be considered but not just the prefrontal cortex; Different depressed subject has individual difference so it is not reasonable to put them together as a whole. In future researches, we will try to put more samples and choose different brain regions so as to provide comprehensive evidences for the acupuncture treatment in depression.

Abbreviations

CUMS: Chronic unpredictable mild stress; AP: Acupuncture; FX: Fluoxetine; OFT: The open field test; EPM: elevated-plus-maze; SPT: Sucrose preference test; WB: Western blot; RT-PCR: Reverse transcription-polymerase chain reaction; BDNF: Brain derived neurotrophic factor; PSD95: Postsynaptic density 95; SYN: Synaptophysin; PKMZ: Protein kinase Mzeta; PFC: Prefrontal cortex.

Declarations

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Data Availability

The original data (including MNOVA, Excel, and USP format) used to support the findings of this study were supplied by Xianjun Meng under license and so cannot be made freely available. Requests for access to these data should be made to Xianjun Meng, mengxianjun@xmu.edu.cn.

Author’s contributions

Xianjun Meng was responsible for the study concept and design. Wenya Huang and Peng Li conducted animal experiment and wrote the manuscript. Wenjing Chen, Yang Huang and Wenjie Chen conducted the experiments and collected the data. Yiping Chen, Xinnan Wu, Yanxun Huang and Yining Yan helped to analyze the data. All authors approved the final version of the manuscript.

Conflict of interest

The authors declare that they have no conflicts of interest.

Consent for publication

Not applicable. The manuscript does not include details, images, or videos relating to individual participants.

Ethics approval and consent to participate

In this study, all rat care and experimental procedures were approved by the Animal Care and Use Committee of Xiamen University (License No. SCXK 2014-0001) and implemented in accordance with the National Institutes of Health Guidelines for the Nursing and Use of Laboratory Animals.

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**Figures**

**Figure 1**

Experimental flow chart. 1 week adaptation, 6 week CUMS procedure, 3 week treatment, behavioral test and prefrontal cortex harvesting at the end of experiment.

The expression of prefrontal cortex BDNF, PSD95 and SYN protein were detected by western blot and their related mRNA were detected by RT-PCR. The number of nerve cells, the length of dendrites, and the density of dendritic spines were observed by Golgi staining.
Figure 2

Schematic diagram of acupuncture and fluoxetine operation in rat. A: Flat needling Shangxing(DU23) toward the direction of the nose, the needle was inserted as deep as 3-5mm; Obliquely stab Daling(PC7) toward the direction of Neiguan(PC6), the needle was inserted as deep as 2-3mm; B: Fluoxetine solution(0.21mg/ml) by gavage (1ml/100g).

Figure 3

Body weight changes in all groups. A: Comparison of rat's body weight growth trend among four groups during the whole experiment; B: Body weight baseline before the experiment; C: Body weight changes at
the end of experiment (21 day). Compared with the Control group, * P < 0.05, ** P < 0.01, *** P < 0.001; compared with the CUMS group, # P < 0.05, ## P < 0.01, ### P < 0.001, Data represent means ± SEM.

**Figure 4**

Behavior changes in all groups. A: Total distance in the open-field test among four groups; B: Time in open arms of the elevated plus maze among four groups; C: The percentage of sucrose consumed by the rat at sucrose concentrations of 1%. Compared with the Control group, * P < 0.05, ** P < 0.01, *** P < 0.001; compared with the CUMS group, # P < 0.05, ## P < 0.01, ### P < 0.001, Data represent means ± SEM.

**Figure 5**
Golgi staining of rat prefrontal cortex [200×(up column) and 1000×(down column)]. A: Control group; B: CUMS group; C: Acupuncture group; D: Fluoxetine group

Figure 6

Morphological analyses of dendritic arborization and spine density of prefrontal cortex. A: Control group; B: CUMS group; C: Acupuncture group; D: Fluoxetine group; E: Intersection per shell of pyramidal neuron arborization; F: Sum of intersections of dendrites; G: Density of dendritic spines. Compared with the Control group, * P < 0.05, ** P < 0.01, *** P < 0.001; compared with the CUMS group, # P < 0.05, ## P < 0.01, ### P < 0.001, Data represent means ± SEM.
Figure 7

The expression of neural plasticity-related proteins in prefrontal cortex measured by western-blot. A: BDNF; B: PSD95; C: SYN; D: PKMz. Compared with the Control group, * P < 0.05, ** P < 0.01, *** P < 0.001; compared with the CUMS group, # P < 0.05, ## P < 0.01, ### P < 0.001, Data represent means ± SEM.
Figure 8

The mRNA level of neural plasticity-related proteins in prefrontal cortex measured by western-blot. A: BDNF; B: PSD95; C: SYN; D: PKMZ. Compared with the Control group, * P < 0.05, ** P < 0.01, *** P < 0.001; compared with the CUMS group, # P < 0.05, ## P < 0.01, ### P < 0.001, Data represent means ± SEM.