Serotonin-Depleted Rats are Capable of Learning in Active Place Avoidance, a Spatial Task Requiring Cognitive Coordination

T. PETRÁSEK, A. STUCHLÍK

Institute of Physiology, Academy of Sciences of the Czech Republic, Prague, Czech Republic

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Summary
Neurotransmitter substrate of spatial cognition belongs to current topics in behavioral neuroscience. The present study examined the effects of serotonin depletion with p-chlorophenylalanine on learning of rats in active place avoidance, a spatial task requiring allothetic mapping and cognitive coordination and highly dependent upon hippocampus. Serotonin depletion transiently increased locomotor activity in response to footshocks, but it did not change the avoidance efficiency measured by three spatial parameters. These results suggest that serotonin neurotransmission is not crucial for cognitive coordination and allothetic learning, i.e. the processes, which are crucial for active place avoidance performance.

Key words
Spatial cognition • Cognitive coordination • Learning • Place avoidance • Serotonin depletion

Corresponding author
A. Stuchlík, Institute of Physiology ASCR, Vídeňská 1083, 14220 Prague, Czech Republic. Fax. +420 241 062 488. E-mail: stuchlik@biomed.cas.cz

Serotonin or 5-hydroxytryptamine (5-HT) is an indolamine neurotransmitter which exerts many physiological functions in the central nervous system (CNS). The functions regulated by serotonin range from feeding and motor regulation to emotion and aggression (Smythies 2005). The vast majority of serotonergic pathways originate in the brainstem nuclei, known as raphe nuclei, from which axons arise to innervate virtually whole CNS (Stanford 2001). The raphe nuclei can be roughly divided to the caudal nuclei projecting to brainstem and spinal cord, dorsal nuclei, which innervate the cerebral cortex, striatum, thalamus and median raphe nuclei providing innervation to hippocampal formation, septum, and neocortical regions, namely frontal, occipital, parietal and cingulated cortices (Stanford 2001, Smythies 2005). Serotonin is synthesized from tryptophan by tryptophan hydroxylase producing 5-hydroxytryptophan, which is decarboxylated to serotonin. Its synaptic actions are mediated by several classes of receptors (5-HT1-7] which are further divided into subgroups. Action of serotonin is terminated by its reuptake from synapses via 5-HT transporter (Povlock and Amara 1997).

The involvement of serotonin neurotransmission in cognition is a matter of intense debate. Although 5-HT has clear effects on cognition in humans (McEntee and Crook 1991) and degeneration of serotonergic (and cholinergic) fibers is found in Alzheimer's disease (Yamamoto and Hirano 1985), results from the animal studies are less conclusive. Blocking of serotonin synthesis did not induce a deficit in the Morris water maze (Richter-Levin and Segal 1989) and reduction of 5-HT levels by 5,7-dihydroxytryptamine (5-DHT) did not also affect spatial learning (Riekkinen et al. 1990). Contrarily, there is evidence that multiple 5-HT receptor subtypes may interact to regulate animal learning. It is often suggested that serotonin modulates cognitive processes especially by the interaction with cholinergic systems (Zarrindast 2006).

Ten years ago, a cognitive task called active place avoidance was designed in our laboratory (Bureš et al. 1998). In this task, rats are trained to actively explore a slowly rotating arena and avoid an unmarked sector,
which is defined as a stable position in the room. Animals must thus localize the shock sector and walk in the safe part of the arena in a direction opposite to arena rotation. It has been shown that in place avoidance tasks, rats use both the intramaze cues (urine, droppings, and scent marks) as well as visual landmarks located in the room. However, both these classes of information are conflicted by arena rotation in this task. Therefore, the animals must segregate spatial cues into representations of the arena and room (Kubík et al. 2006) and select the room frame as the only relevant coordinate system for navigation. This process was described as "cognitive coordination" (Wesierska et al. 2005). The requirement to separate cues into coherent subsets makes the task useful for studying involvement of receptors in the cognition (Stuchlík et al. 2008). Since the active place avoidance task involves higher cognitive functions and is highly dependent upon hippocampus, which receives 5-HT projections from median raphe nucleus, the present study aimed at testing the hypothesis that brain serotonin depletion via systemic injection of \( p \)-chlorophenylalanine will interfere with efficient learning in this task.

Animal procedures were in accordance with the Animal Protection Code of the Czech Republic and with EU directive 86/609/EEC. Twenty-four male Long-Evans rats (4-5 months, 300-450 g), obtained from the Institute's breeding colony, were housed in pairs in 30 x 30 x 40 cm cages in an air-conditioned vivarium with a stable temperature (22 °C) and 12/12 light/dark cycle. Rats were gently implanted with a hypodermic needle, piercing the rat’s skin between its shoulders, and creating a small loop on the needle with tweezers. The loop provided purchase for an alligator clip connecting a shock-delivering cable. Water and food was freely available.

Serotonin depletion was pursued by systemic administration of \( p \)-chlorophenylalanine (\( p \)CPA; Sigma-Aldrich, CR). \( p \)CPA was suspended in the 0.5 % solution of gum arabic in saline (0.9 % NaCl) and administered in two single (500 mg/kg, i.p.) injections dispensed on each of two consecutive days (injected at 9:00). Total volume of each injection was 4 ml/kg. Behavioral training started 72 h after the second injection. We chose this protocol because it was demonstrated to consistently reduce 5-HT levels in brain by 91 % (Dringenberg et al. 1995, Dyer and Cain 2007). All groups consisted of eight animals. Two control groups were used; one untreated (intact controls) and one injected with gum arabic only (sham-injected).

The apparatus was described in detail elsewhere (Bureš et al. 1998, Stuchlík et al. 2008). Briefly, it was a circular elevated arena located in a room containing many visual cues. Every animal was placed onto the arena, which rotated clockwise at one rotation per minute. The animals were to avoid an unmarked, 60° sector defined by its relationships to room cues. A tracking system (iTrack, Biosignal, USA) recorded the rat’s position every 40 ms. Coordinates were stored for off-line analysis (TrackAnalysis; Biosignal, USA). Whenever the rat entered the to-be-avoided sector, the tracking system delivered a mild electric shock (50 Hz, 0.5 s, 0.2-0.8 mA) and counted an entrance. Shocks were delivered through the implanted needle and the grounded arena floor. The current was gradually increased in the beginning of the first session, starting at 0.2 mA, until rapid escape reaction was elicited. Vast majority animals responded appropriately to 0.4 mA, there were also no systematic differences in the threshold current between groups. Four daily sessions of training in active place avoidance task were conducted, each lasting 20 min, and pursued between 9:00 and 12:00.

The following parameters were analyzed. The total distance walked per session reflected the locomotor activity. Two measures of within-session spatial
performance were the number of errors (number of entrances into the shock sector) and maximum time between two errors (maximum time avoided). Latency to the first error was used as a measure of between-session learning. The data were analyzed with a two-way ANOVA (groups x sessions) with repeated measures on sessions and with the Tukey’s HSD post-hoc test (Statistica 7, StatSoft, CR). Significance was accepted at P<0.05.

Visual observation of control rats and animals treated with pCPA did not reveal any signs of stress during and after injections; all basic behaviors were retained. Animals responded to footshocks with mild startles and rapid escapes, suggesting that the perception of shocks was retained despite serotonin depletion. Examples of typical tracks of rats from all three groups are shown in Figure 1.

First, we analyzed the locomotor activity. The quantification of total distance indicated that there was a difference between groups (Fig. 2A). A two-way ANOVA revealed a significant effect of groups (F (2.21) = 3.95; P<0.05), but no effect of 5-HT depletion on the maximum time avoided (mean ± S.E.M.); another measure of within-session learning.

**Fig. 2.** Panel A reveals the effect of 5-HT depletion on locomotor activity measured as total distance (mean ± S.E.M.). 5–HT depleted rats exhibited higher total distances in the beginning of the training, on the last two days of training, all groups walked similar distance. * P<0.05. Panel B shows no effect of 5-HT depletion on the number of entrances (mean ± S.E.M.), a consistent training-induced improvement is seen in all groups. Panel C illustrates the effect of serotonin depletion on the time to first error (mean ± S.E.M.), a measure to between-session learning; no significant between-group differences were found. A two-way ANOVA and a test of linear trend showed that this parameter increased with training (P<0.05), however, no differences were found between groups (P>0.05). Panel D demonstrates no effect of 5-HT depletion on this maximum time avoided (mean ± S.E.M.); another measure of within-session learning.
consistent training-induced improvement in all groups.

Measuring the time to first entrance again revealed the improvement during training, but no between-group differences (Fig. 2C). A two-way ANOVA revealed no effect of groups (F (2.21) = 1.72; P>0.05), but it found a main effect of sessions (F (3.63) = 13.98; P<0.05). No interaction was found (F (6.63) = 2.16; P>0.05), but it revealed a main effect of sessions (F (3.63) = 28.89; P<0.05). Post-hoc analysis of the factor of sessions revealed a consistent learning-induced improvement in this parameter. Finally, analysis of the maximum time avoided again showed no between-group differences (Fig. 2D). A two-way ANOVA failed to reveal a main effect of groups (F (2.21) = 1.72; P>0.05), no interaction (F (6.63) = 2.16; P>0.05), but it revealed a significant main effect of sessions (F (3.63) = 28.89; P<0.05). Post-hoc analysis again revealed that performance measured by this parameter was improving with training in all groups.

Results of the present study show that serotonin depletion induced by the application of 5-HT biosynthesis inhibitor pCPA did not change the spatial efficiency in the active place avoidance task. No alterations in avoidance were measured by any of the spatially sensitive parameters, including the retrieval-sensitive measure of the latency to the first entrance in a session (Wesierska et al. 2005). The only change that we detected was a slight increase in locomotor activity after pCPA treatment, prominent mainly in the beginning of the training.

Regarding locomotor activity, however, it is usually described in the literature that a decrease in the locomotion is observed after serotonin depletion. For example, Dringenberg et al. (1995) described reduction of spontaneous exploratory activity after pCPA treatment in the open-field test. Importantly, this study reported no overt sensorimotor impairments after the same dose of pCPA as used in the present study. This result is in agreement with our present results, since depleted animals were capable of readily learning the task. Our study detected a transient significant increase in locomotion, which seems to contradict the study of Dringenberg et al. (1995), but there are also reports of increased locomotion after pCPA treatment and/or median raphe nuclei lesion (Köhler and Lorens 1978). Since the 5-HT may increase the reaction of animals to stress (Tanke et al. 2008), we propose that transient hyperlocomotion in the present study (most prominent in the initial session) represents an alteration of the locomotion in response to negative reinforcement (footshocks).

Regarding the absence of effect of 5-HT depletion on the spatial learning in this study, our results are in accordance with several other studies. For instance, Beiko et al. (1997) demonstrated no effect of the pCPA application (in the same regimen and doses as in this study) on learning of rats in the water maze. Similar results were obtained by Dyer and Cain (2007), who found no impairment of water maze learning after pCPA as well. However, another study (Hritcu et al. 2007) found impairments in the working memory (WM), but not in the reference memory in the radial maze, after 5-DHT neurotoxic lesion of serotonergic neurons in the rat. Although WM is a function partially involved in the efficient performance of the active place avoidance task (Stuchlík et al. 2008), the main aspect of the task is reference memory (Bureš et al. 1998), because the shock sector location does not change between sessions. Therefore, the overall performance of 5-HT-depleted rats could remain the same as in controls. Moreover, another study (Cassaday et al. 2003) showed that 5-HT depletion may interfere only with acquisition of rules of WM task, but not with the pure WM. In light of their results, it is conceivable that in the place avoidance task, animal learned the rules of the task in the initial sessions and performance on the following days thus remained unaffected. One of the few memory tasks being significantly impaired by 5-HT depletion is object-recognition paradigm (Lieben et al. 2004); however, there was apparent dissociation between the effects of depletion on the object recognition vs. spatial learning, showing no influence upon the latter (Lieben et al. 2004).

To summarize, the present study shows no effect of serotonin depletion upon cognitive functions measured in the place avoidance task, which involves a process of cognitive coordination, and these result support the concept that serotonergic transmission, despite being clearly involved in human learning, is not crucial for spatial cognition in rodents, including spatial mapping and cognitive coordination.

Conflict of Interest
There is no conflict of interest.

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References


STUCHLÍK A, PETRÁSEK T, VALEŠ K: Dopamine D 1 receptors and alpha1-adrenoceptors synergistically modulate locomotion and behavior of rats in a place avoidance task. *Behav Brain Res* **189**: 139-144, 2008.


