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Variations in environmental aversiveness influence emotional memory processes in rats. We have previously shown that cannabinoid effects on memory are dependent on the stress level at the time of training as well as on the aversiveness of the environmental context. Here, we investigated whether the hippocampal endocannabinoid system modulates memory retrieval depending on the training-associated arousal level. Male adult Sprague Dawley rats were trained on a water maze spatial task at two different water temperatures (19°C and 25°C) to elicit either higher or lower stress levels, respectively. Rats trained under the higher stress condition had better memory and higher corticosterone concentrations than rats trained at the lower stress condition. The cannabinoid receptor agonist WIN55212-2 (10–30 ng/side), the 2-arachidonoyl glycerol (2-AG) hydrolysis inhibitor JZL184 (0.1–1 μg/side), and the anandamide (AEA) hydrolysis inhibitor URB597 (10–30 ng/side) were administered bilaterally into the hippocampus 60 min before probe-trial retention testing. WIN55212-2 or JZL184, but not URB597, impaired probe-trial performances only of rats trained at the higher stressful condition. Furthermore, rats trained under higher stress levels displayed an increase in hippocampal 2-AG, but not AEA, levels at the time of retention testing and a decreased affinity of the main 2-AG-degrading enzyme for its substrate. The present findings indicate that the endocannabinoid 2-AG in the hippocampus plays a key role in the selective regulation of spatial memory retrieval of stressful experience, shedding light on the neurobiological mechanisms involved in the impact of stress effects on memory processing.

Key words: cannabinoid receptors; emotional arousal; endocannabinoids; memory for emotional experiences; stress

Significance Statement
Endogenous cannabinoids play a central role in the modulation of memory for emotional events. Here we demonstrate that the endocannabinoid 2-arachidonoylglycerol in the hippocampus, a brain region crucially involved in the regulation of memory processes, selectively modulates spatial memory recall of stressful experiences. Thus, our findings provide evidence that the endocannabinoid 2-arachidonoylglycerol is a key player in mediating the impact of stress on memory retrieval. These findings can pave the way to new potential therapeutic intervention for the treatment of neuropsychiatric disorders, such as post-traumatic stress disorder, where a previous exposure to traumatic events could alter the response to traumatic memory recall leading to mental illness.

Introduction
Extensive evidence indicates that the endocannabinoid system regulates cognitive function (Wotjak, 2005; Campolongo et al., 2009a; Kano et al., 2009; Marsicano and Lafenêtre, 2009; Akirav, 2011). Endocannabinoids, mainly N-arachidonoyl ethanolamine (anandamide, AEA) and 2-arachidonoyl glycerol (2-AG), are synthesized on demand and travel retrogradely to presynaptic sites to bind G-protein-coupled cannabinoid receptors type 1

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The endocannabinoid system plays a key role in the control of emotional responses to environmental challenges (Zanettini et al., 2011; Ruehle et al., 2012; Gunduz-Cinar et al., 2013; Morena and Campolongo, 2014). We have previously reported that cannabinoids induce different effects on memory depending on the aversiveness of the environmental conditions and the stress level at the time of training (Campolongo et al., 2013). Further, we have recently demonstrated that endocannabinoids are physiologically released in limbic brain regions only when animals are trained under high arousal conditions (Morena et al., 2014). Variations in environmental aversiveness differentially influence spatial memory processes in rats (Sandi et al., 1997; Akirav et al., 2004; Salehi et al., 2010). Salehi et al. (2010) induced different stress levels by training rats on a radial-arm water maze task at three different water temperatures (25°C, 19°C, and 16°C). An inverted-U shape effect was found, with animals trained at 19°C showing better memory than animals trained at either the higher (16°C) or lower (25°C) stress condition (Salehi et al., 2010). Once memories are consolidated, the efficacy or accuracy of the information retrieved is vulnerable to stress effects at the time of recall. Considerable evidence points to the hippocampus as a brain structure of crucial importance for retrieval of spatial memory (Hirsh, 1974; Moser and Moser, 1998; Holt and Maren, 1999; Riedel et al., 1999; Eldridge et al., 2000; Brun et al., 2002; Matus-Amat et al., 2004). Stress or exogenous glucocorticoid administration typically impairs memory retrieval of contextual/spatial information in rats and declarative information in humans (de Quervain et al., 1998; Roozaaandael et al., 2003, 2004a; de Quervain et al., 2009). The endocannabinoid 2-AG in the dorsal hippocampus plays an intermediary role in mediating glucocorticoid effects on the impairment of contextual memory retrieval (Atsak et al., 2012).

Here, we investigated whether the hippocampal endocannabinoid system is differentially involved in regulating spatial memory retrieval of rats trained at two experimental conditions that differed with respect to their environment-associated stress levels. Rats were trained on a water maze spatial task at two different water temperatures (19°C and 25°C) to elicit either a higher or lower level of stress. Before retention testing, we pharmacologically manipulated the endocannabinoid system by administering the cannabinoid receptor agonist WIN55212-2, the 2-AG hydrolysis inhibitor JZL184, or the AEA hydrolysis inhibitor URB597 directly into the dorsal hippocampus. Behavioral experiments were paralleled by biochemical measurement of plasma corticosterone levels, hippocampal 2-AG and AEA content, and the cannabionid hydrolytic enzymatic machinery.

Materials and Methods

Animals. Male adult Sprague Dawley rats (320–370 g at the time of behavioral experiments; Charles River Breeding Laboratories) were housed individually in a temperature-controlled (20 ± 1°C) vivarium room and maintained under a 12 h/12 h light/dark cycle (7:00 A.M. to 7:00 P.M. lights on). Food and water were available ad libitum. Training and testing were performed during the light phase of the cycle between 10:00 A.M. and 4:00 P.M. All experimental procedures were in compliance with the European Union Directive on the protection of animals used for scientific purposes (2010/63/EU), the Italian law (D.L. 26/2014), the Declaration of Helsinki, and the Guide for the Care and Use of Mammals in Neuroscience and Behavioral Research (National Research Council, 2004).

Surgery. The rats were anesthetized with sodium pentobarbital (50 mg/kg, i.p.) and given atropine sulfate (0.4 mg/kg, i.p.) to maintain respiration. Subsequently, they were injected with 3 ml of saline (s.c.) to facilitate clearance of these drugs and prevent dehydration. The rats were then positioned in a stereotaxic frame (David Kopf Instruments), and two stainless-steel guide cannulae (23 gauge; 11-mm-long) were implanted bilaterally with the cannula tips 1.5 mm above the CA1 region of the dorsal hippocampus (coordinates: anteroposterior, −3.4 mm; mediolateral, ±1.8 mm; dorsoventral, −2.7 mm) according to the atlas of Paxinos and Watson (2005). The cannulae were affixed to the skull with two anchoring screws and dental cement. Stylets (11-mm-long 00 insect dissection pins) were inserted into each cannula to maintain patency. After surgery, the rats were retained in an incubator until recovered from anesthesia and were then returned to their home cages. Rats were allowed to recover from surgery for 10 d before training and were handled 1 min per day for 3 d before training.

Water maze spatial task and experimental procedures. The experimental apparatus was a circular black galvanized tank, 1.83 m in diameter and 0.6 m in height, filled with water to a depth of 20 cm. The maze was located in a room containing many salient, visual, extra-maze cues. A rectangular platform (20 cm × 25 cm) was placed at a fixed location 25 cm away from the edge of the pool and 2.5 cm below the water surface. A slightly modified procedure of those described by Akirav et al. (2004) and Campolongo et al. (2009b) was used. Separate groups of animals were trained at a water temperature of either 19°C or 25°C, which were previously shown to elicit different stress levels and cognitive performance in rats (Akirav et al., 2004; Salehi et al., 2010). On each day of training, the rats were carried from the vivarium to the experimental room, and training began 120 min later. For spatial training, the rats were given four trials on each daily session for two consecutive days. This relatively small number of trials was chosen such that the animals acquired the task but that their retention performance was moderate to avoid any potential ceiling or floor effects. Before the first training trial, each rat was placed directly onto the submerged platform for 15 s. For each trial, the rat was placed into the tank at one of the four designated starting points and allowed to find the platform. If an animal failed to find the platform within 60 s, it was manually guided to the platform. After mounting the platform, the rat was allowed to remain there for 10 s and was then placed into a holding cage for 25 s until the start of the next trial. The time each rat spent searching for the platform was recorded as the escape latency. Retention of the spatial training was assessed 24 h after the last training session with a 60 s free-swim probe trial under equal temperature conditions (22 ± 0.5°C). During the probe trial, the platform was removed from the tank, and a new starting position was used. Training and probe
trials were videotaped, and an automated tracking system (Smart-B5, Panlab S.L.U., Barcelona, Spain) was used to analyze the swim path of each subject and calculate several corresponding dependent measures, such as time spent in the quadrant containing the platform during training (target quadrant), time spent in the quadrant opposite to the target quadrant (opposite quadrant), initial latency to cross the platform location, number of crossings through the platform location, and total swim distance. The target and opposite quadrants were equidistant from the starting position on the probe trial.

**Drug treatment.** The CB1 receptor agonist WIN55212-2 (10 or 30 ng in 0.5 μl), the 2-AG hydrolysis inhibitor IZL184 (0.1 or 1 μg in 0.5 μl), or the AEA hydrolysis inhibitor URB597 (10 or 30 ng in 0.5 μl) was administered into the dorsal hippocampus 60 min before the probe trial. To examine whether WIN55212-2 and IZL184 effects were mediated via an activation of CB1 receptors, other groups of rats were administered the effective dose of WIN55212-2 (10 ng in 0.5 μl) or IZL184 (1 μg in 0.5 μl) together with the CB1 receptor antagonist AM251 at a dose nonaltering memory performance per se (0.28 ng). Doses were selected on the basis of previous and pilot experiments performed in our laboratory (Morena et al., 2014). All drugs were dissolved in a vehicle containing 5% polyethylene glycol, 5% Tween 80, and 90% saline. Bilateral infusions of drugs or an equivalent volume of vehicle into the dorsal hippocampus were made by using a 30-gauge injection needle connected by polyethylene tubing (PE-20) to a 10 μl Hamilton microsyringe driven by a minipump (KD Scientific). The injection needle protruded 1.5 mm beyond the tip of the cannula, and a 0.5 μl injection volume per hemisphere was infused over a period of 50 s (Roozendaal et al., 2004b; Atsak et al., 2012). The injection needles were retained within the cannulae for an additional 20 s after drug infusion to maximize diffusion and to prevent backflow of drug into the cannulae. All drugs were kindly donated by the National Institute of Mental Health, and drug solutions were freshly prepared before each experiment.

**Histology.** The rats were anesthetized with an overdose of sodium pentobarbital (100 mg/kg, i.p.) and perfused transcardially with 0.9% saline. The brains were then removed and immersed in a 4% formaldehyde solution. At least 48 h before sectioning, the brains were transferred to a 20% sucrose solution in saline for cryoprotection. Coronal sections of 35 μm were cut on a cryostat, mounted on gelatin-coated slides, and stained with cresyl violet. The sections were examined under a light microscope (Microscope Nikon 801), and determination of the location of infusion needle tips within the hippocampus was made according to the standardized atlas plates of Paxinos and Watson (2005) by an observer blind to drug treatment condition. For all experiments, only rats with needle tips terminating within the CA1 region of the hippocampus were included in the data analysis. Approximately 5% of the animals were excluded because of either a missed target or damage to the targeted tissue.

**Plasma corticosterone levels.** Plasma corticosterone levels were determined in parallel groups of rats handled for 3 d and subsequently trained at a water temperature of either 19°C or 25°C. Control animals were handled for 3 d but were not trained or tested. Rats were euthanized 20 min after the two training sessions or immediately after the probe trial. After decapitation, trunk blood was collected and samples were centrifuged at 1900 × g for 20 min at 4°C. Plasma was stored at −20°C and analyzed for corticosterone using an ELISA kit (Assay Designs) according to the manufacturer’s instructions. The kit sensitivity and detection limit were 1.86 and 16.9 pg/ml, respectively. The CV values for intra-assay and interassay precision are 5.2% and 8.7%, respectively.

**Endocannabinoid extraction and analysis.** To examine whether 2-AG and AEA are physiologically released into the hippocampus during memory retrieval, separate groups of nonoperated rats were trained at the two different water temperatures (19°C or 25°C) and euthanized immediately after the probe trial. Control animals were handled but not trained or tested. After rapid decapitation, hippocampi were rapidly dissected and stored at −80°C. The lipid extraction process was performed by using a slightly modified procedure of that described by Dincheva et al. (2015). Brain tissue was weighed and placed into borosilicate glass culture tubes containing 2 ml of acetonitrile with 5 pmol of [3H]AEA and 5 nmol of [3H]2-AG for extraction, and homogenized with a glass rod. Tissue was sonicated for 30 min on ice water and incubated overnight at −20°C to precipitate proteins, then centrifuged at 1500 × g to remove particulates. The supernatants were transferred to a new glass tube and evaporated to dryness under N₂ gas. The samples were reconstituted in 300 μl of acetonitrile and dried again under N₂ gas. Lipid extracts were suspended in 20 μl of acetonitrile and stored at −80°C until analysis. Analysis of AEA and 2-AG was performed by liquid chromatography mass spectrometry analysis as previously detailed (Dincheva et al., 2015).

**Membrane preparation.** Immediately after the probe trial, following rapid decapitation, the hippocampi were dissected from nonoperated rats trained at a water temperature of 19°C or 25°C and from control rats that were handled but not trained or tested. Brain samples were stored at −80°C. All membranes were collected by homogenization of frozen tissue in 10 volumes of TME buffer (50 mM Tris HCl, pH 7.4; 1 mM EDTA, and 3 mM MgCl₂) (Hill et al., 2009). Homogenates were then centrifuged at 18,000 × g for 20 min, and the resulting crude membrane fraction-containing pellet was resuspended in 10 volumes of TME buffer. Protein concentrations were determined using the Bradford method (Bio-Rad). Membranes were used for MAGL and FAAH activity assays.

**MAGL activity assay.** MAGL activity was measured by conversion of 2-oleoylglycerol labeled with [3H] (1-[3H] 2-OG) in the glycerol portion of the molecule to [3H] glycerol preaparations. A slightly modifed procedure of that described by Rademacher et al. (2008) was used. Membranes were incubated in a final volume of 0.5 ml TME buffer (50 mM Tris-HCl, 3.0 mM MgCl₂, 1.0 mM EDTA, and 300 mM URB597, pH 7.4) that contained 1.0 mg/ml fatty acid-free BSA and 100,000 dpm [3H] 2-OG. Isotherms were constructed using six concentrations of 2-OG at concentrations between 10 and 500 μM. Incubation was performed at 30°C, and the enzymatic reaction was stopped by the addition of 2 ml of chloroform/methanol (1:2). After remaining at room temperature for 30 min with frequent mixing, 0.67 ml of chloroform and 0.6 ml of water were added, and the aqueous and organic phases were separated by centrifugation at 1000 rpm for 10 min. The amount of [3H] in 0.5 ml of each of the aqueous and organic phases was determined by liquid scintillation counting and conversion of [3H] 2-OG to [3H] glycerol was calculated. The binding affinity (𝐾m) and maximal hydrolytic activity (𝑉max) values for this conversion were determined by fitting the data to a single-site Michaelis–Menten equation using GraphPad Prism (GraphPad).

**FAAH activity assay.** FAAH activity from hippocampal membranes was measured by conversion of AEA labeled with [3H] in the ethanolamine portion of the molecule to [3H] ethanolamine preparations as reported previously (Hill et al., 2009). The 𝐾m of AEA for FAAH and 𝑉max of FAAH for this conversion were determined by fitting the data to the Michaelis–Menten equation using GraphPad Prism (GraphPad).

**Statistics.** Data are mean ± SEM. Water maze training data were analyzed with one- or two-way ANOVAs with training day as repeated measure. Probe-trial retention data were analyzed with unpaired t tests, one-, two-, or three-way ANOVAs, when appropriate. Plasma corticosterone levels, hippocampal endocannabinoid content, and MAGL and FAAH activity parameters were analyzed with unpaired t tests or one-way ANOVAs. Tukey–Kramer post hoc tests were used to determine the source of the detected significances, when appropriate. p values of <0.05 were considered statistically significant. The number of rats per group is indicated in the figure legends.

**Results**

**Learning under stress enhances probe-trial retention performance**

We first sought to determine whether the different water temperatures at training influenced spatial memory performance of vehicle-treated rats.

Repeated-measures ANOVA for escape latencies to find the hidden platform during training, before drug treatment, revealed a significant effect of training day (𝐹(1,74) = 203.03, 𝑝 <
by significantly longer search times in the vicinity of the platform location (i.e., target quadrant) than in the opposite quadrant \((p < 0.01; \text{Fig. 1D})\). However, rats trained with the lower water temperature spent significantly less time in the opposite quadrant than rats trained with the higher water temperature \((p < 0.01; \text{Fig. 1D})\). No group differences were found for total swim distance on the probe trial \((t_{(174)} = 0.35; p = 0.73; \text{data not shown})\), indicating that the two groups did not differ in motor performance.

**Water maze training under the two stress conditions differentially affects plasma corticosterone levels**

To assess whether the two water temperatures differentially affected plasma corticosterone levels, parallel groups of rats were trained at a water temperature of either 19°C or 25°C, and trunk blood was collected 20 min after either day of training or immediately after the probe trial. Home cage control rats were only handled. As shown in Table 1, one-way ANOVA for plasma corticosterone levels 20 min after the first or second training day revealed a significant experimental condition effect \((\text{training day 1: } F_{(2,20)} = 28.47; p < 0.0001; \text{training day 2: } F_{(2,19)} = 21.41; p < 0.0001)\). Rats trained with either the higher or lower water temperature had higher corticosterone levels than home cage control rats \((p < 0.01 \text{ for both comparisons})\). More importantly, rats trained with the lower water temperature had higher plasma corticosterone levels than those trained with the higher water temperature \((p < 0.05)\), implying that the lower water temperature represented a higher stress condition. As expected, one-way ANOVA for plasma corticosterone levels measured immediately after the 60 s probe trial did not reveal any experimental condition effect \((F_{(2,22)} = 1.29; p < 0.30; \text{Table 1})\) because it is well documented that plasma corticosterone peaks 20 min after the stress onset. Thus, at the time of memory retrieval, both groups of animals presented the same levels of plasma corticosterone.

The cannabinoid receptor agonist WIN55212-2 impairs probe-trial retention performance of rats trained at the higher stress condition

This experiment investigated whether the cannabinoid receptor agonist WIN55212-2 (10 or 30 ng in 0.5 μl) infused into the dorsal hippocampus 60 min before the probe trial would affect retention performance and whether WIN55212-2 effects depend on the stress condition.

**19°C water temperature**

All animals progressively learned to locate the platform across the training sessions, before drug treatment, as indicated by decreasing

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**Figure 1.** Effect of water temperature on training and probe performances of vehicle-treated rats. **A**, Rats trained at either 19°C or 25°C acquired the behavioral task as indicated by a significant difference between mean escape latencies during Training 1 and Training 2. Rats trained at 19°C showed a better memory performance than rats trained at 25°C during Training 2. **\(^{**}p < 0.01\) versus Training 1. **\(^{**}p < 0.01\) mean escape latencies at 19°C versus mean escape latencies at 25°C. **B–D**, The higher stress condition (19°C) enhanced probe-trial retention performance in rats. Rats trained at the higher stress condition (19°C) spent less time to first cross the platform location (B) and performed a higher number of target crossings (C) during probe compared with the group trained under the lower stress condition (25°C). **D**, Both experimental groups spent more time in the target quadrant than in the opposite quadrant in searching of the platform location. Rats trained at the higher stress condition spent less time in the opposite quadrant than rats trained at the lower stress condition. **E**, Illustration of rat coronal sections showing infusion needle termination sites of the rats included in the final analyses. **F**, Representative probe-trial swim paths of the 19°C and the 25°C groups. **\(^{**}p < 0.01\) versus 25°C group. **\(^{**}p < 0.01\) versus the correspondent opposite quadrant. **\(^{**}p < 0.05\) versus opposite quadrant time of the 25°C group. Data are mean ± SEM \((n = 36–40 \text{ per group})\).
mean escape latencies as training progressed ($F_{1,31} = 140.88, p < 0.0001$; data not shown). Figure 2A–C shows that WIN55212-2 administered into the dorsal hippocampus 60 min before the probe trial impaired retention performance of rats trained at this higher stress condition. One-way ANOVAs for crossing latencies and target crossings during the probe trial revealed a significant WIN55212-2 effect (crossing latencies: $F_{2,31} = 3.53, p = 0.04$; target crossings: $F_{2,31} = 5.78, p = 0.007$; Figure 2A,B). Post hoc analyses indicated that the 10 ng dose of WIN55212-2, but not the higher dose, significantly increased crossing latencies ($p < 0.05$; Fig. 2A) and, conversely, decreased the number of crossings through the platform location ($p < 0.01$; Fig. 2B). Two-way ANOVA for quadrant search times during the probe trial showed a significant quadrant effect ($F_{1,62} = 109.23, p < 0.0001$), no WIN55212-2 effect ($F_{1,62} = 9.23, p = 0.56$), and a significant interaction between both factors ($F_{2,62} = 7.37, p = 0.001$). Post hoc analysis revealed that all experimental groups spent more time in the target quadrant than in the opposite quadrant ($p < 0.01$; Fig. 2C), indicating that all groups exhibited memory of the platform location during training. However, WIN55212-2 significantly reduced the time spent searching for the platform location in the target quadrant ($p < 0.05$; Fig. 2C). No WIN55212-2 effect was found for total swim distance during the probe trial ($F_{2,31} = 0.46; p = 0.63$; data not shown).

To investigate whether the CB1 receptor mediates the impairing effect of WIN55212-2, the CB1 receptor antagonist AM251 (0.28 ng in 0.5 µl) was administered together with the effective dose of WIN55212-2 (10 ng) into the dorsal hippocampus 60 min before the probe trial. As shown in Figure 2E, F, two-way ANOVAs for crossing latencies and the number of target crossings during the probe trial revealed a significant main effect of WIN55212-2 (crossing latencies: $F_{1,44} = 8.99; p = 0.004$; target crossings: $F_{1,44} = 5.37; p = 0.03$), no AM251 effect (crossing latencies: $F_{1,44} = 3.43; p = 0.07$; target crossings: $F_{1,44} = 2.81; p = 0.10$), and a significant interaction between WIN55212-2 and AM251 (crossing latencies: $F_{1,44} = 5.59; p = 0.02$; target crossings: $F_{1,44} = 7.22; p = 0.01$). Tukey’s post hoc comparisons showed that WIN55212-2 alone significantly increased the initial time to cross the platform location ($p < 0.01$; Fig. 2E) and decreased the number of target crossings ($p < 0.01$; Fig. 2F). These effects were mediated by CB1 receptor activation because initial crossing latencies of rats given WIN55212-2 together with AM251 were significantly shorter than those of rats given WIN55212-2 alone ($p < 0.05$; Fig. 2E) and equivalent to those of rats given either vehicle or AM251 alone. Consistently, the number of target crossings of rats given WIN55212-2 together with AM251 was significantly higher than those of rats given WIN55212-2 alone ($p < 0.05$; Fig. 2F) and equivalent to those of rats given vehicle or AM251 alone. Three-way ANOVA for quadrant search times during the probe trial revealed no main effect of WIN55212-2 ($F_{1,88} = 1.18; p = 0.28$) but significant AM251 ($F_{1,88} = 4.95; p = 0.03$), quadrant ($F_{1,88} = 191.75; p < 0.0001$), and WIN55212-2 × AM251 × quadrant interaction effects ($F_{1,88} = 4.20; p = 0.04$; Fig. 2G). Rats administered vehicle into the hippocampus spent significantly more time in the target quadrant than in the opposite quadrant ($p < 0.01$; Fig. 2G). As in the previous experiment, WIN55212-2 infused into the hippocampus 60 min before the probe trial reduced the time spent in the target quadrant ($p < 0.01$; Fig. 2G). Administration of AM251 alone into the hippocampus did not affect the time spent in either the target or opposite quadrant, but AM251 blocked the effects induced by WIN55212-2. Rats given combined infusions of AM251 and WIN55212-2 spent significantly more time in the target quadrant than rats given WIN55212-2 alone ($p < 0.01$; Fig. 2G). Two-way ANOVA for total swim distance did not reveal a significant WIN55212-2 ($F_{1,44} = 0.006; p = 0.94$), AM251 ($F_{1,44} = 0.34; p = 0.56$), or WIN55212-2 × AM251 interaction effect ($F_{1,44} = 0.13; p = 0.72$; data not shown).

### 25°C water temperature

WIN55212-2 infused into the hippocampus 60 min before the probe trial did not affect retention performance of rats trained at this lower stress condition (Fig. 2I–K). One-way ANOVAs for initial latency to cross the platform location and the number of crossings through the platform location did not reveal a significant WIN55212-2 effect ($F_{1,32} = 0.50, p = 0.62; F_{1,32} = 0.52, p = 0.60$; respectively, Figure 2I,J). Two-way ANOVA for quadrant search times during the probe trial showed a significant quadrant effect ($F_{1,64} = 59.30, p < 0.0001$) but no WIN55212-2 effect ($F_{1,64} = 0.13, p = 0.88$) or interaction between both factors ($F_{2,64} = 0.65, p = 0.52$; Fig. 2K). Also, no group differences were found for total swim distance during the probe trial ($F_{1,32} = 0.77; p = 0.47$; data not shown).

### 19°C water temperature

All animals progressively learned to locate the platform across the two training sessions, before drug treatment, as indicated by decreasing mean escape latencies as training progressed ($F_{1,26} = 119.11, p < 0.0001$; data not shown). Similar to the effects induced by WIN55212-2, we found that the 2-AG hydrolysis inhibitor JZL184 impaired probe-trial retention performance of rats trained at this higher stress condition. One-way ANOVAs for crossing latencies and target crossings during the probe trial revealed a significant JZL184 effect (crossing latencies: $F_{2,26} = 4.28, p = 0.03$; target crossings: $F_{2,26} = 5.99, p = 0.007$; Figure 3A,B). The 1 µg dose of JZL184, but not the lower dose, significantly increased crossing latencies ($p < 0.05$; Fig. 3A), and both doses of JZL184 decreased the number of crossings through the platform location (both, $p < 0.05$; Fig. 3B). As shown in Figure 3C, two-way ANOVA for quadrant search times during the probe trial showed a significant quadrant effect ($F_{2,52} = 59.30, p < 0.0001$), no JZL184 effect ($F_{2,52} = 0.17, p = 0.84$), and a significant interaction between both factors ($F_{2,52} = 7.05, p = 0.002$). Post hoc analysis indicated that both control rats and rats treated with the lower dose of JZL184 (0.1 µg) spent significantly more time in the target quadrant than in the opposite quadrant ($p < 0.05$; data not shown).
thus indicating that they remembered the platform location. This difference was not found in animals treated with the $1/100$ dose of JZL184 (Fig. 3C), indicating that they did not exhibit memory for the platform location. JZL184 did not affect total swim distance during the probe trial ($F_{(2,26)} = 2.44; p = 0.11$; data not shown).

The next experiment examined whether the JZL184 effect on probe-trial performance is mediated by CB1 receptor activation. To this aim, the CB1 receptor antagonist AM251 (0.28 ng/0.5 μl) and the effective dose of JZL184 (1 μg) were infused together into the dorsal hippocampus 60 min before retention testing. Figure 3E–G shows the effect of JZL184 together with AM251 on probe-trial performance. Figure 2. Intrahippocampal infusions of the cannabinoid agonist WIN55212-2 (WIN) impaired probe-trial retention performance, and the CB1 receptor antagonist AM251 (0.28 ng/0.5 μl) blocked the memory impairing effect induced by WIN (10 ng/0.5 μl; WIN 10), only in the higher stress condition trained group (19°C). In contrast, intrahippocampal infusions of WIN did not affect spatial memory performance at the lower stress condition (25°C). WIN effects on the latency to cross the platform location (A), the number of target crossings (B), and the time spent in the target and opposite quadrants (C) during the 1 min probe trial at 19°C. D. Illustration of rat coronal sections showing infusion needle termination sites of WIN-treated rats and their vehicle control group at the higher stressful condition included in the final analyses. Effects of WIN + AM251 administration on the latency to cross the platform location (E), the number of target crossings (F), and the time spent in the target and opposite quadrants (G) during the 1 min probe trial at 19°C. H. Illustration of rat coronal sections showing infusion needle termination sites of WIN-, WIN + AM251-, AM251-treated rats and their vehicle control group at the higher stressful condition included in the final analyses. WIN effects on the latency to cross the platform location (I), the number of target crossings (J), and the time spent in the target and opposite quadrants (K) during the 1 min probe trial at 25°C. L. Illustration of rat coronal sections showing infusion needle termination sites of WIN-treated rats and their vehicle control group at the lower stressful condition included in the final analyses. M, Representative probe-trial swim paths. * $p < 0.05$ versus the correspondent vehicle (Veh) group. ** $p < 0.01$ versus the correspondent vehicle (Veh) group. *** $p < 0.01$ versus the correspondent opposite quadrant. $p < 0.05$ versus target quadrant time of vehicle group. $p < 0.05$ versus the correspondent WIN + AM251 group. ** $p < 0.01$ versus the correspondent WIN + AM251 group. Data are mean ± SEM ($n = 9–14$ per group).
trial retention performance. Two-way ANOVAs for the crossing latencies and the number of target crossings revealed significant JZL184 (crossing latencies: $F_{(1,41)} = 9.56; p = 0.004$; target crossings: $F_{(1,41)} = 5.90; p = 0.007$), AM251 (crossing latencies: $F_{(1,41)} = 8.24; p = 0.002$; target crossings: $F_{(1,41)} = 4.28; p = 0.04$), and JZL184 × AM251 interaction effects (crossing latencies: $F_{(1,41)} = 9.79; p = 0.003$; target crossings: $F_{(1,41)} = 10.55; p = 0.002$). JZL184 alone significantly increased the initial latency to cross the platform location ($p < 0.01$; Fig. 3E) and reduced the number of target crossings ($p < 0.01$; Fig. 3F). Initial crossing latencies of rats given JZL184 together with AM251 were significantly shorter than those of rats given JZL184 alone ($p < 0.01$;...
Fig. 3) and equivalent to those of rats given either vehicle or AM251 alone. Consistently, the number of target crossings of rats given JZL184 and AM251 was significantly higher than those of rats given JZL184 alone (p < 0.01; Fig. 3F) and equivalent to those of rats given vehicle or AM251 alone. Three-way ANOVA for time spent in the target and opposite quadrant revealed no main effects of JZL184 (F(1,82) = 0.76; p = 0.39) or AM251 (F(1,82) = 1.52; p = 0.22) on quadrant search times during the probe trial (Fig. 3G). However, there was a significant quadrant effect (F(1,82) = 141.89; p < 0.0001) and JZL184 × AM251 × quadrant interaction effect (F(1,82) = 7.18; p = 0.009). Control rats spent significantly more time in the target quadrant than in the opposite quadrant (p < 0.01; Fig. 3G). JZL184 decreased the time spent in the target quadrant (p < 0.05; Fig. 3G). AM251 alone did not affect the time spent in either the target or opposite quadrant but blocked the effects induced by JZL184. Rats treated with AM251 and JZL184 spent significantly more time in the target quadrant (p < 0.01; Fig. 3G) than rats given JZL184 alone and did not differ significantly from those given either vehicle or AM251 alone. Two-way ANOVA for total swim distance did not reveal a significant JZL184 effect (F(1,41) = 0.008; p = 0.93), AM251 (F(1,41) = 1.69; p = 0.20), or JZL184 × AM251 interaction effect (F(1,41) = 1.21; p = 0.28; data not shown).

25°C water temperature

JZL184 administered into the hippocampus 60 min before the probe trial did not affect memory retention of rats trained at this lower stress condition (Fig. 3L–K). One-way ANOVAs did not reveal a significant JZL184 effect for crossing latencies (F(2,36) = 0.24; p = 0.79; Fig. 3J) or target crossings (F(2,36) = 0.37; p = 0.69; Fig. 3J). Two-way ANOVA for quadrant search times indicated a significant quadrant effect (F(1,72) = 47.31, p < 0.0001) but no JZL184 effect (F(2,72) = 0.45; p = 0.64) or interaction between both factors (F(2,72) = 0.19; p = 0.83; Fig. 3K). No group effects were found for total swim distance (F(2,36) = 0.008; p = 0.99; data not shown).

As the effectiveness of JZL184 in rats remains controversial, with some studies reporting unchanged brain 2-AG levels following systemic or local administration of JZL184 (Long et al., 2009; Wiskerke et al., 2012; Kerr et al., 2013) and others studying both biochemical and behavioral effects (Scioliino et al., 2011; Oleson et al., 2012; Seillier et al., 2014), we performed an additional experiment to relate the behavioral effects of JZL184 with any change in endocannabinoid levels. Two parallel groups of rats, handled for 3 d, were bilaterally administered, on day 4, with the behavioral effective dose of JZL184 (1 μg in 0.5 μl) or vehicle into the dorsal hippocampus and killed 60 min later for hippocampal dissection. As shown in Table 2, JZL184 significantly increased 2-AG levels compared with controls (t(15) = 2.41, p = 0.03) but did not affect AEA levels (t(15) = −59, p = 0.56).

The AEA hydrolysis inhibitor URB597 does not alter probe-trial retention performance

We next examined whether an increased endogenous tone of AEA would modulate probe-trial retention performance. To this aim, the AEA hydrolysis inhibitor URB597 was infused bilaterally into the hippocampus 60 min before the probe trial of rats trained with the two water temperatures.

19°C water temperature

Repeated-measures ANOVA for escape latencies to find the hidden platform on the two training days revealed a significant effect of training day (F(1,28) = 84.23, p < 0.0001; data not shown), confirming that rats progressively learned to locate the platform. Figure 4A–C shows that URB597 infused into the hippocampus 60 min before the probe trial did not affect retention of rats trained with this higher stress condition. One-way ANOVA did not reveal any significant drug effect for crossing latencies (F(2,28) = 0.36, p = 0.70; Fig. 4A) or target crossings during the probe trial (F(2,28) = 0.62, p = 0.55; Fig. 4B). Two-way ANOVA for quadrant search times during the probe trial showed a significant quadrant effect (F(1,56) = 82.64, p < 0.0001), but no URB597 (F(2,56) = 0.084, p = 0.92), or quadrant × URB597 interaction effect (F(2,56) = 1.56, p = 0.22). Post hoc analysis revealed that all experimental groups spent more time in the target quadrant than in the opposite quadrant (p < 0.01; Fig. 4C). Also, no group differences were found for total swim distance during the probe trial (F(2,28) = 0.52; p = 0.60; data not shown).

25°C water temperature

URB597 infused into the hippocampus 60 min before the probe trial also did not affect retention of rats trained at the lower stress condition (Fig. 4E–G). One-way ANOVA indicated no significant URB597 effect for initial latency to cross the platform location (F(2,33) = 0.08, p = 0.93; Fig. 4E) or for the number of crossings through the platform location (F(2,33) = 0.69, p = 0.51; Fig. 4F). Two-way ANOVA for quadrant search times during the probe trial showed a significant quadrant effect (F(1,66) = 112.69, p < 0.0001), but no URB597 effect (F(2,66) = 0.16, p = 0.86) or interaction between both factors (F(2,66) = 2.35, p = 0.10; Fig. 4G). No group differences were found for total swim distance (F(2,66) = 0.053; p = 0.95; data not shown).

Water maze training at the higher stress condition increases hippocampal 2-AG levels during probe-trial retention testing

This experiment investigated whether the different stress levels associated to the training procedures induced any alteration in endocannabinoid 2-AG and/or AEA hippocampal levels at the time of retention testing. Parallel groups of rats were trained at a water temperature of either 19°C or 25°C and euthanized immediately after the probe trial for brain tissue dissection. Home cage control rats were handled only. As shown in Figure 5A, one-way ANOVA for hippocampal 2-AG levels indicated a significant experimental condition effect (F(2,24) = 4.68; p = 0.02). Rats trained at 19°C had higher 2-AG levels immediately after the probe trial than rats trained at 25°C or home cage control rats (p < 0.05 for both comparisons; Fig. 5A). Conversely, one-way ANOVA for hippocampal AEA levels did not reveal any group differences (F(2,25) = 0.94; p = 0.40; Fig. 5B).

Water maze training at the higher stress condition alters hippocampal MAGL activity during probe-trial retention testing

This experiment investigated whether the different stress levels associated to the training procedure (19°C or 25°C water temperature) induced any alteration in hippocampal MAGL and FAAH activity at the time of the retention testing. Rats were euthanized immediately after the probe trial for hippocampal tissue dissection.
As shown in Figure 5C, one-way ANOVA for $V_{\text{max}}$ of MAGL did not reveal any significant experimental condition effect ($F_{(2,8)} = 1.12; p = 0.37$). Interestingly, one-way ANOVA for $K_{m}$ of MAGL indicated a significant experimental condition effect ($F_{(2,8)} = 7.04; p = 0.02$; Fig. 5D). Post hoc comparison indicated that rats trained at the higher stress condition had an increase in $K_{m}$ value for MAGL compared with those trained at the lower stress condition or home cage control rats ($p < 0.05$, for both comparisons), indicating that this higher stress condition induced a decrease in the affinity of MAGL for its substrate. No experimental condition effect was found for the $V_{\text{max}}$ ($F_{(2,15)} = 0.85; p = 0.45$; Fig. 5E) or $K_{m}$ ($F_{(2,15)} = 0.28; p = 0.76$; Fig. 5F) of FAAH.

Discussion

The present findings show that the hippocampal endocannabinoid 2-AG selectively modulates spatial memory performance of higher stressful training experience. Indeed, we found that bilateral intrahippocampal infusions of the cannabinoid receptor agonist WIN55212-2 or the 2-AG hydrolysis inhibitor JZL184, but not the AEA hydrolysis inhibitor URB597, impaired spatial memory performance of animals trained under a higher stressful condition (19°C water temperature) in a spatial water maze paradigm, without inducing any effect in animals trained at the lower stressful condition (25°C water temperature). As animals were injected 60 min before memory retention testing, in the time window during which the training performance was retrieved, we demonstrated that boosting 2-AG signaling selectively impaired spatial memory retrieval in rats trained under the higher stress condition through the activation of hippocampal CB1 receptors. Interestingly, vehicle-treated animals trained at the higher stress condition had a better retention (probe) memory performance than animals trained under lower stress condition. Furthermore, only the higher stress training procedure induced an increase in hippocampal 2-AG levels and a reduction of MAGL affinity for its substrate, when assessed at the time of memory retrieval, without altering either AEA levels or FAAH activity in the hippocampus. These findings are in line with evidence reporting increased 2-AG levels after repeated exposure to stressful experiences in limbic brain regions, including the hippocampus (Patel et al., 2005b, 2009; Rademacher et al., 2008; Hill et al., 2010; Dubreucq et al., 2012). Several lines of evidence indicate that the signaling capacity of 2-AG becomes progressively increased under conditions of repeated stress, where it is thought to mediate changes in synaptic transmission and neural excitability (Patel et al., 2005b, 2009; Patel and Hillard, 2008; Hill et al., 2010; Sumislawski et al., 2011; Dubreucq et al., 2012). The present findings add to this notion by demonstrating an increase in 2-AG levels after repeated exposure to the higher stressful training condition and an augmented behavioral response to the increased 2-AG signaling. The mechanisms by which stress increases the signaling capacity of 2-AG remain unknown, deserving further investigation; however, within the amygdala, it has been demonstrated that, following exposure to repeated stress, MAGL expression translocates from the membrane to the cytosol where it would less efficiently catabolize 2-AG (Sumislawski et al., 2011).

Extensive cognitive and neurobiological research on animals, healthy human subjects, and amnesic patients indicates that the hippocampus is an important brain region involved in memory retrieval (Hirsh, 1974; Moser and Moser, 1998; Riedel et al., 1999; Eldridge et al., 2000; Squire et al., 2001; Brun et al., 2002; Matus-Amat et al., 2004) and represents a primary target for stress hor-
mones (Reul and de Kloet, 1985). It is well known that stress and glucocorticoids can facilitate learning performance (Akirav et al., 2001; Salehi et al., 2010) and enhance memory consolidation (Oitzl and de Kloet, 1992; de Kloet et al., 1999; Roozendaal, 2000; Joëls and Baram, 2009). Accordingly, we found higher plasma corticosterone levels than rats trained at 25°C and lower than animals trained at 16°C (Sandi et al., 1997; Akirav et al., 2001, 2004; Salehi et al., 2010). This facilitation of spatial learning by stress, occurring in a water maze at low water temperature, has been shown to be coupled with increased hippocampal synaptic expression of the GluA2 AMPA receptor subunit (Conboy and Sandi, 2010), which has a critical role in controlling various properties of AMPA receptors and in promoting dendritic spine formation and growth (Saglietti et al., 2007) occurring during memory consolidation (Diamond et al., 2006). Interestingly, the increased expression of GluA2 subunits is corticosterone-dependent, and preventing corticosterone release under the higher stress condition also interferes with the stress-induced facilitation of memory (Conboy and Sandi, 2010).

Our findings also show that intrahippocampal infusions of WIN55212-2 or JZL184 impair spatial memory retrieval of rats trained under the higher stress condition and that this effect is mediated by the activation of CB1 receptors. Conversely, the same pharmacological manipulations do not induce any effect on memory of rats trained under the lower stress condition. Therefore, we demonstrate that 2-AG is not only involved in the regulation of this hippocampus-related memory process but also that it can selectively modulate memory retrieval of high stressful experiences. In line with our findings, intrahippocampal infusions of WIN55212-2 have been reported to impair memory retrieval of high arousing test situations (Piri and Zarindast, 2011; Segev and Akirav, 2011; Atsak et al., 2012). A pharmacologically induced enhancement of hippocampal 2-AG exerts a comparable effect (Atsak et al., 2012), whereas exogenous AEA administration in the same brain region has been shown to not alter memory retrieval performance (De Oliveira Alvares et al., 2008).

The finding that stimulating hippocampal CB1 receptors, directly with WIN55212-2 or indirectly with JZL184, impairs memory retrieval only of rats trained at the higher stress condition suggests that there are different circuits implicated in the processing of spatial information depending on the level of stress associated to the learning experience. Previous studies have suggested that spatial learning in a water maze under cold water activates the BLA, whereas increased activation of the hippocampus is detected following spatial learning under warm water (Akirav et al., 2001; Kogan and Richter-Levin, 2008). It is well known that intra-amygdales processes promote memory formation of stressful experiences in other brain regions, including the hippocampus (Roozendaal and McGaugh, 1997; McGaugh, 2000; Packard and Wingard, 2004; Morena et al., 2014). Moreover, it has been demonstrated that, once established, the spatial memory trace formed under high stress condition (i.e., water maze at 19°C) is stronger compared with the trace formed under lower levels of stress (Sandi et al., 1997; Kogan and Richter-Levin, 2010; Salehi et al., 2010). However, this strong memory trace is more complex, thus more vulnerable to interference (Kogan and Richter-Levin, 2010). The dual effect of emotionality may reflect the relative contribution of different neuronal circuits and brain regions recruited to process stressful information (Adolphs et al., 1997;
Hamann et al., 1999; Richardson et al., 2004; Smith et al., 2006). Our findings show that a physiological increase of hippocampal 2-AG levels, likely due to a reduction in the binding affinity of 2-AG for its primary degrading enzyme, selectively promotes the retrieval of stressful experience in nontreated animals. However, following an exogenous manipulation, when the endocannabinoid system is further stimulated with WIN55212-2 or JZL184, memory performance is impaired. It is likely that the memory formation of the stressful experience may have recruited neurons in different circuits, changed the location, and/or altered the functionality of CB1 receptors at cannabinoid-modulated synapses in the hippocampus, rendering this memory process more vulnerable to external manipulation at the time of memory retrieval.

Within the hippocampus, CB1 receptors are primarily located on cholecystokinin-containing interneurons (Katona et al., 1999; Marsicano and Lutz, 1999) surrounding the somata and dendrites of pyramidal neurons (Katona et al., 1999) and compared with cholecystokinin-GABA cells, glutamatergic neurons present a much lower density of CB1 receptors (Domenici et al., 2006; Kawamura et al., 2006). Furthermore, several studies have shown that hippocampal 2-AG, rather than AEA, mediates mechanisms of short- and long-term synaptic plasticity, which are involved in the modulation of memory processes (Makara et al., 2005; Straker and Mackie, 2005; Chevaleyre et al., 2006; Hashimoto et al., 2007; Kano et al., 2009). A growing amount of evidence shows that cannabinoids modulate behavior in an opposite way in regard to the degree of stress to which the subject is exposed (Hill and Gorzalka, 2004; Patel et al., 2005a; Fokos and Panagis, 2010; Morena and Camponolo, 2014). Importantly, both acute and chronic stress has been shown to change the synaptic effects of CB1 receptor activation. Specifically, it has been demonstrated that exposure to stress can induce a switch from CB1 receptor-mediated synaptic inhibition, in the form of LTD, to synaptic enhancement, in the form of LTP (Marsicano and Lutz, 1999). Furthermore, several studies have shown that hippocampal 2-AG, rather than AEA, mediates mechanisms of short- and long-term synaptic plasticity, which are involved in the modulation of memory processes (Makara et al., 2005; Straker and Mackie, 2005; Chevaleyre et al., 2006; Hashimoto et al., 2007; Kano et al., 2009). A growing amount of evidence shows that cannabinoids modulate behavior in an opposite way in regard to the degree of stress to which the subject is exposed (Hill and Gorzalka, 2004; Patel et al., 2005a; Fokos and Panagis, 2010; Morena and Camponolo, 2014). Importantly, both acute and chronic stress has been shown to change the synaptic effects of CB1 receptor activation. Specifically, it has been demonstrated that exposure to stress can induce a switch from CB1 receptor-mediated synaptic inhibition, in the form of LTD, to synaptic enhancement, in the form of LTP (Marsicano and Lutz, 1999).

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