Effects of oral administration of the competitive N-methyl-D-aspartate antagonist, CGP 40116, on passive avoidance, spatial learning, and neuromotor abilities in mice

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ABSTRACT: The effects were investigated of the potent competitive N-methyl-D-aspartate (NMDA) receptor antagonist CGP 40116 [D-(E)-2-amino-4-methyl-5-phosphono-3-pentenoic acid] on the performance of mice in water maze and passive avoidance tasks, and in wire suspension, rotarod, and cage activity tests. The drug was administered per os (p.o.) in its anticonvulsant dose range. CGP 40116 dose-dependently impaired passive avoidance learning when given before, but not when given after training. The antagonist (5, 10, and 20 mg/kg, administered 4 h before each training session) dose-dependently affected water maze acquisition, and impaired retention test performance in both hidden- and visible-platform water maze tasks. In addition, the drug dose-dependently decreased swimming speed during water maze acquisition. Repeated administration of CGP 40116 (20 mg/kg, p.o.) persistently decreased cage activity and wire suspension test performance, whereas motor coordination and equilibrium on the rotarod apparatus remained unimpaired. In our administration protocol, no tolerance was found to the effects of the drug on passive avoidance learning and neuromotor abilities. The parallel effects of CGP 40116 on memory and motor performance are discussed, and it was concluded that the antagonist impairs neuromotor abilities and also induces memory impairments which cannot be entirely reduced to motor interference. © 1999 Elsevier Science Inc.

KEY WORDS: Competitive NMDA antagonist, CGP 40116, Oral administration, Spatial learning, Passive avoidance, Neuromotor performance.

INTRODUCTION

N-Methyl-D-aspartate (NMDA)-type excitatory amino acid (EAA) receptors have been included in a large number of physiological and pathophysiological processes [8]. Although NMDA antagonists impair performance on different learning tasks, spatial learning processes appear to be particularly vulnerable to NMDA receptor blockade [4,5,12,28,35,36]. However, the possibility that the apparent learning impairments are due to other adverse effects of the drug on attention, motivation, or sensory-motor control was not ignored by many authors [20,22,31,50]. More recent studies indeed describe motor impairments caused by systemic administration of competitive and non-competitive NMDA antagonists [1,6]. The effects of NMDA antagonists on neuromotor abilities might interfere with the acquisition or retrieval phase of learning tasks, resulting in decreased performance not caused by memory impairment.

The compound used in this study, CGP 40116, is the active D-enantiomer of the competitive NMDA antagonist, 2-amino-4-methyl-5-phosphono-3-pentenoic acid (CGP 37849, 4-methyl-APPA), and the most potent NMDA receptor ligand presently available. The antagonist was shown to have anticonvulsant activity in several rodent and primate models of epilepsy [7,14,17,19,21,48], to be efficacious in a rat model of depression [42], and to protect against different types of experimentally induced neuronal damage [18,27,38–41,45–47]. Schmutz et al. [47] found that ataxia, muscular hypotonia, and motor incoordination occurred at oral doses well above the anticonvulsant values, but this was later contradicted by studies showing motor impairments in mice within the anticonvulsant dose range [19,48]. Ylinen et al. [51] found that CGP 37849 and its carboxylethyl ester CGP 39551, administered intraperitoneally, protect rats against seizures at doses not impairing spatial learning and swimming speed in the Morris-type water maze. However, their conclusions were based on acquisition trials only, without performance of retention tests, and without any additional behavioural information.

In this study, we have assessed the behavioural effects of orally applied CGP 40116 in its anticonvulsant dose range. With the tests used, we have studied the effect of the compound on neuromotor
abilities as well as its effect on memory performance. We have
tested the compound’s effect on passive avoidance retention when
administered before or after the acquisition trial, and in animals
which had received previous treatment with the antagonist.

The dose-dependent effects on spatial learning were examined
in the Morris-type water maze. Both in the hidden-platform and the
simplified visible-platform condition of the task, the effects of the
compound on the spatial component of the task were examined
using no-platform retention tests (probe trials). Finally, we tested
the effects of the compound on neuromotor performance in cage
activity, wire suspension and rotarod tests, while applying the
same drug-administration protocol as in the water maze experi-
ment.

MATERIALS AND METHODS

Animals and Drug Administration

Seven-week-old male C57BL/6J mice (20–25 g) were used.
The mice were housed in groups of eight under standard laboratory
conditions (food and water *ad libitum*; room temperature of 20–
22°C, and approximate humidity of 55%; 12:12-h light–dark cy-
cles: 0800 h, lights on; 2000 h, lights off). For some of the
experiments, animals were put on reversed light–dark cycle (2000
h, lights on; 0800 h, lights off). Different groups of mice were used
for the passive avoidance and water maze protocols, and neu-
romotor assessment tests. The same animals were never subjected
to different testing procedures as prior handling and/or drug treat-
ment may influence behavioural results.

The competitive NMDA antagonist CGP 40116, d-(E)-2-ami-
no-4-methyl-5-phosphono-3-pentenoic-acid, was kindly provided
by Dr. M. Geelhand (Novartis Belgium). The compound was
dissolved in distilled water, and administered orally using metal
feeding cannulae with plastic protection tip (UNO, Zevenaar, The
Netherlands). In experiments with four treatment groups, a group
treated with sham injections of distilled water and three drug-
treated groups (5, 10, and 20 mg/kg) were considered; in the other
experiments, only the highest dose of CGP 40116 (20 mg/kg) was
applied.

Passive Avoidance Learning

Mice were used after 3 days of reversed light–dark cycle. Passive
avoidance learning was assessed using a step-through
procedure, consisting of a single avoidance training trial (always
performed between 1300 h and 1700 h), 72 h later followed by a
test (retention) trial. For training, mice were placed in the small
(5 × 9 cm²) brightly lit compartment of the box. After 5 s, the
sliding door was opened that led to the big (20 × 30 cm²) dark
compartment of the box (all animals entered within 30 s). Upon
entrance of the dark compartment, the door was closed and the
mice received a slight electric foot shock (Coulbourn Instruments
Small Animal Shocker, Allentown, PA, USA; 0.2 mA, 1–2 s).
Upon shocker activation, the dark compartment cover was lifted
and the response of the mice to the electric foot shock was
confirmed. After 72 h, the mice were placed once more in the small
compartment for their test trial, and latency to enter the dark
compartment was noted up to a maximum of 300 s.

To test the effect of CGP 40116 on avoidance learning, mice
were orally injected with CGP 40116 (0, 5, 10, or 20 mg/kg; 10–15
animals per treatment group) 4 h before training (latency for
maximal anticonvulsant effect according to [47]). To test drug
effect on passive avoidance memory maintenance, an additional
group of mice (*n* = 12) was injected with 20 mg/kg of the drug
immediately after training. To assess the effect of prior treatment,
mice received 8 oral injections of 20 mg/kg CGP 40116 (*n* = 8)
or water (*n* = 7), according to the same daily drug administra-
tion protocol applied for the spatial learning task. After this pre-treat-
ment, mice were injected once more with water or CGP 40116 (20
mg/kg), respectively, 4 h prior to training, and tested 72 h later.

Spatial Learning

For the assessment of spatial learning, we used a Morris-type
water maze consisting of a circular grey plastic pool (150 cm in
diameter, 30 cm high) filled with water made opaque by non-toxic
white paint, and maintained at 26°C. A round Perspex platform (15
cm diameter) was placed inside the pool at the centre of the
north-east quadrant. The platform was placed 1 cm below
the water surface in the hidden-platform condition, but was made
clearly visible with a black pole, in the visible-platform condition.
The protocol consisted of two consecutive sessions of four daily
acquisition trial blocks followed by a 3-day rest period (allowing
96 h without drug application), and a single (retention) probe trial
(also indicated in Fig. 2 as P1 and P2). Acquisition trial blocks
were performed daily, and consisted of four acquisition trials
(15-min intertrial interval) during which the animals were placed
in the water facing the side, in random order at one of the four
starting positions (north, south, east, and west). If an animal could
not find the platform within the maximum swimming time of
120 s, it was placed on the platform and had to stay on it for 15 s
before removal, and a maximal escape latency of 120 s was noted.
Probe trials were performed both in the hidden- and visible-
platform condition. During probe trials, mice had to swim for 100 s
in the pool without a platform. Probe trial 1 (P1) was performed 3
days after the fourth acquisition trial block, and probe trial 2 (P2),
3 days after the eighth trial block (also indicated in Fig. 2).

The effect was examined of CGP 40116 treatment on hidden-
and visible-platform water maze acquisition. Treatment was ad-
ministered 4 h before each acquisition trial block, and consisted of
oral injection of distilled water or CGP 40116 solutions in doses of
5, 10, and 20 mg/kg. Groups of 8–12 mice were considered for
each dose in the hidden- and visible-platform condition.

Neuromotor Abilities

Neuromotor ability tests (cage activity, wire suspension, and
rotarod) were repeatedly performed on the same group of animals
(between 1300 h and 1700 h) using the same drug administration
protocol as used for spatial learning assessment (see above). Cage
activity was measured according to a technique modified from
Crawley and Goodwin [10]. Animals on reversed light–dark cycle,
were placed in transparent mouse cages (20 × 25 cm² floor area),
which were put between three infrared photobeams connected to a
microprocessor counter. Activity (expressed as number of beam
crossings) was measured for 2 h in a dimly lit quiet room. For wire
suspension test of grip strength and endurance, the front paws of
the mouse were placed on a horizontal steel wire (0.6 mm thick)
suspended at a height of 46 cm above tabletop. Mice were allowed
to drop twice during the 60-s assessment period before being
considered to fail the test. Following the wire suspension test,
neuromotor coordination and equilibrium were evaluated on a
lab-build rotarod apparatus [15]. Mice were placed on a rotating
plastic rod (3 cm thick, 15 cm high, 12 rpm) above a glass plate.
There were four trials (10-min intertrial interval), and the latency
before the mice fell off the rod was recorded with a 120-s cut-off.

Treatment consisted of oral injection of distilled water (*n* = 7)
or 20 mg/kg CGP 40116 (*n* = 8). Cage activity was measured 2 h
after drug administration; wire suspension and rotarod tests were
performed after cage activity testing, 4 h after drug administration.
Statistics

Significance of differences between means were assessed using analysis of variance (ANOVA) and two-tailed Student’s *t*-test; differences between proportions were assessed using chi-square analysis and Fisher’s exact test. Level of statistical significance was set at 5%. In the passive avoidance experiments, one-way ANOVA was used to assess the effect of treatment (four treatment groups) on step-through latency. In the water maze acquisition experiments, mean escape latency, path length, and swimming velocity (summed per trial block) were analysed using two-way repeated measures ANOVA with treatment (four groups, *df* = 3), trial block (eight repeated measures, *df* = 7) and interaction (*df* = 21) as sources of variation. One-factor ANOVA and Student’s *t*-tests were used to determine the significance of the effects of treatment on the number of target area entries and time spent in the target (north-east) quadrant during the probe trials. Two-tailed Student’s *t*-test was used for post-hoc pairwise comparison of means.

RESULTS

Effect of Pre-Training and Immediate Post-Training Oral Administration on Passive Avoidance Retention

Figure 1A (left part) shows the effect of p.o. CGP 40116 without pretreatment on step-through latency during training and testing trials. In the water-treated control group, all mice promptly entered the dark compartment during the training trial, whereas none of these animals re-entered during the testing trial, 72 h later. Latency during training was not influenced by CGP 40116 administration (one-way ANOVA; *p* > 0.05). However, the compound dose-dependently reduced step-through latency during the testing trials. One-way ANOVA revealed a highly significant effect of the dose of CGP 40116 on step-through latency during testing (*F*(3,52) = 12.963, *p* < 0.001), and pairwise comparison by post-hoc Student’s *t*-test indicated significantly decreased latencies in the 10 mg/kg and 20 mg/kg groups compared to controls. The drug also dose-dependently decreased the percentage of animals not entering the dark compartment within 300 s, as chi-square analysis indicated a highly significant relationship between the dose and percentage of animals not entering (*p* < 0.001; Fig. 1B, left part). Alternatively, when the drug was administered immediately after the training trial, all animals avoided the dark compartment, and no effect of p.o. CGP 40116 on step-through latency was found (one-way ANOVA; *p* > 0.05; not shown in figure).

Similar impairment of passive avoidance learning was seen in a group pre-treated with eight 20 mg/kg CGP 40116 injections (Fig. 1A, B, right part). Six out of eight CGP 40116-treated animals re-entered the dark compartment during the testing trial, whereas seven water-treated animals did not (*p* < 0.007; Fisher’s exact test), and step-through latency was significantly longer in the control group than in the CGP 40116-treated group (*p* = 0.002; Student’s *t*-test). There was no significant difference in step-through latency and percentage of animals not entering between mice with and those without CGP 40116 pre-treatment (*p* > 0.5; Student’s *t*-test and Fisher exact test, respectively).

Effect on Hidden-Platform Water Maze Acquisition

The four groups (0, 5, 10, or 20 mg/kg) showed a progressive decline in escape latency (Fig. 2A, only sham injection and highest dose displayed) and path length during the 8 training days. Two-way repeated measures ANOVA revealed a significant overall effect of treatment on escape latency (*F*(3,36) = 4.545, *p* = 0.008), whereas the interaction between treatment and training day was not significant. Comparison between treatments showed that 5 mg/kg and 10 mg/kg CGP 40116 did not significantly affect escape latency, whereas mice treated with 20 mg/kg did show significantly increased escape latencies compared to controls (two-way repeated measures ANOVA, *F*(1,20) = 21.255, *p* = 0.001; Fig. 2A). Path length was not significantly different between treatments (not shown in figure).

Swimming velocity was significantly decreased in the CGP 40116-treated animals compared to the sham-injected animals (not shown in figure). Two-way repeated measures ANOVA revealed a significant effect of treatment on swimming speed (*F*(3,36) = 11.151, *p* = 0.001). Analyzed separately, all three doses of CGP 40116 were shown to have a statistically significant effect on
swimming speed [5 mg/kg (p < 0.004), 10 mg/kg (p < 0.001) and 20 mg/kg (p < 0.001)]. Compared to a mean velocity (±SEM) of 21.4 ± 0.6 cm/s in sham-treated animals during the eight acquisition trial blocks, 5, 10, and 20 mg/kg CGP 40116-treated mice showed velocities of 18.4 ± 0.7, 17.2 ± 0.8, and 16.3 ± 0.7 cm/s, respectively.

**Effect on Visible-Platform Water Maze Acquisition**

Two-way repeated measures ANOVA revealed a significant effect of treatment on escape latency in the visible-platform condition [F(3,36) = 2.208, p = 0.022]. Path length was not significantly different between treatments, whereas the interaction between treatment and training day did have a significant effect on path length [F(21,240) = 1.673, p = 0.036].

Like in the hidden-platform condition the drug decreased swimming velocity. Two-way repeated measures ANOVA revealed a significant overall effect of treatment on swimming speed [F(3,35) = 14.666, p = 0.001]. Comparison between treatments showed that doses of 20 mg/kg (p < 0.001) and 10 mg/kg (p < 0.001), but not 5 mg/kg (p > 0.05) had a statistically significant effect on swimming speed. Compared to a mean velocity (±SEM) of 23.6 ± 0.6 cm/s in sham-treated animals during the eight acquisition trial blocks, 10 and 20 mg/kg CGP 40116-treated mice showed velocities of 20.2 ± 0.6 and 18.1 ± 0.7 cm/s, respectively.

**Effect of Administration During Acquisition on Probe Trial Retention**

Probe trials 1 and 2 were performed after the first and the second acquisition session (four acquisition trial blocks), both in the hidden- and visible-platform condition (as indicated in Fig. 2). In contrast to the acquisition trials, there were no significant differences in swimming speed between treatment groups during the probe trials. In the hidden-platform condition, time spent in the different quadrants during Probe 1 indicated a change in spatial search pattern (Fig. 3A); for each of the treatment groups: significant effect of quadrant on the time spent in the different quadrants; one-way ANOVA). However, the results of Probe 1 also demonstrate that treatment with CGP 40116 during the acquisition phase impaired performance during the retrieval phase of the task (Fig. 3A–B). Indeed, while the control group showed a preference to search in the target quadrant, the 20 mg/kg-treated animals spent significantly longer time in the adjacent quadrant compared to the target quadrant (Fig. 3A), and the number of target area entries in the 20 mg/kg group was significantly lower than in the sham-treated group (Student’s t-test, p = 0.01; Fig. 3B).

In the hidden-platform condition Probe 2, the differences between control and CGP 40116-treated animals became even more obvious (Fig. 3C–D). Time spent in the target quadrant decreased with increasing dose of CGP 40116 (Fig. 3C), and one-way ANOVA showed a significant effect of treatment on time spent in the target quadrant [F(3,36) = 2.931, p = 0.047]. A significant difference was found between the 20 mg/kg group and the sham-treated control group (post-hoc Student’s t-test, p = 0.032), but not between sham-treated and respective 5 and 10 mg/kg groups. Also in the 20 mg/kg group, the time spent in the target quadrant during Probe 2 was not significantly longer than the time spent in the other quadrants. Target entries also decreased dose-dependently (Fig. 3D), and one-way ANOVA revealed a significant effect of treatment on target entries [F(3,36) = 4.842, p = 0.006]. Pairwise comparison showed a highly significant difference between the 20 mg/kg-treated animals and the control group (p = 0.001; post-hoc Student’s t-test), but not between sham-treated and respective 5 and 10 mg/kg groups.

Treatment with CGP 40116 also affected the performance in the visible-platform condition probe trials (Fig. 4). The number of target entries decreased dose-dependently (Fig. 4B), and one-way ANOVA revealed a significant effect of treatment on the average number of target entries during Probe 1 [F(3,35) = 5.64, p = 0.003]. Subsequent pairwise analysis revealed a significant difference in the number of target entries between the control group and the respective 10 (p < 0.05) and 20 mg/kg (p < 0.01) groups. However, there was no effect of CGP 40116 treatment on time spent in the target quadrant during visible-platform condition Probe 2 (Fig. 4C), nor on entries in the target quadrant (Fig. 4D).

It is interesting to note that sham-treated animals showed no significant difference between the number of target entries in visible-platform condition Probe 1 and 2 (Fig. 4 B,D), indicating that no additional increase in spatial accuracy was obtained between the two probe trials. In the hidden-platform condition, on the other hand, control animals did show a significantly higher number of target entries during Probe 2 than during Probe 1 (p = 0.026;
Effect on Neuromotor Abilities

During 8 drug-administration days, CGP 40116-treated animals (20 mg/kg) showed persistently decreased cage activity (Fig. 5). Two-way repeated measures ANOVA revealed a highly significant effect of treatment on cage activity \((p = 0.001)\) in an administration schedule parallel to that used in the water maze experiments. The wire suspension test measures muscle strength by examining an animal’s ability to grasp a horizontal wire with its forepaws and remain suspended. A proportion of the mice treated with 20 mg/kg CGP 40116 failed this test, and no indications of improvement were seen during the testing period. All control animals \((n = 7)\) were able to perform the task during the 8 testing days, whereas out of eight CGP 40116-treated mice, seven failed the test on day 5 \((p = 0.001; \text{Fisher’s exact test})\), six on days 2–3 and 6–8 \((p = 0.007)\), five on day 1 \((p = 0.026)\), and four on day 4 \((p = 0.077)\). Finally, in the motor coordination and equilibrium rotarod test, animals of both groups could perform this task without apparent difficulties.

DISCUSSION

This study aimed to investigate the parallel effects of oral administration of the potent competitive NMDA receptor antagonist CGP 40116 on neuromotor and cognitive performance in mice using step-through box, Morris-type water maze and motor assessment protocols. The results of this study are in line with previous findings in suggesting that learning and memory impairments in passive avoidance and water maze tests, following NMDA antag-
onist administration, are not entirely reducible to disruption of sensory-motor processes [12,29,36,37].

We have shown that oral administration of CGP 40116 dose-dependently impairs the acquisition of the passive avoidance response without affecting maintenance or retrieval of stored information. The compound had similar effects in animals which had been pre-treated with the drug. The effects on acquisition are consistent with reported effects of other competitive and non-competitive NMDA receptor antagonists [11,30,35,50]. It has been argued that impaired performance on learning tests could be due to the effect of NMDA antagonists on sensory-motor systems. However, Venable and Kelly [50] obtained no evidence that the drugs decreased shock sensitivity, and in fact found a slight but significant reduction in the shock intensity eliciting vocalization in their mice. In our experiments, the animals’ response to the shock was confirmed during training, and retention was tested when the animals were not under the influence of the drug. The minimal motor requirements of the step-through paradigm minimize the possible contribution of drug-induced motor impairments, and indeed step-through latency during the training phase of the test was not affected by CGP 40116. Our results suggest that CGP 40116 impairs the performance of this task, and affects memory formation in the dose range that was found for the suppression of maximal electroshock-induced seizures in mice [16].

We found that oral administration of CGP 40116 does impair motor performance in the water maze (reduced swimming speed in both hidden- and visible-platform conditions), but in addition appears to decrease dose-dependently the acquisition rate of spatial information in this task. Interestingly, low doses of CGP 37849 have been shown to facilitate learning and memory [23,32,33].
Although we might have observed slightly improved learning ability in the 5 mg/kg treated animals (Fig. 4A), we could not find substantial evidence for such facilitation. Morris and co-workers were the first to show that chronic intraventricular infusion of a competitive NMDA antagonist could block LTP in vivo, and impair spatial learning in the hidden-platform water maze task, while acquisition in the visible-platform task remained unaffected [4,34,35]. It was suggested that this selectivity of the NMDA antagonist-induced impairment indicates that the effect of the antagonist on learning performance is through blockade of NMDA receptor-dependent synaptic plasticity, and not through some gross sensory-motor disturbance [9,35].

In the acquisition of the hidden-platform water maze task, the mice will have to rely mainly on spatial localization strategies. In the visible-platform task, on the other hand, the animals learn to associate the escape platform with the presence of proximal cues. Visible-platform tasks are thus often used as the non-spatial control condition of this test [3]. However, our results in the visible-task probe trials indicate that the animals not only apply cue learning but also spatial search strategies, even when proximal cues are available. This spatial strategy enables them to show a preference for the target quadrant, and to enter the target area when neither the platform, nor the cue are present. This strategy, though apparently not as accurate as in the hidden-platform condition, was shown to be affected by CGP 40116 as well.

The results of Probe 1 indeed indicate that the antagonist affects the spatial component of the visible-platform task. When the drug-treated animals are trained longer, they improve significantly, and reach, in Probe 2, the same spatial accuracy as the control group, whereas the control animals did not show significantly improved spatial accuracy between Probe 1 and 2. The improvement in the drug-treated group might have been a result of tolerance to the effects of the antagonist as tolerance to neuromotor effects of NMDA antagonists has been found in different administration protocols [2,26,44]. However, the neuromotor and passive avoidance results provide no evidence for the development of tolerance within the drug administration protocol applied here. Our findings suggest rather that the antagonist does not abolish the ability for spatial information acquisition altogether, but merely slows down the acquisition process. Slower acquisition rate is likely to explain the difference between control and drug-treated groups in the hidden-platform task also. In this condition, performance of the control group improved significantly between the two probe trials, whereas the difference between probe trial performances in the drug-treated mice was not statistically significant. Acquisition in this more complicated task may be longer than in the visible-platform condition, slowing of acquisition rate would have more profound effects, and this would explain why the effect of the drug is apparent also in Probe 2 of the hidden-platform task.

Cage activity, wire suspension, and rotarod tests were performed using a similar drug administration protocol as during the acquisition of the water maze task. Whereas the wire suspension was clearly impaired in the drug-treated animals, rotarod performance was not different between drug-treated and control animals, indicating that motor coordination and equilibrium were not affected by this kind of administration protocol. Although competitive and non-competitive NMDA antagonists have been shown to exert adverse neuromotor effects in various experimental paradigms, our observations suggest that muscular hypotonia is the main effect following oral administration of competitive NMDA antagonists. The persistent effect of orally administered CGP 40116 on muscle strength or endurance could explain why swimming ability of CGP 40116-treated mice was impaired during 8 trial days of the water maze acquisition test. During the probe trials, which were performed 96 h after drug administration, these effects of the drug were absent. Cage activity was consistently suppressed by CGP 40116 administration. Other studies have reported hyper- as well as hypolactivity as a result of NMDA antagonist administration. Depending on the dose administered, hyperlocomotion or hypolocomotion and ataxia were observed in MK-801-treated kindled and normal rodents and CGP 37849-treated epileptic rats [24,25,49]. Starr and Starr [48], on the other hand, found no hyperactivity in CGP 40116-treated mice, and showed that the drug dose-dependently suppresses motor behavior in mice.

In this study, we have assessed the behavioural effects of oral administration of the potent competitive NMDA receptor antagonist CGP 40116 in mice. CGP 40116 dose-dependently impaired passive avoidance learning when given before, but not when given after training. The antagonist dose-dependently affected water maze acquisition, and impaired retention test performance in both hidden- and visible-platform water maze tasks. In addition, the drug had a dose-dependent effect on swimming speed during water maze acquisition. Repeated administration of CGP 40116 persistently decreased cage activity and wire suspension test performance, whereas motor coordination and equilibrium on the rotarod apparatus remained unimpaired. The present study shows that systemic administration of NMDA receptor antagonists induces both neuromotor- and memory-related impairments.

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