Serotonergic hallucinogens differentially modify gamma and high frequency oscillations in the rat nucleus accumbens

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Abstract

Rationale  The nucleus accumbens (NAc) is a site critical for the actions of many drugs of abuse. Psychoactive compounds, such as N-methyl-D-aspartate receptor (NMDAR) antagonists, modify gamma (40–90) and high frequency oscillations (HFO, 130–180 Hz) in local field potentials (LFPs) recorded in the NAc. Lysergic acid diethylamide (LSD) and 2,5-dimethoxy-4-iodoamphetamine (DOI) are serotonergic hallucinogens and activation of 5HT2A receptors likely underlies their hallucinogenic effects. Whether these compounds can also modulate LFP oscillations in the NAc is unclear.

Objective  This study aims to examine the effect of serotonergic hallucinogens on gamma and HFO recorded in the NAc and to test whether 5HT2A receptors mediate the effects observed.

Methods  LFPs were recorded from the NAc of freely moving rats. Drugs were administered intraperitoneally.

Results  LSD (0.03–0.3 mg/kg) and DOI (0.5–2.0 mg/kg) increased the power and reduced the frequency of HFO. In contrast, the hallucinogens produced a robust reduction in the power of low (40–60 Hz), but not high gamma oscillations (70–90 Hz). MDL 11939 (1.0 mg/kg), a 5HT2A receptor antagonist, fully reversed the changes induced by DOI on HFO but only partially for the low gamma band. Equivalent increases in HFO power were observed after TCB-2 (5HT2A receptor agonist, 0.1–1.5 mg/kg), but not CP 809101 (5H2C receptor agonist, 0.1–3 mg/kg). Notably, hallucinogen-induced increases in HFO power were smaller than those produced by ketamine (25 mg/kg).

Conclusions  Serotonergic hallucinogen-induced changes in HFO and gamma are mediated, at least in part, by stimulation of 5HT2A receptors. Comparison of the oscillatory changes produced by serotonergic hallucinogens and NMDAR antagonists are also discussed.

Keywords  Accumbens · High frequency oscillations · Gamma · Rat · Serotonin receptor · Lysergic acid diethylamide · Hallucinogens · Local field potentials

Introduction

Lysergic acid diethylamide (LSD) and 2,5-dimethoxy-4-iodoamphetamine (DOI) are potent hallucinogens known to produce changes in mood, thought, and sensory perception in humans. LSD is structurally classified as an indoleamine and is a relatively nonselective agonist active at serotonergic receptors as well as on other receptor systems (Halberstadt and Geyer 2011; Marona-Lewicka et al. 2009; Teitler et al. 1990). DOI is a phenethylamine that displays high affinity for 5HT2A/2C receptor subtypes (Appel et al. 1990; Johnson et al. 1987; Krebs-Thomson et al. 1998). Despite differences in chemical structure and pharmacology, these compounds produce relatively similar behavioural effects in humans and rats, much of which appears to be associated with activation of the 5-HT2A receptor subtype; for reviews, see Aghajanian and Marek (1999), Halberstadt and Geyer (2011), Nichols (2004), and Passie et al. (2008).

The nucleus accumbens (NAc) is widely considered to function as a limbic–motor interface, receiving projections from the prefrontal cortex, hippocampus, and amygdala, along with a dense dopaminergic input from the ventral tegmental area (Finch 1996; O’Donnell and Grace 1995;
Oades and Halliday 1987). Altered functioning of these limbic regions may contribute to the effects produced by psychoactive compounds (Jentsch and Roth 1999; Vollenweider and Geyer 2001). In a drug discrimination study, local infusion of LSD to the NAc produced a cueing effect equivalent to that of systemic administration (Nielsen and Scheel-Kruger 1986) suggesting that this structure may directly participate in some of the core effects of hallucinogenic compounds. Accumbal neurons, express 5HT2A and 5HT2C receptors (McKenna et al. 1989; Pazos et al. 1985; Pazos and Palacios 1985) and an in situ hybridization study on rats injected with LSD showed increased c-fos expression in the NAc (Erdtmann-Vourliotis et al. 1999). In vitro, application of 5HT can depolarise accumbal neurons (North and Uchimura 1989) and radiolabelled 125IODI can bind to 5HT2 receptors in the NAc (Appel et al. 1990). Thus, the NAc may be an important region in mediating the effects of serotonergic hallucinogens.

Local field potentials (LFPs) primarily reflect the sum of currents arising predominantly from synchronised synaptic input. Many drugs of abuse, such as ketamine, tetrahydrocannabinol, and amphetamine, have been shown to influence oscillations in different brain areas (Berke 2009; Hunt et al. 2011; Kargieman et al. 2007; Ma and Leung 2000; Pinault 2008; Robbe et al. 2006). Serotonergic hallucinogens produce a general decrease in the power of LFP oscillations in several brain areas of freely moving rats (with exception to alpha where increases are observed; Dimpfel et al. 1989). More recent studies have demonstrated a decrease in the power of cortical delta oscillations in anaesthetised rats (Celada et al. 2008) and a more generalised decrease in frequency bands up to 50 Hz (Palenicek et al. 2012). However, to date, the effects of serotonergic hallucinogens on LFP oscillations recorded from the rat NAc are largely unknown.

In the NAc, gamma can be divided into two frequency bands, low gamma (40–60 Hz) and high gamma (70–90 Hz; for review see, van der Meer et al. 2010). Systemic injection of D-amphetamine and methamphetamine reduce the power of low gamma but increase high gamma power in the NAc of freely moving rats (Berke 2009; Morra et al. 2012). Frequencies above the gamma range, known as high-frequency oscillations (HFO) in this region. Much smaller increases in HFO power are produced by the psychostimulant D-amphetamine (Hunt et al. 2006). Considering that distinct classes of psychoactive compounds can modify LFP oscillations in the NAc, we examined the effect of serotonergic hallucinogens on gamma and HFO in the NAc of freely moving rats.

### Methods

#### Surgery

Thirty-three male Wistar rats (250–350 g) in total were used in this study. Rats were anaesthetized with isoflurane and implanted unilaterally with twisted stainless steel electrodes (110 μm, Science Products, Germany) insulated except at the tip targeted to NAc according to the coordinates of the stereotaxic atlas (Paxinos and Watson 1988) [AP (+1.6 mm) and LM (+1.2 mm) with respect to bregma, DV (-7.1 mm), with respect to cortical surface]. A silver wire connected to a skull screw posterior to the bregma was used as the ground/reference. The locations of the recording electrodes were determined on 40 μm Cresyl violet stained sections post mortem.

#### Experimental procedure

One week after surgery, the animals were handled for approximately 30 min per day, for 2 days. Following this, LFPs were recorded to check the signal quality, habituate the animals to the experimental chamber, and also to the process of connecting/disconnecting to the recording cable. Experiments were performed from 8AM to 7PM. LFPs were recorded using a JFET preamplifier at the head stage, the signals were relayed through a 25-channel commutator (Crist Instruments, USA), amplified ×1,000, filtered 0.1–1000 Hz (A-M Systems, USA), digitised 4 kHz (Micro1401, CED, Cambridge, UK) and stored on a PC for offline analysis. The animals were grouped according to the experimental destinations described in Table 1. Locomotor activity was assessed by beam-breaks (Columbus Instruments, USA). All experiments were performed according to a Latin square design, whereby every animal received each dose of the drug in a pseudorandomised order. The drug washout between consecutive experiments was at least 3 days. All experiments were conducted in accordance with the European Guidelines on the Care and Use of Laboratory Animals (86/609/EEC) and approved by a local ethics committee. All efforts were made to minimise the suffering of the animal and to reduce the number of animals used.

#### Drug treatments

Rats were injected intraperitoneally (i.p.) with different doses of the drug/vehicle as mentioned in Table 1. At least 3 days elapsed between all injections of drug or vehicle. Rats were baselined for 20 min prior to injection and then recorded for at least 60 min thereafter, with exception to group 3. At the end of the main experiment, these rats received injection of ketamine (i.p.). In group 3, every rat
received each of the four combinations of drugs DOI–MDL 11939, DOI–vehicle, saline–MDL 11939, and saline–vehicle. Additionally, after this main experiment, four rats from this group were injected with DOI followed 30 min later by injection of SB 242084. LSD and ketamine were purchased from Sigma-Aldrich (Poznan, Poland), DOI hydrochloride, MDL 11939, SB 242084, TCB-2 and CP 809101 hydrochloride were purchased from Tocris Bioscience (UK).

Data processing

The raw LFPs were cleaned to remove movement large artefacts and then were analysed using Spike2 software. Total power of frequency bands—HFO (typically 130–180 Hz, for TCB-2 and CP 809101 studies 110–180 Hz was used) and gamma (low 40–60 Hz and high 70–90 Hz)—were analysed in 60 s data bins using FFT’s of 4096 points (Spike2). HFO events were extracted by filtering the raw signal with a digital band pass filter at 130–180 Hz and oscillatory episodes greater than 6SD were marked by events, as described previously (Hunt et al. 2008). Inter-event intervals were calculated as an index of HFO frequency.

Statistical analysis

Data were expressed as percentage of baseline values (percent change) and analysed using either, repeated-measures two-way analysis of variance (ANOVA) or one-way ANOVA. Bonferroni post hoc tests were used to find significant differences between doses/drugs. Values were expressed as mean±SEM and were considered to be significant if p<0.05.

Results

Changes in the rat behaviour after injection of serotonergic hallucinogens

Intraperitoneal injection of LSD (0.03, 0.1, and 0.3 mg/kg) or DOI (0.5, 1, and 2 mg/kg) produced similar characteristics of changes in behaviour including, head twitches, wet-dog shakes, and altered posture with the rat often remaining in one corner of the recording chamber and lying flat. For both drugs, these effects were more pronounced at higher doses and typically lasted for the duration of the experiment.

Motor activity was evaluated by analysing the total number of beam breaks in 60 s bins—20 min at baseline and 60 min after injection. Two-way ANOVA with time as the repeated-measure showed a significant effect of LSD on beam-breaks (dose×time F(3,20,2340)=3.20, p=0.0003). Bonferroni post hoc test showed a significant reduction in movement for all doses of LSD versus vehicle, but this was significant for only one time point for all the doses (p<0.01). There were no other significant differences. For the DOI group, we did not observe any significant dose×time interaction (F(3,234,1560)=0.670, p=0.999). At the end of the LSD/DOI studies, all rats received an injection of 25 mg/kg ketamine which produced an immediate increase in the beam breaks lasting around 10 min (LSD, F(3,1950)=1.61, p<0.0001). Bonferroni post hoc test revealed a significant increase in beam breaks for ketamine versus all doses of LSD/DOI and their vehicle (p<0.001).

Serotonergic hallucinogens increase the power of high-frequency oscillations in the LFPs of nucleus accumbens

LFPs were recorded before and at least 1 h after i.p. injection of LSD (0.03, 0.1, and 0.3 mg/kg), DOI (0.5, 1, and 2 mg/kg) or vehicle. Systemic administration of both LSD and DOI increased the power of HFO (130–180 Hz) but decreased the power of the gamma band. Representative time courses of these LFP changes, expressed as spectrograms are shown in Figs. 1a and 2a. Power spectra (10 min) before and after injection of LSD and DOI are shown in Figs. 1b and 2b.

Time courses showing the mean power of HFO (expressed as percent change) for the LSD and DOI studies are shown in Figs. 1c and 2c, respectively. For both LSD and DOI groups, repeated-measures two-way ANOVA revealed a significant effect of dose (LSD, F(3,24)=3.9, p=0.0212; DOI group, F(3,20)=3.77, p=0.0271), time (LSD group, F(79,1896)=10.15,
p<0.0001; DOI group, $F_{78,1560}=5.24, p<0.0001$) and interaction (LSD group, $F_{237,1896}=1.84, p<0.0001$; DOI group, $F_{234,1560}=2.53, p<0.0001$). For the LSD group, Bonferroni post hoc tests revealed significant increases in the power of HFO for 0.1 and 0.3 mg/kg doses compared to vehicle ($p<0.001$, for both), but no difference for 0.03 mg/kg, the lowest dose tested. For the DOI group, Bonferroni post hoc tests revealed significant differences for 1 mg/kg ($p<0.05$) and 2 mg/kg doses ($p<0.001$), compared to vehicle, but no difference for 0.5 mg/kg, the lowest dose tested. The dose of 0.5 mg/kg significantly differed from the 2 mg/kg dose ($p<0.001$).

Additionally, we analysed data for the first 20 min after administration of the drug as the time courses of HFO power showed major effects that occurred during this time. Analysis confirmed that both LSD and DOI produced a significant increase in the power of HFO. Repeated-measures one-way ANOVA revealed a significant effect for both LSD and DOI groups (LSD, $F_{3,18}=9.39, p<0.0006$; DOI, $F_{3,15}=12.12, p<0.0003$, shown alongside the time courses in Figs. 1c and 2c). Bonferroni post hoc test revealed, significant increases in the power of HFO for 0.1 mg/kg ($p<0.01$) dose and 0.3 mg/kg ($p<0.001$) of LSD and for 2 mg/kg ($p<0.001$) dose of DOI group in comparison to the vehicle.
To determine whether serotonergic hallucinogens influence the frequency of HFO, the raw LFPs were digitally 130–180 Hz bandpass filtered and the peaks of oscillations (>6 SD baseline) were marked. The interpeak intervals for the first 20 min after injection of drugs or vehicle were analysed (Figs. 1d and 2d). Repeated-measures one-way ANOVA revealed a significant effect of drug on the inter-peak interval (LSD, $F_{3,18}=3.658$, $p=0.0322$; DOI, $F_{3,15}=19.79$, $p<0.0001$). In both groups, Bonferroni post hoc test revealed an increase in the interpeak interval, with respect to vehicle, that was significant for the highest dose of LSD ($p<0.05$) and for all doses of DOI (at least, $p<0.05$). The raster plot shows individual peaks of HFO (>6 SD) and a gradual increase in the inter-event interval indicative of reduced frequency (Figs. 1d and 2d).

Serotonergic hallucinogens reduce the power of gamma oscillations in the LFPs of nucleus accumbens

Low gamma (40–60 Hz) This band was clearly visible in the baseline power spectra of most rats used in our study and corresponds with the frequency band that has been termed low gamma or gamma-50 in the NAc (Berke et al. 2004; Berke 2009; Kalenscher et al. 2010). Time courses of the total power of low gamma at baseline and after injection of LSD and DOI are shown in Figs. 1e and 2e. For both LSD and DOI groups, repeated-measures two-way ANOVA revealed a significant effect of dose (LSD group, $F_{3,24}=7.83$, $p=0.0008$; DOI group, $F_{3,20}=4.46$, $p=0.015$), time (LSD group, $F_{79,1896}=16.82$, $p<0.0001$; DOI group, $F_{78,1560}=17.04$, shown for the first 20 min post-injection. Raster plot shows the distribution of HFO events across time for DOI 2 mg/kg. e Time courses showing the effect of DOI on the power of gamma in NAc. Mean powers of gamma (60-min average) after injection are shown alongside. f Power of HFO (clear circles) and gamma (filled circles) for individual rats 10 min before and after injection of 2 mg/kg DOI. All animals showed a clear increase in the power of HFO and decrease in the power of gamma when compared to the baseline activity. g Placement of electrodes, $N=6$ rats are indicated by the black dots. Values expressed as mean±SEM, *$p<0.05$, **$p<0.01$, ***$p<0.001$ with respect to vehicle.

Fig. 2 DOI modulates the power of HFO (130–180 Hz) and gamma (40–60 Hz) in the NAc of freely moving rats. a Representative spectrograms showing changes in the power of HFO and gamma across time for vehicle and DOI 2 mg/kg. Black arrows indicate time of injection and the red arrow head points to the HFO band. Colour scales are logarithmic and are the same in both examples. b Power spectra (10-min average) from an example rat at baseline and after injection of 2 mg/kg DOI. e Time courses showing the effect of DOI (0.5, 1, and 2 mg/kg) on the power of HFO. Broken line indicates time of injection. Mean powers of HFO (20-min average) after injection are shown alongside. d Effect of DOI on interpeak interval of HFO events are
p < 0.0001) and interaction (LSD group, \( F_{3,1896} = 1.38, p = 0.0002 \); DOI group, \( F_{3,1560} = 1.53, p < 0.0001 \)). For the LSD group, Bonferroni post hoc tests revealed a significant decrease in the total power of low gamma for 0.1 and 0.3 mg/kg doses \((p < 0.01\) and \(p < 0.05\), respectively) compared to vehicle. For the DOI group, reductions in low gamma power were found for 1 and 2 mg/kg doses \((p < 0.05\) and \(p < 0.01\), respectively) compared to the vehicle. Repeated-measures one-way ANOVA for the entire 60-min post-injection period revealed a significant effect of the drug \((F_{1,14} = 7.45, p = 0.016)\) and interaction \((LSD, F_{3,14} = 7.45, p = 0.016)\) and interaction \((LSD, F_{3,15} = 7.078, p = 0.0035\), shown alongside Figs. 1e and 2e). Bonferroni post hoc revealed significant reductions in the power of low gamma for all doses of LSD \((0.03 \text{ mg/kg, } p < 0.01, 0.1 \text{ and } 0.3 \text{ mg/kg, } p < 0.001\) and DOI \((0.5 \text{ and } 1 \text{ mg/kg, } p < 0.05\) and 2 mg/kg, \(p < 0.01\)) tested when compared to vehicle.

**High gamma (70–90 Hz)** After injection of the highest doses of LSD and DOI, a band in the high gamma range could be discerned in the spectrograms. However, repeated-measures two-way ANOVA did not reveal a significant effect of the dose \((LSD, F_{3,24} = 0.79, p < 0.5119\); DOI group, \( F_{3,20} = 0.86, p = 0.479\) and interaction \((LSD, F_{3} = 0.98, p = 0.5516\); DOI group, \( F_{3,1560} = 0.99, p = 0.5367\)). See also the example spectrograms (shown in Figs. 1 and 2) demonstrating the change in low and high gamma power distribution of across time after LSD and DOI injection. When the power of high gamma was expressed as a percentage of the whole gamma range, we found a significant increase in the proportion of high gamma for the DOI group \((p = 0.001)\) but not for the LSD group \((p = 0.001)\) and interaction \((LSD, F_{3,1580} = 1.48, p < 0.0001\) but not for the LSD group \((p = 0.001)\) and interaction \((LSD, F_{3,1896} = 1.13, p = 0.105\); Electronic supplementary material (ESM) Fig. 1).

Blockade of 5HT2A receptors reverse the increase in HFO power produced by DOI

In a separate study, 30 min after injection of 2 mg/kg DOI, the animals received an i.p injection of the potent 5HT2A receptor antagonist 1 mg/kg MDL 11939 or vehicle. Example spectrograms are shown in Fig. 3a and power spectra in Fig. 3b. Time courses showing the effect of MDL 11939/vehicle after DOI pretreatment on the total power of HFO are shown in Fig. 3c. Repeated-measures two-way ANOVA revealed a significant effect of the drug \((F_{1,14} = 7.45, p = 0.016)\), time \((F_{107,1498} = 24.54, p < 0.0001)\) and interaction \((F_{107,1498} = 10.12, p < 0.0001)\). Bonferroni post hoc tests revealed a significant and sustained reduction in the power of HFO for rats injected with MDL 11939 compared to vehicle \((p < 0.001)\). In the saline-pretreated rats, there was no significant effect of MDL 11939 injection on the power of HFO (ESM Fig. 2). After the main experiment, four out of eight rats were injected with 1 mg/kg SB 242084 30 min after injection of 2 mg/kg of DOI. Time courses showing the effect of SB 242084 on DOI-induced increases in HFO power are shown in Fig. 3f. Although a trend for a reduction in HFO power was observed, repeated-measures two-way ANOVA did not reveal any significant effect of drug \((F_{1,6} = 0.9, p = 0.378)\) or drug \(\times\) time interaction \((F_{102,612} = 1.26, p = 0.0525)\).

It is interesting to note that reductions in the power of HFO after MDL 11939 administration in the DOI pretreated group was immediate and even dropped below the baseline activity for some time but then recovered back to the baseline activity (see Fig. 3a and c). Due to the attenuation of HFO power after injection of MDL 11939, it was not possible to accurately determine the power of dominant frequency in the power spectra or band-pass filtered signals. However, on occasions when bouts of HFO were present in the LFP the frequency did not appear markedly different to the baseline values.

We also analysed the effect of MDL 11939 administration on DOI-induced reductions in low gamma power \((40–60 \text{ Hz}; \text{ Fig. 3d})\). Repeated-measures two-way ANOVA revealed a significant effect of the time \((F_{107,1498} = 7.59, p < 0.0001)\) but not for the drug \((F_{1,14} = 0.64, p = 0.438)\) on the total power of low gamma. Bonferroni post hoc test revealed a short lasting significant increase in the total power of low gamma after MDL 11939 administration \((p < 0.05, p < 0.001, p < 0.0001)\) compared to the vehicle group. In four rats, we also examined the effect of SB 242084 on the gamma changes induced by DOI and compared this to vehicle values. Time courses showing the effect of SB 242084/vehicle on the reduction in low gamma \((40–60 \text{ Hz})\) induced by DOI are shown in Fig. 3g. Repeated-measures two-way ANOVA did not reveal any significant effect of drug \((F_{1,6} = 0.12, p = 0.743)\) or drug \(\times\) time interaction \((F_{102,612} = 0.64, p = 0.997)\).

Activation of 5HT2A receptors modulate the power and frequency of HFO more powerfully than 5HT2C receptors

We performed an additional study using selective agonists at 5HT2A receptors, TCB-2 \((0.1, 0.5, \text{ and } 1.5 \text{ mg/kg})\) and 5HT2C receptors, CP 809101 \((0.1, 1.0, \text{ and } 3 \text{ mg/kg})\). 5HT2A receptor activation, at the highest dose, led to a reduction in the frequency of HFO in four out of six rats (Fig. 4c). Therefore, the range of HFO power was evaluated over a broader range 110–180 Hz. Time courses showing the effect of TCB-2 and CP 809101 on the total power of HFO are shown in Fig. 4a and b. Repeated-measures two-way ANOVA revealed a significant effect of dose \((F_{3,20} = 7.69, p < 0.0013)\), time \((F_{75,1500} = 9.12, p < 0.0001)\) and interaction \((F_{225,1500} = 4.90, p < 0.0001)\) for TCB-2. Bonferroni post hoc
test revealed a significant increase for the 1.5 mg/kg dose compared to vehicle ($p<0.001$) and the other doses; 0.1 mg/kg ($p<0.001$) and 0.5 mg/kg ($p<0.01$). No other differences were found. Repeated-measures two-way ANOVA for HFO power in the CP 809101 experiment revealed an effect of dose ($F_{3,20}=6.87, p=0.0023$), time ($F_{73,1460}=9.31, p<0.0001$), and interaction ($F_{219,1460}=1.69, p<0.0001$). Bonferroni post hoc test revealed a small but significant increase for the highest dose of CP 809101 compared to vehicle ($p<0.001$) and between 0.1 mg/kg ($p<0.001$) and 1 mg/kg doses ($p<0.05$); in both cases, this was significant for only one time point). No other differences were found.

With respect to low gamma power, time-courses showing the effect of TCB-2 and CP 809101 on the total power of gamma are shown in Fig. 4d and e. Repeated-measures two-way ANOVA of the TCB-2 group revealed a significant effect of the dose ($F_{3,20}=3.06, p=0.0519$) and time ($F_{75,1500}=19.10, p<0.0001$) but not interaction ($F_{225,1500}=0.96, p=0.6526$). Bonferroni post hoc test did not reveal any significant differences in the power of low gamma when compared to vehicle. For the CP 809101 group, no significant drug effects or interactions were found for the dose ($F_{3,20}=0.97, p=0.4255$) and interaction ($F_{219,1460}=1.10, p=0.1717$).

Comparison of serotonergic hallucinogens and ketamine-induced changes in HFO and low gamma in the NAc

At the end of the LSD and DOI, studies the rats received an i.p. injection of 25 mg/kg ketamine (Fig. 5). Consistent with our previous findings, ketamine provoked a large increase in the power of HFO (Hunt et al. 2006, 2008).
Fig. 4  Stimulation of 5HT2A (TCB-2), rather than 5HT2C receptors (CP 809101), increase the power of HFO (110–180 Hz) in the NAc of freely moving rats. a, b Time courses showing the effect of TCB-2 (0.1, 0.5, and 1.5 mg/kg) and CP 809101 (0.1, 1, and 3 mg/kg) on HFO power. Mean powers of HFO (20-min average) after injection are shown alongside. c Individual plots showing the dominant frequency within the HFO band after injection for the highest dose of TCB-2 (green circles) or CP 809101 (orange circles) compared to vehicle (Veh). d, e Time courses showing the effect of TCB-2 and CP 809101 on gamma power. Mean powers of gamma (60-min average) post-injection are shown alongside. Broken lines indicate time of injection. f Placement of electrodes for all the rats, N=6 for TCB-2 group are indicated by black asterisks and N=6 for CP 809101 group are indicated by the black dots. Values expressed as mean±SEM, **p<0.01, ***p<0.001 with respect to vehicle.

Fig. 5  Comparison of injection of ketamine and serotonergic hallucinogens on the power of HFO (130–180 Hz) and low gamma (40–60 Hz) in the NAc. a–d Time courses showing, changes in the total power of HFO and low gamma for all doses of LSD, DOI and 25 mg/kg ketamine (data for LSD and DOI are taken from Figs. 1 and 2). Broken lines indicate time of injection. Note the immediate robust increase in the power of HFO lasting for ~10 min after injection of ketamine.
group 1 (LSD group), we observed a long-lasting enhancement of HFO activity after ketamine, different from the rest of the group, and this rat was excluded from the analysis. Time courses of the total power of HFO after ketamine injection for both groups are shown in Fig. 5a, b. Repeated-measures two-way ANOVA for the LSD and DOI groups including their respective ketamine data revealed significant effects of drug×time interaction (LSD group, \( F_{4,25}=4.47, p<0.0073; \) DOI group, \( F_{4,25}=3.93, p=0.0130 \)), time (LSD group, \( F_{79,1975}=5.61, p<0.0001; \) DOI group, \( F_{79,1975}=12.91, p<0.0001 \)) and interaction (LSD group, \( F_{316,1975}=3.18, p<0.0001; \) DOI group, \( F_{316,1975}=7.15, p<0.0001 \)). Bonferroni post hoc test revealed that the ketamine-enhanced HFO power was significantly greater than that produced by all doses of LSD (\( p<0.001 \)) and DOI (\( p<0.001 \)). Similarly, repeated-measures two-way ANOVA for the TCB-2 and CP 809101 groups including their respective ketamine data revealed significant effects of drug×time interaction (TCB-2 group, \( F_{300,1875}=9.98, p<0.0001; \) CP 809101 group, \( F_{292,1825}=21.36, p<0.0001 \)). Bonferroni post hoc test revealed that the ketamine-enhanced HFO power was significantly greater than that produced by all doses of TCB-2 (\( p<0.001 \)) and CP 809101 (\( p<0.001 \)).

Time courses of the total power of gamma after ketamine injection for both groups are shown in Fig. 5c, d. Analysis of low gamma power, with the ketamine data included, revealed drug×time effects for LSD (\( F_{316,1975}=1.80, p<0.0001 \)) and DOI groups (\( F_{312,1975}=2.60, p<0.0001 \)). The power of low gamma for all doses of LSD (\( p<0.01, \) except the lowest dose) and DOI (\( p<0.001 \)) were significantly lower compared to injection of ketamine. Repeated-measures ANOVA for the TCB-2 and CP 809101 studies also revealed significant drug×time interactions (TCB-2 group, \( F_{300,1875}=2.14, p<0.0001; \) CP 809101 group, \( F_{292,1825}=2.15, p<0.0001 \)). The power of low gamma for all doses TCB-2 and CP 809101 (\( p<0.001 \)) were significantly lower compared to injection of ketamine.

**Discussion**

Serotonergic hallucinogens increase the power of HFO recorded in the LFPs of the NAc

LSD is a partial agonist at several 5HT receptor subtypes and also possesses activity on other receptor systems (Halberstadt and Geyer 2011; Marona-Lewicka et al. 2009). DOI is more selective than LSD at 5HT2 receptors with higher affinity for 5HT2A compared to 5HT2C subtypes (\( k_i=0.7 \) and 2.4 nM for human 5HT2A and 5HT2C receptors, respectively). In our study, both drugs produced small but significant increases in the power of HFO in the LFPs recorded from NAc. To try and elucidate the receptor mechanism involved in the hallucination-induced increase in HFO power, we post-treated rats with MDL 11939, a selective 5HT2A receptor antagonist, which fully reversed the DOI-induced increases in HFO power. TCB-2, a potent 5HT2A receptor agonist, induced a robust increase in the power of HFO, implicating a role for 5HT2A receptors in the increases of HFO power we observed in the accumbal LFPs. We observed a small but significant increase in HFO power at the highest dose of CP 809101, suggesting that 5HT2C receptors might also be involved in the generation of HFO but we did not see any significant reductions in the power of DOI-induced HFO after administration of SB 242084, a selective 5HT2C antagonist. Together, these findings indicate that 5HT2A receptors chiefly modulate increases in HFO power induced by serotonergic hallucinogens.

Here, we focussed on the HFO recorded in LFPs from the NAc as we have observed spontaneous HFO in this region and its enhancement after intracccumbal infusion of MK801 (Hunt et al. 2010). Also, our current source density analyses have demonstrated a localisation of HFO after ketamine in this area (Hunt et al. 2011). Activities in the HFO range have been reported in other brain regions after injection of NMDAR receptor antagonists (Nicolas et al. 2011; Phillips et al. 2012). It would be instructive to determine whether, or not, serotonergic hallucinogens produce similar effects on HFO outside the NAc.

LSD, DOI, and TBC-2 not only increased the power of HFO, but decreased their frequency. We have shown before that NMDAR antagonists increase the frequency of accumbal HFO (Hunt et al. 2006). This shows a qualitative difference between these classes of psychoactive compounds. Moreover, we have shown that antipsychotic drugs, particularly the atypicals, in addition to producing a small increase in HFO power, reduced the frequency of both spontaneous and NMDAR induced HFO in the LFPs recorded from NAc (Olszewski et al. 2012). It should be noted that the reductions in frequency produced by antipsychotics are, in general, much larger than those produced by serotonergic hallucinogens. However, how and why antipsychotic and hallucinatory compounds produce similar changes in HFO power and frequency is at present unclear. The mechanisms that underlie the effects of antipsychotics and serotonergic hallucinogens are almost certainly distinct, since hallucinogen mediated effects on HFO appear to be predominantly 5HT2A activation dependent, while both clozapine and risperidone are antagonists at this receptor (Meltzer et al. 1999).

Serotonergic hallucinogens decrease gamma power in the LFPs recorded from NAc

Recordings from freely moving rats have shown that, within the striatum, gamma oscillations appear more dominant in
ventral regions (Berke et al. 2004; Masimore et al. 2005). The gamma band can be divided into two sub-bands, termed low (50 Hz) and high (80 Hz) gamma (van der Meer et al. 2010). In our study, in the majority of rats, we also observed two discrete bands of gamma activity, corresponding to those reported previously (Berke et al. 2004; Berke 2009; Kalenscher et al. 2010), with the power of low gamma being the more dominant of the two. In the presence of hallucinogens, small bouts of low gamma still occurred but these were far less frequent, compared to vehicle controls, and in many rats we observed almost complete cessation of gamma activity which typically lasted for the entire time course. In this respect, our findings are similar to those of Berke (2009) who demonstrated that systemic injection of amphetamine or apomorphine reduced the power of low gamma in the ventral striatum (see Fig. 3 of Berke (2009) and our study). However, he observed decreased low gamma was also associated with an increase in the power of high gamma. A very recent study examining the effect of methamphetamine on accumbal gamma also demonstrated an increase in the power of high gamma, but reported no significant effects on the power of low gamma (Morra et al. 2012). In contrast to psychostimulants, serotonergic hallucinogens did not increase the absolute power of high gamma in the NAc indicating fundamental differences between these drugs. However, at the highest doses, a band in the high gamma frequency range could be seen in the spectrograms and a corresponding peak in the power spectra. Analysis of the proportion of high gamma in the total gamma range did yield a significant difference for DOI, but not for the LSD group. This indicates that serotonergic hallucinogens have some impact on the high gamma frequency band albeit to a far lesser degree than the low gamma band.

Activation of 5HT2A or 5HT2C receptors alone do not appear to adequately account for the reduction in gamma observed after serotonergic hallucinogens. Firstly, blockade of 5HT2A receptors, produced only a short-lasting reversal of low gamma and secondly, the reductions in gamma were not reproduced fully using the selective 5HT2A agonist TCB-2. However, we should point out that, in our study we found relatively low doses of DOI can produce profound reductions in the power of gamma. Considering that the MDL 11939 studies were carried out in the presence of DOI 2 mg/kg, we cannot exclude the possibility that the short lasting effect was due to excess competition by DOI at the 5HT2A receptor site, displacing the already bound antagonist.

Several in vivo studies have shown that putative fast spiking interneurons, and to a lesser extent medium spiny neurons, can phase lock to either low or high gamma but not both (Berke 2009; Hernandez and Cheer 2012; Kalenscher et al. 2010; van der Meer and Redish 2009). This indicates that the generation of gamma in the NAc may be caused by distinct networks arising from different afferent input or possibly generated by distinct types of interneuron. Cross-structural synchrony of low gamma in the NAc has been observed in the amygdala (Popescu et al. 2009) and ventral tegmental area (Dzirasa et al. 2011) both of which project to the NAc. Indeed, the changes we recorded in the NAc may be secondary to effects on one or more of the accumalb affrent sites. Behaviourally, changes in the power of low gamma in the striatum has been related to reward-associated tasks (Berke 2009; Hernandez and Cheer 2012; Kalenscher et al. 2010; van der Meer and Redish 2009), movement initiation (Masimore et al. 2005), and anxiety-related behaviours (Dzirasa et al. 2011). Consequently, the reduction in low gamma power produced by serotonergic hallucinogens may impair functional coupling between mesolimbic regions leading to behavioural disturbances.

The mechanism of gamma generation in the NAc is only partially understood and there is much debate about the locality of gamma oscillations in the striatum and the extent to which they reflect truly local activity or volume-conducted currents, such as from the piriform cortex (Berke 2005). While a proportion of low gamma recorded in the NAc may originate from the piriform cortex, findings that the firing of both interneurons and medium spiny neurons can cohere with this oscillation indicates that a proportion of low gamma does represent authentic local processing (Berke 2009; Dzirasa et al. 2011; Hernandez and Cheer 2012; Kalenscher et al. 2010; van der Meer and Redish 2009).

Comparison of the changes in accumbal HFO and gamma with NMDAR antagonists

The most obvious difference between serotonergic hallucinogens and NMDAR antagonist-enhanced HFO is mainly quantitative in nature, with NMDAR antagonists producing much greater increases in power. This was borne out in our current study where, at the end, all rats were injected with ketamine 25 mg/kg and we observed several fold greater increases in power compared to the hallucinogens (see Fig. 5). The maximal doses of LSD and DOI used in this study were towards the top end of those typically used by other groups (Bishop et al. 2004; Erdtmann-Vourliotis et al. 1999; Gaggi et al. 1997; Nielsen and Scheel-Kruger 1986). Thus, the relatively weaker effect on HFO power, in comparison to NMDAR antagonists, is unlikely to be related to the doses we used. Indeed, at the medium and high doses of LSD and DOI, we observed clear classical behavioural effects such as wet-dog shakes and postural changes, as reported by others (Gaggi et al. 1997). Even at the lowest dose, we found a clear reduction in gamma power (see Figs. 1e and 2e) demonstrating that the low doses of hallucinogens do affect the electrical activity of the NAc, but that
larger doses are required to influence the power of HFO. Also, the smaller increases in HFO power produced by LSD and DOI cannot be attributed to their partial agonist activities at 5HT receptors, since TCB-2, is a full agonist at 5HT2A receptors and produced a quantitatively similar increase. Thus, hallucinogens do not merely produce a “weakened” increase in HFO power, but rather modulate HFO in a qualitatively and quantitatively distinct manner. MDL 11939 did not influence the power of spontaneous or MK801-enhanced HFO (unpublished observation) indicating that 5HT2A receptors do not underlie the generation of HFO per se, but rather mediate increases in power produced by serotonergic hallucinogens. In freely moving rats, systemic injection of hallucinogens produced a dramatic reduction in the power of low gamma in the NAc, indicating potent effects of these compounds on gamma oscillations. In general, NMDAR antagonists increase the power of gamma in several brain regions, including the hippocampus, cortex and a variety of other areas. A recent study by Nicolas et al. (2011) demonstrated that increases in both low and high gamma occur in several subcortical areas. In contrast in the NAc, a different pattern emerges. We found using bipolar recordings that ketamine produces a small but significant decrease in the power of “broadband” gamma in the NAc, but simultaneously increases the power of gamma in the hippocampus (Hunt et al. 2011). Reductions in gamma power were also found after administration of a moderated dose 0.5 mg/kg of MK801, which produced a sustained decrease in broadband gamma power in the NAc (Hunt et al. 2010). In the current study, we focussed on low gamma and found it to be relatively unaffected after injection of ketamine.

We summarise this aspect of our study as follows: systemic administration of serotonergic hallucinogens and NMDAR antagonists influence the generation of HFO and gamma in the NAc, but with different orders of potency [changes in HFO power: NMDAR antagonists->hallucinogens, changes in gamma power: hallucinogens->NMDAR antagonists].

Conclusions

The NAc has long been considered one of the core structures involved in integration of corticolimbic signalling, in part, due to its anatomical location and ability to gate afferent information. These findings add to a growing body of literature suggesting that the NAc is an important target for the actions of psychoactive compounds. Given the importance of fast oscillations in higher cognitive function and binding alterations, these electrophysiological changes may either impose, or reflect, altered information processing in this structure.

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