Adjunctive treatment with mianserin enhances effects of raclopride on cortical dopamine output and, in parallel, its antipsychotic-like effect

Charlotte Wiker
Love Linnér
Marie-Louise Wadenberg
Torgny H Svensson

Section for Neuropsychopharmacology, Department of Physiology and Pharmacology, Karolinska Institutet, Stockholm, Sweden

Abstract: Clinical studies indicate that adjunctive treatment with the antidepressant drug mianserin, a 5-hydroxytryptamine (5-HT)2A/C receptor antagonist and an α2- and α1-adrenoceptor antagonist, may enhance the effect of conventional antipsychotic drugs in schizophrenia, in particular on negative symptoms such as withdrawal retardation, akathisia, and some aspects of cognitive impairment. Here, we have examined the effect of mianserin in combination with the selective dopamine (DA) D2/D3 receptor antagonist raclopride on conditioned avoidance response (CAR), a preclinical test of antipsychotic efficacy with high predictive validity; catalepsy, a preclinical test of extrapyramidal side effect liability; and DA output in the medial prefrontal cortex (mPFC) and the nucleus accumbens (NAC), respectively. Mianserin (5 mg/kg intraperitoneal) significantly enhanced the suppressant effect of a low dose of raclopride (0.1 mg/kg subcutaneous) on CAR without any increase in catalepsy. Administration of raclopride to rats pretreated with mianserin resulted in a large enhancement of DA output in the mPFC and, at the same time, a small but significant reduction in the raclopride-induced DA output in the NAC. These experimental results indicate that adjunctive treatment with mianserin to a typical D2 antagonist generates an atypical antipsychotic profile.

Keywords: adjunctive mianserin, typical antipsychotic drugs, dopamine, prefrontal cortex, atypicality

Introduction

The clinical effect of conventional (typical) antipsychotic drugs (APDs) has been largely ascribed to their dopamine (DA) D2 receptor blocking action (Carlsson 1988). However, although these drugs, eg, haloperidol, are usually effective against positive symptoms, they show less efficacy against negative and cognitive symptoms (Carpenter 1996; Volavka et al 1996; Leucht et al 1999). Clinically effective doses of typical APDs generate about 70%–85% D2 receptor occupancy in brain, but a D2 receptor occupancy above 80% has been associated with a relatively high risk of extrapyramidal side effects (EPS) (Farde et al 1992). In contrast, some, but not all, of the so-called second generation (atypical) APDs may be clinically effective at lower D2 occupancy in brain, with clozapine around 45% (Nordstrom 1995), and are associated with a significantly reduced, and in the case of clozapine essentially abolished, EPS liability (Safferman et al 1991). Yet, clozapine shows superior efficacy in treatment-resistant schizophrenia, and moreover, several atypical APDs have been claimed to show improved efficacy against negative and cognitive symptoms (Meltzer 1995; Davis et al 2003). Clozapine possesses a complex pharmacology, acting as an antagonist not only at several DA receptors (ie, D2, D3, and D4 receptors) but also at,
addition of ritanserin, a 5-HT 2A/C antagonist, was found to contribute to the atypical APD profile (Svensson 2003). By contrast, the alternative D_2_ receptor “fast-off” hypothesis of Kapur and Seeman (2001) applies only to clozapine and quetiapine and is inconsistent with the “slow” off-rate of most atypical APDs (Meltzer et al 2003).

Previous clinical augmentation trials involving conventional APDs may help elucidate this issue. Thus, the addition of ritanserin, a 5-HT 2A/C antagonist, was found to enhance the antipsychotic effect of typical APDs, in particular negative symptoms (Gelders et al 1986, Gelders 1989). Other clinical studies have shown that adjunctive treatment with idazoxan, a selective α_2-adrenoceptor antagonist, can enhance the efficacy of conventional APDs in treatment-resistant schizophrenia (Litman 1993; Litman et al 1996), including both positive and negative symptoms and total Brief Psychiatric Rating Scale score. Moreover, mianserin, a clinically effective antidepressant drug, which acts as an antagonist at 5-HT 2A/C receptors, α_2- and α_1- adrenoceptors, and muscarinic and histaminergic (H_1) receptors (Pinder 1991), has also been found to enhance the clinical effect of typical APDs, particularly concerning negative symptoms such as withdrawal retardation, akathisia, and some aspects of cognitive dysfunction (Itil et al 1974; Mizuki et al 1990, 1992; Poyurovsky 1999; Grinshpoon et al 2000; Shiloh et al 2002; Poyurovsky et al 2003). This generates a clozapine-like clinical effect (Lindstrom 2000) as well as a combined receptor binding profile relatively similar to that of clozapine (cf Pinder 1991; Bymaster et al 1996).

Several experimental studies may contribute to clarify the neurobiological mechanisms of particular importance in this respect. Thus, clozapine as well as other atypicals have, in contrast to typical APDs, been found to preferentially enhance DA outflow in the prefrontal cortex (Imperato and Angelucci 1989; Moghaddam and Bunney 1990; Nomikos et al 1994; Kuroki et al 1999; Westerink et al 2001). As previous experimental results using both a neurodevelopmental model and the phencyclidine model of schizophrenia inter alia have indicated an impaired prefrontal DA projection in this disease (see Weinberger and Lipska 1995; Svensson et al 1995; Egan and Weinberger 1997; Egan et al 2001), the selective augmentation of prefrontal DA output by clozapine and other atypicals may allow for restoration of a pathophysiological deficit in schizophrenia, tentatively related to cognitive and/or negative symptoms (see Svensson 2003). Both the addition of ritanserin and idazoxan to the selective D_2/3 antagonist, raclopride, produced an enhanced suppression of the conditioned avoidance response (CAR) in rodents, ie, an antipsychotic-like effect, without any concomitant increase in catalepsy (Wadenberg et al 1996; Hertel et al 1999), as well as a preferential enhancement of prefrontal DA output not seen with the D_2_ antagonist alone (Andersson et al 1995; Svensson et al 1995; Hertel et al 1999). Since subsequent behavioral (Wadenberg et al 1998) and biochemical (Liégeois et al 2002) studies using the selective 5-HT 2A receptor antagonist M100907 produced similar results, whereas 5-HT 2C receptor blockage activates both cortical and subcortical DA projections (Di Matteo et al 2000; Gobert et al 2000), the preferential enhancement of prefrontal DA efflux by atypicals may be associated with blockade of 5-HT_2A_ rather than 5-HT_2C_ receptors. Thus, clinical as well as preclinical, correlative behavioral, and biochemical studies indicate that an enhanced effect of typical D_2_ antagonists in schizophrenia can be obtained by adding 5-HT_2A_ and/or α_2- receptor blockage and that this effect is indeed associated with enhanced prefrontal DA output.

Here, we have experimentally investigated the effect of combining raclopride and mianserin, which when given alone increases DA output both in the medial prefrontal cortex (mPFC; Tanda et al 1996) and in the nucleus accumbens (NAC; Di Matteo et al 2000) on DA efflux in both terminal areas of the mesocorticolimbic system, as well as by correlative behavioral studies using the CAR paradigm and additional assessment of catalepsy scores.

### Materials and methods

#### Animals

Adult male BK1:WR (Wistar) rats weighing 300–390 g (CAR), 270–310 g (catalepsy), or 230–390 g (microdialysis) were used in all experiments. Animals arrived at least five days prior to experimental use and were housed (4 per cage [Makrolon IV]) in the animal facility under standard laboratory conditions with a 12-h light/dark cycle. Rats designated for microdialysis had lights on at 6.00 am, whereas animals designated for behavioral experiments were subjected to a reversed light/dark cycle, ie, lights off at 6.00 am. All experiments were performed between 8.00 am and 6.00 pm. Food and water were available ad lib.
experiments were approved by, and conducted in accordance with, the local Animal Ethics Committee (Stockholms Norra och Södra Försöksdjursetiska Kommitéer) (permit numbers N216/00, N11/00).

Conditioned avoidance response
A shuttle-box (530 mm × 250 mm × 225 mm) divided into two compartments by a partition was used. The rats were free to move from one compartment to the other via an opening (75 mm × 75 mm) in the partition. Upon presentation of the conditioned stimulus (CS), 80 dB white noise (White Noise Generator, Lafayette 1501, Lafayette, IN, USA), the rat had 10 s to avoid the unconditioned stimulus (USC), an intermittent electric shock in the grid floor of approximately 0.2 mA (intershock interval 2.5 s, shock duration 0.5 s), by moving into the opposite compartment. The following behavioral variables were recorded: (1) avoidance (response to CS within 10 s); (2) escape (response to CS + USC); (3) escape failure (if the rat was unable to respond to the shock within 50 s the trial was terminated). The animals were trained for 5 consecutive days, and were adapted to the shuttlebox 5 min before the training session started. Each training session consisted of 20 trials randomly distributed over 15 min. All subsequent pre-tests and experimental test sessions consisted of 10 trials randomly distributed over 7.5 min. Experimental manipulations were always preceded by a pre-test. The same animals were tested repeatedly according to a change-over design (Li 1964) serving as their own controls. Only rats that showed at least 90% avoidance on the last day of training were included in the experiments.

Catalepsy
Animals were placed on an inclined (60°) grid and, excluding the first 30 s, the time the rat remained in the same position was measured for a maximum of 2.5 min. The catalepsy was scored from 0–5 according to the time (square root transformation) the animal remained immobile (min): 0 = 0–0.08, 1 = 0.09–0.35, 2 = 0.36–0.80, 3 = 0.81–1.42, 4 = 1.43–2.24, 5 = ≥ 2.25 min; ie, if the rat remained immobile for ≥2.25 min it was scored 5, etc (Ahlenius and Hillegaart 1986).

Microdialysis
The probe implantation and dialysis procedure, as well as the biochemical analyses, were similar to those previously described (Hertel et al 1996). Anesthetized male BK1:WR (Wistar) rats (B&K Universal, Sollentuna, Sweden; sodium pentobarbital, 60 mg/kg, intraperitoneal [IP]) were implanted with dialysis probes in the mPFC or NAC (AP: +2.6, +1.4; ML: −0.6, −1.4; DV: −5.2, −8.2), respectively, relative to bregma and dural surface (Paxinos and Watson 1998). Dialysis occurred through a semipermeable membrane (AN69 Hospal) with an active surface length of 4 and 2.25 mm for mPFC and NAC, respectively. Dialysis experiments were conducted approximately 48 h after surgery in freely moving rats. The dialysis probe was perfused with a physiological perfusion solution (147 mmol/L sodium chloride, 3.0 mmol/L potassium chloride, 1.3 mmol/L calcium chloride, 1.0 mmol/L magnesium chloride, and 1.0 mmol/L sodium phosphate, pH 7.4) at a rate of 2.5 µL/min set by a microinfusion pump (Harvard Apparatus, Holliston, MA). Online quantification of DA in the dialysate was accomplished by high pressure liquid chromatography coupled to electrochemical detection. The detection limit for DA was approximately 0.2 fmol/min. The placement of the probe was later verified in slices stained with neutral red.

Drugs
Mianserin (Sigma-Aldrich, Stockholm, Sweden) and raclopride tartrate (Astra Zeneca, Södertälje, Sweden) were dissolved in saline. Mianserin was subsequently adjusted to pH ~6 (at a higher pH the drug precipitated) and raclopride was adjusted to physiological pH (7.0–7.3). Mianserin or saline was administered IP (1.0 mL/kg) and raclopride or saline was administered subcutaneously (SC) (1.0 mL/kg). The dose of mianserin was chosen based on previous studies with mianserin in microdialysis (Tanda et al 1996; Di Matteo et al 2000). The doses of raclopride used (0.05–0.1 mg/kg SC) were chosen based on earlier experiments, where no or only moderate effects on CAR (Hertel et al 1999) and low, subtherapeutic (≤65%) D₂ receptor occupancy (Wadenberg, Kapur, et al 2000) have been observed. During microdialysis, administration of drugs or vehicle was performed after stable outflow (<10% variation) of DA was obtained.

Statistics
Behavioral experiments. Statistical evaluation was performed by means of the Friedman two-way analysis of variance (ANOVA), followed by the Wilcoxon matched-pairs signed-ranks test (CAR) or the Kruskal-Wallis
one-way ANOVA, followed by the Mann-Whitney U-test (catalepsy) (Siegel and Castellan 1988).

**Microdialysis.** Data were calculated as percent changes of basal DA output over time. Baseline (= 100%) was defined as the average of the last two (mPFC) or four (NAC) preinjection values. Data were statistically evaluated using two-way (treatment × time) ANOVA for repeated measures followed by the Newman-Keuls test for multiple comparisons.

**Results**

**Conditioned avoidance response**

Raclopride (0.05 or 0.1 mg/kg SC) produced a statistically significant suppression of CAR 20 minutes after administration, but not at later observation times (90 and 240 minutes). Pretreatment with mianserin (5 mg/kg IP 30 min before) produced a significant enhancement of the suppression of CAR induced by raclopride (0.1 mg/kg) compared with animals treated with raclopride alone. Mianserin by itself had no effect on CAR. No significant effects were observed at later observation times, and no escape failures were recorded under any treatment condition; ie, a decrease in avoidance responses was always accompanied by a corresponding increase in escape responses (Figure 1).

**Catalepsy**

Raclopride (0.1 mg/kg SC) alone, or in combination with mianserin (5 mg/kg IP), did not produce any significant catalepsy at any of the observation times. The administration of a higher dose of raclopride (1.0 mg/kg SC) resulted in a cataleptic response that peaked at 60 minutes after administration. At this time point and dose of raclopride, pretreatment with mianserin (5 mg/kg IP – 30 min) caused a consistent, albeit not significant, decrease in catalepsy scores compared with animals treated with raclopride alone (Figure 2).

**Microdialysis**

The mean baseline concentration (fmol/min ± SEM) of DA in the mPFC and the NAC was 0.44 ± 0.09 (n = 24) and 4.41 ± 0.61 (n = 24) fmol/min, respectively, (data not corrected for in vitro dialysis probe recovery). Administration of mianserin (5 mg/kg IP) alone, or in combination with raclopride (0.1 mg/kg SC), resulted in a significant increase in DA output in the mPFC, which was significantly larger in the combination-treatment group (Figure 3a). Administration of raclopride (0.1 mg/kg SC) alone had no significant effect on mPFC DA output but resulted in a significant increase in DA output in the NAC. Pretreatment with mianserin (5 mg/kg IP) caused a small, but significant reduction of the effect of raclopride (0.1 mg/kg SC) on NAC DA output. Administration of saline or mianserin (5 mg/kg IP) alone had no effect on NAC DA output (Figure 3b).

![Figure 1](image-url)

*A Figure 1 Effects of saline or mianserin (5 mg/kg IP) pre-treatment (30 min) on saline– or raclopride (0.05 and 0.1 mg/kg SC)–induced effects in the conditioned avoidance response test (a) 20 min, (b) 90 min, and (c) 240 min after saline or raclopride administration. Each bar represents the median avoidance % (± semi-interquartile range; n = 9 in all groups). Statistical evaluation was performed by means of the Friedman two-way analysis of variance (ANOVA), followed by the Wilcoxon matched-pairs signed ranks test. ANOVA Chi² (df = 5) = 35.5, *p < 0.001, **p < 0.05, ***p < 0.01 compared with respective control (saline/saline or mianserin/saline). †p < 0.05 for comparisons between saline/raclopride and mianserin/raclopride treatment group.*
Discussion

The major finding of the present study is that the addition of mianserin to the selective D<sub>2/3</sub> receptor antagonist raclopride causes a significant enhancement of the raclopride-induced suppression of CAR without increasing catalepsy scores, providing experimental evidence for an enhanced antipsychotic efficacy of the drug combination compared with raclopride alone, yet without any increased EPS liability. The relatively low doses of raclopride used, that did not cause an effective antipsychotic-like effect when given alone has been observed to generate a central D<sub>2</sub> occupancy of about 50%–65% (Wadenberg, Kapur, et al 2000). Notably, about 80% D<sub>2</sub> occupancy is necessary to produce a suppression of the CAR of such magnitude that it would predict sufficient antipsychotic activity, although at this level of D<sub>2</sub> occupancy catalepsy also begins to occur (Wadenberg, Kapur, et al 2000). Thus, the adjunctive treatment with mianserin allowed for a sufficient
significant suppression of the raclopride-induced increase
Green 1996; Harvey 1998). A potential cognitive enhancing effect of adding mianserin also in the parietal and occipital cortex (Valentini et al 2004).

In contrast to ritanserin or idazoxan, the addition of mianserin to raclopride also caused a slight, but still significant suppression of the raclopride-induced increase in DA outflow in the subcortical brain region studied; ie, NAC, an effect that in itself is secondary to the D2 receptor blockade (see Carlsson 1988). Previous studies show that both the α1-adrenoceptor antagonist prazosin and the selective 5-HT2A receptor antagonist M100907 inhibit the D2 receptor antagonist-induced increase in accumbal DA release (Andersson et al 1995; Liégeois et al 2002) and, moreover, that prazosin (Wadenberg, Hertel, et al 2000) in similarity with M100907 (cf Introduction) also enhances the CAR suppression induced by D2 antagonists. Thus, both the 5-HT2A receptor antagonistic action and the α1-adrenoceptor antagonistic effect of mianserin may, in principle, contribute to the slight inhibition of the raclopride-induced accumbal DA release observed in the present study and, by inference, to the suppression of the CAR. Such a mechanism may also have bearing on the overall antipsychotic efficacy of combining mianserin and typical D2 receptor antagonists in schizophrenia, albeit not necessarily related to relief of negative symptoms. This notion gains further support from both preclinical and clinical observations. Thus, both prazosin (Mathé et al 1996) and M100907 (Schmidt and Fadayel 1995) act to suppress evoked, but not basal DA release in the NAC of experimental animals, and clinical studies indicate that schizophrenic patients display an enhanced evoked, but not basal subcortical DA release (Laruelle et al 1996; Breier et al 1997).

In conclusion, several pharmacological properties of mianserin, namely blockade of 5-HT2A receptors, α2- as well as α1- adrenoceptors may contribute to its clinical effect when added to typical D2 receptor antagonists in the treatment of schizophrenia, which appears to include not only an effect on negative symptoms, but also a reduction in dysphoria, anxiety, akathisia, and potentially cognitive dysfunction. Yet, the sedative effect of mianserin may, as in the case of clozapine, largely be correlated with its histamine H1 receptor antagonistic action, which may contribute to weight gain. Several of the new, second generation APDs are relatively expensive in comparison with conventional APDs, as pointed out by Lindstrom (2000). Combining typical APDs with mianserin might thus represent a more affordable treatment strategy in some areas of the world. In addition, by using a combination of mianserin and typical D2 antagonists one serious side-effect of clozapine, namely agranulocytosis, can be avoided. Tentatively, also mirtazapine, which has a receptor affinity profile similar to mianserin, may be used for the same purpose (Berk et al 2001).
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**References**

Ahlenius S, Hillegaart V. 1986. Involvement of extrapyramidal motor mechanisms in the suppression of locomotor activity by antipsychotic drugs: a comparison between the effects produced by pre- and post-synaptic inhibition of dopaminergic neurotransmission. *Pharmacol Biochem Behav*, 24:1409–15.

Andersson JL, Nomikos GG, Marcus M, et al. 1995. Risperidone potentiates the stimulatory effects of raclopride on neuronal activity and dopamine release selectivity in the mesolimbic dopaminergic system. *Naunyn Schmiedebergs Arch Pharmacol*, 352:374–85.

Berk M, Ichim C, Brook S. 2001. Efficacy of mirtazapine add on therapy to haloperidol in the treatment of the negative symptoms of schizophrenia: a double-blind randomized placebo-controlled study. *Int Clin Psychopharmacol*, 16:87–92.

Breier A, Su TP, Saunders R, et al. 1997. Schizophrenia is associated with elevated amphetamine-induced synaptic dopamine concentrations: evidence from a novel positron emission tomography method. *Proc Natl Acad Sci U S A*, 94:2569–74.

Bystmister FP, Calligaro DO, Falcone JF, et al. 1996. Radioreceptor binding profile of the atypical antipsychotic olanzapine. *Neuropsychopharmacology*, 14:87–96.

Carlsson A. 1988. The current status of the dopamine hypothesis of schizophrenia. *Neuropsychopharmacology*, 1:179–86.

Carpenter Jr WT. 1996. The treatment of negative symptoms: pharmacological and methodological issues. *Br J Psychiatry Suppl*, 29:17–22.

Davis JM, Chen N, Glick ID. 2003. A meta-analysis of the efficacy of second-generation antipsychotics. *Arch Gen Psychiatry*, 60:553–64.

Di Matteo V, Di Mascio M, Di Giovanni G, et al. 2000. Acute administration of amitriptyline and mianserin increases dopamine release in the rat nucleus accumbens: possible involvement of serotonin2C receptors. *Psychopharmacology (Berl)*, 150:45–51.

Egan MF, Goldberg TE, Kolachana BS, et al. 2001. Effect of COMT Val108/158 Met genotype on frontal lobe function and risk for schizophrenia. *Proc Natl Acad Sci U S A*, 98:6917–22.

Egan MF, Weinberger DR. 1997. Neurobiology of schizophrenia. *Curr Opin Neurobiol*, 7:701–7.

Farde L, Nordstrom A-L, Wiesel F-A, et al. 1992. Positron emission tomographic analysis of central D1 and D2 dopamine receptor occupancy in patients treated with classical neuroleptics and clozapine. Relation to extrapyramidal side effects. *Arch Gen Psychiatry*, 49:538–44.

Gelders YG. 1989. Thymosthenic agents, a novel approach in the treatment of schizophrenia. *Br J Psychiatry Suppl*, 5:33–6.

Gelders Y, Vandeven Bussche G, Reyntjens A, et al. 1986. Serotonin-S2 receptor blockers in the treatment of chronic schizophrenia. *Clin Neuropharmacol*, 9:325–7.

Gobert A, Rivet JM, Lejeune F, et al. 2000. Serotonin(2C) receptors tonically suppress the activity of mesocortical dopaminergic and adrenergic, but not serotonergic, pathways: a combined dialysis and electrophysiological analysis in the rat. *Synapse*, 36:205–21.

Green MF. 1996. What are the functional consequences of neurocognitive deficits in schizophrenia? *Am J Psychiatry*, 153:321–30.

Grinshpoon A, Valevski A, Moskowit M, et al. 2000. Beneficial effect of the addition of the 5-HT2A/2C and alpha2 antagonist mianserin to ocura N, Imai T, et al. 1990. Effects of mianserin on negative symptoms in schizophrenia. *Int Clin Psychopharmacol*, 5:83–95.

Harvey PD. 1998. Symptoms, cognitive functioning, and adaptive skills in geriatric patients with lifelong schizophrenia: a comparison across treatment sites. *Am J Psychiatry*, 155:1080–6.

Hertel P, Fagerquist MV, Svensson TH. 1999. Enhanced cortical dopamine output and antipsychotic-like effects of raclopride by alpha2 adrenoceptor blockade. *Science*, 286:105–7.

Hertel P, Nomikos GG, Ihrlo M, et al. 1996. Risperidone: regional effects in vivo on release and metabolism of dopamine and serotonin in the rat brain. *Psychopharmacology (Berl)*, 124:74–86.

Imperato A, Angelucci L. 1989. The effects of clozapine and flupenthixol on the in vivo release and metabolism of dopamine in the striatum and in the prefrontal cortex of freely moving rats. *Psychopharmacol Bull*, 25:383–9.

Itil TM, Polvan N, Dinmchen K, et al. 1974. New drug developments in the Netherlands. *Dis Nerv Syst*, 35:10–17.

Kapur S, Seeman P. 2001. Does fast dissociation from the dopamine d2 receptor explain the action of atypical antipsychotics?: a new hypothesis. *Am J Psychiatry*, 158:360–9.

Kuroki T, Meltzer HY, Ichikawa J. 1999. Effects of antipsychotic drugs on extracellular dopamine levels in rat medial prefrontal cortex and nucleus accumbens. *J Pharmacol Exp Ther*, 288:774–81.

Laruelle M, Abi-Dargham A, van Dyck CH, et al. 1996. Single photon emission computerized tomography imaging of amphetamine-induced dopamine release in drug-free schizophrenic subjects. *Proc Natl Acad Sci U S A*, 93:9235–40.

Leucht S, Pitschel-Walz G, Abraham D, et al. 1999. Efficacy and extrapyramidal side-effects of the new antipsychotics olanzapine, quetiapine, risperidone, and sertindole compared to conventional antipsychotics and placebo. A meta-analysis of randomized controlled trials. *Schizophr Res*, 35:51–68.

Li CC. 1964. Introduction to experimental statistics. New York: McGraw-Hill. p 207–26.

Li Z, Ichikawa J, Dai J, et al. 2004. Aripiprazole, a novel antipsychotic drug, preferentially increases dopamine release in the prefrontal cortex and hippocampus in rat brain. *Eur J Pharmacol*, 493:75–83.

Lieseos JS, Ichikawa J, Meltzer HY. 2002. 5-HT(2A) receptor agonist potentiates haloperidol-induced dopamine release in rat medial prefrontal cortex and nucleus accumbens. *Psychopharmacology (Berl)*, 150:45–51.

Lindstrom LH. 2000. Schizophrenia, the dopamine hypothesis and alpha2-adrenoceptor antagonists. *Trends Pharmacol Sci*, 21:198–9.

Littman RE. 1993. Idazoxan, an alpha 2 antagonist, augments fluphenazine in schizophrenic patients: a pilot study. *J Clin Psychopharmacol*, 13:264–7.

Littman RE, Su T-P, Potter WZ, et al. 1996. Idazoxan and response to typical antipsychotics and placebo. A meta-analysis of randomized controlled trials. *Schizophr Res*, 35:51–68.

Meltzer HY. 1995. Role of serotonin in the action of atypical antipsychotic drugs. *Clin Neurosci*, 3:64–75.

Meltzer HY, Li Z, Kaneda Y, et al. 2000. Serotonin receptors: their key role in drugs to treat schizophrenia. *Prog Neuropsychopharmacol Biol Psychiatry*, 24:1159–72.

Mizuki Y, Kajimura N, Imai T, et al. 1990. Effects of mianserin on negative symptoms in schizophrenia. *Int Clin Psychopharmacol*, 5:83–95.

Mizuki Y, Kajimura N, Kai S, et al. 1992. Effects of mianserin in chronic schizophrenia. *Prog Neuropsychopharmacol Biol Psychiatry*, 16:517–28.

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Moghaddam B, Bunney BS. 1990. Acute effects of typical and atypical antipsychotic drugs on the release of dopamine from prefrontal cortex, nucleus accumbens, and striatum of the rat: an in vivo microdialysis study. *J Neurochem*, 54:1755–60.

Nomikos GG, Iurlo M, Andersson JL, et al. 1994. Systemic administration of amperozide, a new atypical antipsychotic drug, preferentially increases dopamine release in the rat medial prefrontal cortex. *Psychopharmacology (Berl)*, 115:147–56.

Nordstrom AL. 1995. D1, D2, and 5-HT2 receptor occupancy in relation to clozapine serum concentration: a PET study of schizophrenic patients. *Am J Psychiatry*, 152:1444–9.

Paxinos G, Watson C. 1998. The rat brain in stereotaxic coordinates. 4th ed. San Diego: Academic Pr.

Pinder RM. 1991. Mianserin: pharmacological and clinical correlates. *Nordisk Psychiatrik Tidsskrift*, 45:13–26.

Poyurovsky M. 1999. Treatment of neuroleptic-induced akathisia with the 5-HT2 antagonist mianserin. Double-blind, placebo-controlled study. *Br J Psychiatry*, 174:238–42.

Poyurovsky M, Koren D, Gonopolsky I, et al. 2003. Effect of the 5-HT2 antagonist mianserin on cognitive dysfunction in chronic schizophrenia patients: an add-on, double-blind placebo-controlled study. *Eur Neuropsychopharmacol*, 13:123–8.

Safferman A, Lieberman JA, Kane JM, et al. 1991. Update on the clinical efficacy and side effects of clozapine. *Schizophr Bull*, 17:247–61.

Schmidt CJ, Fadayel G. 1995. The selective 5-HT2A receptor antagonist, MDL 100,907, increases dopamine efflux in the prefrontal cortex of the rat. *Eur J Pharmacol*, 273:273–9.

Schotte A, Janssen PFM, Gommeren W, et al. 1996. Risperidone compared with new and reference antipsychotic drugs: in vitro and in vivo receptor binding. *Psychopharmacology (Berl)*, 123:57–73.

Shiloh R, Zemishlany Z, Aizenberg D, et al. 2002. Mianserin or placebo as adjuncts to typical antipsychotics in resistant schizophrenia. *Int Clin Psychopharmacol*, 17:59–64.

Siegel S, Castellan NJ Jr. 1988. Nonparametric statistics for the behavioral sciences. New York: McGraw-Hill.

Svensson TH. 2003. Preclinical effects of conventional and atypical antipsychotic drugs: defining the mechanisms of action. *Clin Neurosci Res*, 3:34–46.

Svensson TH, Mathe JM, Andersson JL, et al. 1995. Mode of action of atypical neuroleptics in relation to the phencyclidine model of schizophrenia: role of 5-HT2 receptor and alpha 1-adrenoceptor antagonism [corrected]. *J Clin Psychopharmacol*, 15:115–18.

Tanda G, Bassareo V, Di Chiara G. 1996. Mianserin markedly and selectively increases extracellular dopamine in the prefrontal cortex as compared to the nucleus accumbens of the rat. *Psychopharmacology (Berl)*, 123:127–30.

Valentini V, Frau R, Di Chiara G. 2004. Noradrenaline transporter blockers raise extracellular dopamine in medial prefrontal but not parietal and occipital cortex: differences with mianserin and clozapine. *J Neurochem*, 88:917–27.

Volavka J, Cooper TB, Czobor P, et al. 1996. Effect of varying haloperidol plasma levels on negative symptoms in schizophrenia and schizoaffective disorder. *Psychopharmacol Bull*, 32:75–9.

Wadenberg ML, Hertel P, Fernholm K, et al. 2000. Enhancement of antipsychotic-like effects by combined treatment with the alpha1-adrenoceptor antagonist prazosin and the dopamine D2 receptor antagonist raclopride in rats. *J Neural Transm*, 107:1229–38.

Wadenberg ML, Hicks PB, Richter JT, et al. 1998. Enhancement of antipsychoticlike properties of raclopride in rats using the selective serotonin2A receptor antagonist MDL 100,907. *Biol Psychiatry*, 44:508–15.

Wadenberg ML, Kapur S, Soliman A, et al. 2000. Dopamine D2 receptor occupancy predicts catalepsy and the suppression of conditioned avoidance response behavior in rats. *Psychopharmacology (Berl)*, 150:422–9.

Wadenberg ML, Salmi P, Jimenez P, et al. 1996. Enhancement of antipsychotic-like properties of the dopamine D2 receptor antagonist, raclopride, by the additional treatment with the 5-HT2 receptor blocking agent, ritanserin, in the rat. *Eur Neuropsychopharmacol*, 6:305–10.

Weinberger DR, Lipska BK. 1995. Cortical maldevelopment, antipsychotic drugs, and schizophrenia: a search for common ground. *Schizophr Res*, 16:87–110.

Westerink BH, Kawahara Y, De Boer P, et al. 2001. Antipsychotic drugs classified by their effects on the release of dopamine and noradrenaline in the prefrontal cortex and striatum. *Eur J Pharmacol*, 412:127–38.