Effects of selective antagonism or depletion of the cholinergic system on visual discrimination performance in rats

W.H.I.M. Drinkenburg,¹H.N.P.M. Sonntag,¹C.J.H. Coenders,¹J.S. Andrews² and J.M.H. Vossen¹

¹Department of Comparative and Physiological Psychology, University of Nijmegen, The Netherlands; and ²Scientific Development Group, Organon International B.V., The Netherlands

Correspondence to: W.H.I.M. Drinkenburg, Nijmegen Institute of Cognition and Information, Department of Comparative and Physiological Psychology, University of Nijmegen, P.O. Box 9104, 6500 HE Nijmegen, The Netherlands, E-mail: DRINKENBURG@NICI.KUN.NL

A two-lever simultaneous visual discrimination task was used to study the effects on performance in Long-Evans rats of the muscarinic antagonists scopolamine (0.0125, 0.05, 0.2 and 0.8 mg/kg s.c.), the M₁ antagonist pirenzepine, the M₂ antagonist AF-DX 116, the M₃ antagonist UH-AH 37 (each 3.2, 10, 32 μg/rat, i.c.v.) and the cholinergic depleting agent, hemicholinium-3 (0.04, 0.2, 1.0 and 5.0 μg/rat i.c.v.). Scopolamine dose-dependently decreased accuracy, increased the number of trials on which the rats failed to respond, and significantly lengthened latency to respond. Only the highest doses of hemicholinium-3, pirenzepine and AF-DX 116 reduced accuracy and increased errors of omission as well as response latency. UH-AH 37 reduced overall task performance at 10 and 32 μg, suggesting that antagonism of both M₁ and other muscarinic receptors (including M₃) had a greater effect on performance than selective antagonism of the M₃ or M₂ receptors. These data indicate that the disruptive effects of cholinergic antagonism on attentionally demanding tasks are strengthened by activity at multiple subtypes of the receptor.


INTRODUCTION

Age-related memory deficits, especially those found in Alzheimer's disease, have been attributed to loss of cholinergic function (Davies and Maloney, 1976; Coyle et al., 1983; Perry, 1986). Accordingly, the effects on learning and memory of substances which induce cholinergic hypofunction have been extensively investigated both in humans and animals (for reviews see Kopelman (1986), Hagan and Morris (1988)). The muscarinic receptor antagonist scopolamine has been widely used to induce amnestic effects in many species from rat to human (Hagan and Morris, 1988; Rupniak et al., 1990; Rusted et al., 1991). However, at certain doses of scopolamine, discriminability rather than memory is disturbed (Warburton and Brown, 1971; Ksir and Slifer, 1982; Kirk et al., 1988). Indeed, a broad range of non-specific effects on for example vigilance or attention, have been proposed as possible confounding factors for the putative direct effects on memory (Andrews et al., 1992; Dunnett et al., 1991; Rupniak et al., 1991).

At least five different subtypes of muscarinic cholinergic receptors can be identified within the central nervous system (Levey, 1993). The M₁, M₂ and M₃ receptors have received the most attention from pharmacologists, and consequently little is known about M₄ and M₅ receptors. The postsynaptic M₁ receptors are primarily located in the cerebral cortex and hippocampus (Mash et al., 1985; Araujo et al., 1988; Levey, 1993), whereas M₂ receptors, representing a class of autoreceptors, can be found in the cerebral cortex, various midbrain regions and brainstem nuclei (Messer et al., 1989; Levey, 1993). Lastly, the M₃ receptors are located in the hippocampus as well as in other structures, but are less common in the central nervous system (CNS) than either the M₁ or M₂ receptors. Research into cholinergic based treatments for Alzheimer's disease has concentrated on the M₁ receptor, because this subtype appears to be preserved during the progression of the disease (Mash et al., 1985; Araujo et al., 1988). However, the exact role for the
muscarnic subtypes in cognitive performance is still largely unclear.

Studies on the effects of specific muscarinic antagonists on cognitive performance are rare and studies on discriminability even rarer. Most studies have been confined to the effects of the intracerebroventricular (i.c.v.) administration of the M₁ antagonist pirenzepine in spatial learning tasks (Hagan et al., 1987; Hunter and Roberts, 1988; Messer et al., 1990; Sala et al., 1991). Andrews et al. (1994) recently studied the effects of pirenzepine, AF-DX 116 and UH-AH 37 after i.c.v. administration in a delayed matching to position (DMTP) task and found clear differences in the effects of these selective ligands. The M₁ receptor antagonist pirenzepine disrupted accuracy in a delay-, but not dose-related manner. After injection of pirenzepine into the hippocampus, Messer et al. (1990) also found a decrease in performance on a representational memory (non-matching-to-sample, T-maze) task. Less agreement exists about the consequences of antagonising the M₂ receptor. Andrews et al. (1994) found no disruptions after central injection of the muscarinic M₂ antagonist AF-DX 116. However, other studies reported a decrease in representational memory performance after hippocampal injection of AF-DX 116 (Messer et al., 1990). To date, only one study used the M₁ antagonist UH-AH 37: in a delayed matching to position task UH-AH 37 was found to disrupt performance severely even in the zero delay condition (Andrews et al., 1994).

An alternative method of inducing cholinergic hypo-funcion is to deplete the cholinergic neurons of acetylcholine. To this end, i.c.v. injections of hemicholinium-3 (HC3), a high-affinity choline uptake blocker (Russell and Macri, 1978), have been used to induce a reversible cholinergic depletion. Although at high dosages behavioural disruptions have been reported (Freeman et al., 1975; Jenden et al., 1977), low doses of HC3 can induce cognitive deficits without accompanying motor disturbances (Ridley et al., 1984, 1987; Hagan et al., 1989; Muir et al., 1992; Andrews et al., 1994).

Given the knowledge that a generalised cholinergic blockade results in many non-specific, and probably non-cognitive, disruptive effects on behaviour, it is important to determine the effects on performance of cholinergic depletion as well as of antagonising cholinergic receptors. Therefore, the present experiment investigated the contribution of the muscarinic cholinergic system to task performance, by general and selective antagonism as well as by cholinergic depletion. The compounds used were chosen because their effects in a DMTP task are known and provide a reference for effects on delay-dependent and delay-independent performance. An operant two-lever visual discrimination task was chosen, as a similar task has proven useful in elucidating the effects of psychoactive drugs and lesions on performance (Andrews and Sahgal, 1984; Andrews and Holtzman, 1988; Evenden et al., 1989). These experiments help to establish whether blockade of a particular muscarinic subtype can disrupt visual discrimination in a similar manner to that repeatedly observed after general muscarinic blockade (e.g. Andrews et al., 1992), cholinergic lesions (e.g. Dunnett et al., 1991) or cholinergic depletion (e.g. Muir et al., 1992).

MATERIALS AND METHODS

Subjects and surgery

Subjects were male Long-Evans rats (supplied by HAR-LAN, Zeist), weighing approximately 275 g at the beginning of the experiment. Rats were individually housed in standard Makrolon cages and maintained on a 12:12 h light:dark cycle, with lights on at 08.00 h. A restricted feeding schedule (down to 90% of free-feeding weight) was introduced three days before training, but access to tap water was free at all times. After training the animals to criterion (see below) animals were allowed to free-feed for three days before surgery. Each rat was implanted under anaesthesia (Nembutal, 60 mg/kg i.p.) with a stainless steel guide cannula (Organon B.V., The Netherlands), aimed at the right lateral ventricle: coordinates from Bregma, AP = −0.8 mm, Lat = −1.5 mm, HV = 3.0 mm (Paxinos and Watson, 1982). I.c.v. injections of dye histologically verified correct cannula positioning post mortem. Food deprivation was reinstated one week after surgery.

Apparatus

Eight operant chambers (L 27 x W 25 x H 24 cm), equipped with two retractable levers, a centrally placed food tray and pellet dispenser (delivering 45 mg pellets), red house light, and two cue light displays above the levers, were connected to Skinner Box Controllers and controlled by an Apple Macintosh SE 30. Boxes, controllers and software were developed by the Electronic and Computer Engineering Department and the Mechanical Engineering Department of the Psychological Laboratory, University of Nijmegen. The cue light consisted of 64 green LEDs (PD 1167, Siemens), together forming a 25.4 mm square light.

Visual discrimination learning

Rats were initially trained to press the levers on an FR1 schedule of food reinforcement using an autoshaping schedule. Subsequently, both levers were permanently available in the chamber; when a cue light was presented,
the rat was required to press the lever directly under the stimulus in order to obtain a food pellet and progress to the next trial. As soon as this task had been adequately mastered, training on the discrimination proper started. In the discrimination task, each trial began with the presentation of a light over one of the two levers, while at the same time both levers were inserted into the chamber. Upon pressing the correct lever, both levers were withdrawn, the cue light was switched off, the tray light was illuminated and a food pellet delivered. The next trial followed after a variable interval of 5 s mean duration (range 3.5–6.5 s). If the rat failed to respond within 5 s or responded on the incorrect lever, the cue light was extinguished, the levers were withdrawn and no food pellet was delivered. In these cases the next trial began after a variable time-out interval of mean 10 s duration (7.5–12.5 s). Each session consisted of 100 trials, in which the stimulus (left or right) occurred 50 times in a random order; the order of presentation of the stimuli was given in a different random pattern each day. Each animal received one session a day, five days a week. Over a period of four weeks the cue light duration was reduced in a step-wise manner from 5.0 s to 0.3 s to prevent ceiling effects in accuracy scores; if the rat responded correctly on at least 70% of all responses in one session, the stimulus duration was halved for the next session. The animals reached criterion when they completed three successive sessions using the final parameters with a minimum of 75% correct responses. Animals were then implanted with a cannula during a two-week period of surgery and recovery; thereafter, all animals were re-trained to the pre-operation criterion. Parameters used for the final training session were also used for the drug testing session.

**Drug testing**

Separate groups of rats were used to study each dose of each drug in one-session studies. Rats were randomised prior to testing and the overall accuracy of the different groups evaluated; for each experiment the baseline performance of each group was equivalent on the training day immediately before testing. At least one week of drug-free training was allowed before further injection of compounds. Scopolamine (placebo, 0.0125, 0.05, 0.2 and 0.8 mg/kg, injection volume 1 ml/kg; \( n = 8 \) /group) was injected s.c. (neck) 30 min prior to test. HC3 (placebo, 0.04, 0.02, 1.0 and 5.0 µg/ rat in 5 µl of merlysin vehicle; \( n = 8 \) /group) was infused into the ventricle over 30 s, 60 min prior to test, using a CMA micropump. The same method was used to inject rats i.c.v. with pirenzepine and the structural analogues AF-DX 116 or UH-AH 37 (doses 0.0, 3.2, 10 and 32 µg, in 5 µl of merlysin over 30 s; \( n = 10 \) /group) immediately before testing.

**Data analysis**

Accuracy as measured by percentage correct responding (calculated as the number of trials correct/number of trials on which a response occurred × 100), errors of omission (missed trials), and response latency were analysed using a one-factor independent groups analysis of variance; followed, when significance was achieved, by Scheffe F post hoc testing. Rats which responded on fewer than ten trials were excluded from the analysis in order to prevent an extreme bias in the calculation of the errors of commission.

**RESULTS**

**Pirenzepine**

A significant drug effect was found for the percentage correct responses \([F(3,36) = 4.56, \ p < 0.01]\), for latency to respond \([F(3,36) = 11.23, \ p < 0.01]\), and for missed trials \([F(3,36) = 6.60, \ p < 0.01]\). Post hoc testing revealed a decrease in percentage correct and an increase in response latency and missed trials only at the highest dose (Fig. 1, top panel).

**AF-DX 116**

Five rats injected with the highest dose (32 µg) were discarded because they responded on fewer than ten trials. A significant drug effect was found for all parameters: the percentage correct responses \([F(3,31) = 6.26, \ p < 0.01]\), latency to respond \([F(3,31) = 4.67, \ p < 0.01]\) and missed trials \([F(3,31) = 4.59, \ p < 0.01]\). Post hoc testing revealed that only the highest dose decreased accuracy, and increased response latencies and failures to respond (Fig. 1, middle panel).

**UH-AH 37**

Five rats, which received the highest dose (32 µg), and one rat injected with 10 µg were discarded for failing to respond on the minimally required number of trials. A significant drug effect was found for percentage correct responses \([F(3,30) = 6.49, \ p < 0.01]\), missed trials \([F(3,30) = 7.54, \ p < 0.01]\) and latency to respond \([F(3,30) = 8.80, \ p < 0.01]\). Post hoc testing revealed a decrease in accuracy and an increase in response latencies at 10 and 32 µg UH-AH 37. The highest dose also increased the number of missed trials (Fig. 1, bottom panel).

**Scopolamine**

Scopolamine dose-dependently decreased accuracy \([F(4,34) = 14.4, \ p < 0.01]\), and increased the number of missed trials \([F(4,34) = 34.5, \ p < 0.01]\) and response...
FIG. 1. Effects of the M₁ antagonist pirenzepine, the M₂ antagonist AF-DX 116, and the M₃ antagonist UH-AH 37 on visual discrimination performance in the rat. Doses (in µg per rat i.c.v.) are represented on the horizontal axis. Left panels: effects on accuracy as measured by percentage correct responses; centre panels: mean number of missed trials from a session of 100 trials; right panels: mean latency to respond in seconds. Values are means and standard deviations; * indicates a significant difference from vehicle (0).
latencies \( [F(4,34) = 28.9, \ p < 0.01] \) (Fig. 2, upper panel).
All but one rat from the 0.05 mg/kg scopolamine group responded adequately; this rat was removed from the analysis.

**Hemicholinium-3**

HC3 decreased correct responding \( [F(4,35) = 10.9, \ p < 0.01] \), and increased failures to respond \( [F(4,35) = 15.1, \ p < 0.01] \) and response latencies \( [F(4,35) = 25.2, \ p < 0.01] \). Post hoc testing showed that only the highest dose (5 \( \mu \)g) had effects on any aspect of performance (Fig. 2, lower panel). All rats responded adequately following HC3 administration.

**DISCUSSION**

The highest doses of all compounds enhanced errors of commission (decreased accuracy) and errors of omission (missed trials), as well as increased the latency to respond. In general, the effects of each compound were similar in pattern, although the magnitude differed considerably between compounds. No compound unambiguously disrupted accuracy without affecting latency to respond or trials completed. In the concentrations given, scopolamine and UH-AH 37 induced the most severe effects on all aspects of performance; pirenzepine and HC3 the weakest.

Several studies have indicated effects of pirenzepine on spatial learning tasks and although effective doses differ between studies, the general pattern of results is similar.
accuracy is affected before speed or frequency of responding (e.g. Andrews et al., 1994; Bymaster et al., 1993; Hagan et al., 1987). An earlier study had shown that pirenzepine was able to disrupt performance in the DMTP task at a 10-fold lower dose without corresponding effects on ability to respond (Andrews et al., 1994). It should be noted that the effects of pirenzepine in both this and the DMTP study are small and indicate some caution in ascribing a major role for M1 receptors in memory and attention without substantially more data than is available at the current time.

The effects of AF-DX 116 were surprising; apart from greater M2 activity in vitro, AF-DX 116 is rather less potent at other muscarinic receptors than the other analogues tested (Doods et al., 1994). To date, M2 antagonists have been reported to have no effect or a facilitating effect on mnemonic performance in rats (Andrews et al., 1994; Doods et al., 1993); there are no comparable studies on visual attention or discrimination. Messer et al. (1990) reported a decrement in performance following intrahippocampal injections of AF-DX 116 in their representational memory task. In the same experiment, pirenzepine was observed to disrupt memory at lower doses and the difference between the effective doses of the two compounds appeared to reflect relative potencies at the M1 receptor. A similar explanation is not possible here: the effect of AF-DX 116 was greater than that of pirenzepine, and contrasts strongly with results obtained using similar doses and strain of rats in a DMTP task (Andrews et al., 1994). It seems unlikely that AF-DX 116 could exert a large and selective effect on visual discrimination performance without causing some decrements in complex and demanding procedures such as DMTP. Thus, there is no immediately convincing explanation for these large discrepancies and further research is required to evaluate all the possibilities.

Of the three antagonists tested i.c.v., UH-AH 37 had the most potent effects on performance. The differences between pirenzepine and its close structural analogue UH-AH 37 are potentially the most interesting. Pirenzepine had only a small effect on accuracy at the highest dose, whereas UH-AH 37 had a greater effect on accuracy at both 10 and 32 μg doses and suppressed responding more than all other compounds apart from scopolamine.

The present data are consistent with a previous study involving DMTP, indicating a more potent disruption of performance by UH-AH 37 than by pirenzepine (Andrews et al., 1994). A number of studies have indicated that in functional assays pirenzepine and UH-AH 37 differ mostly in their effects on M1 mediated responses (Hagan et al., 1988; Kilbinger et al., 1991; Wess et al., 1991; Ten Berge et al., 1993; Doods et al., 1994). Nevertheless, other differences between pirenzepine and UH-AH 37 in both binding and functional essays, reflecting activity at other muscarinic subtypes, have also been reported (Eberlein et al., 1989; Wess et al., 1991; Bogner et al., 1992; Doods et al., 1994). Therefore, it is debatable whether the difference between the effects of UH-AH 37 and pirenzepine can be solely attributed to the differences in activity at M1 and M2 receptors. UH-AH 37 also shows high affinity for M4 receptors (Doods et al., 1994) and despite the fact that these receptors are much more prevalent in the brain than M1 (but not M4) receptors (Levey, 1993), little is known concerning their pharmacology. To resolve the question as to the relative importance of M3 receptors in performance in the visual discrimination task requires the use of a more selective M1 antagonist than pirenzepine and a more selective M3 antagonist than UH-AH 37.

The effects of all compounds apart from scopolamine can be assumed to be strictly centrally mediated. Despite the possible involvement of peripheral cholinergic receptor antagonism in the action of scopolamine, the similarities in effects to the other compounds indicate a strong influence of central muscarinic receptors on performance. Dissociating the peripheral and central effects of scopolamine has proven to difficult: qualitatively similar effects of methyl scopolamine and scopolamine on operant performance are often reported: e.g. Andrews et al., 1992; Dunnett et al., 1989; Hudzik and Wenger, 1993 (see Moore et al. (1992) for a discussion). However, the greater suppression of overall responding in the scopolamine-treated animals may be due to the additional antagonism of peripheral cholinergic receptors. Accordingly, it is of interest that scopolamine significantly affected frequency to respond at a lower dose than accuracy, whereas UH-AH 37 had a significant effect on accuracy at a lower dose than on trials completed.

Although HC3 has been available for more than 30 years, the number of studies involving complex behaviours remains small. In studies to date the active dose range appears to be 1–5 μg per rat i.c.v.; doses higher than this do not have any greater effects on acetylcholine levels in the brain (Freeman et al., 1979). Within this range, effects are seen on acquisition of a lever press (Andrews, unpublished observations), on spatial memory as measured in the swim maze, and in DMTP tasks (Hagan et al., 1989; Andrews et al., 1994), and on aspects of visual attention as measured here and in a serial reaction time task (Muir et al., 1992). Earlier studies using monkeys have suggested a separation of mnemonic and perceptual or motor effects (Ridley et al., 1984, 1987). The effects here and in the earlier DMTP study (Andrews et al., 1994) may indicate that, although the effects on attentional and perceptual factors are close, they are perhaps separable: 1 and 5 μg HC3 had similar effects on DMTP, but only 5 μg had any effect on the current task.
Interestingly, a recent study has indicated that attentional tasks involving visual discrimination, such as the five-choice serial reaction time task, may be less sensitive to HC3 than DMTP (Kirkby et al., 1994). However, other factors such as minor strain differences in the response to HC3 may account for some of the small differences between the studies mentioned above (see also Andrews et al., 1995). The advantages of HC3 over systemically administered antagonists such as scopolamine remain: central activity only; and as the effect is one of cholinergic deficiency and not receptor blockade, receptors can be stimulated directly without first competing with a high affinity antagonist.

To summarise, in agreement with previous studies, these data indicate an important role for the cholinergic system in attentional processes (Dunnett et al., 1991; Rupniak et al., 1991; Muir et al., 1992). It is noteworthy that no compound had an effect on accuracy independent of changes in other aspects of performance. With the exception of AF-DX 116, the relative disruptive effects of the compounds tested were broadly in line with the effects observed in an earlier experiment using a DMTP task (Andrews et al., 1994). The effects of pirenzepine were relatively small and occurred at doses higher than required to disrupt DMTP performance, suggesting some functional selectivity in its cognitive effects. UH-AH 37 induced more potent disruptive effects on visual discrimination performance than pirenzepine, a result consistent with a previous study involving DMTP (Andrews et al., 1994).

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