RESEARCH ARTICLE

Effects of Pubertal Cannabinoid Administration on Attentional Set-Shifting and Dopaminergic Hyper-Responsivity in a Developmental Disruption Model of Schizophrenia

Felipe V. Gomes, PhD; Francisco S. Guimarães, PhD; Anthony A. Grace, PhD

Department of Pharmacology, Medical School of Ribeirão Preto, University of São Paulo, Brazil (Drs Gomes and Guimarães); Center for Interdisciplinary Research on Applied Neurosciences, University of São Paulo, Brazil (Drs Gomes and Guimarães); Departments of Neuroscience, Psychiatry and Psychology, University of Pittsburgh, A210 Langley Hall, Pittsburgh, PA 15260 (Dr Grace).

Correspondence: Felipe V. Gomes, MD, Department of Pharmacology, Medical School of Ribeirão Preto, University of São Paulo, 3900 Bandeirantes Ave, Ribeirão Preto, SP, 14049–900, Brazil (gomesfv@usp.br).

Abstract

Background: Adolescent exposure to cannabinoids in vulnerable individuals is proposed to be a risk factor for psychiatric conditions later in life, particularly schizophrenia. Evidence from studies in animals has indicated that a combination of repeated pubertal cannabinoid administration with either neonatal prefrontocortical lesion, isolation rearing, or chronic NMDA receptor antagonism administration induces enhanced schizophrenia-like behavioral disruptions. The effects of adolescent exposure to CB1 receptor agonists, however, have not been tested in a developmental disruption model of schizophrenia.

Methods: This was tested in the methylazoxymethanol (MAM) model, in which repeated treatment with the synthetic cannabinoid agonist WIN 55,212-2 (WIN; 1.2 mg/kg) was extended over 25 days throughout puberty (postnatal days 40–65) in control and MAM rats. The rats received 20 injections, which were delivered irregularly to mimic the human condition. Adult rats were tested for attentional set-shifting task and locomotor response to amphetamine, which was compared with in vivo recording from ventral tegmental area (VTA) dopamine (DA) neurons.

Results: MAM-treated rats showed impairment in the attentional set-shifting task, augmented locomotor response to amphetamine administration, and an increased number of spontaneously active DA neurons in the VTA. Interestingly, pubertal WIN treatment in normal animals induced similar changes at adulthood as those observed in MAM-treated rats, supporting the notion that adolescence exposure to cannabinoids may represent a risk factor for developing schizophrenia-like behavioral signs at adulthood. However, contrary to expectations, pubertal WIN administration did not exacerbate the behavioral and electrophysiological changes in MAM-treated rats beyond that observed in WIN-treated saline rats (Sal). Indeed, WIN treatment actually attenuated the locomotor response to amphetamine in MAM rats without impacting DA neuron activity states.

Conclusions: Taken together, the present results indicate that the impact of cannabinoids during puberty/adolescence on schizophrenia models is more complex than may be predicted.

Keywords: adolescence, cannabinoid, dopamine, MAM treatment, schizophrenia
Introduction

Early insults during brain development have been associated with increased risk of schizophrenia (Harrison and Weinberger, 2005; Rapoport et al., 2005). Evidence for a neurodevelopmental disruption is largely based on follow-back, cohort, and population studies in which the pre-morbid history is associated with the presence of subtle prenatal perturbations that may interact with genetic predisposition to result in the schizophrenia phenotype (van Os et al., 2010). This finding is central to the development of animal models which utilize perinatal insults to produce a behavioral phenotype as adult.

One model that has substantial face validity utilizes the administration of the DNA methylating agent methylazoxymethanol acetate (MAM) to pregnant dams on gestational day (GD) 17 (Moore et al., 2006). This model utilizes a non-selective developmental disruption with no selective genetic manipulation or loss of a specific brain structure, and has findings consistent with those seen in schizophrenia patients. Furthermore, the deficits observed in this model parallel those observed in schizophrenia patients, including anatomical changes (Moore et al., 2006), behavioral deficits (Talamini et al., 2000; Flagstad et al., 2004; Moore et al., 2006), and disruption of rhythmic activity in the frontal cortex (Goto and Grace, 2006). In addition, the MAM model also shows pharmacological validity, with typical and atypical antipsychotics being able to reverse MAM-induced behavioral and electrophysiological changes (Pen et al., 2010; Valenti et al., 2011; Belujon et al., 2013) at a time course consistent with schizophrenia in humans (Agid et al., 2003; Valenti et al., 2011).

In addition to the perinatal phase, adolescence is also a period extremely vulnerable to disruption by environmental influence. This period is characterized by cognitive, emotional, and social maturation. Moreover, other important changes observed in adolescence are risk taking and novelty seeking (Kelley et al., 2004). Besides their adaptive benefits, these behaviors also render adolescents more vulnerable to pathology. For example, epidemiological studies showed an increased risk for drug abuse during adolescence (Fried et al., 2001; Martin et al., 2002). Among these drugs, epidemiological data indicate a causal association between early cannabis abuse and development of psychiatric conditions later in life, including schizophrenia (Arsenault et al., 2004; Degenhardt and Hall, 2006; Fergusson et al., 2006; van Laar et al., 2007). Adolescents initiate cannabis use before consuming other illicit drugs (Cleveland and Wiebe, 2008) and, although the majority of people who experience this drug during adolescence do not develop a psychiatric condition later in life (Gregg et al., 2007; Dekker et al., 2009; Kolliakou et al., 2011), genetic and environmental factors, such as childhood trauma or psychosocial stress, may predispose them to be particularly vulnerable to the effects of cannabis (Caspil et al., 2005; Arsenault et al., 2011; Kuepper et al., 2011). Therefore, adolescent exposure to cannabinoids (that is, CB₁ receptor agonists) in vulnerable individuals is proposed to act as a risk factor for inducing behavioral disturbances (Casadio et al., 2011). This conclusion seems to be more credible when the two-hit hypothesis of schizophrenia is taken into account. In this hypothesis, genetic or environmental factors disrupt early central nervous system development, producing vulnerability to a “second hit” that then may lead to the onset of schizophrenia symptoms. In fact, a combination of a neonatal prefrontocortical lesion with repeated pubertal cannabinoid (CB₁ receptor agonist WIN55,212-2) administration leads to greater impairments in social behavior (Schneider and Koch, 2005) and object recognition memory (Schneider and Koch, 2007), suggesting that pubertal cannabinoid administration in vulnerable individuals might induce enhanced behavioral disturbances. Furthermore, pubertal exposure to Δ⁹-tetrahydrocannabinol (THC), the major psychotomimetic compound present in cannabis, worsened disruption of prepulse inhibition induced by isolation rearing (Malone and Taylor, 2006) and impairment in the object recognition test, induced by chronic administration of phencyclidine (Vigano et al., 2009), an NMDA receptor antagonist. The effects of pubertal exposure to CB₁ receptor agonists, however, have not been tested in neurodevelopmental disruption models of schizophrenia which, due to their delayed onset, would be expected to have greater interaction with adolescent cannabis use.

MAM-treated animals show increased dopamine (DA) neuron population activity in the ventral tegmental area (VTA) that correlates with the enhanced locomotor response to amphetamine (Lodge and Grace, 2007), indicating an enhanced activity in the mesolimbic DA system. Cannabis, like most drugs of abuse, causes an increase in extracellular DA levels (Gardner, 2005). In addition, repeated use of cannabis in adolescence could lead to sensitization of the mesolimbic DA system, a fact that would help to explain why cannabis use during adolescence may facilitate the development of schizophrenia (Stefanis et al., 2004). Moreover, altered DA function is proposed to make schizophrenia patients more vulnerable to the effects of CB₁ receptor agonists (Abi-Dargham, 2004).

Based on this evidence, the effects of pubertal exposure to the CB₁ receptor agonist WIN55,212-2 (WIN) were tested on the behavioral changes and VTA DA neuronal activity observed in MAM-treated rats.

Material and Methods

Animals and MAM Treatment

All experiments were conducted according to the guidelines established by the National Institutes of Health Guide for the Care and Use of Laboratory Animals and were approved by the Institutional Animal Care and Use Committee of the University of Pittsburgh. Pregnant female Sprague-Dawley rats were obtained from Hilltop Lab Animals on GD 15 and individually housed in ventilated plastic breeding tubs. MAM (20 mg/kg, i.p.; Midwest Research Institute) was administered on GD 17. Control dams received injections of saline (1 mL/kg, i.p.). Male pups were weaned on postnatal day (PD) 21 and housed in groups of two to three with littermates approximately 3–4 months of age, at which time they were used for behavioral and electrophysiological experiments. All experiments were performed on multiple litters of MAM- and saline-treated rats.

Experimental Design

Pregnant rats were administered MAM or saline on GD 17. Male offspring of both groups were then administered either the CB₁ receptor agonist WIN55,212-2 (Sigma-Aldrich) or vehicle (Veh). WIN was emulsified in 0.5% Tween 80 and then diluted in saline (0.9%). The drug was administered intraperitoneally at a dose of 1.2 mg/kg in a volume of 1 mL/kg. The treatment with either the synthetic cannabinoid WIN or Veh lasted 25 days, from PD40 to PD65. This period corresponds to the pubertal phase in male rats. It should be noted that puberty and adolescence are
overlapping time periods with puberty being a part of adolescence (Schneider, 2013).

During the treatment period the rats received either one or two injections daily or no injection at all (10 times one injection, 5 times two injections, and 10 times no injection per day, for a total of 20 injections). This protocol was chosen in order to mimic the irregular consumption practice in humans.

As adults, the animals were submitted to the behavioral (PD85–PD100) and electrophysiological tests (PD100–PD125; Figure 1). The WIN dose and the experimental design were based on previous studies (Schneider and Koch, 2002, 2003, 2005, 2007; Schneider et al., 2005; Schneider et al., 2008; Du and Grace, 2013; Zimmerman et al., 2013). All experiments were performed with investigators blinded to treatment.

The rats showed a normal increase in body weight (measured every other day from PD40 to PD80) that was independent of prenatal treatment (MAM or saline) or pubertal drug administration (WIN or Veh; data not shown).

**Attentional Set-Shifting Task**

Both five days before the attentional set-shifting task and during testing, rats were food restricted to approximately 85% of their body weight. The attentional set-shifting task, designed to evaluate extra-dimensional shift as a rodent analog of the Wisconsin Card Sort Test (Tait et al., 2014), was conducted in a white test box (L70 x W40 x H30 cm³) in which a wood panel was used to divide one-third of the box length into two equal sections, forming the choice chambers to which access could be blocked via removable doors. During behavioral testing, one ceramic bowl (diameter of 8 cm and depth of 4 cm) was placed in each choice chamber. Food rewards were one-third pieces of Honey Nut Cheerios (General Mills), which were placed in one bowl per trial and covered with digging media. The media varied by odor and/or texture to provide two different stimulus dimensions to guide choice behavior. Testing was performed according to a modified version of the protocol previously described (Birrell and Brown, 2000; Gastambide et al., 2012). Rats were habituated to the testing box and then initially trained to dig in bowls filled with cage bedding to retrieve food rewards. Once habituated, rats were trained on two simple discriminations (SDs): one based on odor (mustard vs. celery) and one based on texture (shredded paper vs. styrofoam). SD order and reinforced stimuli were pseudorandomly chosen per rat, but counterbalanced across the rat groups. These odor and texture stimuli were not used again in later phases of the experiment. The purpose of this preliminary phase was to acquaint rats with the basic discrimination learning process, as well as to encourage attention to the two different dimensions of the digging media that could be relevant for subsequent stages of discrimination learning. The following day, rats were given a series of seven discriminations (Table 1): a simple discrimination (SD); a compound discrimination (CD) in which digging media differed according to both odor and texture, but with correct and incorrect exemplars remaining similar to the preceding SD; a reversal (Rev1) in which the reward contingency of the CD exemplars is reversed; an intra-dimensional shift (IDS) in which a novel discrimination is learned with new stimuli, the new correct exemplar being of the same dimension as before; a second reversal (Rev2); and an extra-dimensional shift (EDS), in which another discrimination with new stimuli is learned, but in this case the correct exemplar is now from the other previously irrelevant dimension; and finally a third reversal. For each discrimination stage, testing continued until rats reached a criterion level of six correct consecutive trials. The procedure was the same for each stage: a trial was initiated by raising the removable doors to give rats access to the two digging bowls, one of which was baited. The first four trials of each discrimination stage were deemed discovery trials, where rats were permitted to dig in both bowls if they chose the incorrect bowl first. An error was recorded if rats dug first in the un baited bowl. On subsequent trials, if rats started to dig in the unbaited bowl, an error was recorded and the trial was terminated. If rats did not dig at all in either bowl within 3 min, the trial was aborted, recorded as an omission, and reintiated. The number of errors made to reach criterion was recorded per rat for each stage of the test.

**Locomotor Response to Amphetamine**

Adult rats were tested in an open-field chamber (Coulbourn Instruments) in which locomotor activity was determined by beam breaks and recorded with TruScan software (Coulbourn Instruments). All experiments were conducted at the same time each day. Spontaneous activity was recorded for 30 min. After that, rats were injected with D-amphetamine sulfate (0.5 mg/kg, Gastambide et al., 2012).

### Table 1. Example of Order of Exemplar Exposure in the Attentional Set-Shifting Task.

<table>
<thead>
<tr>
<th>Discrimination</th>
<th>Odor Pair</th>
<th>Medium Pair</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simple discrimination</td>
<td>cumin/cinnamon</td>
<td>beddig</td>
</tr>
<tr>
<td>Compound discrimination</td>
<td>cumin/cinnamon</td>
<td>cedar shavings/boxo</td>
</tr>
<tr>
<td>Reversal 1</td>
<td>cumin/cinnamon</td>
<td>cedar shavings/boxo</td>
</tr>
<tr>
<td>Intra-dimensional shift</td>
<td>thyme/cloves</td>
<td>fine shavings/paperchips</td>
</tr>
<tr>
<td>Reversal 2</td>
<td>thyme/cloves</td>
<td>fine shavings/paperchips</td>
</tr>
<tr>
<td>Extra-dimensional shift</td>
<td>oregano/paprika</td>
<td>Cat litter/ground cat litter</td>
</tr>
<tr>
<td>Reversal 3</td>
<td>oregano/paprika</td>
<td>Cat litter/ground cat litter</td>
</tr>
</tbody>
</table>

The table shows a possible order of exposure to the exemplar, where the rat must shift its attention from odor to digging medium at the extra-dimensional shift acquisition. The rewarded exemplar is indicated by the underlined words. Presented with either of the two exemplars from the irrelevant dimension, so that during each trial within a discrimination, all four exemplars are present. Rats were counterbalanced so that 50% underwent odor to medium shift.
i.p.; Sigma-Aldrich) and their locomotor activity was recorded for another 90 min.

**In Vivo Recording from VTA DA Neurons**

Rats were anesthetized with chloral hydrate and mounted on a stereotaxic frame (Kopf). The body temperature was maintained at 37°C using a thermostatically-controlled feedback heating pad (Fintronics). A burr hole was drilled in the skull overlying the right VTA. Extracellular recording microelectrodes were pulled from Omegadot 2.0 mm glass tubing on a Narishige P-5 vertical electrode puller, the tip broken back under microscopic control, and filled with 2M NaCl containing 2% Pontamine Sky Blue dye. The impedance of the electrodes ranged from 6 to 15 MΩ. The stereotaxic coordinates for the VTA were 5.3 mm posterior from bregma, 0.6 mm lateral to the midline, and 6.5–9.0 mm ventral from the brain surface. Single-unit activity was filtered using a highpass filter at 30 Hz and lowpass at 10 kHz. All data analysis was performed using custom software (Neuroscope). Only neuronal activity with a signal-to-noise ratio greater than 3:1 and at least 1–3 min of stable spontaneous activity was used. Six to nine vertical tracks, separated by 200 mm, were sampled in a predetermined pattern within the VTA of each rat. DA neurons were identified according to well-established electrophysiological features (Grace and Bunney, 1983; Ungleess and Grace, 2012), which included the following criteria: (1) location; (2) an action potential duration > 2.2 ms with variable waveform within a train; (3) slow firing rate (1–10 Hz); and (4) irregular and burst firing patterns, with the start of burst characterized by interspike interval < 80 ms, and the end of burst characterized by inter-spike interval > 160 ms. The activity of each identified DA neuron was recorded for 1–3 min. Three parameters of the DA neuron activity were analyzed: (1) the number of spontaneously active DA neurons per electrode track; (2) average firing rate; and (3) the percentage of spikes that occurred in bursts. At the end of recordings, the recording sites were marked via electrophoretic ejection of Pontamine Sky Blue dye from the tip of the electrode (20 μA constant negative current, 30 min). Rats were euthanized by an overdose of anesthetic; the brains were removed, fixed for at least 48 h in 8% paraformaldehyde, cryoprotected in 25% sucrose, and sectioned for histological confirmation of the electrode sites.

**Statistical Analysis**

The attentional set-shifting task was analyzed using repeated-measures 3-way ANOVA with condition (prenatal treatment, MAM, or Sal) and pubertal treatment (Veh or WIN) as the main independent factors, and discrimination type as a repeated measurement followed by Bonferroni post hoc tests. Locomotor activity was analyzed by TruScan software and compared using repeated-measures 3-way ANOVA with condition and treatment as the main independent factors, and time as a repeated measurement. As the 3-way ANOVA indicated a significant effect of condition versus treatment versus discrimination type (F6,174 = 2.5, p < 0.05), but no effects for interaction measurements, planned comparison analyses were conducted which compared groups during each stage of discrimination. MAM:Veh rats (n = 8) required a significantly greater number of trials to reach criterion during the three reversals discriminations and the IDS discrimination compared to Sal:Veh rats (n = 8; p < 0.05, Bonferroni post hoc test; Figure 2). A similar pattern was observed during two of the three reversal discriminations (Rev1 and Rev2) and IDS discrimination, with MAM:WIN animals (n = 9) requiring more trials to reach criterion compared to Sal:Veh rats (p < 0.05, Bonferroni post hoc test; Figure 2). Moreover, pubertal WIN treatment in normal rats (Sal:WIN rats, n = 8) required a significantly shorter time to reach criterion compared to Sal:Veh rats (p < 0.05, Bonferroni post hoc test; Figure 2). Concerning the omitted trials, although no significant difference was observed, the animals were most likely to stop digging at Rev1 (Sal:Veh, -2.9 ± 0.7; Sal:WIN, -3.6 ± 0.6; MAM:Veh, -2.7 ± 0.4; MAM:WIN, -3.2 ± 0.5), while fewer stopped digging at the IDS or Rev2, and very few stopped digging after that.

**Results**

**Effects of Pubertal WIN Exposure on the Behavioral Flexibility in the Attentional Set-Shifting Task in Normal and MAM-Treated Rats Tested as Adults**

During the habituation and training sessions, all rats learned to dig in bowls to retrieve the food reward and perform the SDs. During the test session, Sal:WIN, MAM:Veh, and MAM:WIN rats required significantly more trials to reach criterion than the Sal:Veh group. The repeated-measures 3-way ANOVA indicated significant effects of condition (F1,29 = 18.3, p < 0.0001), treatment (F1,29 = 8.9, p < 0.01), and discrimination type (F6,174 = 68.7, p < 0.0001). There were also significant condition versus treatment (F1,29 = 5.4, p < 0.05), condition versus discrimination type (F6,174 = 3.1, p < 0.05), and treatment versus discrimination type interactions (F6,174 = 2.5, p < 0.05), but no effects for interaction among condition versus treatment versus discrimination type (F6,174 = 1.7, p > 0.05).

All rats required more trials to learn the reversals than they required for either initial acquisition (SD and CD stages) or the IDS (discrimination type: F6,174 = 68.7, p < 0.0001). Moreover, Sal:Veh animals made significantly more errors to reach the criterion of six consecutive correct trials in the EDS than in the IDS (p < 0.05), demonstrating that they had formed an attentional set towards the relevant dimension before the EDS stage (Birrell and Brown, 2000; Gastambide et al., 2012). Concerning the omitted trials, although no significant difference was observed, the animals were most likely to stop digging at Rev1 (Sal:Veh, -2.9 ± 0.7; Sal:WIN, -3.6 ± 0.6; MAM:Veh, -2.7 ± 0.4; MAM:WIN, -3.2 ± 0.5), while fewer stopped digging at the IDS or Rev2, and very few stopped digging after that.

**Figure 2.** Sal:WIN, MAM:Veh, and MAM:WIN treated rats exhibited deficits in the attentional set shifting task (n = 8–9/group). Graph bars represent the mean ± SEM of the number of errors made to reach the criterion of six consecutive correct trials in each test discrimination. *p < 0.05 vs. Sal:Veh rats; repeated measures of 3-way ANOVA followed by Bonferroni post hoc test. SD: simple discrimination; CD: compound discrimination; IDS: intra-dimensional shift; EDS: extra-dimensional shift; Rev: discriminations requiring a reversal learning.
greater number of trials to reach criterion during the Rev1 and Rev2 (p < 0.05, Bonferroni post hoc test; Figure 2). No differences were observed among Sal:WIN, MAM:Veh, and MAM:WIN during any discriminations.

**Effects of Pubertal WIN Exposure on Amphetamine-Induced Hyperlocomotion in Normal and MAM-Treated Rats Tested as Adults**

Both Sal:WIN and MAM:Veh rats exhibited greater locomotor response to amphetamine compared to Sal:Veh rats. However, the MAM:WIN rats showed significantly less amphetamine-induced locomotor activity than either Sal:Win or MAM:Sal rats and were not significantly different from controls. The repeated-measures 3-way ANOVA indicated no significant effects of condition (F<sub>1,23</sub> = 1.3, p > 0.05) or treatment (F<sub>1,23</sub> = 0.22, p > 0.05); however, there was a significant effect of time (F<sub>23,667</sub> = 52.4, p < 0.001) and an interaction between condition and treatment (F<sub>3,667</sub> = 6.5, p < 0.05). There were also significant condition versus time (F<sub>23,667</sub> = 2.1, p < 0.05), treatment versus time (F<sub>23,667</sub> = 1.9, p < 0.05), and condition versus treatment versus time interactions (F<sub>23,667</sub> = 2.4, p < 0.01). Although the 3-way ANOVA did not indicate any effect of condition and treatment, the 2-way ANOVA showed significant effects of group (Sal or MAM + Veh or WIN; F<sub>1,60</sub> = 3.5, p < 0.05), time (F<sub>23,667</sub> = 53.12, p < 0.001), and interaction between group and time (F<sub>23,667</sub> = 2.2, p < 0.001).

Consistent with previous studies showing that rats treated with MAM on GD17 exhibited an enhanced locomotor response to amphetamine (Flagstad et al., 2004; Moore et al., 2006), MAM:Veh rats (n = 8) showed significantly higher levels of locomotor activity in response to amphetamine administration (0.5 mg/kg, i.p.) compared to controls (Sal:Veh, n = 8; p < 0.05 at 5, 10, and 15 min after amphetamine, Bonferroni post hoc test; Figure 3). Likewise, pubertal WIN treatment in normal rats (Sal:WIN, n = 8) produced a significant enhancement in amphetamine-stimulated locomotion compared to Sal:Veh rats (p<0.05 at 5, 10, 20, and 40 min after amphetamine, Bonferroni post hoc test; Figure 3).

Surprisingly, WIN treatment in MAM rats (MAM:WIN, n = 9) induced a significantly lower level of amphetamine-stimulated locomotion compared to MAM:Veh rats (p < 0.05 at 10 and 15 min after amphetamine, Bonferroni post hoc test; Figure 3), and was not significantly different from Sal:Veh rats (p > 0.05, Bonferroni post hoc test). The locomotor activity before amphetamine administration did not differ significantly among all four groups (p > 0.05, Bonferroni post hoc test; Figure 3).

Based on the opposite effects induced by the chronic pubertal treatment with WIN in normal and MAM-treated rats, we tested the effects of a WIN administration given once on PD65 to test whether repeated administration would be required to induce altered locomotor responses (Supplementary Figure 1). Indeed, no effect in the amphetamine-stimulated locomotion induced by the single WIN injection was observed in saline-or MAM-treated rats tested on PD65–PD90 (Supplementary Figure 2).

**Effects of Pubertal WIN Exposure on VTA DA Neuron Activity in Normal and MAM-Treated Rats Tested as Adults**

Both MAM groups as well as the Sal:WIN group demonstrated significant increases in the number of DA neurons firing spontaneously compared to Sal:Veh rats. The number of spontaneously active DA neurons was significantly affected by prenatal MAM (condition: F<sub>2,13</sub> = 4.4, p < 0.05) and pubertal WIN administration (treatment: F<sub>1,23</sub> = 12.9, p < 0.001), but no interaction was observed (F<sub>2,29</sub> = 2.0, p > 0.05; 2-way ANOVA). Consistent with what has been reported previously (Lodge and Grace, 2007, 2009), recordings from MAM:Veh rats (n = 7 rats, 78 neurons) showed a significantly greater number of spontaneously active DA neurons per electrode track (1.6 ± 0.1 cells/track, p < 0.05, Bonferroni post hoc test) compared to Sal:Veh rats (n = 6 rats, 46 neurons, 0.99 ± 0.2 cells/track; Figure 4A).

Compared to Sal:Veh rats, Sal:WIN rats (n = 7, 92 neurons) showed a significantly greater number of spontaneously active DA neurons (1.9 ± 0.2 cells/track, p < 0.05, Bonferroni post hoc test). These changes required repeated WIN exposure, given that no effect was observed after a single WIN injection on PD65 (Supplementary Figure 3).

Similar to the MAM:Veh rats, the numbers of DA neurons firing in the MAM:WIN (n = 7, 90 neurons; 1.9 ± 0.3 cells/track) was greater than the Sal:Veh rats (p < 0.05, Bonferroni post hoc test). Importantly, despite differences in amphetamine-induced locomotion, no difference was observed between MAM:Veh and MAM:WIN rats with respect to DA neuron activity (p > 0.05, Bonferroni post hoc test; Figure 4A). The firing rate and percentage of spikes in bursts did not differ significantly across all four groups (p > 0.05 by two-way ANOVA; Figure 4B and 4C).

**Discussion**

Consistent with previous studies (Featherstone et al., 2007; Lodge and Grace, 2007; Gastambide et al., 2012), MAM-treated rats showed an impairment in the attentional set-shifting task, augmented locomotor response to amphetamine administration, and an increased number of spontaneously active DA neurons in the VTA. Importantly, despite differences in amphetamine-induced locomotion, no difference was observed between MAM:Veh and MAM:WIN rats with respect to DA neuron activity (p > 0.05, Bonferroni post hoc test; Figure 4A). The firing rate and percentage of spikes in bursts did not differ significantly across all four groups (p > 0.05 by two-way ANOVA; Figure 4B and 4C).
locomotion in the MAM rats without altering the increase in DA neuron activity.

The results of the present study are consistent with recent findings showing that repeated pubertal cannabinoid treatment induces lasting behavioral changes in adulthood, including sensorimotor gating impairment, abnormal social behavior, and anhedonia (Schneider and Koch, 2003, 2005, 2007; Schneider et al., 2008). Moreover, rodents chronically treated during different periods of adolescence with other CB1 receptor agonists (CP55,940 and THC) and tested as adults also exhibited deficits in sensorimotor gating, object recognition, and spatial working memory (O’Shea et al., 2004; Quinn et al., 2008; Rubino, Realini, Braida, Alberio, et al., 2009; Rubino, Realini, Braida, Guidi, et al., 2009; Gleason et al., 2012). The current study is the first time that the long-term effects of cannabinoid exposure during adolescence have been evaluated in the attentional set-shifting task.

In the set-shifting task, rats are required to solve a series of discriminations by attending to a particular perceptual dimension of a multidimensional stimulus. A critical discrimination occurs when rats are required to shift to an alternate perceptual dimension after having acquired an attentional set to the previous dimension (Tait et al., 2013). The neural substrates of set shifting and reversal learning are reasonably well-defined. Lesions of the monkey lateral prefrontal cortex (Dias et al., 1997) and the equivalent prelimbic and infralimbic regions of the rat medial prefrontal cortex (Birrell and Brown, 2000) disrupt attentional set-shifting ability, whereas lesions of the orbitofrontal cortex selectively impair reversal learning in both species (Dias et al., 1997; McAlonan and Brown, 2003). Impaired reversal learning has recently been highlighted to occur reliably in schizophrenia patients. Indeed, Leeson et al. (2009) found in a large group of first-episode schizophrenia patients that although they were impaired on set shifting, they also exhibited small but consistent deficits in reversal learning. Moreover, chronic cannabis use in adolescence appears to be associated with overall less efficient executive function and attention (Abdullaev et al., 2010).

Consistent with previous studies (Featherstone et al., 2007; Gastambide et al., 2012), MAM rats exhibited a variety of cognitive impairments, including reversal learning and attentional set shifting, as defined by requiring a greater number of trials than controls to successfully learn to shift cognitive set between stimuli belonging to the same perceptual dimension (IDS), and having difficulties in learning to reverse a previously acquired discrimination. Interestingly, the pubertal WIN treatment in normal rats also resulted in a significantly greater number of trials to reach criterion during two of the three reversal learning discrimination trials. Thus, the deficits observed in reversal learning, together with the preservation of ability to shift strategy, indicated that MAM-treated rats and the pubertal WIN exposure in normal animals induced an increased rigidity in the processes required to update responses based on affective associations between stimuli and reward presentation, but did not affect ability for higher order attentional flexibility (EDS). This suggests that the deficits in the reversal learning and IDS were not due to a generalized performance or cognitive impairment. Although it has been suggested that adolescent cannabinoid exposure in vulnerable individuals might induce even more pronounced behavioral disturbances, and studies with animals have shown an enhanced cognitive impairment observed after the combination of pubertal exposure to cannabinoids with neonatal prefrontocortical lesions, social isolation, or chronic administration of phencyclidine (Schneider and Koch, 2005, 2007; Malone and Taylor, 2006; Vigano et al., 2009), no significant difference was observed between Veh- and WIN-treated MAM rats in the attentional set-shifting task.

Similar to MAM-treated rats, WIN administration during puberty induced an augmented locomotor response to amphetamine and an increased number of spontaneously active DA neurons in the VTA in normal animals as adults. Previous studies have shown that acute CB1 receptor activation increases mesolimbic DA activity (French, 1997; Tanda et al., 1997; Wu and French, 2000). However, this is the first study showing persistent long-term changes in mesolimbic DA activity induced by pubertal cannabinoid exposure.

Prenatal MAM administration and pubertal cannabinoid exposure have been shown to induce similar changes in GABA neurons of adult rats. Zamberletti et al. (2014) observed that adolescent THC exposure in mice reduced GAD67 expression in interneurons containing the calcium binding protein parvalbumin (PV) within the adult prefrontal cortex. Moreover, repeated CB1 receptor activation in adolescence elicited an enduring state of prefrontal cortex disinhibition due to a developmental
impairment of prefrontal GABAergic transmission (Cass et al., 2014). Decreased GABAergic signaling is among the most robust postmortem pathological changes observed in schizophrenia (Reynolds et al., 2002; Lewis et al., 2005). Specifically, a decrease in GAD67 protein is observed postmortem throughout the cortex of schizophrenia patients (Hashimoto et al., 2003) that are largely restricted to the GABAergic PV-positive interneurons (Lewis et al., 2005). Interestingly, a decrease in PV-containing interneurons is also a consistent observation in a diverse variety of animal models of schizophrenia, including the MAM model (Penschuck et al., 2006; Lodge et al., 2009; Gill and Grace, 2014).

It has been suggested that the augmented DA neuron activity and hyper-responsivity to psychomotor stimuli observed in MAM-treated rats results from an increased activity within ventral regions of the hippocampus due to a loss of PV interneurons (Lodge et al., 2009). In addition to the ventral hippocampus, adult MAM-treated rats also display specific reductions in the number of PV-positive interneurons throughout the medial prefrontal cortex (Penschuck et al., 2006; Lodge and Grace, 2009) and the orbitofrontal cortex (Gastambide et al., 2012), the main brain structures involved in set-shifting and reversal learning, respectively (Birrell and Brown, 2000; Alfonso and Brown, 2003). Thus, based on the evidence indicating that cannabinoid exposure during adolescence may reduce the number of PV-positive interneurons (Zambrerletti et al., 2014), changes in the PV expression could be involved in the dopaminergic dysfunction and in the impairment observed during the attentional set-shifting task in WIN-treated normal, similar to that observed with MAM-treated rats.

No additive or synergic effect, however, was found in MAM-treated rats that received WIN. Although this could reflect a ceiling effect, it is not possible to rule out the involvement of at least partially distinct (and parallel) mechanisms for the attentional set-shifting task impairment induced by these two treatments.

The most intriguing finding of our study is that, although WIN-treated MAM rats showed an enhanced VTA DA neuronal spontaneous activity that was similar to Veh-treated MAM rats, pubertal WIN exposure in MAM rats decreased amphetamine-induced hyperlocomotion. The reason for this attenuation is unclear, particularly given that increases in DA neuronal activity in the VTA, such as those observed after either MAM (Lodge and Grace, 2007) or WIN administration, along with other models (e.g., amphetamine sensitization [Lodge and Grace, 2012], temporal lobe epilepsy [Cifelli and Grace, 2012]) have consistently shown parallel changes between the number of DA neurons firing spontaneously and amphetamine-induced locomotor activity. This suggests that the pubertal exposure to WIN may have induced compensatory changes in MAM rats that are downstream from DA neuron activity. Thus, it is known that exogenous cannabinoids affect the function of the endocannabinoid system. Indeed, pubertal cannabinoid exposure can change the expression of components of the endocannabinoid system in brain structures related to motivation and motor control (Marco et al., 2007; Ceci et al., 2014). Therefore, plastic changes in the endocannabinoid system induced by repeated CB1 receptor agonist administration could lead to plastic changes in the CB1 receptor modulation of GABA- or glutamate-mediated neurotransmission (Wilson and Nicoll, 2002; Fernandez-Ruiz et al., 2010) in key brain structures related to hyperlocomotion, such as the nucleus accumbens, thereby compensating for the modifications in other brain areas (for example, the ventral subiculum) induced by MAM. Interestingly, Spano et al. (2013) observed that both WIN self-administration and passive WIN administration (i.v.) over 14 days attenuated hyperlocomotion in response to an acute phencyclidine (PCP) challenge in adult rats treated chronically with PCP, a model of schizophrenia based on the N-methyl-D-aspartate receptor hypofunction.

Alternately, a decrease in the DA transporter seen in the caudate nucleus of schizophrenia patients was not observed in patients who had used cannabis (Dean et al., 2003), leading the investigators to suggest that THC might reverse the decreases in DA transporter expression associated to schizophrenia.

In conclusion, these results are consistent with the notion that adolescent exposure to cannabinoids may represent a risk factor for developing schizophrenia-like signs at adulthood. However, contrary to our hypothesis that pubertal MAM-treated rats would be more susceptible to the cannabinoid exposure, WIN administration did not exacerbate the behavioral and electrophysiological changes in MAM-treated rats, and in fact prevented the augmentation of the locomotor response to amphetamine. While several epidemiological studies have clearly shown an association between cannabis use and susceptibility to schizophrenia (Arseneault et al., 2004; Casadio et al., 2011), it is not possible to evaluate if subgroups of patients that may develop schizophrenia later in life are protected by cannabis use. Therefore, it is clear that the relationship between cannabinoid exposure and susceptibility to disease states is more complex than may be predicted by correlative epidemiological studies.

Supplementary Material
For supplementary material accompanying this paper, visit http://www.ijnp.oxfordjournals.org/

Acknowledgments
We thank Niki MacMurdo for her technical assistance. This work was supported by the Brazilian Federal Agency for Support and Evaluation of Graduate Education, CAPES/PDSE (10865/13–7, FVG) and USPHS MH57440 (AAG). Drs Gomes and Guimarães are, respectively, recipients of a FAPESP doctoral (2010/17343-0) and of a CNPq researcher fellowship.

Statement of Interest
Drs Gomes and Guimarães declare no conflict of interest. Dr Grace has received funds from Johnson & Johnson, Lundbeck, Pfizer, GSK, Merck, Takeda, Dainippon Sumitomo, Otsuka, Lilly, Roche, Asubio, and Abbott.

References


Lewis DA, Hashimoto T, Volk DW (2005) Cortical inhibitory neurons and schizophrenia. Nat Rev Neurosci 6:312–324.

Lodge DJ, Behrens MM, Grace AA (2009) A loss of parvalbumin-containing interneurons is associated with diminished oscil-


Gomes et al. | 9

by guest on October 7, 2016 http://ijnp.oxfordjournals.org/ Downloaded from