

Cannabinoids Prevent the Effects of a Footshock Followed by Situational Reminders on Emotional Processing

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Posttraumatic stress disorder (PTSD) can develop following exposure to a traumatic event. Hence, what we do in the first few hours after trauma exposure may alter the trajectory of PTSD. We examined whether cannabinoids can prevent the effects of a single footshock followed by situational reminders (SRs) on emotional processing. Rats were exposed to a footshock (1.5 mA, 10 s) on day 1 followed by exposure to SRs of the shock on days 3 and 5. The CB1/2 receptor agonist WIN55,212-2 or vehicle were injected intraperitoneally 2 h after the shock. After 1 week, PTSD-like symptoms were examined. Exposure to SRs exacerbated the effects of the shock as rats exposed to shock and SRs, but not shock alone, showed impaired extinction of the traumatic event, impaired plasticity in the hippocampal-accumbens pathway, enhanced latency to startle, and altered expression of CB1 receptors (CB1r) and glucocorticoid receptors (GRs) in the CA1, basolateral amygdala (BLA) and prefrontal cortex (PFC). WIN55,212-2 prevented the effects of the shock and SRs on extinction, plasticity, and startle response. WIN55,212-2 normalized the shock/SR-induced upregulation in CB1r in the PFC, and CA1 and GRs in the CA1, with no effect on BLA downregulation of CB1r and GRs. Shock and SRs caused lasting (1 week) alterations in emotional processing associated with changes in GR and CB1r expression in brain areas related to PTSD. WIN55,212-2 administered after trauma exposure prevented these alterations via PFC- and CA1-CB1r and CA1-GRs.

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INTRODUCTION

Posttraumatic stress disorder (PTSD) is an anxiety disorder that can develop following exposure to traumatic life events. PTSD is different from other psychiatric disorders, in that it has a very clear point of onset. Hence, what we do in the first few hours after trauma exposure may alter the trajectory of PTSD (Zohar *et al*, 2011).

The endocannabinoid system consists of cannabinoid receptors (CB1 and CB2), their endogenous lipid ligands, and the enzymatic machinery for endocannabinoid synthesis and degradation (Kogan and Mechoulam, 2006).

Recent clinical and preclinical evidence suggests that this system is a therapeutic target for the treatment of stress- and anxiety-related disorders such as PTSD (Marsicano *et al*, 2002; Fraser, 2009; Ganon-Elazar and Akirav, 2009, 2012, 2013; Moreira and Wotjak, 2010). We found a relatively broad therapeutic time window in the aftermath of trauma exposure for preventive treatment with cannabinoids. Rats were exposed to the single-prolonged stress (SPS) model of PTSD (restraint, forced swim, and sedation), and after an undisturbed period of 7 days, rats showed impaired extinction, enhanced acoustic startle response (ASR), and

exaggerated negative feedback on the hypothalamic-pituitary-adrenal (HPA) axis (Ganon-Elazar and Akirav, 2012). The CB1/2 receptor agonist WIN55,212-2 (0.5 mg/kg) administered intraperitoneally (i.p.) 2 or 24 h after SPS prevented the trauma-induced alterations in extinction, ASR potentiation, and HPA axis inhibition.

However, it has become increasingly clear that the consequences of exposure to trauma are affected not only by aspects of the event itself, but also by the frequency and severity of trauma reminders. Places and situations are the most frequent trauma reminders (Pynoos *et al*, 1996; Louvart *et al*, 2005; Corral-Frias *et al*, 2013). Hence, in this study, rats were placed in the light side of the light-dark apparatus and once entering the dark side, they received a single inescapable footshock (day 1), followed by two short reexposures to the light side (situational reminders (SRs)) on days 3 and 5 (Pynoos *et al*, 1996; Louvart *et al*, 2005; Corral-Frias *et al*, 2013). We aimed to assess the effects of posttrauma exposure on SRs and to examine whether cannabinoid receptor activation 2 h after exposure to trauma would prevent the effects of the shock and SRs on PTSD-like behavioral, physiological, and biochemical measures.

Impaired extinction of fear memories is thought to contribute to the development and persistence of the unrelenting memories of the trauma and to one of the hallmark symptoms in PTSD, that is, avoidance (Milad *et al*, 2007; Sripada *et al*, 2013). The exposure to trauma and SR model allows measuring avoidance behavior directed at the

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context of the trauma as well as the extinction of the traumatic event itself, and not the extinction of conditioned fear that is not directly associated with the traumatic event, as was previously studied in humans and animal models (Milad *et al*, 2007; Milad *et al*, 2008; Ganon-Elazar and Akirav, 2012, 2013; Sripada *et al*, 2013).

Maladaptive plasticity processes in response to specific external challenges are believed to underline disorders such as PTSD (Nathan *et al*, 2011). Growing attention has been focused on plasticity in the ventral subiculum (vSub)-nucleus accumbens (NAc) pathway (O'Donnell and Grace, 1995; Abush and Akirav, 2010, 2013; Segev *et al*, 2013), as there is growing evidence for a role of the NAc in regulating mood and motivation (for review: Nestler and Carlezon, 2006), and in modulating fear-related behaviors after stress exposure that affects vSub and amygdala inputs to the NAc (Muschamp *et al*, 2011; Gill and Grace, 2013).

Finally, PTSD is associated with alterations in CB1 receptors (CB1r) and glucocorticoid receptors (GRs). Yehuda *et al* (2006) reported on a greater ACTH decline in response to hydrocortisone in PTSD patients, implying that central GRs are more responsive. Neumeister *et al* (2013) found that anandamide concentrations were reduced in PTSD patients relative to non-PTSD patients and suggested that abnormal CB1r-mediated anandamide signaling is implicated in the etiology of PTSD. Hence, we also measured the expression of CB1r and GRs in the fear circuit, which is considered dysfunctional in PTSD (that is, hippocampus, basolateral amygdala (BLA), and prefrontal cortex (PFC); Heim and Nemeroff, 2009) and in the NAc.

MATERIALS AND METHODS

Subjects

Male Sprague–Dawley rats (60 days old, ~250 g; Harlan, Jerusalem, Israel) were caged together at $22 \pm 2^\circ\text{C}$ under 12-h light/dark cycles (lights turned on at 07:00). Rats were allowed water and laboratory rodent chow *ad libitum*. The experiments were approved by the University of Haifa Ethics and Animal Care Committee, and adequate measures were taken to minimize pain or discomfort.

Drug Treatment

WIN55,212-2 (WIN; 0.5 mg/kg) and the CB1 receptor antagonist AM251 (0.3 mg/kg; Cayman Chemicals) were dissolved in 1% dimethylsulfoxide (DMSO), 1% Tween-80, and 98% saline. Controls were given the vehicle (Veh) only. The selective serotonin reuptake inhibitor (SSRI) sertraline hydrochloride (SH; 10 mg/kg; Toronto research chemicals) was dissolved in 5% DMSO, 5% Tween-80, and 90% saline freshly every day. Drugs (vehicle, WIN, AM251, or SH1) were injected *i.p.* 2 h after the shock. An additional group was injected with SH once daily for 7 days (SH7; Matar *et al*, 2006).

Trauma and SRs

Animals were placed in an inhibitory avoidance apparatus (50 cm \times 25 cm \times 30 cm), divided into two equal-size compartments, and separated by an automatic guillotine door.

Except for the sensory-motor sequence tests, every rat underwent one behavioral or electrophysiological test to prevent carryover effects due to multiple behavioral tests.

Shock. On day 1, each rat was placed in the light compartment and after 2 min of exploration, the door was raised allowing access to the dark compartment. Thirty seconds after the rat entered the dark compartment, the door closed and the rat received an inescapable 1.5 mA shock for 10 s. The rat remained in the dark side for an additional 20 s, after which it was returned to the home cage. The no-shock groups received the same treatment, but with the shock mechanism inactivated.

SRs. Rats were placed in the light compartment on days 3 and 5 for 1 min with the gate closed to prevent them from entering the shock compartment (to avoid extinction).

Avoidance and extinction. On day 8, rats were submitted to a non-reinforced test trial every 24 h for 4 days (extinction 1–extinction 4). Each rat was placed in the light side of the box, and the time elapsed until it crossed over to the dark side (that is, latency) was measured. If, after 300 s, the rat did not cross over on its own, the experimenter gently guided it to the dark side. The opening between the two sides of the shuttle was then blocked and the rat was allowed to freely explore the dark side for 180 s, after which it was removed back to the home cage.

Elevated Platform

A different set of rats were placed on an elevated platform (EP; 11 cm \times 11 cm) 80 cm above the ground for 30 min in a brightly lighted room on days 3 and 5. This procedure elicits stress responses in the form of behavioral 'freezing', defecation, and urination, and impairs extinction (Maroun and Akirav, 2008; Ganon-Elazar and Akirav, 2009).

Electrophysiology

Electrophysiology was as described in Abush and Akirav (2013) and Segev *et al* (2013). Briefly, rats were anesthetized (40% urethane, 5% chloral hydrate in saline; 4 ml/1 kg, *i.p.*) and placed in a stereotaxic frame. A recording microelectrode was inserted into the NAc shell (anteroposterior, +1.6 mm; lateral, \pm 1.0 mm; ventral, –5.5 mm) and a bipolar 125 μm stimulating electrode into the vSub (anteroposterior, –6.5 mm; lateral, \pm 5.0 mm; ventral, –6.0 mm). Evoked field potentials (EFPs) were digitized (10 kHz) and analyzed using Cambridge Electronic Design (Cambridge, UK). Offline measurements were made of the amplitude and slope of the EFPs using averages of 30 successive responses to a given stimulation intensity applied at 0.1 Hz. Test stimuli (monopolar pulses, 100 μs duration) were delivered at 0.1 Hz. After positioning the electrodes, the rats were left for 1 h before commencing the experiment.

LTP was induced by high-frequency stimulation (HFS; three sets of 10 trains; each train consisting of 10 pulses at 200 Hz; inter-train interval, 200 ms; inter-set interval, 1 min) to the vSub.

Acoustic Startle Response

Rats were placed in $8 \times 8 \times 16$ cm open animal holder that restricted locomotion but did not immobilize the animal (Coulbourn Instruments). Chambers were calibrated for both sensitivity to movement and sound level to ensure consistency between chambers and experiments. The animals were placed in the holder and allowed a 5-min acclimatization period with background noise only. Following the 5-min acclimatization period, 30 acoustic startle trials (120 dB white noise; 50 ms duration; 20–40 s intertrial interval) were presented over the 68 dB white noise background. Three measures of behavior were assessed: (1) mean startle amplitude; (2) mean latency to reach peak; and (3) an habituation index (a block of the last five responses/ a block of the first five responses).

Sensory-Motor and Anxiety Tests

Sensory-Motor and Anxiety Tests were as described in Abush and Akirav (2010). Briefly, rats were placed in a darkened open field (OF; $40 \times 40 \times 40$ cm) for 5 min. After 24 h, rats were placed in the elevated plus maze (EPM) for 5 min. The EPM consisted of four arms (55 cm each) on a stand 80 cm high. Data for the OF and EPM were analyzed using the behavioral tracking system Ethovision \times T 8.5 (Noldus). After 24 h, pain sensitivity (PS) was assessed by applying continuously ascending mild electric footshock (beginning at 0.0 mA and ending as soon as the animal flinched or vocalized). Two observers scored flinch and vocalization thresholds.

Western Blotting

Rats were killed and brain tissues of the infralimbic PFC, NAc shell, BLA, and CA1 were collected and homogenized in buffer. We concentrated on the IL-PFC as previous studies suggested the IL as involved in the enhancement of extinction learning (Milad and Quirk, 2002; Mueller *et al.*, 2008).

Protein levels were determined by the bicinchoninic acid Protein Assay Kit (Pierce) according to the manufacturer's protocol. The samples were then diluted in SDS sample buffer, boiled (100 °C) for 5 min, and stored at -80 °C. Aliquots were subjected to SDS-PAGE (10% polyacrylamide) and immunoblot analysis. Blots were incubated with the GR/CB1 antibody overnight at 4 °C (1:100, Pierce Antibodies), followed by washing and 1 h incubation with an HRP-linked secondary antibody at room temperature (goat anti-rabbit IgG; Jackson ImmunoResearch Laboratories, 1:10 000). Blots were visualized by enhanced chemiluminescence with ECL (Biological Industries) and quantified with an XRS charge-coupled device camera (Bio-Rad Laboratories) and Quantity One software. All protein samples were standardized with β -actin (1:5000, polyclonal goat antibody; Santa Cruz Biotechnology, USA).

Statistical Analysis

The results are expressed as means \pm SEM. For statistical analysis, one-way ANOVA, two-way ANOVA, mixed design three-way ANOVA, and one-sample *t*-test were used as

indicated. All *post hoc* comparisons were made using the least significant difference multiple-comparison test.

RESULTS

WIN55,212-2 Prevents the Effects of Shock and SRs on Extinction

Mixed design three-way ANOVA (shock \times SR \times extinction day ($2 \times 2 \times 4$)) on the latency to enter the dark side indicated a significant effect for shock ($F_{(1,35)} = 57.43$, $p < 0.001$; Figure 1a), day ($F_{(3,105)} = 16.51$, $p < 0.001$), and the following interactions: shock \times SR ($F_{(1,35)} = 4.28$, $p < 0.05$), day \times shock ($F_{(3,105)} = 11.86$, $p < 0.001$), day \times SR ($F_{(3,105)} = 6.59$, $p < 0.001$), and day \times shock \times SR ($F_{(3,105)} = 5.51$, $p < 0.001$).

Post hoc comparison revealed that on extinction 1, the shock groups demonstrated increased latency compared with no-shock groups ($p < 0.001$) and the shock–no SR group demonstrated increased latency compared with the shock–SR group ($p < 0.05$). On extinction 2–4, the shock–SR group demonstrated increased latency compared with all groups (no shock–no SR: extinction 2: $p < 0.01$, extinction 3–4: $p < 0.001$; no shock–SR: $p < 0.001$; shock–no SR: extinction 2–3: $p < 0.05$, extinction 4: $p < 0.001$). Hence, the shock groups avoided the dark side (extinction 1), but only the shock–SR group demonstrated impaired extinction kinetics.

Next, we examined whether WIN55,212-2 can prevent the effects of trauma and SRs on extinction. A mixed design three-way ANOVA (SR \times drug \times extinction day ($2 \times 2 \times 4$)) indicated significant effects for SRs ($F_{(1,34)} = 9.24$, $p < 0.01$), drug ($F_{(1,34)} = 15.65$, $p < 0.001$), and day ($F_{(3,102)} = 17.27$, $p < 0.001$; Figure 1b) with no significant interactions. Although the interactions were not significant, there was an important effect for drug. Hence, we continued with one-way ANOVA and *post hoc* comparisons on each extinction day. *Post hoc* comparison revealed that on extinction 2, the shock–SR Veh group showed increased latency compared with the shock–no SR WIN group ($p < 0.05$). On extinction 3–4, the shock–SR Veh group showed increased latency compared with all groups (extinction 3: no SR groups: $p < 0.01$, shock–SR WIN: $p < 0.001$; extinction 4: no SR groups: $p < 0.001$, shock–SR WIN: $p < 0.05$). Hence, WIN55,212-2 prevented the effects of shock and SRs on extinction.

As WIN55,212-2 is a CB1/CB2 agonist, we examined whether the preventing effects of WIN55,212-2 on extinction are mediated by the CB1 receptor. A mixed design two-way ANOVA (group \times extinction day (4×4)) on the latency to enter the dark side did not reveal significant effects (Figure 1c). Hence, injecting a low dose of AM251 after the shock had no effect on extinction. AM251 partially prevented the effects of WIN55,212-2 on extinction as there was no significant increase in the latency from extinction 1 to extinction 4 in the shock–SR WIN + AM251 group.

To further validate our model, we added two control experiments. In the first experiment, rats were exposed to the shock, and on days 3 and 5, were exposed to an out-of-context stressor (that is, the EP stress). These rats (shock–EP) were compared with shocked rats that were exposed to SRs (shock–SR) or not (shock–no SR). A mixed design two-way ANOVA (group \times extinction day (3×4)) on the

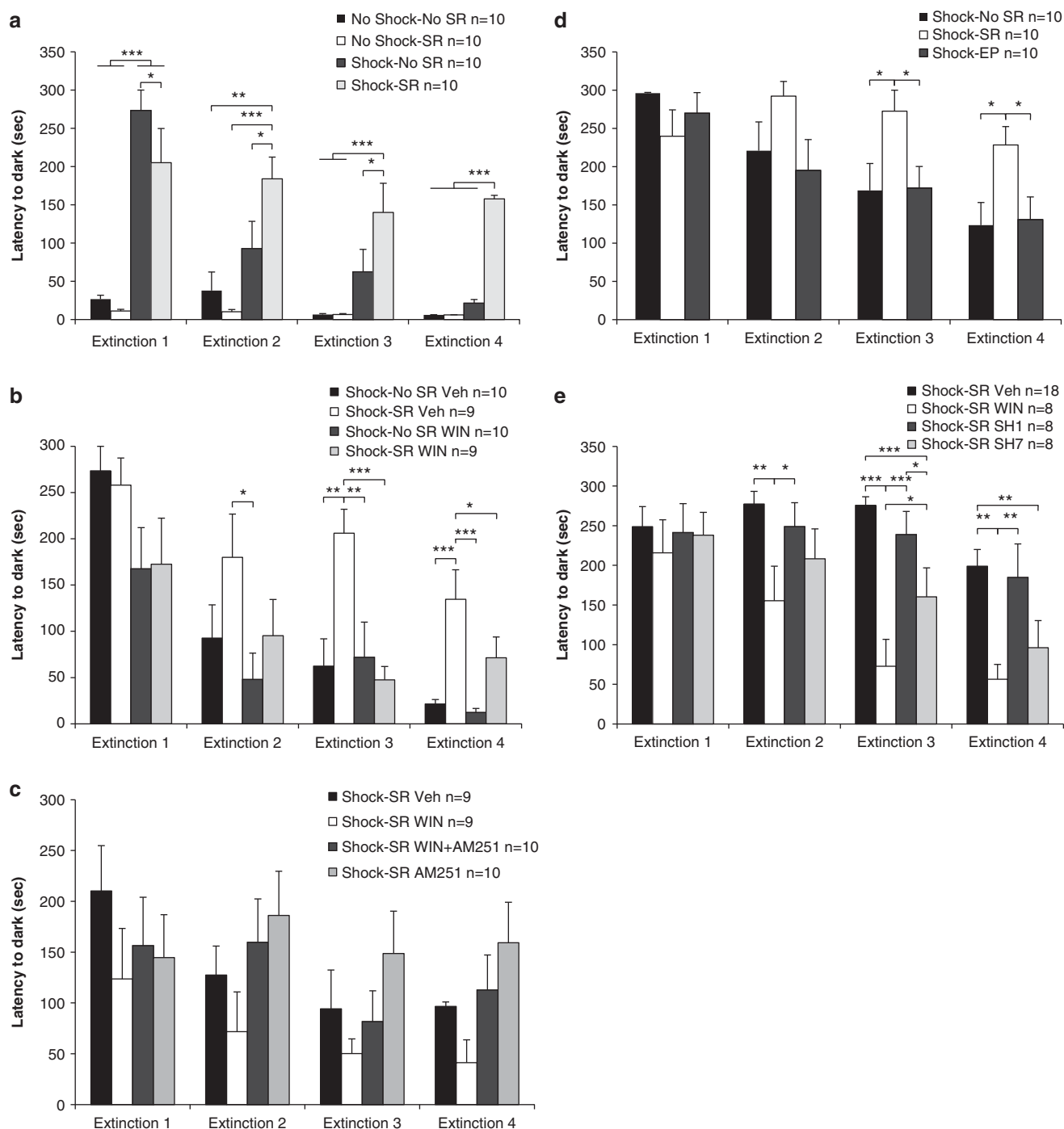


Figure 1 WIN55,212-2 prevents the effects of shock and SRs on extinction (Ext). (a) On Ext 1, the shock groups (shock–no SR, shock–SR) demonstrated increased latency compared with the no-shock groups. On Ext2–4, the shock–SR group demonstrated increased latency compared with all groups ($*p < 0.05$; $**p < 0.01$; $***p < 0.001$). (b) When WIN55,212-2 was injected 2 h after the shock, the shock–SR Veh group showed increased latency compared with the shock–no SR WIN group on Ext2. On Ext3 and 4, the shock–SR Veh group showed increased latency compared with all groups ($*p < 0.05$; $**p < 0.01$; $***p < 0.001$). (c) When the antagonist AM251 was injected, no significant differences were observed between the groups. (d) The shock–SR group demonstrated increased latency to enter the dark side on Ext3 and 4, compared with the shock–no SR group and with rats exposed to shock and placed on the EP on days 3 and 5 after the shock (shock–EP; $*p < 0.05$). (e) Rats were injected with sertraline once (shock–SR SH1) or once daily for 7 days (shock–SR SH7). The shock–SR WIN group demonstrated reduced latency compared with the shock–SR Veh and shock–SR SH1 groups on Ext2, Ext3, and Ext 4. On Ext3, the WIN group also demonstrated increased latency compared with the SH7 group. The SH7 group demonstrated reduced latency compared with the Veh ($p < 0.001$) and SH1 ($p < 0.05$) groups on Ext3 and compared with the Veh group on Ext4 ($*p < 0.05$; $**p < 0.01$; $***p < 0.001$).

latency to enter the dark side revealed a significant effect for group ($F_{(1,27)} = 3.99$, $p < 0.05$; Figure 1d), day ($F_{(3,81)} = 9.56$, $p < 0.001$), and a significant day \times group interaction

($F_{(6,105)} = 2.58$, $p < 0.05$). *Post hoc* comparison revealed that on extinction days 3 and 4, the shock–SR group demonstrated increased latency compared with all groups

($p < 0.05$). Hence, the shock-EP group demonstrated the same extinction kinetics as the shock-no SR group.

In the second control experiment, we tested whether a classic SSRI can prevent the effects of shock and reminders on extinction. Rats were exposed to the shock and injected with SH, which was previously shown to reduce PTSD-like symptoms in rats when injected for 7 days starting after the trauma (Matar *et al*, 2006). Repeated measures ANOVA (drug \times extinction day (4×4)) revealed a significant effect for drug ($F_{(3,38)} = 7.43$, $p < 0.001$), day ($F_{(3,114)} = 14.77$, $p < 0.001$), and a day \times drug interaction ($F_{(9,114)} = 2.17$, $p < 0.05$; Figure 1e). *Post hoc* comparison revealed that on extinction day 2, shock-SR WIN showed reduced latency compared with the shock-SR Veh ($p < 0.01$) and shock-SR SH1 ($p < 0.05$; injected once with SH) groups. On extinction 3, the shock-SR WIN group showed reduced latency compared with all groups (shock-SR Veh SH1: $p < 0.001$; shock-SR SH7: $p < 0.05$). The shock-SR SH7 group (injected for 7 days) showed reduced latency compared with the shock-SR Veh ($p < 0.001$) and shock-SR SH1 ($p < 0.05$) groups. On extinction 4, the shock-SR WIN group showed reduced latency compared with the shock-SR Veh and SH1 groups ($p < 0.01$). The shock-SR SH7 group showed reduced latency compared with the shock-SR Veh group ($p < 0.01$). Hence, treating rats with SH for 7 days after the trauma facilitated extinction compared with vehicle-treated rats. Nonetheless, the effects of SH were not as robust as the effects of WIN55,212-2 on extinction. The acute SSRI treatment had no effect on extinction.

Repeated measures ANOVA (treatment \times extinction day (2×4)) did not reveal differences between the vehicle-treated rats (1 vehicle vs 7 vehicle injections; $F_{(1,16)} < 1$, NS), hence they were grouped together to a control group (shock-SR Veh, $n = 18$).

WIN55,212-2 Prevents the Effects of Shock and SRs on Synaptic Plasticity

Post-HFS analysis using mixed design three-way ANOVA (shock \times SR \times time ($2 \times 2 \times 12$)) on EFPs' amplitude (Figure 2a) and slope (Figure 2b) levels indicated a significant main effect for shock (amplitude: $F_{(1,28)} = 9.78$, $p < 0.01$; slope: $F_{(1,28)} = 11.69$, $p < 0.01$) and SR (amplitude: $F_{(1,28)} = 4.78$, $p < 0.05$; slope: $F_{(1,28)} = 5.13$, $p < 0.05$). *Post hoc* analysis revealed a significant difference between the shock-SR group and the other groups ($p < 0.01$). Hence, exposure to shock and SRs impaired LTP. Mixed design three-way ANOVA on amplitude and slope pre-HFS (drug \times stress \times time ($2 \times 2 \times 5$)) did not reveal any significant effects.

To examine the effects of shock and SRs on baseline synaptic activity, input-output measurements were taken. Analysis revealed a significant effect for stimulation intensity ($F_{(1,28)} = 134.15$, $p < 0.001$). Stimulation of the vSub input into the NAC with different stimulus intensities did not result in any other significant effects on basal EFPs (Figure 2c).

When WIN55,212-2 was injected, post-HFS analysis using mixed design three-way ANOVA (drug \times SR \times time ($2 \times 2 \times 12$)) on EFPs' amplitude (Figure 2d) and slope (Figure 2e) indicated a significant main effect for shock (amplitude: $F_{(1,25)} = 5.37$, $p < 0.05$; slope: $F_{(1,25)} = 4.25$,

$p < 0.05$), SR (amplitude: $F_{(1,25)} = 4.15$, $p < 0.05$; slope: $F_{(1,25)} = 4.05$, $p < 0.05$), and drug \times SR interaction (amplitude: $F_{(1,25)} = 4.48$, $p < 0.05$; slope: $F_{(1,25)} = 4.68$, $p < 0.05$). *Post hoc* analysis revealed a significant difference between the shock-SR Veh group and the other groups ($p < 0.01$). Hence, WIN55,212-2 prevented the impairing effects of the shock and SRs on LTP.

To examine the effects of WIN55,212-2 on baseline synaptic activity, input-output measurements were taken. Mixed design three-way ANOVA on amplitude and slope pre-HFS (drug \times SR \times time ($2 \times 2 \times 5$)) did not reveal any significant effects. Analysis of input-output curve revealed a significant effect for stimulation intensity ($F_{(1,25)} = 104.51$, $p < 0.001$; Figure 2f).

When AM251 was injected, post-HFS analysis using mixed design ANOVA (group \times time (4×12)) on EFPs' amplitude (Figure 2g) and slope (Figure 2h) indicated a significant main effect for group (amplitude: $F_{(3,24)} = 11.46$, $p < 0.001$; slope: $F_{(3,24)} = 15.18$, $p < 0.001$). *Post hoc* analysis revealed a significant difference between the shock-SR WIN group and the other groups ($p < 0.001$). Hence, AM251 prevented the effects of WIN55,212-2 on LTP.

Mixed design three-way ANOVA on amplitude and slope pre-HFS (group \times time (2×5)) did not reveal any significant effects. Analysis of input-output curve revealed a significant effect for stimulation intensity ($F_{(1,24)} = 134.62$, $p < 0.001$) and stimulation intensity \times group interaction ($F_{(3,24)} = 3.33$, $p < 0.05$; Figure 2i).

WIN55,212-2 Prevents the Effects of Shock and SRs on Startle Response

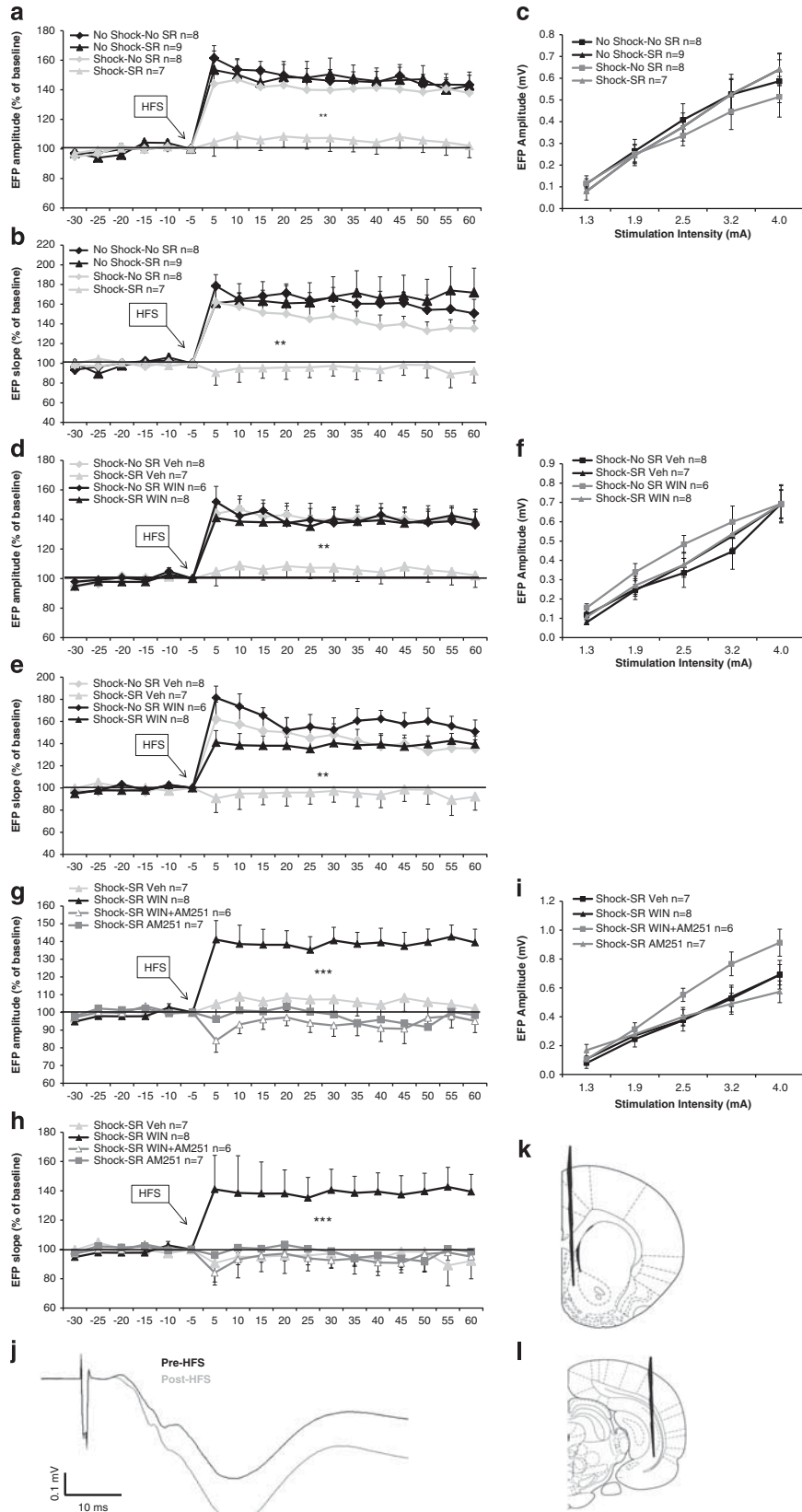
Two-way ANOVA (shock \times SR (2×2)) on the latency to startle revealed a significant effect for SRs ($F_{(1,37)} = 12.69$, $p < 0.001$) and shock \times SR interaction ($F_{(1,37)} = 6.43$, $p < 0.05$; Figure 3a). *Post hoc* comparison revealed that the shock-SR group demonstrated a significant reduction in the latency to startle compared with all groups (no shock-no SR: $p < 0.01$; no shock-SR: $p < 0.05$; shock-no SR: $p < 0.001$).

When WIN55,212-2 was injected, two-way ANOVA (drug \times SR (2×2)) revealed a significant effect for SRs ($F_{(1,36)} = 7.87$, $p < 0.01$) and the interaction drug \times SR ($F_{(1,36)} = 12.07$, $p < 0.001$; Figure 3b). *Post hoc* comparison revealed that the shock-SR Veh group demonstrated a significant reduction in the latency to startle compared with all groups ($p < 0.01$). Hence, WIN55,212-2 prevented the shock/SR-induced decrease in the latency to startle.

When the antagonist AM251 was injected, one-way ANOVA revealed a significant effect for drug ($F_{(3,36)} = 8.76$, $p < 0.001$). *Post hoc* comparison revealed that the shock-SR WIN group demonstrated increased latency to startle compared with all groups ($p < 0.001$; Figure 3c). Hence, AM251 prevented the effects of WIN55,212-2 on the latency to startle.

There were no effects on startle amplitude or the habituation index throughout the experiments, and no correlation was found between these measures and the latency to startle.

To further validate our model, a different set of rats were exposed to the shock, and on days 3 and 5, were exposed to an out-of-context stressor (that is, the EP stress). One-way ANOVA revealed a significant effect for group



($F_{(2,27)} = 3.94, p < 0.05$). *Post hoc* comparison revealed that the shock-SR group demonstrated increased latency to startle compared with all groups ($p < 0.05$; Figure 3d). Hence, the latency to startle in the shock-EP group was similar to the latency observed in the shock-no SR group.

The Effects of Trauma Exposure, SRs, and WIN55,212-2 on Anxiety and Sensory-Motor Measures

In the OF (Table 1, experiment 1), two-way ANOVA (shock \times SR (2×2)) revealed a significant effect for shock on the total distance covered ($F_{(1,35)} = 39.15, p < 0.001$), freezing levels ($F_{(1,35)} = 20.17, p < 0.001$), and time in the center ($F_{(1,35)} = 22.45, p < 0.001$). *Post hoc* comparison revealed that compared with the no-shock groups, the shock groups traveled less distance ($p < 0.001$), demonstrated increased freezing ($p < 0.01$), and avoided the center of the arena ($p < 0.01$).

When WIN55,212-2 was injected (Table 1, experiment 2), one-way ANOVA did not reveal a significant difference between the groups in the distance traveled, freezing levels, or time in the center. Hence, WIN55,212-2 did not prevent the shock/SR-induced increase in anxiety levels.

In the EPM (Table 1, experiment 1), two-way ANOVA (shock \times SR (2×2)) revealed a significant effect for shock on the total distance covered ($F_{(1,36)} = 13.98, p < 0.001$) and the percent of time the rats spent in the open arms ($F_{(1,36)} = 9.98, p < 0.01$). *Post hoc* comparison revealed that the shock groups traveled less distance than the no shock-no SR group ($p < 0.01$), and the shock-no SR group traveled less than the no shock-SR group ($p < 0.01$). Further, the shock groups spent more time in the open arms than the no shock-no SR group ($p < 0.01$).

When WIN55,212-2 was injected (Table 1, experiment 2), one-way ANOVA did not reveal significant differences between the groups.

In the PS test (Table 1, experiment 1), two-way ANOVA (shock \times SR (2×2)) revealed a significant effect for shock in the PS threshold ($F_{(1,37)} = 31.41, p < 0.001$). *Post hoc* comparison revealed that the shock groups demonstrated higher threshold than the no shock-no SR ($p < 0.001$) and the no shock-SR ($p < 0.01$) groups.

When WIN55,212-2 was injected (Table 1, experiment 2), one-way ANOVA revealed a significant difference between the groups ($F_{(3,37)} = 6.45, p < 0.001$). *Post hoc* comparison revealed that the WIN-treated rats demonstrated lower threshold ($p < 0.01$) than the vehicle-treated rats. Hence,

WIN55,212-2 prevented the shock/SR-induced increase in PS threshold.

When AM251 was injected (data not shown), one-way ANOVA revealed a significant difference between the groups ($F_{(3,35)} = 3.88, p < 0.05$). *Post hoc* comparison revealed that the shock-SR WIN group demonstrated lower threshold compared with all groups ($p < 0.01$). Hence, AM251 prevented the effects of WIN55,212-2 on PS threshold.

The Effects of Trauma Exposure, SRs, and WIN55,212-2 on the Expressions of CB1r and GRs

Finally, we aimed to examine whether the preventive effects of WIN55,212-2 after shock and SRs are mediated by CB1r and GR in the fear circuit.

CB1r and GR levels were analyzed using one-way ANOVA in all areas tested. To confirm equal protein loading, the same blots were re-hybridized with antibodies specific for β -actin. No significant difference in β -actin levels was observed between the groups in any of the brain areas tested, suggesting that β -actin levels were not affected by the treatment.

In the *IL-PFC*, a significant group effect was found for CB1r ($F_{(2,27)} = 4.62, p < 0.05$; Figure 4a) but not for GR (Figure 4b). *Post hoc* comparison revealed that the shock-SR Veh group expressed significantly more CB1r than all groups ($p < 0.05$).

In the *BLA*, a significant group effect was found for CB1r ($F_{(2,27)} = 4.46, p < 0.05$; Figure 4d) and GR ($F_{(2,27)} = 6.53, p < 0.01$; Figure 4e). *Post hoc* comparison revealed that the no shock-no SR Veh group expressed more CB1r ($p < 0.01$; $p < 0.05$) and GR ($p < 0.05$; $p < 0.01$) than the shock-SR Veh and the shock-SR WIN groups, respectively.

In the *CA1*, a significant group effect was found for CB1r ($F_{(2,27)} = 4.65, p < 0.05$; Figure 4g) and GRs ($F_{(2,27)} = 5.04, p < 0.05$; Figure 4h). *Post hoc* comparison revealed that the shock-SR Veh group expressed significantly more CB1r ($p < 0.05$) and GRs ($p < 0.01$) than all groups.

In the *NAc*, no significant differences were observed between the groups in CB1r (Figure 4j) and GR expression (Figure 4k).

We added a control experiment in which rats were injected with vehicle or WIN55,212-2 and 1 week later decapitated (with no shock or SRs). We found no differences in the expression of CB1r and GRs between the groups (PFC: CB1r ($t_{(12)} = 0.25, NS$), GR ($t_{(12)} = 0.1, NS$);

Figure 2 The effects of shock, SRs and WIN55,212-2 on LTP. (a) When tested on day 8, the shock-SR group demonstrated significantly reduced amplitude compared with all groups post-HFS (** $p < 0.01$). (b) When tested on day 8, the shock-SR group demonstrated significantly reduced slope compared with all groups post-HFS (** $p < 0.01$). (c) Input-output curve: no significant differences between the groups were found in NAc EFP amplitudes after stimulation of vSub input with different stimulus intensities. (d) When the agonist WIN55,212-2 was injected, the shock-SR Veh group demonstrated significantly reduced amplitude compared with all groups post-HFS (** $p < 0.01$). (e) When the agonist WIN55,212-2 was injected, the shock-SR Veh group demonstrated significantly reduced slope compared with all groups post-HFS (** $p < 0.01$). (f) Input-output curve: no significant differences between the groups were found in NAc EFP amplitudes after stimulation of vSub input with different stimulus intensities. (g) When the antagonist AM251 was injected, the shock-SR WIN group demonstrated significantly increased amplitude compared with all groups post-HFS (** $p < 0.001$). (h) When the antagonist AM251 was injected, the shock-SR WIN group demonstrated significantly increased slope compared with all groups post-HFS (** $p < 0.001$). (i) Input-output curve: no significant differences between the groups were found in NAc EFP amplitudes after stimulation of vSub input with different stimulus intensities. (j) Representative signal trace in the NAc taken before and 1 h after HFS to the vSub. (k) Schematic of coronal sections of the rat brain representing the recording site in the nucleus accumbens shell (coordinates: 1.6 mm anterior, 1 mm lateral, and 5 mm ventral to bregma). (l) Schematic of coronal sections of the rat brain representing the stimulating site in the ventral subiculum (coordinates: 6.5 mm posterior, 5 mm lateral, and 6 mm ventral to bregma).

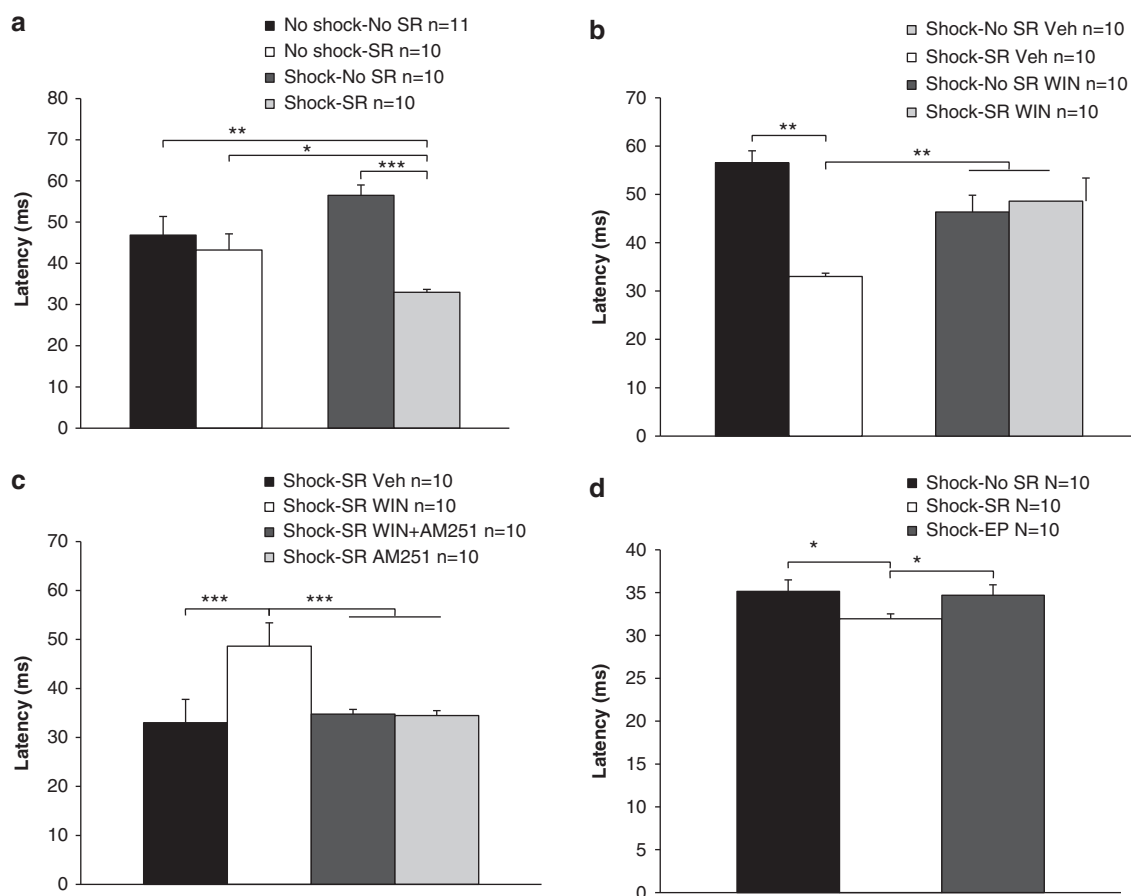


Figure 3 WIN55,212-2 prevents the effects of shock and SRs on startle response. (a) The shock–SR group demonstrated a significant reduction in the latency to startle compared with all groups ($*p < 0.05$; $**p < 0.01$; $***p < 0.001$). (b) When WIN55,212-2 was injected 2 h after the shock, the shock–SR Veh group demonstrated a significant reduction in the latency to startle compared with all groups ($***p < 0.01$). (c) When the antagonist AM251 was injected, the shock–SR WIN group demonstrated increased latency to startle compared with all groups ($***p < 0.001$). (d) The shock–SR group demonstrated increased latency to startle compared with the shock–no SR group and with rats exposed to shock and placed on the EP on days 3 and 5 after the shock (shock–EP) ($*p < 0.05$).

BLA: CB1r ($t_{(12)} = 0.78$, NS), GR ($t_{(12)} = 0.81$, NS); CA1: CB1r ($t_{(12)} = 0.53$, NS), GR ($t_{(12)} = 0.82$, NS); NAc: CB1r ($t_{(11)} = 0.1$, NS), GR ($t_{(12)} = 0.81$, NS).

DISCUSSION

Exposure to shock and SRs resulted in changes in emotional processing observed 1 week after the initial stress. Rats exposed to intense shock and SRs avoided the shock context and showed impaired extinction of the traumatic event, showed impaired plasticity in the vSub–NAC pathway, and enhanced latency to startle. Exposure to shock by itself was not sufficient to provoke all of these changes and rats exposed to shock (with or without the following exposure to SRs) demonstrated avoidance, increased anxiety, and hypoalgesia. This suggests that exposure to SRs exacerbated the effects of the shock on behavior and physiology. Rats exposed to shock and SRs also demonstrated alterations in the expression of CB1r and GRs in brain areas found to be dysfunctional in PTSD (that is, BLA, PFC, and CA1).

Notably, administering the cannabinoid agonist WIN55,212-2 2 h after shock exposure prevented the effects of the shock and SRs on extinction, plasticity, startle response, and PS. These findings are particularly encouraging as the available drugs (for example, SSRIs and serotonin–norepinephrine reuptake inhibitors) provide some benefit in the management of PTSD symptoms, but effect sizes are small and in some cases ineffective (Stein *et al*, 2006).

The agonist WIN55,212-2 also prevented the shock/SR-induced upregulation in CB1r in the PFC and CA1 and upregulation in GRs in the CA1, suggesting that these receptors mediate the therapeutic effects of WIN55,212-2 on behavior and physiology. WIN55,212-2 did not prevent the shock/SR-induced downregulation of CB1r and GRs in the BLA.

In general, a low dose of the CB1 antagonist AM251 administered after the shock had no effect on behavior or plasticity by itself, but when co-administered with WIN55,212-2 it blocked the preventive effects of WIN55,212-2 administered after the shock on plasticity, startle response, and PS. Hence, the preventing effects of

Table 1 The Effects of Shock, SRs and WIN55,212-2 on Sensory-Motor and Anxiety Tests

Group	Open field			Elevated plus maze		Pain sensitivity mA to flinch or vocalization
	Total distance moved (cm)	Freezing (s)	Time in center (% of total)	Total distance moved (cm)	Time in open arms (%)	
<i>Experiment 1:</i>						
No shock–no SR Veh	2317 ± 377	49 ± 8	42 ± 10	1788 ± 101	28 ± 8	0.24 ± 0.01
No shock–SR Veh	2080 ± 330	55 ± 28	38 ± 7	1596 ± 138	44 ± 7	0.26 ± 0.01
Shock–no SR Veh	1387 ± 424**	98 ± 46**	7 ± 2**	1181 ± 102**	59 ± 11**	0.33 ± 0.02***
Shock–SR Veh	1430 ± 434**	111 ± 40**	15 ± 1**	1269 ± 138**	62 ± 5**	0.32 ± 0.01***
<i>Experiment 2:</i>						
Shock–no SR Veh	1387 ± 424	98 ± 46	7 ± 2	1181 ± 102	59 ± 11	0.33 ± 0.02
Shock–SR Veh	1430 ± 434	111 ± 40	15 ± 1	1269 ± 138	62 ± 5	0.32 ± 0.01
Shock–no SR WIN	1308 ± 355	121 ± 28	18 ± 3	1455 ± 139	68 ± 5	0.26 ± 0.01**
Shock–SR WIN	1666 ± 575	91 ± 51	27 ± 7	1139 ± 100	68 ± 6	0.27 ± 0.01**

Abbreviation: SR, situational reminder.

In the OF (experiment 1), the shock groups (shock–SR and shock–no SR) traveled less distance, spent more time freezing, and spent less time in the center than the no-shock groups (no shock–no SR and no shock–SR; ** $p < 0.01$).

When WIN55,212-2 was injected (experiment 2), there was no significant difference between the groups in the distance traveled, freezing levels, or time in the center. In the EPM (experiment 1), the shock groups (shock–SR and shock–no SR) traveled less distance and spent more time in the open arms than the no-shock groups (no shock–no SR and no shock–SR; ** $p < 0.01$).

When WIN55,212-2 was injected (experiment 2), there was no significant difference between the groups in the distance traveled or time spent in the open arms. In the PS test (experiment 1), the shock groups (shock–SR and shock–no SR) were less sensitive to shock intensity than the no-shock groups (no shock–no SR and no shock–SR; *** $p < 0.001$).

When WIN55,212-2 was injected (experiment 2), the WIN groups (shock–SR WIN and shock–no SR WIN) were more sensitive than the vehicle groups (shock–SR WIN and shock–no SR WIN; ** $p < 0.01$).

WIN55,212-2 on behavior and physiology were mostly mediated by CB1r.

However, the blocking effect of AM251 was not observed when extinction was tested. WIN55,212-2 is a nonselective cannabinoid agonist that activates CB1r, CB2r, TRPV1, and so on (Breivogel *et al*, 2001; Pertwee *et al*, 2010). Specifically, CB2r was found to be involved in the consolidation of fear memory (García-Gutiérrez *et al*, 2013), which might explain why the CB1 antagonist AM251 administered after the conditioned shock did not block the effects of WIN55,212-2 on extinction.

We have previously found that AM251 blocked the effects of WIN55,212-2 on inhibitory avoidance and contextual extinction (Ganon-Elazar and Akirav, 2012, 2013). However, here we measured the extinction of the traumatic event itself, and not the extinction of conditioned fear that is not directly associated with the traumatic event (Ganon-Elazar and Akirav, 2012, 2013).

Avoidance and Extinction

Shocked rats exposed to SRs persistently avoided the dark chamber, although they were repeatedly exposed to this chamber under safe conditions (that is, with no shock exposure). Interestingly, shocked rats not exposed to SRs demonstrated increased avoidance on the first extinction trial, but their extinction kinetics was intact. Pamplona *et al* (2011) examined the emergence of avoidance behavior in the aftermath of a trauma in mice and found that fear

incubation after a highly aversive experience induces time-dependent quantitative and qualitative change in avoidance behavior, mirroring fear incubation in human PTSD. Specific conditioned avoidance to shock-paired odor occurred shortly after trauma (2 days) and turned into generalized avoidance after 28 days of fear incubation. They suggested that generalized avoidance results from a mixture of both sensitized and conditioned fear responses and that generalized avoidance reduced through both habituation to the test environment (that is, safety learning) and post-incubation extinction of contextual fear (Pamplona *et al*, 2011). In our study, 1 week after the shock, extinction training reduced avoidance in rats exposed to shock with no SRs, whereas shock–SR rats continued to avoid the dark chamber.

The relationship between fear extinction deficits and avoidance symptoms in PTSD might be bidirectional. Previous studies suggest that extinction deficits can lead to the development of avoidance symptoms, and conversely that pre-existent 'higher avoidance' can be a contributor to extinction deficits. In support of the first hypothesis, greater fear in response to aversive stimuli was associated with greater levels of subsequent avoidance in rats (Chen *et al*, 2012) and pre-trauma deficits in extinction learning were associated with greater risk for developing PTSD after trauma in humans (Lommen *et al*, 2013).

In contrast to previous studies in rats and humans in which the deficits in extinction were to a fear-conditioning paradigm not directly associated with the initial trauma (Milad *et al*, 2007, 2008; Ganon-Elazar and Akirav, 2012,

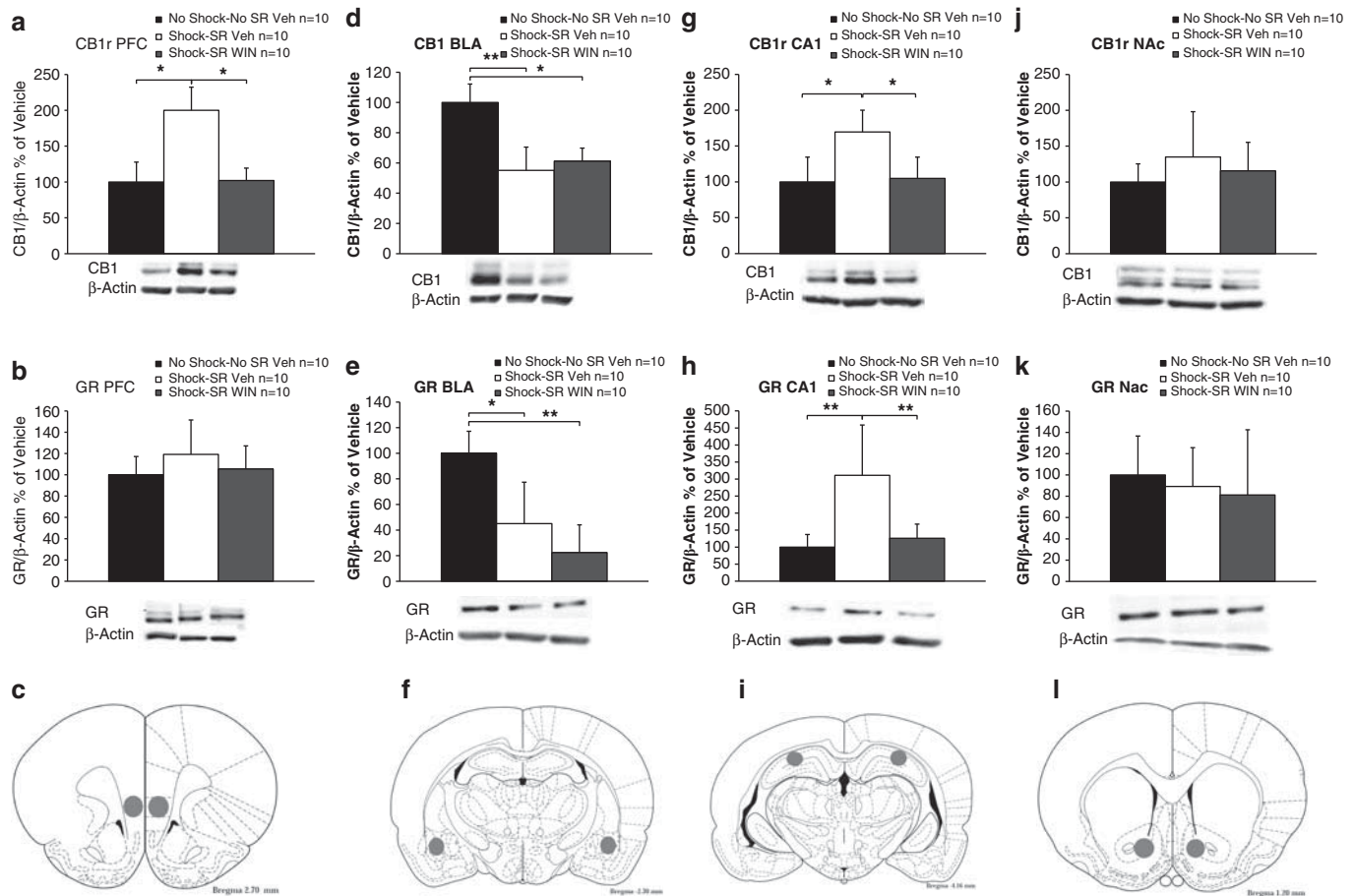


Figure 4 CB1 receptor and GR expression in the fear circuit. Representative bands for the expression of GR/CB1r and β -actin (lower panel). (a) In the IL-PFC, the shock–SR Veh group expressed significantly more CB1r compared with all groups ($*p < 0.05$). (b) In the IL-PFC, no significant difference was observed in GRs between the groups. (c) Brain sites from where the tissue samples were extracted. The IL-PFC was obtained by punches (1 mm diameter) bilaterally. The numbers refer to the distance from Bregma (based on Paxinos and Watson, 1996). (d) In the BLA, the no shock–no SR Veh group expressed more CB1r compared with all groups ($*p < 0.05$; $**p < 0.01$). (e) In the BLA, the no shock–no SR Veh group expressed more GRs compared with all groups ($*p < 0.05$; $**p < 0.01$). (f) The BLA was obtained by punches (1 mm diameter) bilaterally. (g) In the CA1, the shock–SR Veh group expressed significantly more CB1r compared with all groups ($*p < 0.05$). (h) In the CA1, the shock–SR Veh group expressed more GR compared with all groups ($**p < 0.01$). (i) The CA1 was obtained by punches (1 mm diameter) bilaterally. (j) In the NAc, no significant effect was found for CB1r. (k) In the NAc, no significant effect was found for GR. (l) The NAc was obtained by punches (1 mm diameter) bilaterally.

2013; Sripada *et al*, 2013), here extinction training was directed at extinguishing the traumatic event. This is important as abnormalities in extinction may be particularly salient for the persistence of fear memories in PTSD (Milad *et al*, 2008).

We found that exposure to SRs exacerbated the effects of the shock. To further validate the model, we added a control experiment in which rats were exposed to shock on day 1, and then exposed to a stressor on days 3 and 5 (30 min on the EP). Rats exposed to shock and EP (with no SRs) demonstrated increased latency to enter the dark side and reduced latency to startle compared with rats exposed to shock and SRs. Hence, exposure to repeated stress did not exacerbate the effects of the shock as seen with SR exposure. This suggests that the effects of exposure on SRs could not be mimicked by repeated out-of-context stress.

We also added a control experiment in which a ‘classical’ PTSD drug (that is, sertraline) was injected acutely 2 h after the shock (similar to the WIN55,212-2 experiment) or for once daily for 7 days. We found that the acute treatment

had no effect on extinction and that the 1-week treatment facilitated extinction compared with a vehicle-treated group. This corroborates with a previous study demonstrating that administration of sertraline for 7 days immediately post stress exposure (that is, predator scent stress) reduced anxiety-like and avoidant behavior compared with the saline controls (Matar *et al*, 2006). This further validates the trauma and reminders’ model as a PTSD model. Nonetheless, the effects of sertraline, even when administered for 7 days, were not as robust as the effects of WIN55,212-2 on extinction in our model (the sertraline-treated rats showed reduced latency on the third and fourth extinction days).

Plasticity

Shock–SR rats showed impaired NAc LTP 1 week after shock exposure and the agonist WIN55,212-2 prevented this impairment. Recently, we found that exposure to 3 weeks of chronic mild stress impaired NAc LTP tested 48 h, but not 1 week after the stressor ended (Segev *et al*, 2013). Gill and

Grace (2013) demonstrated that following repeated restraint, but not after acute restraint, HFS of the vSub failed to potentiate the vSub-NAc pathway. Hence, it seems that the joint effects of the shock and SRs were significant enough to result in a lasting impairment effect on NAc LTP.

Together with data from the literature, our results suggest that the NAc is embedded within the brain circuit that regulates fear (Schwienbacher *et al*, 2004; Muschamp *et al*, 2011). It has been shown that inactivating the NAc blocked conditioned fear (Schwienbacher *et al*, 2004) and that CREB activation within the NAc produced multiple behavioral signs (anhedonia, impaired extinction) characteristic of experience-dependent psychiatric conditions such as depression and PTSD (Muschamp *et al*, 2011).

Startle Response

Exposure to shock and SRs, but not shock alone, enhanced the latency to startle, suggesting that exposure to SRs exacerbated the effects of the trauma. It is notable that the agonist WIN55,212-2 prevented this shock/SR-induced enhancement because abnormal startle response is a noninvasive translational tool of research that bridges the gap between animal and human investigations and it is a recognized symptom of human PTSD (Grillon and Baas, 2003).

However, evidence is conflicting regarding the question whether the auditory startle reflex is in fact increased in patients with PTSD and other anxiety disorders (Grillon and Baas, 2003). Some studies have shown increased startle responses in PTSD (Grillon *et al*, 1998; Holstein *et al*, 2010), whereas other studies found no difference (Lipschitz *et al*, 2005; Jovanovic *et al*, 2009), or even smaller responses (Ornitz and Pynoos, 1989). Also there are no consistent findings on habituation (Grillon *et al*, 1996; Metzger *et al*, 1999). As far as the increased latency of the startle response, there is evidence that PTSD patients show increased latency to startle (Vrana *et al*, 2013).

Sensory-Motor Tests and Unconditioned Anxiety

In the OF test, shocked rats (with or without SR exposure) demonstrated decreased locomotion, increased freezing, and a decrease in time spent in the center of the OF, indicating enhanced anxiety 1 week after exposure to an intense shock. The agonist WIN55,212-2 did not prevent the shock-induced anxiety, corroborating with a previous study demonstrating that WIN55,212-2 can prevent PTSD-like symptoms (such as enhanced startle response, impaired extinction, and alterations in the HPA axis), but it has no effect on unconditioned anxiety levels (Ganon-Elazar and Akirav, 2012). This suggests that the effects of WIN55,212-2 in preventing PTSD-like symptoms are not due to a general 'relaxation effect' or to an erasure of the stressful event, as rats injected with WIN55,212-2 after the shock still exhibit increased unconditioned anxiety (Ganon-Elazar and Akirav, 2012).

In the EPM, shocked rats also demonstrated decreased locomotion but surprisingly spent more time in the open arms, which usually suggests less anxiety. A possible explanation to this puzzling effect is that shocked rats generalized from the dark chamber where they received the

shock to the dark arms of the EPM, hence resulting in spending more time in the 'safe' open arms. In support, we found that shocked animals demonstrated increased freezing in the open arms compared with nonshocked rats (data not shown). Hence, they stayed in the open arms for longer duration. This generalization effect was observed in other studies as well. Olson *et al* (2011) demonstrated that mice exposed to shock and SRs that were tested in the light-dark test as a second measure of fear response spent less time in the dark chamber than control mice.

In the PS test, shocked rats (with or without SRs) were less sensitive to shock intensity, suggesting an effect that was specifically associated with the shock. Assessment of PS in PTSD patients suggests that PTSD can be associated with both increases and decreases in experimental PS (Asmundson and Katz, 2009; Moeller-Bertram *et al*, 2012). Hypoalgesia was observed in other animal models for PTSD such as SPS (Imanaka *et al*, 2006; Takahashi *et al*, 2006). In human studies, PTSD veterans demonstrated a 30% decrease in reported pain intensity ratings of standardized heat stimuli after the combat videotape suggesting hypoalgesia in response to combat-related cues (Pitman *et al*, 1990).

CB1r and GRs

Fear-conditioning experiments have delineated an amygdala-hippocampal-cortico-striatal circuit as a key brain circuit responsible for processing and storing fear-related memories and for coordinating fear-related behaviors (Neumeister *et al*, 2013).

Shock and SR exposure upregulated CB1r in the PFC and CA1, and WIN55,212-2 normalized this effect. Our data are consistent with other studies showing that CB1r expression is sensitive to stressful experiences, as animals submitted to a fear-conditioning paradigm presented CB1r upregulation in the PFC (Lisboa *et al*, 2010). Campos *et al* (2012) on the other hand have shown that 7 days after a single predator exposure, CB1 mRNA expression was downregulated in the frontal cortex. This discrepancy may result from the different stress paradigms that were used. CB1r are also modulated in the hippocampus by chronic stress; a significant reduction in CB1 receptor binding in the dentate gyrus with a parallel increase in the CA3 region was observed (Hill *et al*, 2009).

Shock and SR exposure also upregulated GRs in the CA1 and WIN55,212-2 prevented this effect. We have previously shown that exposure to the SPS model of PTSD upregulated GRs in the CA1, BLA, and PFC, and WIN55,212-2 administered 2 h after SPS exposure prevented the CA1 and BLA upregulation (Ganon-Elazar and Akirav, 2013). The differences may stem from the different stress paradigm used; the SPS paradigm involves 2 h of restraint followed by forced swim and sedation, and then the animals are left undisturbed for 1 week. In the current paradigm, an intensive footshock is applied, followed by SRs that exacerbate the negative effects of the shock.

In the BLA, shock and SR exposure downregulated CB1r and GRs, and WIN55,212-2 did not normalize this effect. It is possible that exposure to an intense shock caused immediate changes in the BLA, such as increased release of endocannabinoids and glucocorticoids that may result in lasting downregulation of CB1r and GRs in this brain area. In support,

it has been shown that glucocorticoids released during stress lead to an increase in endocannabinoids in the amygdala (Hill *et al*, 2005) and that exposure to shock or tone that was associated with shock increased endocannabinoid release in the amygdala (Marsicano *et al*, 2002; Morena *et al*, 2012). It should be noted that WIN55,212-2 did not normalize the unconditioned anxiety observed in the OF test, suggesting that these effects are mediated by the BLA.

In the NAc, no alterations in CB1r or GRs were observed. The NAc integrates limbic and cortical inputs arising from monosynaptic glutamatergic projections that originate in the vSub, BLA, and PFC (O'Donnell and Grace, 1995; French and Totterdell, 2003). Each of these regions is believed to supply a different mode of input to the NAc, with the BLA involved in affective responses, the vSub in context dependency, and the medial PFC in behavioral flexibility.

Studies reported a low level or even lack of CB1r in the NAc, whereas other brain regions, such as the PFC, the hippocampus, and the amygdala, which densely innervate the NAc, show moderate to very high CB1r levels (Tsou *et al*, 1997). Nevertheless, it has been argued that CB1-expressing neurons in the NAc, although sparse, appear to be critical for emotional and motivational responses. Specifically, it has been suggested that manipulation of CB1 signaling within the NAc triggers robust emotional/motivational alterations related to drug addiction and other psychiatric disorders, and that these effects cannot be exclusively attributed to CB1 located at afferents to the NAc (Winters *et al*, 2012). Taken together, this might explain why we have robust effects on plasticity in the vSub-NAc pathway, but not on the expression of CB1r in this area.

A similar picture emerges with GRs; GRs are highly expressed in the CA1 and moderately expressed in the BLA and PFC, whereas the NAc shell expresses low density (Ahima and Harlan, 1990). It has been suggested that the BLA, hippocampus, and PFC are likely candidates for conveying the indirect effects of cannabinoids on dopamine release within the NAc, thereby contributing to reward processes (Katona *et al*, 2001).

Studies indicate a bidirectional, functional relationship between glucocorticoids and the endocannabinoid system (for review, see Akirav, 2013) and this interaction can modulate memory consolidation of emotionally arousing experiences (Campolongo *et al*, 2009; Hill and McEwen, 2009; Ganon-Elazar and Akirav, 2009, 2013; Atsak *et al*, 2012; de Oliveira Alvares *et al*, 2010; de Bitencourt *et al*, 2013; Segev *et al*, 2013).

Stress and glucocorticoids can trigger endocannabinoid synthesis to constrain HPA axis activity under acute conditions in the brain's fear circuit (that is, BLA, hippocampus, and medial PFC) (Marsicano *et al*, 2002; Rademacher *et al*, 2008; de Oliveira Alvares *et al*, 2010; Hill *et al*, 2011; Wang *et al*, 2011). In the hippocampus and PFC, the recruitment of endocannabinoids by glucocorticoids likely increases the outflow of the principal neurons of these regions to contribute to termination of the stress response (de Oliveira Alvares *et al*, 2010; Hill *et al*, 2011). In the BLA, the endocannabinoids bind to presynaptic CB1r on GABAergic terminals, which rapidly suppress the release of GABA. One model suggests that inhibition of GABA release disinhibits norepinephrine release and increases norepinephrine activation of postsynaptic β -adrenoreceptors,

increasing the consolidation of emotionally aversive memories (Campolongo *et al*, 2009; Hill and McEwen, 2009; Atsak *et al*, 2012). Another model suggests that inhibition of GABA release in BLA interneurons reduces their inhibition of the GABAergic neurons of the intercalated nuclei, which, in turn, increases their inhibition of the pyramidal neurons of the central amygdala (Katona *et al*, 2001). The end result of the reduction in inhibitory tone may be reduced HPA axis activity and attenuation of the effects of stress on memory (Katona *et al*, 2001; Akirav, 2013).

Taken together, it can be suggested that in our study, endocannabinoids release was not sufficient to constrain HPA axis activity after exposure to shock and SRs, and administering WIN55,212-2 compensated for this reduced constrain and prevented the effects of shock and SRs on extinction, startle, and plasticity via CB1r in the PFC, and CB1r and GRs in the CA1. In previous studies, we found that intra-BLA WIN55,212-2 prevented the effects of stress on emotional memory (Ganon-Elazar and Akirav, 2009, 2012); however, in this study, WIN55,212-2 was injected systemically and this probably results in a different effect on emotional processing than a specific intra-BLA injection. Another explanation is that the effects of an intense shock on amygdala processing are different than the effects of exposure to other stressors that are not associated with shock or fear (that is, the SPS model). In support, it has been shown that restraint stress decreased (Hill *et al*, 2009), whereas shock or tone associated with the shock increased (Marsicano *et al*, 2002; Morena *et al*, 2012) anandamide levels in the amygdala.

In conclusion, shock and SR exposure in rats caused lasting alterations in emotional processing associated with changes in GR and CB1r expression in brain areas dysfunctional in PTSD. WIN55,212-2 administered after trauma exposure prevented these alterations via CB1r in the PFC, and CA1 and GRs in the CA1. This suggests that the connectivity in the fear circuit is altered following trauma and CB1r activation and thus may result in lasting effects on emotional processing.

FUNDING AND DISCLOSURE

The authors declare no conflict of interest.

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