Effects of quetiapine and sertindole on subchronic ketamine-induced deficits in attentional set-shifting in rats

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Abstract

Rationale Prefrontal cortical dysfunctions, including an impaired ability to shift perceptual attentional set, are core features of schizophrenia. Nevertheless, the effectiveness of second-generation antipsychotic drugs in treating specific prefrontal dysfunctions remains equivocal.

Objectives To model schizophrenia-like cognitive inflexibility in rats, we evaluated the effects of repeated administration of ketamine, the noncompetitive antagonist of the N-methyl-D-aspartate receptor, after a washout period of 14 days in the attentional set-shifting task (ASST). Next, we investigated whether the atypical antipsychotics quetiapine and sertindole would alleviate the ketamine-induced set-shifting impairment.

Methods Ketamine (30 mg/kg) was administered intraperitoneally to rats once daily for 5 or 10 consecutive days to assess its efficacy in producing cognitive impairment. The ASST was performed 14 days following the final drug administration. Quetiapine (0.63, 1.25 or 2.5 mg/kg) or sertindole (2.5 mg/kg) was administered per os 120 min before testing.

Results The results of the present study demonstrate that ketamine treatment for 10 but not 5 days significantly and specifically impaired rats’ performance in the extradimensional shift (EDs) stage of the ASST. This cognitive inflexibility was reversed by acute administration of sertindole or quetiapine. Quetiapine also promoted set-shifting in cognitively unimpaired control animals.

Conclusion The data presented here show that subchronic administration of ketamine induces cognitive inflexibility after a washout period. This cognitive deficit likely reflects clinically relevant aspects of cognitive dysfunction encountered in schizophrenic patients. The beneficial effects of quetiapine on set-shifting may have therapeutic implications for the treatment of schizophrenia and other disorders associated with frontal-dependent cognitive impairments.

Keywords Cognitive flexibility · Attentional set-shifting · Ketamine · Quetiapine · Sertindole · Schizophrenia · Animal models · Prefrontal cortex

Introduction

Deficits of prefrontal cortical function are prominent features of schizophrenia. These neurocognitive dysfunctions include reduced flexibility in modifying behavior in response to the altering relevance of stimuli. This aspect of executive function is commonly assessed in humans using the Wisconsin Card Sorting Test (WCST) (Grant and Berg 1948) and its modified version, the Intradimensional/Extradimensional Shift (ID/ED) task, developed by Roberts et al. (1988). In fact, a poor WCST and/or ID/ED performance, which is an impairment reminiscent of that observed in patients with frontal lobe damage (Pantelis et al. 1999), is the key cognitive symptom of schizophrenia (Elliott et al. 1998).

Cognitive flexibility may also be assessed in the rodent version of the ID/ED task, i.e., in the attentional set-shifting task (ASST) (Birrell and Brown 2000). In this paradigm, rats must select a bowl containing a food reward based on the ability to discriminate the odors and the media covering the bait. The ASST requires rats to initially learn a rule and form an attentional “set” within the same stimulus...
dimensions. At the extra-dimensional shift (EDs), animals must switch their attention to a new, previously irrelevant stimulus dimension and, for example, discriminate between the odors and no longer between the media covering the bait. The EDs phase, regarded as an index of cognitive flexibility, is impaired by lesions of the medial prefrontal cortex (mPFC) (Birrell and Brown 2000). Hence, the ASST measures specific frontal-dependent cognitive functions in a way homologous to human tests and therefore represents a useful translational approach from animal models to the clinic (Keeler and Robbins 2011).

Noncompetitive antagonists of the N-methyl-D-aspartate receptor (NMDAR), such as ketamine and phencyclidine (PCP), produce a behavioral syndrome in healthy humans that closely resembles the symptoms of schizophrenia (Lahti et al. 1999). Therefore, NMDAR-based models are commonly used to mimic a schizophrenia-like state in laboratory animals. Interestingly, ketamine administration to healthy volunteers produced inflexible responding in the WCST, as revealed by an increase in perseverative errors (Krystal et al. 1994). In line with clinical findings, our previous study demonstrated that acute administration of ketamine to rats also impaired their cognitive flexibility during the EDs phase of the ASST (Nikiforuk et al. 2010).

However, although acute administration of NMDAR antagonists evokes a broad range of schizophrenia-like symptoms, experimental data suggest that repeated dosing protocols might represent a more appropriate preclinical approach for modeling neurochemical changes relevant to the pathophysiology of this disorder. Specifically, the repetitive administration of NMDAR antagonists reduced the expression of glutamic acid decarboxylase 67 (GAD67) and the calcium-binding protein, parvalbumin, in cortical aminobutyric acid-ergic (GABAergic) interneurons (Abdul-Monim et al. 2007). This GABAergic impairment has consistently been shown to reverse cognitive inflexibility in both nonhuman primates and rodents (Jentsch et al. 1997; Jentsch et al. 1998). Because these enduring neurochemical and metabolic alterations in the PFC are thought to underlie cognitive impairments (Dawson et al. 2010; Abdul-Monim et al. 2007; Jentsch et al. 1997), NMDAR antagonist-based preclinical models may represent a useful tool for assessing the pro-cognitive efficacy of antipsychotic drugs.

The first aim of the present study was to investigate the impact of subchronic (5 or 10 consecutive days) administration of the subanesthetic dose (30 mg/kg) of ketamine on rats' ASST performance assessed after 2 weeks of drug withdrawal. This protocol has been effective for inducing behavioral changes mimicking the positive and negative symptoms of schizophrenia, as revealed by potentiating dizocilpine-induced hyperlocomotion and by the decrease in social behavior, respectively (Becker et al. 2003). Likewise, cognitive deficits were observed in tasks assessing spatial working memory (Enomoto and Floresco 2009) and reversal learning (Floresco et al. 2009). Additionally, second-generation antipsychotic drugs (i.e., clozapine and risperidone) but not typical neuroleptics such as haloperidol ameliorated ketamine-induced impairment of social interaction, suggesting predictive validity of this subchronic treatment paradigm (Becker and Grecksch 2004). In line with persistent behavioral abnormalities, this ketamine dosing schedule also induced dysfunction in GABAergic interneurons (Keilhoff et al. 2004). Therefore, repeated ketamine administration represents a validated model in preclinical schizophrenia research.

Although atypical antipsychotic medications appear to demonstrate a promising effect on cognitive functioning in schizophrenia patients (Harvey and Keefe 2001), several specific dysfunctions including cognitive inflexibility are not normalized by these treatments (Goldberg et al. 2007). Accordingly, atypical antipsychotics including clozapine, olanzapine and risperidone were largely ineffective in normalizing set-shifting deficits in preclinical schizophrenia models (Rodefer et al. 2008; Goetghebeur and Dias 2009; but see also McLean et al. 2008). To date, only sertindole has consistently been shown to reverse cognitive inflexibility (Broberg et al. 2009; Goetghebeur and Dias 2009; Kos et al. 2011; Nikiforuk et al. 2010; Rodefer et al. 2008).

Clinical data indicate that among the newer atypical drugs, quetiapine may possess a distinctive cognitive-enhancing action (Riedel et al. 2010). The pro-cognitive efficacy of quetiapine has been demonstrated in several cognitive domains known to be affected in schizophrenia, including executive functions (Kopala et al. 2006; Purdon et al. 2001; Voruganti et al. 2007). Therefore, the second goal of the present study was to evaluate the effects of quetiapine on the ASST performance of ketamine-treated and control rats; sertindole was used as a positive control.

**Materials and methods**

**Animals**

Male Sprague–Dawley rats (Charles River, Germany) weighing 250–280 g on arrival were used in this study. They were initially group-housed (5 rats/cage) in a temperature- (21±1°C) and humidity-controlled (40–50%) colony room under a 12/12-h light/dark cycle (lights on at
0600 hours). Rats were allowed to acclimatize for at least 7 days before the start of the experimental procedure. For 1 week prior to testing, rats were individually housed with mild food restriction (15 g of food pellets per day) and ad libitum access to water. Behavioral testing was performed during the light phase of the light/dark cycle. The experiments were conducted in accordance with the NIH Guide for the Care and Use of Laboratory Animals and were approved by the Ethics Committee for Animal Experiments, Institute of Pharmacology.

Attentional set-shifting: apparatus

Testing was conducted in a modified wire rat housing cage (length × width × height: 42 × 32 × 22 cm) with a white plywood wall dividing half of the length of the cage into two sections (Nikiforuk et al. 2010). During testing, one ceramic digging pot (an internal diameter of 10.5 cm and a depth of 4 cm) was placed in each section. Each pot was defined by a pair of cues along with two stimulus dimensions. To mark each pot with a distinct odor, 5 μl of a flavoring essence (Dr. Oetker®, Poland) was applied on a piece of blotting paper fixed to the external rim of the pot immediately prior to use. A different pot was used for each combination of digging medium and odor; only one odor was ever applied to a given pot. The bait (one-third of a Honey Nut Cheerio, Nestle®) was placed at the bottom of the “positive” pot and buried in the digging medium.

Attentional set-shifting: procedure

The procedure was adapted from Birrell and Brown (Birrell and Brown 2000) and entailed 3 days for each rat.

Day 1, habituation: rats were habituated to the testing area and trained to dig in the pots filled with sawdust to retrieve the food reward. Rats were transported from the housing facility to the testing room where they were presented with one unscented pot (filled with several pieces of Cheerios) in their home cages. After the rats had eaten the Cheerio from the home cage pot, they were placed in the apparatus and given three trials to retrieve the reward from both of the sawdust-filled baited pots. With each exposure, the bait was covered with an increasing amount of sawdust.

Day 2, training: rats were trained on a series of simple discriminations (SD) to a criterion of six consecutive correct trials. For these trials, rats had to learn to associate the food reward with an odor cue (e.g., arrack vs. orange, both pots filled with sawdust) and/or a digging medium (e.g., plastic balls vs. pebbles, no odor). All rats were trained using the same pairs of stimuli. The positive and negative cues for each rat were presented randomly and equally. These training stimuli were not used again in later testing trials.

Day 3, testing: rats performed a series of discriminations in a single test session. The first four trials at the beginning of each discrimination phase were a discovery period (not included in the six criterion trials). In subsequent trials, an incorrect choice was recorded as an error. Digging was defined as any distinct displacement of the digging media with either the paw or the nose; the rat could investigate a digging pot by sniffing or touching without displacing material. Testing was continued at each phase until the rat reached the criterion of six consecutive correct trials, after which testing proceeded to the next phase.

In the simple discrimination (SD) involving only one stimulus dimension, the pots differed along one of two dimensions (i.e., a digging medium). For the compound discrimination (CD), the second (irrelevant) dimension (i.e., an odor) was introduced, but the correct and incorrect exemplars of the relevant dimension remained constant. For the reversal of this discrimination (Rev 1), the exemplars and relevant dimension were unchanged, but the previously correct exemplar was now incorrect and vice versa. The intradimensional (ID) shift was then presented, comprising new exemplars of both the relevant and irrelevant dimensions, with the relevant dimension remaining the same previously. The ID discrimination was then reversed (Rev 2) so that the formerly positive exemplar became the negative one. For the extra-dimensional (ED) shift, a new pair of exemplars was again introduced, but this time a relevant dimension was also changed. Finally, the last phase was the reversal (Rev 3) of the ED discrimination problem. The exemplars were always presented in pairs and varied so that only one animal within each treatment group received the same combination. The following pairs of exemplars were used: pair 1: odor: lemon vs. almond, medium: cotton wool vs. crumpled tissue; pair 2: odor: spicy vs. vanilla, medium: metallic filler vs. shredded paper; and pair 3: odor: rum vs. cream, medium: clay pellets vs. silk. Our previous study demonstrated that there were no differences in performance of rats shifted from odor to medium and from medium to odor (Nikiforuk et al. 2010). Therefore, in an attempt to simplify the experimental design, the order of discrimination was always the same (i.e., from digging medium to odor). The assignment of each exemplar in a pair as being positive or negative at a given phase and the left–right positioning of the pots in the test apparatus on each trial were randomized.

Drugs

Ketamine (aqueous solution (115.34 mg/ml), Biowet, Pulawy, Poland) was diluted in sterile physiological saline. Sertindole (Sigma Aldrich, Poznan, Poland) was dissolved in a minimum amount of 0.1-M hydrochloric acid and diluted with saline, and quetiapine (AstraZeneca) was suspended in 0.5% methylcellulose.

Ketamine (30 mg/kg) was administered intraperitoneally to rats once daily for 5 or 10 consecutive days. The ASST
was performed 14 days following the final drug administration. Animals were left undisturbed during the washout period except for the last 2–3 days when habituation and training were performed. Control animals received vehicle (physiological saline) according to the same experimental schedule. Because no differences were observed between animals receiving vehicle injections for 5 or 10 days, these subgroups were collapsed into one vehicle-treated group for further statistical analysis (Fig. 1).

Sertindole and quetiapine were given per os (PO) 120 min before testing. Drugs or physiological saline were administered in a volume of 1 ml/kg of body weight.

The ketamine treatment regime and doses of antipsychotics (i.e., sertindole and quetiapine) were based on previously reported studies (Nikiforuk et al. 2010; Becker et al. 2003; Swerdlow et al. 1996).

Data analysis

The number of trials required to achieve the criterion of six consecutive correct responses was recorded for each rat and for each discrimination phase. Additionally, the time required by the rats to complete all discriminations was recorded because the compounds could nonspecifically affect performance. The effect of ketamine administration was assessed using two-way ANOVAs with ketamine treatment (0, 5 and 10 days) as a between-subject factor and discrimination phase (SD, CD, Rev 1, ID, etc.) as a within-subject factor. Data from experiments evaluating the effects of antipsychotic drugs were calculated using three-way ANOVAs with one within-subject factor, i.e., discrimination phase, and two between-subject factors: ketamine treatment (0 and 10 days) and a respective antipsychotic drug treatment (sertindole: 0 and 2.5 mg/kg, or quetiapine: 0, 0.63, 1.25 and 2.5 mg/kg). Post-hoc comparisons were performed using the Newman–Keuls test. The alpha value was set at $p<0.05$. The data fulfilled the criteria for normal distribution. Statistical analyses were performed with the use of Statistica 7.0 for Windows.

Results

Effect of repeated ketamine administration on rats’ performance on the attentional set-shifting task

As illustrated in Fig. 1, there was a statistically significant interaction between ketamine treatment and the discrimination phase: two-way ANOVA interaction factor $F(12, 126)=18.26$, $P<0.0001$. Post-hoc analysis revealed that vehicle-treated rats required significantly more trials to reach the criterion on the ED than the ID stage of the task, indicating that they had formed an attentional set towards the relevant dimension before the ED discrimination stage. Furthermore, repeated ketamine administration (30 mg/kg) for 10 (but not 5) consecutive days significantly and specifically impaired rats’ performance in the ED stage of the ASST, as indicated by an increased number of trials to criterion during this phase. There was no significant drug effect during any other discrimination stage.

Corresponding to the increased number of trials during the ED phase, 10-day ketamine administration also prolonged the time to complete this discrimination stage (vehicle: 13.5±0.9 min and ketamine×10 days: 26.3±0.9; ANOV A interaction: $F(12, 126)=9.71$, $P<0.0001$; data not shown).

Effect of sertindole on the set-shifting ability of vehicle- and ketamine-treated rats

As illustrated in Fig. 2, there was a statistically significant interaction between ketamine treatment, sertindole treatment and discrimination phase: three-way ANOVA interaction factor $F(6, 120)=3.73$, $P<0.05$. Acute administration of sertindole (2.5 mg/kg, PO) reversed the 10-day ketamine-administration-induced deficit in ED set-shifting performance. In addition, sertindole administration to vehicle-treated controls did not produce differences across any of the discrimination phases.

The sertindole-induced facilitation of EDs performance in ketamine-treated rats was associated with reduced time to complete this phase (ketamine–vehicle: 21.3±1.2 min and ketamine–sertindole: 13.0±1.6 min; ANOVA interaction: $F(6, 120)=2.43$, $P<0.05$; data not shown).
Effect of quetiapine on the set-shifting ability of vehicle- and ketamine-treated rats

As illustrated in Fig. 3, there was a statistically significant interaction between the ketamine treatment, quetiapine treatment and discrimination phase: three-way ANOVA interaction factor $F_{(18,318)}=1.79$, $P<0.05$. Acute administration of quetiapine (1.25 and 2.5 mg/kg, PO) reversed the subchronic ketamine-induced deficit in ED set-shifting performance. Ketamine-treated rats that received quetiapine at a dose of 2.5 mg/kg also achieved the criterion during the ED stage in fewer trials compared to the vehicle-treated control group. Additionally, quetiapine administration (1.25 and 2.5 mg/kg) improved ED set-shifting in vehicle-treated rats. Quetiapine did not affect any other discrimination phase in either vehicle- or ketamine-treated rats.

The improved EDs performance after quetiapine administration to ketamine-treated and control rats was accompanied by the shortened duration of this discrimination phase (vehicle–vehicle: 15.3±1.1, vehicle–quetiapine 1.25: 9.4±1.2 and vehicle–quetiapine 2.5: 9.2±0.7; ketamine–vehicle: 23.5±1.5, ketamine–quetiapine 1.25: 14.2±1.5 and ketamine–quetiapine 2.5: 11.45±1.6; ANOVA interaction: $F_{(18,318)}=1.87$, $P<0.05$; data not shown).

Discussion

The present study reveals that the subchronic administration of ketamine specifically impairs rats’ performance on the ED stage of the attentional set-shifting task. Furthermore, this ketamine-induced persistent cognitive inflexibility was reversed by acute quetiapine and sertindole treatment. Quetiapine also facilitated ED set-shifting in cognitively unimpaired control animals.

The present demonstration of ketamine-induced ED set-shifting impairment is in line with studies employing repetitive dosing regimens of another NMDAR antagonist, PCP. Indeed, PCP administered twice daily for 7 consecutive days has been consistently demonstrated to impair rats’ performance in the ED stage of the ASST following a 10-day washout period. The present study reveals that the subchronic administration of ketamine specifically impairs rats’ performance on the ED stage of the attentional set-shifting task. Furthermore, this ketamine-induced persistent cognitive inflexibility was reversed by acute quetiapine and sertindole treatment. Quetiapine also facilitated ED set-shifting in cognitively unimpaired control animals.

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day washout period (Rodefer et al. 2005; Rodefer et al. 2008; Goetghebeur and Dias 2009; McLean et al. 2008). Additionally, Egerton et al. (2008) demonstrated selective impairment of ED set-shifting occurring 72 h after the last PCP administration following subchronic (5 days) and chronic intermittent treatment regimes. In another study, chronic (14-day) infusion of PCP via osmotic mini-pumps to rats followed by a 7-day drug-free phase also produced selective deficits in the ED stage (Pedersen et al. 2009). The fact that similar results were obtained in neurodevelopmental schizophrenia models, i.e., early post-natal PCP administration (Broberg et al. 2008) or neonatal ventral hippocampal lesions (Marquis et al. 2008), further supports the utility of the ASST in preclinical schizophrenia research.

In contrast to the present findings, using a similar schedule of ketamine administration, Floresco et al. (2009) showed drug-induced deficits in reversal learning during the instrumental and maze-based versions of the set-shifting task. This discrepancy is not surprising in light of the available data. It is known that NMDAR antagonist-induced impairments are common in tasks specifically designed to assess reversal learning ability at the level of changing stimulus–reward contingencies (Abdul-Monim et al. 2007). Conversely, no effect on reversal learning as assessed in the ASST procedure has been demonstrated after either the repeated administration of PCP (Goetghebeur and Dias 2009; Rodefer et al. 2008; Egerton et al. 2005) or acute ketamine treatment (Nikiforuk et al. 2010). As discussed by Floresco et al. (2009), these discrepancies might be due to procedural differences between these paradigms. In contrast to the maze- and operant-based set-shifting protocols that involve extended training, in the ASST, animals usually achieve CD phase in six consecutive trials; this may prevent the formation of a stable stimulus–reward association. Consequently, reversal learning discrimination seems to be relatively easy for animals, as indicated by the high basal level of performance, and is therefore less prone to disruption. However, the results of Floresco et al. (2009) also demonstrated that repeated ketamine administration improved rats’ set-shifting performance due to the reduction in perseverative errors. This pattern of responding was similar to that previously noted in rats with inactivation of the nucleus accumbens shell and interpreted as an inability to learn the irrelevance of stimuli during the initial discrimination (Floresco et al. 2006a). Therefore, it is likely that the current protocol is sensitive to different aspects of ketamine-evoked set-shifting disruption as compared with the maze and instrumental box protocols.

Alternatively, the differences in ketamine administration protocols may contribute to these discrepancies. Ketamine (30 mg/kg) was administered twice daily for 5 days in the Floresco et al. study and once daily for 10 days in the present experiments. However, Enomoto and Floresco (2009) reported that 5 days of ketamine administration was insufficient to induce PFC-dependent working memory deficits; these occurred only after 10 days of drug administration. Likewise, 5-day ketamine administration did not affect ASST performance in the present study, while the same drug administration protocol has been previously shown to disrupt rats’ social behavior and latent inhibition (Becker et al. 2003). Thus, it appears likely that to disrupt mPFC-dependent functions, ketamine should be administered for more than 5 days.

Finally, the potential limitation of the current experiments is the fact that flexibility in shifting attentional set was assessed only in one direction, i.e., from digging medium to odor. However, an initial relevant dimension (odor or medium) did not affect ED set-shifting (Nikiforuk et al. 2010), and it cannot be excluded that ED performance of animals shifted from medium to odor may be more susceptible to the disrupting effects of ketamine.

Sertindole-induced inhibition of the ASST deficit agrees with previous data demonstrating its efficacy in ameliorating ED deficits induced by either subchronic (Broberg et al. 2009; Goetghebeur and Dias 2009; Rodefer et al. 2008) or post-natal (Broberg et al. 2009) PCP administration in rats. Similar beneficial effects of sertindole have been shown in rats (Nikiforuk et al. 2010) and mice (Kos et al. 2011) treated with ketamine acutely, confirming sertindole-induced improvement of WCST performance in patients with schizophrenia (Gallohfer et al. 2007). Thus, the effectiveness of sertindole in reversing the subchronic ketamine-induced attentional set-shifting deficit may support the predictive validity of the applied model in finding novel compounds that purportedly improve cognitive functioning.

The main finding of the current study is quetiapine-induced facilitation of ED set-shifting; this is the first demonstration of the effectiveness of this drug for modulating prefrontal functions assessed in the ASST. The beneficial effect of quetiapine on cognitive flexibility in ketamine-treated animals may corroborate clinical data, as quetiapine has been demonstrated to improve WCST performance in schizophrenic patients (Kopala et al. 2006; Purdon et al. 2001; Voruganti et al. 2007). However, the lack of drug effects on this specific aspect of executive skills has also been noted (Good et al. 2002; Keefe et al. 2007).

Although quetiapine’s effects on set-shifting have not yet been addressed in preclinical studies, existing data suggest its effectiveness in other cognitive domains. For example, Tanibuchi et al. (2009) demonstrated that 2-week administration of quetiapine reversed the deficits caused by repeated PCP administration on a novel object recognition task in mice. Chronic treatment with quetiapine also counteracted the impairment caused by acute PCP administration on rats’
reference memory in the radial arm maze (He et al. 2006a). Additionally, acute administration of quetiapine restored prepulse inhibition in PCP- and ketamine-treated rats (Swerdlow et al. 1996; Swerdlow et al. 1998). In contrast to its beneficial effects on preattentive information processing, quetiapine treatment did not reverse PCP-induced impairments on attention and impulsivity as assessed in the 5-choice serial reaction time task (Amiati and Markou 2009). Nevertheless, the reported pro-cognitive action of quetiapine was not restricted to schizophrenia-like states. Accordingly, quetiapine treatment ameliorated learning and spatial memory impairments in rats exposed to enhanced single prolonged stress, an animal model of post-traumatic stress disorder (Wang et al. 2010). Quetiapine also alleviated spatial working and reference memory deficits in a transgenic mouse model of Alzheimer’s disease (He et al. 2009). Beneficial effects of quetiapine on cognitive functioning were also noted in a cerebral ischemia model (Yan et al. 2007) and in a methamphetamine-induced neurotoxicity model (He et al. 2006b). Correspondingly, the present demonstration of quetiapine-induced facilitation of set-shifting in cognitively unimpaired animals may predict its efficacy in disorders other than schizophrenia that are also characterized by disturbances in frontal-dependent cognitive flexibility.

The cognitive benefits of quetiapine in the clinic have been partially ascribed to its relatively low affinity for dopamine D2 receptor together with fast dissociation from these receptors and to the lack of affinity for M1 muscarinic cholinergic receptors (Keefe et al. 2007; Kopala et al. 2006; Riedel et al. 2007; Voruganti et al. 2007). As suggested by Floresco et al. (2006b), D2 receptors may play an important role in regulating set-shifting; the authors demonstrated that the infusion of the D2 receptor antagonist eticlopride into the rat mPFC disrupted set-shifting ability, as indicated by the increase in perseverative errors (Floresco et al. 2006b). Likewise, systemic administration of the D2 antagonist sulpiride impaired attentional set-shifting in humans (Mehta et al. 2004). Another line of research showed that administration of the muscarinic antagonist scopolamine also impaired rats’ performance on the ASST (Chen et al. 2004). Altogether, the unique receptor profile of quetiapine may have an advantage over other antipsychotics.

In contrast to other novel generation antipsychotics (e.g., clozapine) that are characterized by a complex multi-receptor profile, quetiapine displays marked affinity only for 5-HT2A, D2, 5-HT1A, α1-adrenergic and α2-adrenergic receptors (Richelson and Souder 2000). Some data indicate that α1-adrenoreceptors might be involved in the mechanism of action of quetiapine. Thus, quetiapine’s effectiveness against PCP-induced cognitive deficits in mice (Tanibuchi et al. 2009) and PCP-induced deficits in prepulse inhibition (Bakshi and Geyer 1999) may have resulted from the drug’s antagonism at α1-adrenoreceptors.

In contrast to this potential therapeutic implication of α1-adrenergic antagonism, the results of Lapiz and Morilak (2006) suggested that α1-receptor stimulation rather than blockade may play a role in facilitating the ASST responding. Thus, it remains questionable if the blockade of α1-adrenergic receptors could improve prefrontal functions specifically related to set-shifting capacity.

Recent data have indicated that the active metabolite of quetiapine, N-desalkylquetiapine, may be involved in its actions (Jensen et al. 2008). This compound was characterized as a potent inhibitor of the noradrenaline reuptake transporter (NET) (Jensen et al. 2008) and showed relatively high affinity for the serotonin 7 (5-HT-7) receptor. The pharmacological profile of N-desalkylquetiapine, already proposed to explain quetiapine’s antidepressant actions, may also account for the drug’s pro-cognitive effects. Indeed, NET inhibitors may represent one of the strategies destined to improve prefrontal functions in schizophrenic patients (Apud and Weinberger 2007). In our study, desipramine, a NET inhibitor, facilitated rats’ performance in ED set-shifting (Nikiforuk and Popik 2011). Moreover, the selective blockade of 5-HT7 receptors has been shown to be effective for ameliorating cognitive deficits in an NMDAR-based animal model (McLean et al. 2009) and for improving cognitive flexibility in rats (our unpublished data). Therefore, one can assume that the activities of N-desalkylquetiapine at the NET and/or 5-HT7 receptors may contribute to the pro-cognitive action of quetiapine in the ASST.

It has been proposed that the ability of atypical antipsychotic drugs to increase catecholamine levels in the prefrontal cortex might underlie their potential pro-cognitive action. Likewise, quetiapine has been demonstrated to increase dopamine and noradrenaline levels in the rat mPFC (Ichikawa et al. 2002; Pira et al. 2004; Yamamura et al. 2009). However, the lack of an effect of quetiapine on PFC dopamine release was also noted (Denys et al. 2004). Because medications enhancing frontal dopaminergic and noradrenergic neurotransmission were shown to facilitate ED set-shifting (Nikiforuk et al. 2010; Nikiforuk and Popik 2011; Tunbridge et al. 2004), the quetiapine-evoked increase in catecholamine levels could account for its beneficial action in the ASST. However, it should be noted that novel generation antipsychotic drugs (i.e., risperidone and olanzapine) increased the prefrontal dopamine level but had no effect on the set-shifting task (Rodefer et al. 2008), suggesting that the mere enhancement of dopamine release is insufficient to produce facilitation of set-shifting. Furthermore, similarly to the actions of sertrindole but not risperidone (Mork et al. 2009), quetiapine enhances the cortical glutamate level (Yamamura et al. 2009). The role of glutamatergic transmission in modulating cognitive flexibility has previously been suggested (Nikiforuk...
et al. 2011). Finally, elevation of the acetylcholine level in the mPFC was also noted after quetiapine administration (Ichikawa et al. 2002). Nevertheless, it remains equivocal whether cholinergic transmission plays a critical role in the modulation of attentional set-shifting because acetylcholine depletion in the rat mPFC did not affect performance on that task (McGaughy et al. 2008).

In summary, the present demonstration of the subchronic ketamine-induced attentional set-shifting deficit reflects clinically relevant aspects of cognitive dysfunction encountered in schizophrenic patients. Furthermore, the effectiveness of sertindole in reversing ketamine-induced cognitive inflexibility may support the predictive validity of the applied model. Finally, the beneficial effects of quetiapine on ASST performance may have therapeutic implications for the treatment of schizophrenia and other disorders associated with frontal-dependent cognitive impairments.

Acknowledgments This study was supported by the Statutory Funds of the Institute of Pharmacology, the Polish Academy of Sciences and project “Prokog,” UDA-POIG.01.03.01-12-063/09-00, co-financed by the European Union from the European Fund of Regional Development.

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