

Reduced Synaptic Plasticity Contributes to Resistance Against Constant-Stimulus Electroconvulsive Treatment in a Rat Model of Stress-Induced Depression

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Purpose: Depression is a common mood disorder in humans worldwide. Electroconvulsive therapy (ECT) remains the most effective treatment for patients with drug-resistant or severe depression; however, during ECT, electrical resistance can occur, antagonizing ECT efficacy. We aimed to investigate how depressed patients develop resistance to electric shocks during ECT.

Methods: Rats exposed to chronic unpredictable stress exert similar impairments in hippocampal synaptic plasticity as those in depressed humans, including hippocampal neuronal atrophy and reduced synaptic function and synapse-related proteins. Therefore, a rat model was used to model depressive-like behaviors in the current study. Depression-like behavior was stimulated in Sprague Dawley (SD) rats that were then randomized into six groups: control group (C); a rat model of stress-induced depression group (D); and four groups in which a rat model of stress-induced depression received one, three, five, or seven electroconvulsive shocks (ECS; DE1, DE3, DE5, and DE7). The sucrose preference test (SPT) and Morris water maze (MWM) were utilized to evaluate anhedonia and spatial learning and memory in rats, respectively. Synaptic plasticity was recorded electrophysiologically in terms of field excitatory postsynaptic potential (fEPSP) and long-term potentiation (LTP).

Results: The rat model of stress-induced depression triggered a decrease in the sucrose preference percentage (SPP) and the baseline fEPSP slope relative to those observed for the C group, and these changes were significantly rescued by ECT in a shock number-dependent manner within five shocks. However, the rat model of stress-induced depression displayed an increase in the escape latency and a decrease in space exploration time, in addition to decreased LTP relative to those in the C group, which was further augmented by ECT in a shock number-dependent manner within five shocks.

Conclusion: Changes in synaptic plasticity might be responsible for the development of resistance against constant-stimulus ECT in a rat model of stress-induced depression.

Keywords: depression, electroconvulsive shocks, electrical resistance, synaptic plasticity

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Introduction

Depression has been predicted to be the second most frequent mood disorder through 2020 by the World Health Organization (WHO).¹ Over 350 million people are estimated to suffer from depression, which is associated with over half of the 800,000 suicides/year worldwide.²

Our previous study indicated that electroconvulsive therapy (ECT), which remains the most effective therapeutic method for patients with drug-resistant or severe depression, is ineffective in up to 30% of depressed patients³ due to inefficacy caused by electrical resistance during ECT.^{4,5} As ECT progresses, the seizure threshold increases by 40–125%; consequently, the electrical charge and number must also increase to maintain the anti-depressant effects, particularly in older patients,⁴ which can severely damage the learning and memory functions in treatment-resistant depression patients.⁶

However, the underlying mechanisms that are responsible for the development of electrical resistance against ECT are complex. One possible reason is synaptic plasticity, which is altered in depression and may underlie the ability of ECT to relieve depression in patients.^{7,8} For instance, similar to depressed patients, rats exposed to chronic unpredictable stress present with impaired hippocampal synaptic plasticity, as indicated by the atrophy of hippocampal neurons and reduced synaptic function and synapse-related protein expression,^{9,10} consequently, the rat model was used to model depressive-like behaviors in the current study. In addition, repeated ECSs have been shown to enhance the synaptic function of the hippocampus, as reflected in increased nerve fiber growth, synapse numbers, and electrophysiological function observed in animal models.^{11,12} Altogether, synaptic enhancement may at least partially underlie the anti-depressive effects of ECT.

Here, we explored whether synaptic plasticity contributes to the development of electrical resistance in ECT using a rat model of stress-induced depression.

Materials and Methods

Rats

Adult male (2–3 months, 200–250 g) Sprague Dawley (SD) rats were maintained in the Laboratory Animal Centre of the First Affiliated Hospital of Chongqing Medical University (Chongqing, China) under controlled conditions of 22–24 °C, 60–64% humidity, and a 12-h day/night cycle, with lights on/off at 8:00/20:00. One week before the experiments, the animals were adapted to the new environment. The procedures of the current study were approved by the Ethics Committee of the First Affiliated Hospital of Chongqing Medical University and were performed in accordance with the US National Institutes of Health Guide for the Care and Use of

Laboratory Animals. All efforts were made to minimize the suffering and the number of rats used in the current study.

Stress-Induced Depression

Rats exposed to chronic unpredictable stress present with similar hippocampal synaptic plasticity impairments as are observed in depressed patients, including the atrophy of hippocampal neurons and reductions in synaptic function and synapse-related proteins,^{9,10,13} consequently, a rat model of chronic unpredictable mild stress, as described in our previous study, was used to model depressive-like behaviors in the current study.¹⁴ The stress procedure is comprised of the following 9 types of stressors: 1) swimming for 5 min in 4 °C cold water; 2) swimming for 5 min in 45 °C hot water; 3) pinching the tail for 60 sec; 4) food deprivation for 1 day; 5) water deprivation for 1 day; 6) shaking for 20 min; 7) continuous lighting for 1 day; 8) being caged in a cage containing damp sawdust for 1 day; 9) being in a cage tilted 45° from horizontal for 1 day. Animals were housed one rat per cage and were randomly subjected to one of the described stressors once/day for 28 consecutive days. The randomization was applied individually to each rat. The same stressor was not used on successive days to ensure the unpredictability of the stimulation.

Electroconvulsive Shock (ECS)

Shocks were delivered under propofol anesthesia (10 mg/mL, 9 mL/kg, intraperitoneal; FX061, AstraZeneca, UK), as described in our previous report.¹⁴ Shocks featuring bidirectional square wave pulses of 120 mC were delivered once/day for seven consecutive days by ear clip electrodes on the Nivique ECT system (Nivique Meditech, Bangalore, India). Sham shocks were delivered as described above, except without the current. Oxygen was supplied to rats before, during and after the shocks, and before the recovery of body movement, the blood oxygen saturation (SpO₂) was monitored to maintain at least 95%. During the whole process of ECS and the recovery period after ECS, there was no obvious inhibition of respiration. As described by our previous report,¹⁵ all rats in the current study also experienced tonic-clonic seizures for at least 10 s in each group after treatment with ECS, indicating the success of ECS treatment. Tonic-clonic seizures in rats weighing over 300 grams can cause spinal cord injuries (SCIs), resulting in bilateral lower-limb palsy. Fortunately, none of the rats (n =15 per group) developed SCI in the present study.

Grouping

The SD rats were randomly allocated to six groups ($n = 15$ per group). The rats in the C group were subjected to neither chronic unpredictable stress nor ECS, whereas those in the D group were subjected to chronic unpredictable stress and sham shocks (as described above, except without the current), and those in the DE1, DE3, DE5, and DE7 groups were subjected to chronic unpredictable stress, followed by the delivery of one, three, five, or seven ECSs. All rats were exposed to oxygen during the experiment. In brief, at the end of the experimental period, nine rats were subjected to behavioral tests, and six rats underwent electrophysiological measurements.

The experimental timeline of the procedure was similar to that used in our previous study¹⁶ and is summarized in Figure 1. Starting the day after the completion of the chronic unpredictable stress model (day 29), baseline measurements were established in nine rats using the sucrose preference test (SPT), and the Morris water maze test (MWM) was performed on days 30–35. Subsequently, the rats received one, three, five, and seven ECS once per day, on days 36, 36–38, 36–40, and 36–42, respectively. The rats were subjected to SPT the day after the last ECS was administered and subjected to the MWM for six consecutive days, starting two days after the last ECS.

Sucrose Preference Test (SPT)

SPT was used to assess anhedonia, as we previously described.¹⁶ On the first day, all rats were supplied with two bottles of 1% (w/w) sucrose to learn how to drink. On the second day, one bottle of sucrose was replaced with sterile water. On the third day, rats were deprived of water and food for 23 h and then provided with access to one bottle containing 1% sucrose and one bottle containing sterile water for 1 h. All rats were free to drink for 30 min, then the position of two bottles were exchanged to prevent position preference. The weight of each bottle was weighed before and after the experiment to determine the sucrose preference of each rat. The SPP was calculated using the following equation: $SPP = \{ \text{sucrose consumption (mL)} / [\text{sucrose consumption (mL)} + \text{sterile water consumption (mL)}] \} \times 100\%$.

Morris Water Maze (MWM)

The MWM is widely used to evaluate spatial learning and memory in rodents, which depends on activity in the hippocampus and N-methyl-D-aspartate (NMDA) receptor-mediated Hebbian plasticity. The MWM was performed as we previously reported.¹⁷ The MWM was conducted in a circular water tank with a diameter of 150 cm, divided into four quadrants (SE, SW, NW, and NE). Water at a temperature of $22 \pm 1^\circ\text{C}$ was rendered opaque using dark, nontoxic, and washable paint. The ZH0065 (Zhenghua Instruments, China) was utilized to record the swimming velocity, trajectory, and time required to reach the platform and the time spent in each quadrant.

Rats were released in each quadrant and allowed as long as 60 sec to find a platform with a diameter of 11 cm that was placed 1.5 cm beneath the water surface in the center of the NE quadrant. Rats were subjected to four trials in different quadrants daily (one from each quadrants) for five consecutive days and placed onto the platform for 15 sec after each trial. For those rats that successfully reached the platform within 60 sec, the time necessary was recorded as the escape latency, which was each day's average of all four trials. For those rats that failed to find the platform within 60 sec, the escape latency was recorded as 60 sec after rats were guided onto the platform.

On the sixth day, the platform in the center of NE was removed. Rats were released to SW and allowed to swim for 60 sec. The time spent swimming in NE was recorded as space exploration time.

Electrophysiology

At 24 h after the completion of ECS treatment, the rats were anesthetized by intraperitoneal injection of sodium pentobarbital (2%, 50 mg/kg). Afterward, cardiac perfusion was performed with artificial cerebrospinal fluid (ACSF) at $0-4^\circ\text{C}$, as we previously reported.¹⁷ The perfused brain was placed into ice-cold ACSF equilibrated with 95% O_2 and 5% CO_2 for 2 min. Next, brain slices were transferred to a perfusion trough and immobilized in the center of the visual field using nylon thread. The hippocampal CA3 and CA1 regions were placed under the microscope, and bipolar stimulator electrodes were placed in the Schaffer collaterals of the CA3 region.¹⁸ We

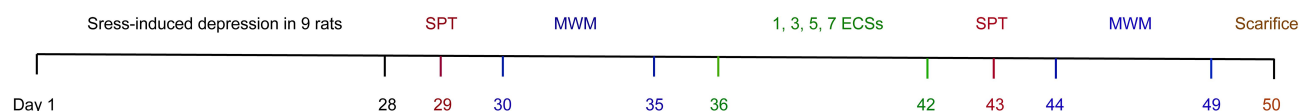


Figure 1 Experimental timeline.

performed the electrophysiology experiments in the rats from one group per day.

Tissues were stimulated by a 0.2-msec square wave with gradually increasing power until the induced fEPSP no longer rose. The stimulation power was adjusted to 50% of this maximum power to induce a baseline fEPSP. When the induced fEPSP waveform and slope were stable, the baseline fEPSP was recorded every 2 min. Three traces were recorded with an interval of 28 sec. The baseline fEPSP was recorded for half an hour, during which time the average slope was recorded as the baseline fEPSP value.

LTP was stimulated by high-frequency stimulation of 200 pulses at 100 Hz. Post-fEPSP was recorded every 2 min for 60 min. Three traces were recorded with an interval of 28 sec. After 30 min of high-frequency stimulation, the average fEPSP slope was recorded as the post-fEPSP value. The magnitude of LTP was calculated using the formula: $LTP (\%) = [(mean \text{ slope post-fEPSP} - mean \text{ slope baseline fEPSP}) / mean \text{ slope baseline EPSP}] \times 100\%$.

Statistical Analysis

Data were analyzed by SPSS 22.0 (IBM, Chicago, IL, USA) and are presented as the mean $\pm$ SD. All data were tested for normal distribution and homogeneity of variance. Intergroup differences in escape latency were assessed by repeated-measures analysis of variance (ANOVA). Post-mortem variables were evaluated using the Student–Newman–Keuls (SNK)-q test, while differences in SPP, space exploration time, baseline fEPSP, and LTP were analyzed by one-way

ANOVA, and the groups were compared using the SNK-q test. $P < 0.05$ indicated a significant difference.

Results

Sucrose Preference Test (SPT)

As shown in Figure 2 and Table 1, before ECT treatment in SD rats, the SPP differed significantly among the C, D, DE1, DE3, DE5, and DE7 groups ($F = 19.33$, $P < 0.001$). SPP was significantly reduced by approximately 30% in the D, DE1, DE3, DE5, and DE7 groups compared with that in the C group ($P < 0.01$). In addition, no significant difference was observed between the D group and any of the DE groups ($P > 0.05$).

After the ECT treatment of rats exposed to chronic unpredictable stress, SPP differed significantly among the six groups ($F = 22.52$, $P < 0.001$), which was reversed by ECS application in a shock number-dependent manner; it was significantly higher in the DE5 group and DE7 groups ($P < 0.01$) compared with the D group, but no significant difference was observed between the D group and the DE1 or DE3 groups ($P > 0.05$). Additionally, no significant differences in SPP were observed between the DE1 and DE3 groups ($P > 0.05$) or between the DE5 and DE7 groups ($P > 0.05$).

Morris Water Maze (MWM)

No significant difference was observed in the swimming speeds among the six groups ($P > 0.05$, Figure 3A).

As shown in Figure 3B, the escape latency for each group decreased over the course of the five-day training

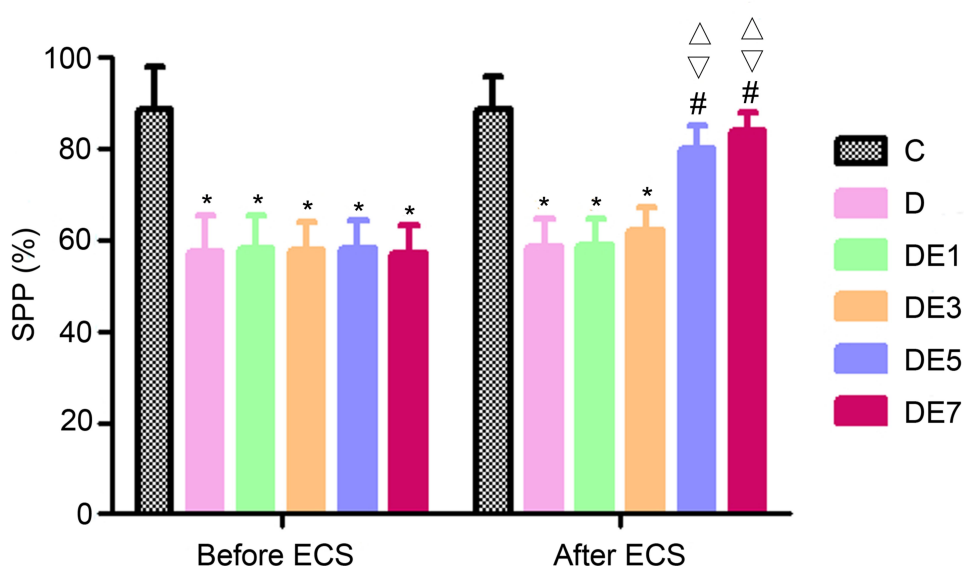


Figure 2 SPP before and after ECSs. * $P < 0.05$ vs. C. # $P < 0.05$ vs. D. ^ $P < 0.05$ vs. DE1. ^ $P < 0.05$ vs. DE3.

Table 1 SPP of Each Rat Before and After ECS

	Rats/Groups	Rat 1	Rat 2	Rat 3	Rat 4	Rat 5	Rat 6	Rat 7	Rat 8	Rat 9	Mean $\pm$ SD	P
Before ECS	C	81.87	86.97	92.32	92.00	87.88	70.33	91.51	102.31	94.56	88.90 $\pm$ 8.95	
	D	63.24	68.24	57.25	60.37	52.62	63.65	58.25	47.77	45.44	57.43 $\pm$ 7.57	a
	DE1	48.89	62.28	51.23	61.03	64.23	58.80	68.73	59.59	50.51	58.37 $\pm$ 6.79	a
	DE3	57.76	59.55	67.66	57.76	57.76	59.80	57.70	57.76	44.13	57.76 $\pm$ 6.03	a
	DE5	55.57	58.39	61.50	47.36	58.39	56.97	70.57	58.39	58.41	58.39 $\pm$ 6.02	a
	DE7	58.30	46.16	60.09	57.58	51.75	62.01	52.04	61.73	64.82	57.16 $\pm$ 6.02	a
After ECS	C	81.64	82.96	82.13	94.23	94.27	94.00	92.84	81.85	98.21	89.13 $\pm$ 6.79	
	D	54.31	65.79	60.01	62.17	52.07	59.87	65.32	60.45	48.71	58.74 $\pm$ 5.89	a
	DE1	61.91	63.80	52.40	52.69	67.37	63.37	55.45	56.10	61.28	59.37 $\pm$ 5.40	a
	DE3	55.31	55.39	63.25	66.14	67.70	63.11	66.00	58.21	67.07	62.46 $\pm$ 4.94	a
	DE5	78.29	78.10	81.68	80.22	73.36	91.47	79.65	80.79	78.38	80.22 $\pm$ 4.80	bcd
	DE7	87.84	78.74	87.24	81.55	80.05	87.27	83.21	86.74	86.95	84.40 $\pm$ 3.55	bcd

Notes: ^a $P < 0.05$ vs C. ^b $P < 0.05$ vs D. ^c $P < 0.05$ vs DE1. ^d $P < 0.05$ vs DE3.

period and differed significantly among the six groups ($F = 8.58$, $P < 0.001$), with significantly longer escape latencies recorded for the D group than for the C group ($P < 0.05$). In rats exposed to chronic unpredictable stress and treated with ECT, the escape latency was significantly lengthened by ECS applications in a shock-number-dependent manner.

The escape latency increased significantly in the DE5 and DE7 groups compared with that in the D group ($P < 0.05$), but these increases were not significant for the DE1 or DE3 groups ($P > 0.05$). No significant differences in escape latency were observed between the DE1 and DE3 groups ($P > 0.05$) or between the DE5 and DE7 groups ($P > 0.05$).

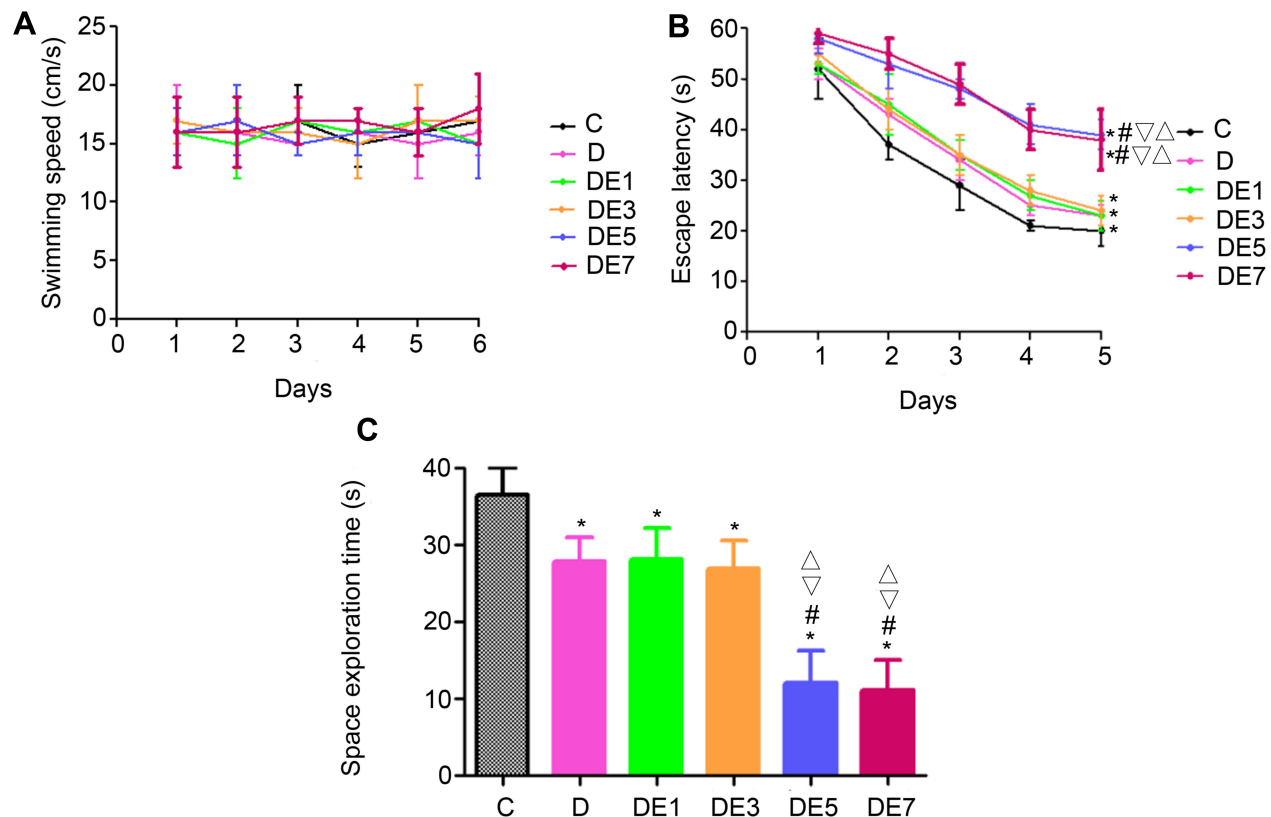


Figure 3 MWM outcomes for different groups. (A) Swimming speed. (B) Escape latency. (C) Space exploration time. * $P < 0.05$ vs C. # $P < 0.05$ vs D. $\nabla P < 0.05$ vs DE1. $\Delta P < 0.05$ vs DE3.

On day 6 of the MWM, the space exploration time was measured (Figure 3C and Table 2), which differed significantly among the six groups ($F = 10.229$, $P < 0.001$). The space exploration time was significantly shorter in the D group compared with that in the C group ($P < 0.05$), which was further significantly shortened by ECS application when using the increased shock numbers in the DE5 and DE7 groups ($P < 0.05$) but was not significantly shortened in the DE1 or DE3 groups ($P > 0.05$) compared with that in the D group. No significances in space exploration time were found between the DE1 and DE3 groups ($P > 0.05$) or between the DE5 and DE7 groups ($P > 0.05$).

Constant-Stimulus ECSs and Baseline Field Excitatory Postsynaptic Potential (fEPSP) in the Hippocampal SC-CA1 Pathway

Diagrammatic representation of a hippocampal cross-section, showing the placement of the stimulating and recording electrodes, is presented in Figure 4A.

As exhibited in Figure 4B and C and Table 3, baseline fEPSP differed significantly among the six groups ($F = 66.127$, $P < 0.05$). Baseline fEPSP was significantly lower in the D group than in the C group ($P < 0.05$), which was rescued by ECS application in a shock number-dependent manner in the DE3, DE5, and DE7 groups ($P < 0.01$) but not in the DE1 group ($P > 0.05$) in comparison with the D group. Moreover, the baseline fEPSPs in DE5 and DE7 groups ($P < 0.05$) but not in the DE3 group ($P > 0.05$) were significantly higher than that in the C group. No significance was observed in baseline fEPSP between the DE5 and DE7 groups ($P > 0.05$).

Constant-Stimulus ECSs and Long-Term Potentiation (LTP) in the Hippocampal SC-CA1 Pathway

As shown in Figure 5A and B and Table 4, LTP was stimulated using a high-frequency stimulation procedure, which resulted in significantly different outcomes among

the six groups ($F = 43.522$, $P < 0.05$). LTP was significantly reduced in the D group compared with the C group ($P < 0.05$), which was further lowered with the use of increasing ECS numbers in the DE3, DE5, and DE7 groups ($P < 0.05$) but not DE1 group ($P > 0.05$). Additionally, no significant differences in LTP were found between the DE5 and DE7 groups ($P > 0.05$).

Discussion

Until now, synaptic plasticity, which is altered during depression, has been considered to underlie the ability of ECT to relieve depression and represent a potential factor in the development of electrical resistance against ECT,^{7,8} although the precise underlying mechanisms remain undiscovered. To the best of our knowledge, the present study provides the most detailed insights thus far into the potential mechanisms underlying electrical resistance against ECT in a rat model of stress-induced depression, which was manifested in the progressively improved SPP and the increased synaptic function (fEPSP) observed with the increase from one to five shocks, peaking at five shocks and not increasing further at seven shocks. In addition, MWM and LTP gradually worsened with increasing shocks, likely reflecting the increasingly severe damage to learning and memory functions.

The current study utilized a chronic, unpredictable, mild stress paradigm to stimulate depression-like symptoms in SD rats, which is a model of “reactive” depression that presents with similarities to depression observed in humans.¹³ Anhedonia remains the core symptom of depression,¹⁹ which was assessed using the SPT in the current study and identified a significantly lower SPP among the rats in the D group compared with the rats in the C group, indicating the successful establishment of a rat model of stress-induced depression. In addition, multiple ECSs significantly increased SPP, indicating the ability of ECS applications to produce anti-depressive effects, consistent with the existing global literature and our

Table 2 Space Exploration Time of Each Rat

Rats/Groups	Rat 1	Rat 2	Rat 3	Rat 4	Rat 5	Rat 6	Rat 7	Rat 8	Rat 9	Mean $\pm$ SD	P
C	30.35	36.28	39.72	38.71	38.53	35.19	35.07	33.55	41.56	36.55 $\pm$ 3.45	
D	28.95	23.59	30.73	27.86	25.21	31.08	28.12	27.89	28.18	27.96 $\pm$ 3.01	a
DE1	25.62	32.12	23.99	25.25	32.47	30.55	24.01	32.59	28.40	28.33 $\pm$ 3.59	a
DE3	30.75	22.46	29.30	24.23	29.89	24.97	23.97	29.49	27.32	26.93 $\pm$ 3.52	a
DE5	6.92	15.18	8.68	17.36	10.95	13.55	11.06	13.17	12.09	12.11 $\pm$ 4.40	abcd
DE7	8.25	14.29	16.03	12.02	9.91	6.89	9.79	10.96	12.23	11.15 $\pm$ 3.89	abcd

Notes: ^a $P < 0.05$ vs C. ^b $P < 0.05$ vs D. ^c $P < 0.05$ vs DE1. ^d $P < 0.05$ vs DE3.

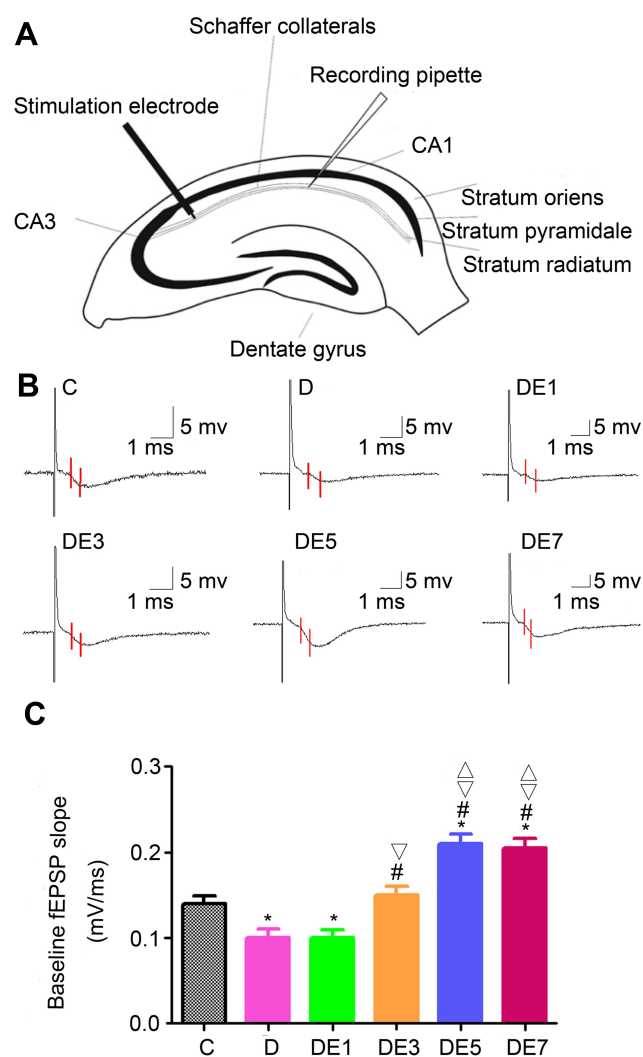


Figure 4 Baseline fEPSP in the hippocampal SC-CA1 pathway. (A) Diagrammatic representation of a hippocampal cross-section showing the stimulating and recording electrodes. (B) Field potentials recorded from hippocampal SC-CA1. The baseline fEPSP slope was measured for the region between the red lines. (C) Comparison of baseline fEPSP slopes. * $P < 0.05$ vs C. # $P < 0.05$ vs D. ∇ $P < 0.05$ vs DE1. $^{\Delta}P < 0.05$ vs DE3.

previous report.^{17,20,21} Consistently, the present study indicated that the use of an increasing number of ECSs resulted in an increased anti-depressive effect, as observed by the upregulation of SPP with the increasing number of

shocks, although the effect peaked at five shocks and did not further increase at seven shocks.

As we previously reported, MWM is commonly used for the examination of hippocampus-dependent spatial learning and memory functions in rodents.¹⁷ Consistently, in the current study, damage to the hippocampus was indicated by longer escape latencies and shorter space exploration times. In rats exposed to chronic unpredictable stress, the escape latency was significantly prolonged by ECS application for up to five shocks, with no further prolongation observed at seven shocks. The space exploration times were dramatically shortened in rats exposed to chronic unpredictable stress and treated with five ECSs, but no further shortening was observed at seven ECSs. According to the results of the current study, the application of ECSs using five and seven shocks reduced the symptoms in a rat model of stress-induced depression by reducing anhedonia, which was accompanied by impaired learning and spatial memory.

In animal models exposed to chronic unpredictable stress, ECS application promotes synaptic growth in the hippocampus and induces changes that resemble functional LTP.^{22,23} Moreover, synaptic changes have been suggested to represent a potential mediator of the anti-depressant effects of ECSs.^{7,8} In the rat model of stress-induced depression used in the present study, the baseline fEPSP slope significantly increased with the increased number of shocks, peaking at five shocks; analogously, LTP was remarkably decreased with increasing shocks, reaching a minimum value at five shocks. These results further suggested that reduced synaptic plasticity is positively associated with electrical resistance against anti-depressive ECT at five shocks, and no significant difference was observed between the DE5 and DE7 groups. Moreover, ECS application using five and seven shocks to a rat model of stress-induced depression reduced anhedonia, as measured by SPT, while simultaneously exhibiting impaired learning and spatial memory, as measured by the

Table 3 Baseline fEPSP of Each Rat

Rats/Groups	Rat 1	Rat 2	Rat 3	Rat 4	Rat 5	Rat 6	Mean $\pm$ SD	P
C	0.14	0.15	0.13	0.14	0.14	0.14	0.14 $\pm$ 0.01	
D	0.10	0.09	0.10	0.08	0.12	0.11	0.10 $\pm$ 0.01	a
DE1	0.10	0.09	0.09	0.08	0.09	0.10	0.10 $\pm$ 0.01	a
DE3	0.14	0.14	0.15	0.15	0.15	0.15	0.15 $\pm$ 0.01	bc
DE5	0.19	0.23	0.21	0.21	0.20	0.21	0.21 $\pm$ 0.01	abcd
DE7	0.21	0.17	0.21	0.20	0.19	0.19	0.20 $\pm$ 0.02	abcd

Notes: $^aP < 0.05$ vs C. $^bP < 0.05$ vs D. $^cP < 0.05$ vs DE1. $^dP < 0.05$ vs DE3.

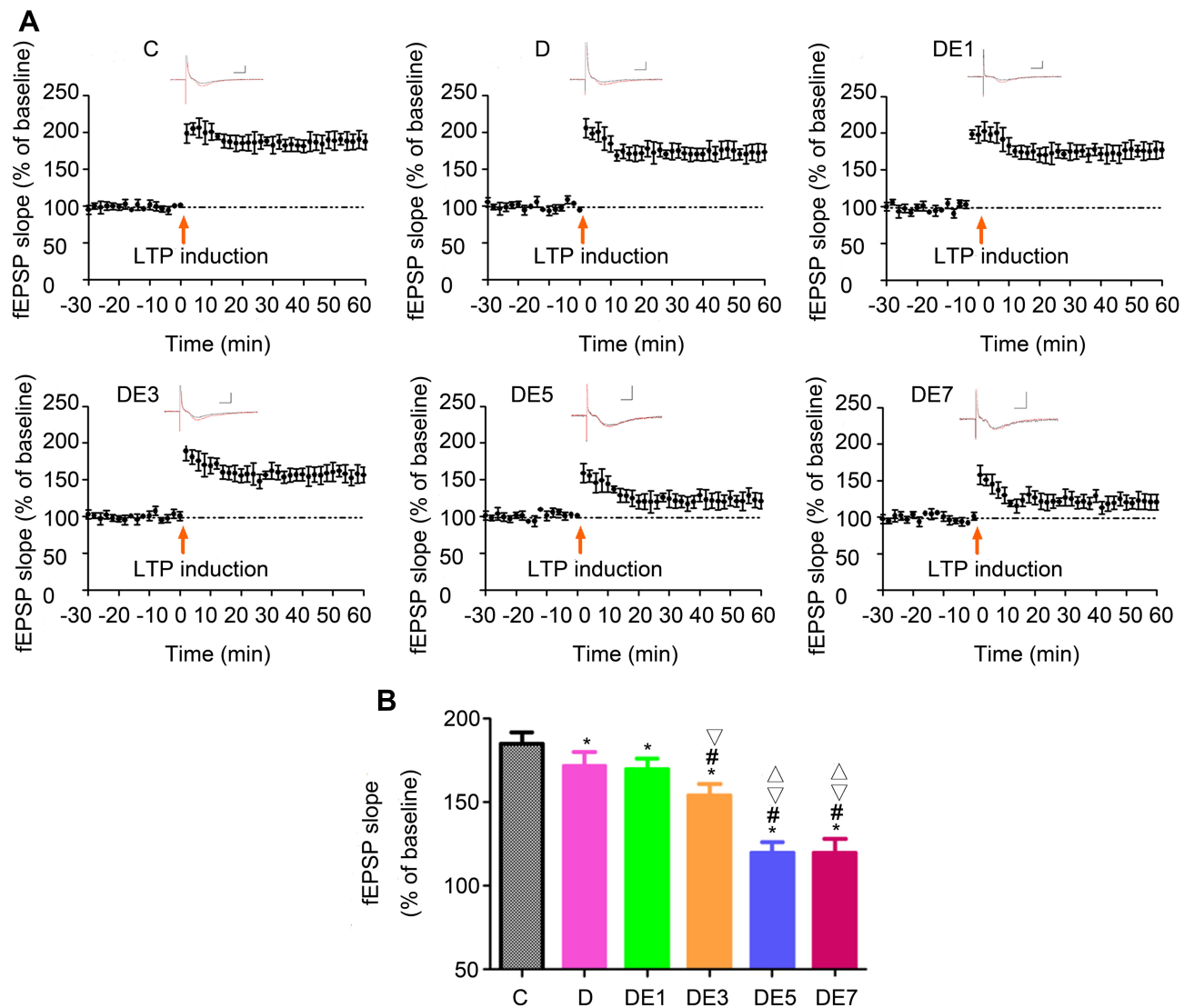


Figure 5 LTP in the hippocampal SC-CA1 pathway. (A) Post-fEPSP after high-frequency stimulation. The black trace indicates the pre-stimulation baseline; the red trace indicates the post-stimulation recording. Horizontal scale bar, 5 ms; vertical scale bar, 1 mV. Arrows indicate the stimulus (200 pulses at 100 Hz). (B) Comparison of post-fEPSP. * $P < 0.05$ vs C. # $P < 0.05$ vs D. Δ $P < 0.05$ vs DE1. Δ $P < 0.05$ vs DE3.

MWM, which corresponds well with the observed reduction of LTP magnitude.

Taken together, these results indicated the exiting of a “ceiling effect” of five ECSs applied with constant power

in the present study, which was observed in the SPT, fEPSP, LTP, and MWM analyses and mirrors the electrical resistance against ECT observed in patients with depression, as reported by previous studies^{4,5} Furthermore, the

Table 4 fEPSP of Each Rat

Rats/Groups	Rat 1	Rat 2	Rat 3	Rat 4	Rat 5	Rat 6	Mean ± SD	P
C	176.32	189.46	189.31	179.30	192.45	188.28	185.85 ± 6.45	
D	166.85	181.39	180.55	162.30	176.27	166.60	172.33 ± 8.11	a
DE1	162.82	174.64	175.62	172.20	171.86	164.10	170.21 ± 5.43	a
DE3	154.23	155.50	154.23	146.50	148.68	166.26	154.23 ± 6.88	abc
DE5	118.96	121.59	111.23	121.94	116.58	129.66	119.99 ± 6.15	abcd
DE7	122.58	132.59	118.30	123.51	109.60	116.52	120.52 ± 7.73	abcd

Notes: ^a $P < 0.05$ vs C. ^b $P < 0.05$ vs D. ^c $P < 0.05$ vs DE1. ^d $P < 0.05$ vs DE3.

findings of the present study were consistent with a previous study which demonstrated the limited ability of ECSs to amplify the progenitors required to increase the generation of neurons when applied at 10 shocks, without observable effects on depression-like behaviors.²⁴ In addition, another report showed significantly more apoptotic cells in animals treated with 10 ECSs compared with those in sham animals, but no significant difference was observed in the numbers of apoptotic cells between animals treated with 20 ECSs and sham animals.²⁵

In the current study, ECS application exhibited a shock number-dependent effect on the alleviation of a rat model of stress-induced depression up to five shocks, with larger numbers of ECSs associated with a higher SPP, longer escape latency, shorter space exploration time, higher baseline fEPSP slope, and reduced induction of LTP in rats exposed to chronic unpredictable stress. In addition, in the current study, the rats in the C and D groups were not exposed to propofol. In our previous study, propofol was not found to significantly affect brain plasticity in rats exposed to chronic unpredictable stress (16). In the current study, we aimed to demonstrate the effects of different numbers of ECSs on synaptic plasticity among the DE1, DE3, DE5, and DE7 groups, with only one variable: the number of ECS applied. Therefore, we acknowledge that the impacts of propofol were not considered; however, the findings in the current study support our view. We provide strong evidence indicating that reduced synaptic plasticity contributes to resistance against ECS during constant-stimulus ECT in a rat model of stress-induced depression, as no significant differences were observed between the DE5 and DE7 groups.

However, three limitations exist in the current study: 1) the changes in the body weight in each group were not measured; 2) although the rat model used in this study resembles the human model of depression, these findings must be applied with caution due to rheological differences in depression between rodents and humans; and 3) the rats exposed to chronic unpredictable stress in our study were initially healthy, whereas depression in humans is often comorbid with other diseases; therefore, our findings should be verified and extended in studies with a larger number of animals and in preclinical and human trials in the future.

Disclosure

The authors report no conflicts of interest in this work.

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