Timosaponin derivative YY-23 acts as a non-competitive NMDA receptor antagonist and exerts a rapid antidepressant-like effect in mice

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Aim: N-methyl-D-aspartic acid (NMDA) receptor modulators have shown promising results as potential antidepressant agents, whereas timosaponins extracted from the Chinese herb Rhizoma Anemarrhenae exhibit antidepressant activities. In the present study we examined whether YY-23, a modified metabolite of timosaponin B-III, could affect NMDA receptors in rat hippocampal neurons in vitro, and evaluated its antidepressant-like effects in stressed mice.

Methods: NMDA-induced currents were recorded in acutely dissociated rat hippocampal CA1 neurons using a whole-cell recording technique. C57BL/6 mice were exposed to a 6-week chronic mild stress (CMS) or a 10-d chronic social defeat stress (CSDS). The stressed mice were treated with YY-23 (20 mg·kg$^{-1}$·d$^{-1}$) or a positive-control drug, fluoxetine (10 mg·kg$^{-1}$·d$^{-1}$) for 3 weeks. Behavioral assessments were carried out every week.

Results: In acutely dissociated rat hippocampal CA1 neurons, YY-23 selectively and reversibly inhibited NMDA-induced currents with an EC$ _{50} $ value of 2.8 μmol/L. This inhibition of NMDA-induced currents by YY-23 was non-competitive, and had no features of voltage-dependency or use-dependency. Treatment of the stressed mice with YY-23 not only reversed CMS-induced deficiency of sucrose preference and immobility time, and CSDS-induced reduction of social interaction, but also had faster onset as compared to fluoxetine.

Conclusion: YY-23 is a novel non-competitive antagonist of NMDA receptors with promising rapid antidepressant-like effects in mouse models of CMS and CSDS depression.

Keywords: Rhizoma Anemarrhenae; timosaponin; YY-23; depression; NMDA receptor antagonist; chronic mild stress; chronic social defeat stress; fluoxetine

Introduction
The incidences of psychiatric diseases preceded by depressive disorder have been rising annually as the pressure of life in modern society grows. In response, the development of antidepressants has been ongoing for decades[1]. Tricyclic antidepressants and monoamine oxidase inhibitors were the earliest two types of antidepressants developed, represented by imipramine and iproniazid, respectively, and are still widely used clinically. According to the mechanism of their antidepressant-like effects, the ‘monoamine deficiency hypothesis’ holds the view that a deficiency of synaptic monoamines (e.g., dopamine, serotonin and norepinephrine) was the main cause of depression[2]. However, the clinical application of these agents was limited because of their side effects, which resulted from poor selectivity; therefore, selective serotonin reuptake inhibitors (SSRIs) became the main therapeutic drugs that were widely applied in clinical settings. The most common SSRI is fluoxetine, which has been widely reported to reverse or prevent the depressive-like behaviors induced by chronic mild stress (CMS)[3, 4]. However, the problems related to the slow onset, long timescale and low curative rate of these drugs have not been sufficiently resolved. Therefore, there is an urgent to elucidate the pathogenesis of depressive disease and to develop a new generation of antidepressants with rapid action and fewer side effects.

For the past few years, the emerging role of the N-methyl-D-aspartic acid (NMDA) receptor in psychiatric diseases has gradually attracted the attention of a growing number of researchers. Evidence from post-mortem and in vivo brain imaging studies implicated abnormalities in glutamate signaling in patients with depression[5]. One of the breakthrough discoveries was the observation that the administration of an NMDA receptor antagonist, ketamine, produced a fast
antidepressant-like effect\textsuperscript{[6–7]}. Moreover, the underlying signaling pathways of ketamine also revealed novel targets for antidepressant discovery and development\textsuperscript{[6–11]}. Several NMDA receptor modulators have been developed and currently show promise as potential antidepressant agents with favorable effects and rapid onset in both clinical settings and animal models of depression\textsuperscript{[5]}. The results from these ketamine studies and the preclinical reports on a variety of other agents acting on the NMDA receptor have aroused increasing interest in the development of NMDA modulators as novel therapeutic agents to treat mood disorders\textsuperscript{[12, 13]}. The combination of NMDA receptor antagonists with first-line antidepressants has the potential to become a more effective and efficient therapeutic strategy for treatment-resistant patients or those with suicidal tendencies\textsuperscript{[14, 15]}. Although its beneficial effects were promising, the clinical use of ketamine has been hampered by its psychotomimetic character and toxicity due to drug abuse\textsuperscript{[16]}

The psychotropic effects of natural products have been of interest to many clinicians and scientists due to their lack of side effects and toxicity\textsuperscript{[17, 18]}. Compelling research studies have shown that many types of effective components from natural products were effective agents against central nervous system diseases. For example, huperzine A and l-stepholidine (l-SPD), which have been proven to be efficient at reversing cognitive deficits, appear to be good candidates for the treatment of Alzheimer’s disease and schizophrenia, respectively\textsuperscript{[19, 20]}. Recent research suggests that timosaponin, which is derived from \textit{Rhizoma Anemarrhenae}, has unique sedative action on mice with neural dysfunction caused by stress. Many preclinical studies also indicated that timosaponin may have the potential to play an essential role in both depression and anxiety treatment\textsuperscript{[18, 21]}. Specifically, timosaponin B-II and B-III are the two main bioactive constituents that are thought to improve learning and memory. Moreover, B-III and its derivatives were reported to be important for the main antidepressive activity of timosaponin\textsuperscript{[22, 23]}. Although its detailed mechanism of action remains unknown, the therapeutic potential of timosaponin is obvious, and its mechanism of psychotropic action deserves further exploration.

We have found that there were several metabolites of B-III present in the brain areas that are related to mood disorders after intragastric administration of B-III in both mice and rats. A structurally modified derivative of one of the primary metabolites, YY-23 (Figure 1), elicited significant improvements in stress-related behaviors in our preliminary study\textsuperscript{[24]}. In the present study, we used electrophysiological methods combined with behavioral studies to explore the psychotropic mechanisms of YY-23.

**Materials and methods**

**Animals**

All Sprague–Dawley rats and C57BL/6 mice used in our experiments were purchased from Shanghai SLAC Laboratory Animal Co Ltd and kept in the animal center of the Shanghai Institute of Materia Medica at the Chinese Academy of Science (SIMM, CAS). The protocols were approved by the Institutional Animal Care and Use Committee, and the experiments were carried out in accordance with the EU Directive 2010/63/EU on the protection of animals used for scientific purposes.

**NMDA-current recording**

**Preparation of dissociated hippocampal neurons**

Experiments were carried out according to methods previously described\textsuperscript{[25, 26]}, with slight modifications. In brief, dissociated hippocampal neurons were prepared from 10–14-d-old Sprague–Dawley rats. The whole brain was removed and dissected in oxygenated ice-cold dissociated solution that consisted of the following (in mmol/L): Na\textsubscript{2}SO\textsubscript{4} 82, K\textsubscript{2}SO\textsubscript{4} 30, MgCl\textsubscript{2} 5, HEPES 10, NaPy 1 and glucose 20 (pH=7.4 with NaOH). Hippocampal slices (500 μm) were cut with a vibratome (VT1000S, Leica, Germany), and the CA1 region was dissected out. The CA1 pieces were treated with dissociated solution containing 3 mg/mL protease XXIII (Sigma, USA) at 32°C for 8 min, washed three times with the dissociated solution and then incubated in dissociated solution containing 1 mg/mL bovine serum albumin (Sigma) and 1 mg/mL trypsin inhibitor type II-S (Sigma, USA) for 1 h. Neurons were dissociated with a series of fire-polished Pasteur pipettes by waiting for the neurons to adhere to the bottom of the recording dishes and then replacing the solution with an Mg\textsuperscript{2+}-free external solution that omitted Mg\textsuperscript{2+} and consisted of the following (in mmol/L): NaCl 140, KCl 3, CaCl\textsubscript{2} 2, HEPES 10 and glucose 10 (pH=7.4 with NaOH). Pyramidal neurons with large pyramidal-shaped cell bodies and thick apical dendritic stumps were chosen for this study.

**Whole-cell voltage-clamp recording**

Patch pipets (tip resistance 6–8 MΩ) were pulled with borosilicate glass pipets (SUTTER P-97, SUTTER INSTRUMENT Co, USA) and filled with internal solution containing the following (in mmol/L): KCl 140, HEPES 10, CaCl\textsubscript{2} 1 and EGTA 10 (pH=7.4 with KOH). Current responses were recorded under a whole-cell voltage-clamp configuration using an Axonpatch

![Figure 1. The chemical structure of YY-23.](image)
200B amplifier (Axon Instruments, USA). The membrane potential was held at -60 mV, except when the voltage-depen- 
dency of YY-23 action was tested. The data were filtered at 1 kHz and then acquired online by a computer through an Axon DigiData-1440A interface using the Clampex 10.3 Data Acqui-
sition Module (Molecular Devices, USA).

**Drug application**

NMDA, α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA), glycine and dizocilpine (MK-801) were pur-
chased from Sigma. YY-23 was provided by the Department of Phytochemistry at the Shanghai Institute of Materia Medica, 
dissolved in dimethylsulfoxide (DMSO, Sigma) and then diluted to a series of concentrations with Mg2+-free external 
solution. The DMSO content was maintained below 0.1%, 
except when the efficacy of 15 μmol/L YY-23 was tested, and 
the DMSO control was carried out on this occasion. Other 
drugs were directly dissolved in the Mg2+-free external solu-
tion. The solutions were applied to the recorded neuron through the Rapid Solution Changer (BioLogic RSC-200) with a 10-tube head; 50 μmol/L AMPA, or 100 μmol/L NMDA in 
the presence of 2 μmol/L glycine was administered to induce 
current responses. When tested, YY-23 was applied 10 s prior to 
and during NMDA and AMPA application.

**CMS induction and drug evaluation**

We used the same method described in our previous work24, 
according to the naturalistic rodent modeling procedure of depression reported in previous studies27, 28. Briefly, each 
mouse, with the exception of the control mice, was caged alone and exposed to a series of chronic, unpredictable, mild 
stressors, including 4 min of cold swimming, 12 h of food/water deprivation, 12 h of a soiled cage, 12 h of cage rotation, 
alteration of the normal light and dark cycle, and 12 h of white 
noise (65 dB SPL). These stressors were randomly applied 
every week to avoid habituation and maximize unpredictabil-
ity. C57BL/6 male mice (weighing 20–25 g) were randomly 
divided into different groups depending on their treatment. 
Seven control mice (CON) were kept under standard condi-
tions, in which they were caged together and intragastrically (ig) administered vehicle, 0.5% sodium carboxymethylcel-
lulose (CMC-Na), as a solvent control from week 1 to week 3. 
Seven CMS-induced mice (CMS) were maintained under CMS 
conditions, in which each mouse was exposed to chronic mild 
stressors, caged alone and administered vehicle, 0.5% CMC-
Na (ig), from week 1 to week 3. Six YY-23-treated mice (YY-
23) were maintained under CMS conditions and administered 
20 mg/kg YY-23 (ig) (YY-23 was suspended in 0.5% CMC-Na) 
per day for the last 3 weeks. Six fluoxetine-treated mice (FLX) 
were maintained under CMS conditions and intraperitoneally (ip) administered 10 mg/kg fluoxetine (fluoxetine was dis-
solved in water) per day for the last 3 weeks.

**Tail suspension test (TST)**

We evaluated mice using the tail suspension test each week 
using the method described in a previous study24. Briefly, the 
mice were suspended by placing their tails on the edge of a 
shelf at a height of 80 cm above the floor for 6 min. A blinded 
experiment was conducted to record the immobility time dur-
ing the last 4 min of the testing period, excluding the first 2 
min.

**Sucrose preference test (SPT)**

We evaluated the mice using a weekly sucrose preference test. 
The mice were allowed to choose freely between two bottles 
during 24 h: one bottle contained 1% sucrose solution, and 
the other contained normal drinking water. After the first 
12 h, the position of each bottle was switched to prevent the 
effects of position preference. The two bottles were weighed 
before and after the test. Sucrose preference was calculated as 
follows: sucrose preference (%)=(sucrose intake (g)+sucrose 
intake (g)+water intake (g))×100.

**Open field test (OFT)**

As previously described29–31, the total distance travelled 
and the walking speed of the mice in an open field test were 
measured as indicators of locomotor activity. The mice were 
placed into the arena (40×40×35 cm) and allowed to explore 
freely for 5 min. Locomotor activity was recorded and ana-
lyzed using a Mouse Spontaneous Activity Video Analysis 
System (JLBehv-LAG-4, Shanghai Jiliang Software Technology 
Co, Ltd).

**Chronic social defeat stress (CSDS) induction and behavioral 
testing**

The CSDS model and social interaction test were performed 
according to published protocols32, 33. In brief, after screen-
ing to identify aggressive CD-1 mice, the test C57BL/6 mice 
were exposed to a different CD1 aggressor mouse each day 
for 10 min over a total of 10 d. During this exposure, all of the 
test mice showed signs of stress and subordination, includ-
ing vocalization, flight response and a submissive posture. 
After 10 min of contact, the test mice were separated from the 
aggressor. The test mice were placed in an adjacent compart-
ment of the same cage, separated by a plastic divider with 
holes, where they were exposed to chronic stress in the form 
of threat for the next 24 h. The control mice were housed in 
equivalent cages but with members of the same strain, which 
changed daily. Twenty-four hours after the last session, we 
tested the behavioral consequences of the chronic social defeat 
stress and selected all of the susceptible mice according to the 
published method32. Briefly, the social interaction ratio (SI ratio) 
was obtained by dividing the time spent in the interac-
tion zone when the target was present by the time spent in the 
interaction zone when the target was absent. An SI ratio of 
less than 1, indicating that less time was spent in the presence 
versus absence of a social aggressor, was used as the threshold 
for identifying susceptible mice. All mice were then housed 
individually for 3 weeks, and all of the 21 susceptible mice 
were randomly divided into three groups: the DEFEAT group 
was administered 0.5% CMC-Na (ig) per day, whereas the 
YY-23 group was treated with 20 mg/kg YY-23 (ig) per day,
and the FLX group was treated with 10 mg/kg fluoxetine (ip) per day. Eight control mice were administered 0.5% CMC-Na (ig) per day. We obtained the SI ratio weekly using the same method as previously described.

Statistical analysis
The data are expressed as the mean±SEM. GraphPad Prism 5.0 software (for Mac, US) was used to carry out the statistical analysis, and a value of $P<0.05$ was considered to be significant. For the electrophysiological studies, Student’s two-tailed $t$-test was used to determine statistical significance. The EC$_{50}$ values for NMDA and the IC$_{50}$ values for YY-23 were obtained using GraphPad Prism 5.0 software (for Mac, US). For the tail suspension test, sucrose preference test, open field test and social interaction test, statistical significance was evaluated by two-way repeat measures ANOVA, and post-hoc analyses were performed with the Bonferroni test.

Results
YY-23 selectively inhibited NMDA-induced current and showed features as a non-competitive antagonist
For the purpose of investigating the target mechanism of YY-23, an NMDA current was induced on dissociated hippocampal CA1 neurons using a method previously described$^{[25, 26]}$, with modifications. The application of 3 μmol/L YY-23 reduced the NMDA-induced current by more than 50% ($n=5$, $P<0.01$ vs CON); we observed no effect on the holding current and complete recovery by washing (Figure 2A). The dose-response curve of YY-23 gave an IC$_{50}$ value of 2.8 μmol/L, and the 95% confidence intervals ranged from 1.8 to 4.2 μmol/L (Figure 2B). In contrast, no effect of the same concentration was detected on the current induced by 50 μmol/L AMPA (Figure 1A and 1C, $n=7$, $P>0.05$ vs CON). Next, we examined the inhibitory effect of YY-23 on different concentrations of NMDA (Figure 3A). The NMDA dose-response curve was constructed in the presence and absence of 3 μmol/L YY-23. The pooled data from four neurons demonstrated that the maximal current response to NMDA showed a 40%–50% reduction in the presence of 3 μmol/L YY-23 (Figure 3B). We also found no prominent change in the EC$_{50}$ (46.7±6.70 μmol/L in the control mice; 49.9±9.47 μmol/L with YY-23, $n=4$, $P=0.90$) (Figure 3C), illustrating the non-competitive antagonist characteristic of YY-23.

YY-23 inhibited NMDA-induced current in a voltage-independent or ‘use-independent’ manner
To explore the antagonistic manner of YY-23, we examined the effects of membrane potential on YY-23 inhibition. The current induced by NMDA was detected in the presence and absence of YY-23 when the membrane potential of the neuron was held at −30, −60 and −90 mV (Figure 4A). The application of 3 μmol/L YY-23 inhibited the NMDA-induced currents equivalently at three different holding potentials, and similar results were observed in five neurons (Figure 4B). We observed no significant differences among groups (One-way ANOVA, $F_{(2,14)}=0.038$, $P=0.962$), indicating that the YY-23 inhibitory effect on the NMDA-induced current was voltage-independent.

Figure 2. The effect of YY-23 on NMDA and AMPA-induced current. (A) YY-23 (3 μmol/L) inhibited the current induced by NMDA (100 μmol/L with glycine 2 μmol/L) but had little effect on the current induced by AMPA (50 μmol/L) of representative neurons. The inserted long and short horizontal lines represent the presence of YY-23 and NMDA (or AMPA), respectively. (B) The dose-response curve of the inhibitory effect of YY-23 on NMDA-induced current; 5–6 neurons were detected for every concentration. (C) The statistical diagram of the inhibitory effect of 3 μmol/L YY-23 on NMDA or AMPA-induced current. The data are represented as the mean±SEM. $n=6$ per group. *$P>0.05$, †$P<0.01$ vs CON.
To further understand whether the blocking effect of YY-23 on the NMDA receptor was ‘agonist-dependent’ or ‘use-dependent,’ we examined whether the NMDA-induced currents could be affected by the sustained stimulation of 3 μmol/L YY-23. One representative test of six repetitions was applied, as shown in Figure 4C. The results showed that the inhibitory strength with rapid onset was almost the same with repeated applications of YY-23 and vanished immediately following the washout of YY-23, indicating that the YY-23 inhibition of the NMDA-induced current was not occurring in a ‘use-dependent’ manner.

The antagonistic effect of YY-23 on the NMDA receptor was partially blocked by MK-801 and Mg²⁺

In an effort to understand YY-23’s binding characteristics and to confirm the NMDA receptor antagonistic effect of YY-23, the classic antagonist, dizocilpine (MK-801), and the endogenous blocker, Mg²⁺, were administered, respectively[34]. Dosages of 120 nmol/L for MK-801 and 0.5 mmol/L for Mg²⁺ were selected as working concentrations, based on previous studies[34-36]. The results suggested an evident reduction in the NMDA-induced current under these conditions (Figures 5A and 5B). In the presence of MK-801 and Mg²⁺, the inhibitory effect of 3 μmol/L YY-23 on the NMDA-induced current was attenuated and found to be significantly less than the inhibitory effect of 3 μmol/L YY-23 alone, which elicited more than a 50% reduction (Figure 5C). These results demonstrated that the proven blockers such as MK-801 and Mg²⁺ could partially neutralize the antagonistic effect of YY-23 on the NMDA receptor, proving the NMDA receptor antagonist activity of YY-23.

YY-23 exhibited a prominent antidepressant-like effect on CMS and a social defeat depression model with rapid onset

Because many types of NMDA receptor antagonists or modulators exhibit prominent antidepressant-like effects with rapid onset, we investigated the psychotropic outcomes of YY-23 as a potentially effective agent against affective disorders. Two animal models were used to validate its efficacy on different aspects of mood disorders. First, an unpredictable CMS model was established using the method described in previ-
The NMDA receptor antagonistic effect of YY-23 was partially neutralized by proven blockers. (A) The NMDA-induced current of one of the representative neuron showed a reduction after the sustained application of 120 nmol/L MK-801, and the antagonistic effect of YY-23 (3 μmol/L) was partially neutralized by pre-blocked MK-801. (B) The NMDA-induced current of one of the representative neuron exhibited reduction after pre-applied 0.5 mmol/L Mg²⁺, partially blocking the antagonistic effect of YY-23 (3 μmol/L). The inserted long and short horizontal lines in (A) and (B) represent the presence of YY-23, MK-801, Mg²⁺ and NMDA. The steady status of NMDA current is noted by arrows in (A) and (B). (C) Different combinations of applied drugs inhibited the NMDA-induced current to different extents. The ‘+’ and ‘-’ below the statistical diagram represent the presence and absence of drugs and NMDA. The data are represented as the mean±SEM. *P<0.01 vs CON (only NMDA was applied). †P<0.05 vs MK-801 (NMDA and MK-801 were applied). ‡P<0.05 vs Mg²⁺ (NMDA and Mg²⁺ were applied). n=5–7 neurons for every combination of applied drugs.

Figure 5. The NMDA receptor antagonistic effect of YY-23 was partially neutralized by proven blockers. (A) The NMDA-induced current of one of the representative neuron showed a reduction after the sustained application of 120 nmol/L MK-801, and the antagonistic effect of YY-23 (3 μmol/L) was partially neutralized by pre-blocked MK-801. (B) The NMDA-induced current of one of the representative neuron exhibited reduction after pre-applied 0.5 mmol/L Mg²⁺, partially blocking the antagonistic effect of YY-23 (3 μmol/L). The inserted long and short horizontal lines in (A) and (B) represent the presence of YY-23, MK-801, Mg²⁺ and NMDA. The steady status of NMDA current is noted by arrows in (A) and (B). (C) Different combinations of applied drugs inhibited the NMDA-induced current to different extents. The ‘+’ and ‘-’ below the statistical diagram represent the presence and absence of drugs and NMDA. The data are represented as the mean±SEM. *P<0.01 vs CON (only NMDA was applied). †P<0.05 vs MK-801 (NMDA and MK-801 were applied). ‡P<0.05 vs Mg²⁺ (NMDA and Mg²⁺ were applied). n=5–7 neurons for every combination of applied drugs.
the control mice (post hoc Bonferroni test, all $P<0.001$ vs CON and $P>0.05$ vs DEFEAT). These promising results showed that a 3-week treatment with YY-23 significantly reversed this social avoidance behavior. In addition, YY-23 triggered the rapid onset of an antidepressant-like effect, causing a reversal in the SI ratio at week 1 (post hoc Bonferroni test, all $P>0.05$ vs CON, $P<0.01$ vs DEFEAT for week 1, $P<0.001$ vs DEFEAT for week 2, $P<0.05$ vs DEFEAT for week 3), whereas FLX failed to exert a reversal effect on the SI ratio within 3 weeks ($P<0.001$ vs CON for week 1 and week 3, $P<0.01$ vs CON for week 2, $P<0.05$ vs DEFEAT for week 1, $P>0.05$ vs DEFEAT for week 2 and week 3).

These results suggested that YY-23 could act as a potential antidepressant, significantly reversing depressive behavior in both CMS and CSDS depression models and exhibiting faster onset than FLX treatment.

Discussion

YY-23 acts as a non-competitive NMDA receptor antagonist with fast blocking and unblocking characteristics

In the present study, YY-23 acted as an NMDA receptor antagonist and showed no effect on AMPA-induced current at a concentration demonstrated a remarkable inhibition of NMDA-induced current. Therefore, YY-23 could act as a selective NMDA receptor antagonist. The NMDA dose-response curve indicated that in the presence of YY-23, the maximal response to NMDA was reduced by 40%-50%, with little change in the EC$_{50}$. This result suggested a non-competitive feature of YY-23, indicating that it acted on the allosteric modulatory site of the NMDA receptors, rather than on the glutamate recognition site. Furthermore, the inhibitory effect of YY-23 on the NMDA-induced current was not affected by membrane potential, and sustained administration of YY-23 inhibited
all subsequent responses to the same extent and returned to control levels following the washout of YY-23. These results revealed that the inhibition of the NMDA-induced current by YY-23 showed neither ‘voltage-dependency’ nor ‘use-dependency’ and demonstrated that YY-23 exhibited the binding feature of fast blocking and unblocking. This suggested that YY-23 did not act at a site within the NMDA receptor-channel lumen[25, 34]. This is entirely in contrast to the findings for the non-competitive antagonist, ketamine, as well as the uncompetitive channel blockers, memantine and MK-801[35, 36, 40-42]. MK-801 is poorly tolerated clinically due to its slow unblocking kinetics[34, 43]. YY-23 would be expected to be safer for clinical use because it exhibits fast action with rapid blocking and unblocking rates at low micromolar concentrations.

To validate the direct inhibitory effect of YY-23 on the NMDA receptor, as shown in Figure 5A, 120 nmol/L MK-801 was applied according to specifications in a previous study[35]. The presence of 120 nmol/L MK-801 partially blocked the inhibitory effect of YY-23 on NMDA-induced current. This result demonstrated that the semi-blocked status of the channel could affect the interaction of YY-23 with the channel. The physiological levels of $\text{Mg}^{2+}$ (0.5–1 mmol/L) were a natural blocker of the NMDA receptor, causing a substantial reduction in NMDA-induced current at a resting potential[36]. In the present study, YY-23 still produced an inhibition of NMDA-induced current in the presence of 0.5 mmol/L $\text{Mg}^{2+}$, but this was weaker than the level of inhibition that was observed in the presence of 3 μmol/L YY-23 alone (Figure 5B). This result suggested that when $\text{Mg}^{2+}$ produced an ionic blockade of the channel[36], it affected the interaction of YY-23 with the channel. All of these findings suggested that when the channel of the NMDA receptor was blocked by other blockers, such as MK-801 and $\text{Mg}^{2+}$, the gating status of the channel was changed. Based on this, the inhibitory effect of YY-23 on the NMDA receptor could be directly influenced by an altered channel status. Therefore, we can conclude that YY-23 acts directly on the NMDA receptor, rather than through a non-specific effect, or through intracellular signaling mechanisms, illustrating that YY-23 likely acts as an NMDA receptor antagonist.

All of these results demonstrated that YY-23 may act as a selective, non-competitive NMDA receptor antagonist. In addition, YY-23 may be a safer agent than other traditional NMDA receptor antagonists and blockers.

**YY-23 showed antidepressant potential with rapid onset in CMS and CSDS models**

Many types of animal models have been developed to imitate the clinical syndrome of depression in patients. An animal model induced by CMS was developed and confirmed to be the most similar to the symptoms of clinical depression due to the construct validity, face validity and predictive validity[28, 44, 45]. In the present study, we investigated the effects of YY-23 and FLX on depressive behavior caused by CMS stressors, including alterations in the day/night cycle, rotated or soiled cages, isolated housing and food/water deprivation. A 3-week chronic treatment with YY-23 and FLX reversed the reduced sensitivity to reward (anhedonia) and the reduced desire to struggle (despair). Beyond that, the most striking finding was that despite the same onset as FLX in anhedonia that was indicated by SPT, YY-23 exhibited the rapid ability to reverse the depressive behavior 1 week earlier than the SSRI FLX, which had a delayed onset of 3 weeks. These results demonstrated that YY-23 elicited an antidepressant effect on a CMS depression model with faster onset than the SSRIs that are widely used clinically.

Social avoidance is also the main symptom of clinical depression[46]; moreover, social stressors always control affective-like behavioral responses across a wide variety of mammalian species[32]. Therefore, an animal model reflecting a deficit of social behavior is necessary to explore the pathological mechanism of depression and antidepressant action. CSDS is an increasingly popular model that exploits the ethological relevance of territorial aggression. It is considered to have construct, face, and predictive validity[32, 47] for the assessment of social interaction. The reversal of defeat-induced social avoidance in mice by antidepressants also suggests that this behavioral pathology may be relevant to human depression[33, 47, 48]. Multiple antidepressant agents, such as NMDA receptor antagonists and standard antidepressants, have been validated as being capable of reversing the social interaction effect of social defeat, but long-term treatment is required for the effectiveness of standard antidepressants[30, 48]. Therefore, to further understand the antidepressant-like effect of YY-23, we established the CSDS model using a published protocol and explored the effect of YY-23 and FLX on social avoidance in the present study. Our findings showed that YY-23 prominently reversed the decreased social interaction ratio that was induced by social defeat stress, with a very rapid onset of 1 week. However, FLX failed to rescue the deficit in social interaction within 3 weeks. This result was consistent with those of previous studies, in which FLX showed a slow onset time of 4 weeks[33, 48]. Therefore, these results demonstrated that YY-23 was absolutely effective on social behavior, exerting an antidepressant-like effect in a CSDS model with significantly faster onset than standard antidepressants.

All of these observations provided powerful evidence suggesting that YY-23, acted as a selective antagonist of NMDA receptor and may be more suitable for use as a rapid-acting antidepressant than the SSRIs with slow onset that are used clinically.

**YY-23 possesses a distinct possibility of being a rapid antidepressant-like agent**

The antidepressants used clinically function primarily by elevating the concentrations of synaptic monoamines, such as serotonin, but are associated with various side effects and a long timescale of action. On this occasion, the NMDA receptor antagonist displayed advantageous characteristics, such as rapid action. Various compounds with NMDA antagonist actions are awaiting testing in clinical trials[33]. The hypothesis that NMDA antagonists are effective antidepressants was based on the observation that interfering with NMDA...
antagonists such as MK-801, 1-aminocyclopentane-carboxylic acid (a partial agonist at glycine B receptors) and D,L-2-amino-7-phosphonoheptanoic acid (a competitive antagonist) would mitigate the behavioral deficits induced by inescapable stressors in rodents[59, 60]. Multiple NMDA receptor antagonists were reported to mimic the effects of antidepressants in more complex models, including chronic mild stress and olfactory bulbectomy[51-53]. This clinical and preclinical evidence linked major depressive disorder to dysregulated glutamate neurotransmission. In our previous study, similar observations were made. CMS could cause a deficit in spontaneous-burst firing in PFC pyramidal cells, and YY-23 treatment may enhance glutamate neurotransmission based on reversing the spontaneous-burst firing and could also rescue the CMS-induced decrease in BDNF levels[24]. All of these processes are the pillars of the glutamatergic theory of depression, demonstrating the rapid antidepressant potential of NMDA receptor antagonists. Moreover, the antagonistic effect on the NMDA receptor affords YY-23 the capability to act as a rapid-acting antidepressant.

The possible mechanism involved in the antidepressant-like effect of YY-23

In recent years, the response to intravenous ketamine infusion has now been replicated in multiple studies, including in depressed patients with suicidal ideation, treatment-resistant depression and bipolar disorder[54, 55]. The rapid and robust antidepressant effects of the NMDA receptor antagonists ketamine and traxoprodil provided target validation for the family of ionotropic glutamate receptors[50, 54]. In the present study, we observed that YY-23 could act as a non-competitive NMDA receptor antagonist; moreover, YY-23 indeed exhibited rapid antidepressant-like effects on both CMS and CSDS models. Based on these observations and the research results of other NMDA receptor antagonists, we speculated that some detailed mechanisms may be involved in the antidepressant-like effects of YY-23. With the capacity to block the NMDA receptor, YY-23 could likely enhance neurotransmission by one of two ways. First, if YY-23 blocked the extra-synaptic NMDA receptors, it could reduce signaling via elongation factor-2, thereby disinhibiting BDNF levels[56, 57]. An alternative possibility is that YY-23 acts on synaptic NMDA receptors, stimulating synaptic glutamate release and enhancing synaptic AMPA receptor downstream signaling[6, 8]. Both mechanisms lead to dendritic spine growth, followed by enhanced synaptic plasticity, neurotransmission and functional connectivity[56]. This speculation that YY-23 may enhance synaptic neurotransmission was not only supported by numerous observations related to the effects of other NMDA receptor antagonists[58-60] but was also confirmed by our previous work, which showed the significant ability of YY-23 to reverse the decreased burst firing induced by CMS[24]. Regarding the downstream mechanism, we have noted interesting changes in some of the signaling pathways downstream of the glutamate receptors after treatment with YY-23, but additional work is needed to draw definitive conclusions regarding these changes.

Conclusions

The present study highlights that YY-23, which is derived from timosaponin B-III, could act as a selectively non-competitive NMDA receptor antagonist that does not exhibit voltage-dependency or use-dependency. Behavioral studies aiming to reveal the psychotropic outcomes of YY-23 noted the rapid antidepressant-like effects on both the CMS and the CSDS models. Although further studies will be necessary to elucidate the neurotrophic molecular expression (or signal transduction pathways) and detailed mechanisms of rapid action by YY-23, these results unveiled the target mechanism of the metabolite of timosaponin B-III and confirmed the essential roles of NMDA receptor antagonists in antidepressant treatments. Moreover, we provided evidence of the therapeutic potential of timosaponin for the treatment of depression and demonstrated the more rapid onset of the effectiveness of the compound compared with SSRIs.

Acknowledgements

This work was supported by the National Natural Science Foundation (31128009 and 31171011 to Yang LI), the National Science & Technology Major Project ‘Key New Drug Creation and Manufacturing Program’ (2014ZX09102-001-005 to Cheng-gang HUANG), and the Ministry of Science and Technology (2013CB910601 to Yang LI).

Author contribution

Qi ZHANG performed the electrophysiological experiments; Qi ZHANG, Fei GUO and Bing ZHANG performed the behavioural experiments and analysed the data; Zhi-wen FU and Cheng-gang Huang contributed the new compounds; Qi ZHANG, Cheng-gang HUANG and Yang LI designed the project and wrote the paper.

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