Involvement of NMDA-AKT-mTOR Signaling in Rapid Antidepressant-Like Activity of Chaihu-jia-Longgu-Muli-tang on Olfactory Bulbectomized Mice

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Background: Fast-onset antidepressants are urgently needed. Chaihu-jia-Longgu-Muli-tang (CLM), a classic Chinese herbal medicine, has been used for antidepressant treatment with long history. Olfactory bulbectomy (OB) model is validated for identification of rapid antidepressant efficacy. Here we used OB model for investigating the rapid onset activity of CLM in mice, and also tested the involvement of prefrontal Akt-mTOR and associated AMPA/NMDA receptors as well as hippocampal BDNF in the rapid antidepressant-like effect of CLM.

Methods: The OB model was first characterized with depression-like behaviors and the time course changes of the behaviors. The fast onset of antidepressant effect of CLM was evaluated using sucrose preference test, tail suspension test and forced swim test in OB mice after a single administration. The expression of synaptic proteins of AMPA and NMDA subunits as well as Akt/mTOR signaling in the prefrontal cortex, and hippocampal BDNF was evaluated with the immunoblotting method.

Results: A single dose of CLM significantly improved the deficiency in the sucrose preference and decreased the immobility time in the tail suspension test in OB mice. In the prefrontal cortex (PFC) in OB mice, there was lower expression level of the AMPA receptor subunit GluR1, rescued by a single dose of CLM. Additionally, the expression of NMDA subunit NR1 was up-regulated in OB mice, whereas mTOR and its upstream Akt signalings were both down-regulated. These deficiencies were reversed by a single dose of CLM. The CLM treatment also attenuated the expressions of NMDA receptor subunits NR2A and NR2B, which did not change in OB mice. In the hippocampus, expressions of GluR1 and brain derived neurotrophic factor (BDNF) were both up-regulated in OB mice, although CLM increased GluR1, but not BDNF.
INTRODUCTION

Major depressive disorder (MDD) is one of the most common diseases of persistent emotion (feeling) stepping down, inhibition of thought and psychomotor retardation. It has been becoming the leading cause of disability and a major contributor to the disease burden of the world's population (Smith, 2014). Serotonin selective reuptake inhibitors (SSRIs) represent the first-line antidepressants. However, there are several major disadvantages for SSRIs, including the long lag-time of therapeutic action, an appreciate number of non-responsive MDD patients, and adverse reactions (Bull et al., 2002). Therefore, development of fast onset, safe and effective antidepressants is urgently needed.

Recently, ketamine (KET), a N-methyl-D-aspartic acid receptor (NMDAR) antagonist, is found to elicit fast-onset and long-lasting antidepressant effects: a single dose of KET quickly alleviates the depressive symptoms, and the effect may last for several days in both MDD patients and various animal models of depression (Zarate et al., 2006; aan het Rot et al., 2010; Li et al., 2010; Autry et al., 2011; Duman et al., 2012). In the past few years, studies continue to discover the novel mechanisms underlying the rapid antidepressant effects. Emerging evidence support that rapid and persistent improvement of neural plasticity is essential for mediating antidepressant effects (Tsankova et al., 2006; Martinowich et al., 2007). Experimental studies also suggest that the rapid enhancement of BDNF expression in the hippocampus is responsible for the rapid antidepressant-like effect of KET (Li et al., 2010; Autry et al., 2011). The NMDA-KET-AMPA signaling in the PFC and BDNF expression in the hippocampus are probably responsible for the rapid antidepressant-like effect.

Although antidepressant efficacy of KET was well-established experimentally, clinical use of KET is challenged by the potential toxic and addictive effects. Thus, efforts have been made to discover fast-onset and safer antidepressant drugs, including Chinese herbal medicine that has been used clinically in a safe manner. For example, a herbal medicine Yueju pill, has been found to elicit rapid antidepressant-like effect, similar to KET in various animal models in mice (Xue et al., 2013; Tang et al., 2015; Xue et al., 2016). Chaihu-jia-Longgu-Muli-tang (CLM) is also a traditional Chinese herbal formulation in Chinese medicine, Doctor Zhang Zhongjing, 1800 years ago. "Shang Han Lun" written by the founding theorist of traditional Chinese medicine, Doctor Zhang Zhongjing, 1800 years ago. The therapeutic effects and mechanisms of CLM on depression and insomnia have been reported previously (Tsujimura et al., 2015; Xue et al., 2016). CLM also suggest that the rapid enhancement of BDNF expression in the hippocampus are probably responsible for the rapid antidepressant-like effect of CLM following repeated administration linked to reversal of the reduction of dopaminergic and serotonergic transmission, normalization of the dysfunctional hypothalamo-pituitary-adrenal system, and modulation of the glucocorticoid secretion system (Sasaki et al., 1998; Mizoguchi et al., 2003). Clinical observations also suggested a relatively fast antidepressant action of CLM, which, however, has not been scientifically investigated using appropriate animal models.

Olfactory bulbectomy (OB) is a validated model accurately predicting onset time of classical antidepressants or KET-like rapid antidepressant reagents (Ramaker and Dulawa, 2017). After surgically removing bilateral olfactory bulbs, rats or mice demonstrates steady depression-like behaviors, including reduced preference of sucrose, mimicking the core symptom of depression, anhedonia (Redmond et al., 1997; Song and Leonard, 2005). Many neuromolecular features are shared with other depression models, except for increased BDNF expression in the hippocampus in contrast to decreased level in most other models (Angelucci et al., 2004, 2005; Greenwood et al., 2007). It

**Conclusion:** CLM elicited rapid antidepressant-like effects in the OB model mice, and CLM reversal of the abnormality in PFC expression of AMPA and NMDA receptors and associated Akt-mTOR signaling may underlie the effects.

**Keywords:** rapid antidepressant, NMDA receptor, Akt-mTOR signaling, Chaihu-jia-Longgu-Muli-tang, olfactory bulbectomized mice
is believed that OB led to dysregulated NMDA-AMPA and Akt-mTOR signaling in PFC similar to other models, which, however, remains to be investigated.

Here, using a mouse OB model, we investigated whether a single dose of CLM was able to elicit rapid antidepressant-like activity; furthermore, we tested the association of NMDA-AMPA receptors and related Akt-mTOR signaling in PFC or increase in hippocampal BDNF with the rapid antidepressant-like action of CLM in OB mice.

MATERIALS AND METHODS

Animals

Kunming mice weighing between 20 and 25 g were purchased from China Academy of Military Medical Sciences (Beijing). Mice aged approximately 6 weeks old and were housed in cages for 7 days before behavioral testing and OB. Animals were maintained in a temperature and humidity-controlled environment, (temperature 22 ± 2°C and room humidity, 50 ± 10%) under a 12: 12 h light/dark cycle. All the mice were offered ad libitum access to food and water. All procedures in this study conformed to the Guide for the Care and Use of Laboratory Animals and were approved by the Institutional Animal Care and Use Committee at Nanjing University of Chinese Medicine.

Drugs

The medicinal plants used to prepare CLM are Bupleuri radix (Chai Hu), Fossilia oissis mastoid (Long Gu), Scutellariae radix (Huang Qin), Zingiberis rhizoma (Sheng Jiang), Ginseng radix (Fu Ling), Pinelliae tuber (Ban Xia), Radix et Rhizoma Rhei (Da Huang), Osttreae testa (Mu Li), Zizyphi fructus (Da Zao). All the medicinal plants were purchased from Nanjing GuoYi Clinical, Medicinal Material Department (Nanjing, China). CLM was supplied in the form of a water-extracted cream that was manufactured from a mixture of the crude drugs according to a fixed ratio listed in Table 1, and the quality control of different batches of CLM was evaluated using HPLC method (Supplementary Figure S1 and Supplementary Table S1). Briefly, these 11 dried herbal and mineral drugs were soaked in water for 30 min, and then were decocted two times with boiling water in the ratio of 1:8 for 2 h. After collecting all the solution, placed in a water bath at 60°C for 2 h. After collecting all the solution, placed in a water bath at 60°C and evaporated to a target concentration for frozen storage. The yield of the preparation was about 42%. The solutions of the herb preparation and vehicle were diluted to 210 mg/ml.

Behavioral Tests

Open Field Test (OFT)

Open field test estimated locomotor activity and anxiety-like behavior. Mice were freely explore for 5 minutes in a well-illuminated (~300lux) plastic cage (40 cm × 40 cm × 40 cm). A camera was placed in the top of the box for recording the activity. Both the distance traveled (cm) and time spent in central area(s) were analyzed by a computer-based tracking system.

| Plant name              | Composition (g) | Major components                      |
|-------------------------|-----------------|---------------------------------------|
| Bupleuri radix          | 6               | Saikosaponin a, c, d, e               |
| Fossilia oissis mastoid | 2.25            | Calcium base                          |
| Scutellariae radix      | 2.25            | Baccalin, Wogonin                     |
| Zingiberis rhizoma      | 2.25            | Gingerol, Shogaol                     |
| Ginseng radix           | 2.25            | Ginsenoside                           |
| Cinnamomi cortex        | 2.25            | Cinnamitic aldehyde                   |
| Hoelen                  | 2.25            | Eburicic acid                         |
| Pinelliae tuber         | 3               | Homogenistic acid                     |
| Radix et Rhizoma Rhei   | 3               | Anthraquinone                         |
| Osttreae testa          | 2.25            | Calcium base                          |
| Zizyphi fructus         | 3               | Zysynus saponin, betulinic acid       |

These 11 dried herbal and mineral drugs were soaked in water for 30 min, and then were decocted two times with boiling water in the ratio of 1:8 for 2 h. After collecting all the solution, placed in a water bath at 60°C and evaporated to a target concentration for frozen storage. The yield of the preparation was about 42%. The concentration of the solutions for administration was diluted to 210 mg/ml.
Testing apparatus was thoroughly cleaned with 75% ethanol between each test.

**Sucrose Preference Test (SPT)**
The SPT followed a procedure published before with minor modifications. Briefly, mice were housed and exposed to a sucrose solution (2% in tap water) for 72 h, followed by 24 h of water deprivation and mice were offered two individual bottles containing either tap water or 2% sucrose solution for 2 h. Liquid consumption from each bottle was measured by comparing the differences in bottle weight before and after a 2hr exposure. Sucrose preference was measured by the percentage of sucrose solution intake versus total (water + sucrose liquid) intake.

**Tail Suspension Test (TST)**
The apparatus is consisted of 6 chambers which allowed 6 animals to be videotaped and tested at one time. In a chamber which is isolated acoustically and visually, an individual mouse was individually suspended 1cm from the tip of the tail to the vertical bar with adhesive tape. A camera positioned in front of the TST box was used to record the animal’s behavior for 6 min. Total immobility time during the last 4 min was analyzed by ANY-maze software. No animal climbed its tail during the test.

**Western Blot**
After the mouse behavioral test was finished, the brain was taken on ice. The entire hippocampus and PFC were lysed in RIPA buffer containing protease inhibitors and phosphatase inhibitors. Protein concentration was determined colorimetrically by BCA assay (Pierce, Rockford, IL, United States). Protein lysates were separated by SDS-PAGE electrophoresis and were transferred onto polyvinylidene difluoride (PVDF) membranes. After blocking with 1% BSA for 1 h, the membranes were incubated with primary antibodies. BDNF (Santa Cruz Biotechnology, sc-546, 1 : 400), GluR1(Cell Signaling Technology, #13185, 1:1000), NMDAR1 (Cell Signaling Technology,S104s, 1 : 1000), NR2A (Cell Signaling Technology, #4205, 1 : 1000), NR2B (Cell Signaling Technology, 4212s,1 : 1000), p-mTOR (Cell Signaling Technology, #2971, 1:1000), mTOR (Cell Signaling Technology, #2972, 1:1000), phosphor-Akt (Cell Signaling Technology, #4060, 1:1000), Akt (Cell Signaling Technology, #9272, 1 : 1000), and tubulin (Proteintech, 10094-1-AP , 1 : 2000) were used at 4°C overnight. The next day, blots were washed three times in TBST and incubated with horseradish peroxidase conjugated anti-mouse or anti-rabbit secondary antibody (1:3000) for 1 h. After final three washes with TBST, the bands were visualized using the Super Signal West Pico Chemiluminescent Substrate (Thermo Fisher Scientific Inc.). BDNF, mTOR, Akt, NR1, NR2A, NR2B and GluR1 were all normalized to β-Tubulin bands, and pro-BDNF, mature BDNF, p-mTOR, p-Akt bands were normalized to total protein levels then expressed as a percentage. All experiments were performed 3 times.

**FIGURE 1** | Behavioral effects at different time points after a single ketamine and CLM treatment on OB mice. Control animals (CTL) received vehicle treatment, and animals exposed to olfactory bulbectomy received a single administration of vehicle (OB), Chaihu-jia Longgu-Muli decoction (CLM) or ketamine (KET). (A) Sucrose preference test at 1 week, 2 weeks, 3 weeks, and 4 weeks after olfactory bulbectomy surgery (7-test, *p < 0.05 and n = 6–10/group). (B) TST (ANOVA, F(3,30) = 25.85, p < 0.0001, and n = 7–8/group) was tested at 24 h. (C) SPT (ANOVA, F(3,35) = 57; p < 0.0001, and n = 9/group) was tested at 48 h. (D) TST (ANOVA, F(3,32) = 4.735; p < 0.01, and n = 8–9/group) was tested at 72 h. *p < 0.05, **p < 0.01, ***p < 0.001, compared to CTL; #p < 0.05, ##p < 0.01, ###p < 0.001, compared to Veh. (E,F) OFT was performed 24 h post administration of CLM, KET or saline [F(3,23) = 5.591, p < 0.05 for total distance, ANOVA, F(3,23) = 1.225, p > 0.05 for central time, and n = 6/group].
**Statistical Analysis**

All data are presented as means ± SEM. Two-sample comparisons were carried out using two-tailed Student’s t-test. Others was performed by one-way analysis of variance (ANOVA) followed by the Bonferroni multiple comparison tests. A value of $p < 0.05$ was considered statistically significant.

**RESULTS**

**A Single Dose of CLM Elicited Rapid Antidepressant-Like Activity in OB Mice**

Following the surgery for OB, SPT was performed weekly for 4 weeks to monitor the time course of depression-like response (Figure 1A). OB resulted in a significant decrease in sucrose preference at week 1, 2, and 3 ($p < 0.05$ vs. CTL). By week 4 post surgery, OB group no longer differed from the control group ($p > 0.05$), indicating that the OB surgery induced a relatively long but reversible depression-like behavior. Thus, in the following experiments, animals were treated at week 2 post surgery with a single dose of CLM or ketamine (KET). Then, different behavioral tests were performed at 24, 48, and 72 h post administration. Although the immobility time in OB mice did not differ from the non-OB control group, a single administration of CLM and KET both significantly decreased the immobility time in TST in OB mice at 24 hr, compared to vehicle control ($p < 0.01$, Figure 1B) and at 72 h ($p < 0.05$, Figure 1D). CLM or KET increased sucrose preference in SPT ($p < 0.001$, Figure 1C) post a single administration. By 2 weeks after surgery, mice
showed increased locomotor activity ($p < 0.05$, Figure 1E) in the OFT. Administration of CLM or KET did not affect the time spent in central area or total distance in the OFT (Figures 1E,F).

**CLM Normalized AMPA and NMDA Receptor Expression in PFC of OB Mice**

The expression of NMDA and AMPA receptor subunits in PFC were examined across groups at 24 h post a single dose of CLM...
CLM Activated AKT-mTOR Signaling in the PFC

In the PFC, OB mice showed significantly lower level of total Akt and pAkt \((p < 0.05, \text{Figures 3A–C})\), without change in the ratio of pAkt to Akt. A single dose of CLM or KET improved the deficits in total Akt and pAKT, without effect on pAkt/Akt. Furthermore, total mTOR, pmTOR as well as pmTOR/mTOR were all reduced in the OB mice \((p < 0.05, \text{Figures 3D–F})\), which were reversed by a single dose of CLM \((p < 0.001)\), similar to KET \((p < 0.001, \text{Figures 3D–F})\).

CLM Increased GluR1 but Not BDNF in Hippocampus in OB Mice

The rapid enhancement of BDNF is also a neuroplastic mechanism of rapid antidepressant effect. Interestingly, the expression of BDNF and GluR1 was up-regulated at 24 hr in the veh group, compared to the CTL group \((p < 0.05, \text{Figures 4A, B})\). Both CLM and KET group increased the GluR1 expression \((p < 0.001 \text{ for CLM}, p < 0.001 \text{ for KET, Figure 4C})\), but they didn’t have any difference in mature and pro-BDNF expression compared to Veh \((p > 0.05, \text{Figures 4A, B})\).

DISCUSSION

We aimed to test a traditional Chinese herbal medicine CLM for a rapid antidepressant activity and to examine associated neuromolecular substrate. We found a single dose of CLM instantly induced an antidepressant-like action, which lasted for 3 days. Using the OB depression model in mice, we found the deficits in sucrose preference in OB mice, which was reversed by a single dose of CLM, similar to ketamine. Moreover, in the PFC, mice with OB showed decreased level of expression of GluR1 and increased level of NR1, which was reversed by a single dose of CLM. There was also remarked deficiency in Akt and mTOR signaling regulating neural plasticity in OB mice, which was ameliorated by a single dose of CLM. These results suggested that normalization of NMDA and AMPA receptors and related Akt-mTOR signaling may account for the rapid antidepressant-like activity of CLM.

Rapid Antidepressant-Like Effect of CLM in OB Mice

In OB model, a period of 2 weeks is suggested to be optimal for the development of the bullectomy syndrome, including hyperactivity and alterations in exploration and social behavior (van Riezen and Leonard, 1990). Consistent with these results, we found OB mice were hyperactive in OFT at post surgery week 2, and displayed a robust depression-like phenotype on the basis of the performance in SPT at week 1, but it was reversible by week 4. In this model, it requires 14 days for SSRIs to improve depressive behaviors, recapitulating a minimum of 2–4 weeks of continuous treatment for onset of antidepressant effects of SSRIs in humans (Redmond et al., 1997). The present findings showed that, similar to ketamine, only a single dose of CLM was capable to reverse the deficits in SPT, and the effect lasted for 3 days. Additionally, the immediate and persistent antidepressant effect of CLM was also evident in the test of TST in both OB model and non-OB conditions. As OB chronic depression model is one of the best to be used for testing the rapid antidepressant potential after a single or a few number of dosing (Ramaker and Dulawa, 2017), the results from this study suggest the fast-onset antidepressant-like effect of CLM.

Akt-mTOR Signaling Pathway Was Involved in the Rapid Antidepressant Effect of CLM

We found the level of phosphorylation of AKT and mTOR decreased in PFC of OB mice, which was increased post a single administration of CLM. Akt-mTOR signaling is important for neural plasticity for rapid antidepressant effects. The activation of Akt-mTOR signaling is responsible for persistent increase of expression of synaptic proteins such as GluR1 (Li et al., 2010). In the current study, we found for the first time that the phosphorylation of Akt and mTOR as well as the expression of GluR1 decreased in PFC of OB mice, consistent with the results in the chronic mild stress model and the postpartum depression model (Li et al., 2011; Tang et al., 2015; Xia et al., 2016). A single administration of CLM rescued the phosphorylated Akt, phosphorylated mTOR and expression of synaptic protein GluR1, similar to ketamine. It is worth noting that, the expression of total mTOR or total Akt did not alter in other chronic depression models, but it was down-regulated in OB mice, rescued by a single dose of CLM here. These results may indicate some distinct molecular deficits in the OB model (Chandran et al., 2013). Nonetheless, these molecular deficits were still repairable by a single dose of CLM or ketamine.

The Regulation of Synapse Protein Related to CLM’s Rapid Antidepressant Activity

The current study showed that, contrary to GluR1, NR1 was overexpressed in PFC of OB mice, which was reversed by CLM and ketamine. The expression of NR1 in the PFC or hippocampus was up-regulated in mice of chronic mild stress, learned helplessness and pre-pregnancy stress depression models.
acute administration and the effect lasted for 3 days. In OB of CLM elicited an antidepressant effect on OB mice post an
like activity of CLM. An equivalent to half of clinical dose
The present study demonstrated the fast-onset antidepressant-like activity of CLM. An equivalent to half of clinical dose of CLM elicited an antidepressant effect on OB mice post an
easy administration and the effect lasted for 3 days. In OB
mice, a single dose of CLM rescued the deficient Akt-mTOR
signaling which controls the expression of synaptic proteins
including GluR1. Additionally, the dysregulations of the subunits
of NMDA was also reversed by CLM. These findings suggest
that the activation of Akt-mTOR signaling and normalization
of AMPAR/NMDAR ratio in the PFC may underlie the rapid
antidepressant-like effect of CLM in OB mice.

CONCLUSION
The present study demonstrated the fast-onset antidepressant-like activity of CLM. An equivalent to half of clinical dose of CLM elicited an antidepressant effect on OB mice post an
cutaneous and the effect lasted for 3 days. In OB

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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