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Metyrapone blunts stress-induced hyperthermia and increased locomotor activity independently of glucocorticoids and neurosteroids

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KEYWORDS
Locomotor activity; Stress-induced hyperthermia; Corticosterone; Neuroactive steroids; EEG; Arousal

Summary Metyrapone, a cytochrome P450 inhibitor used to inhibit corticosterone synthesis, triggers biological markers of stress and also reduces stress-induced anxiety-like behaviors. To address these controversial effects, 6 separate investigations were carried out. In a first set of investigations, abdominal temperature (Tabd) and spontaneous locomotor activity (AS) were recorded in freely moving rats treated with either saline or 150 mg kg−1 metyrapone. An increase in Tabd and AS occurred in saline rats, while metyrapone rats exhibited an immediate decrease, both variables returning to basal values 5 h later. Concomitantly, the EEG spectral power increased in the gamma and beta 2 bands and decreased in the alpha frequency band, and the EMG spectral power increased. This finding suggests that metyrapone depressed stress-induced physiological response while arousing the animal. In a second step, restraint stress was applied 5 h after injection. Metyrapone significantly blunted the stress-induced Tabd and AS rise, without affecting the brain c-fos mRNA increase. Corticosterone (5 and 40 mg kg−1) injected concomitantly to metyrapone failed to reverse the observed metyrapone-induced effects in Tabd and AS. Finasteride (50 mg kg−1), which blocks neurosteroid production, was also unable to block these effects. In conclusion, metyrapone acutely reduced stress-induced physiological response in freely behaving rats independently from glucocorticoids and neurosteroids.

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1. Introduction

Metyrapone is a cytochrome P<sub>450</sub> inhibitor (Williamson and O’Donnell, 1969) that blocks the 11β-hydroxylation of deoxycorticosterone (DOC) into corticosterone in the adrenal cortex (Jenkins et al., 1958). It has been extensively used in rodents to study the role of glucocorticoids in stress processes (Mousa et al., 1981; Haleem et al., 1988; Calvo et al., 1998; Mikics et al., 2005). However, metyrapone administration leads to apparently contradictory findings.

Metyrapone limits stress-induced behaviors. Metyrapone reduces immobility time when administrated one (Healy et al., 1999) or 3 h (Baez and Volosin, 1994) prior to forced swim test. Its administration 3 h before inescapable foot-shock exposure also decreases the percentage of inactive trials during the test (Baez et al., 1996). These behavioral changes occurring during stress exposure are likely to reflect a decrease in stress-induced anxiety. The subsequent anxiety-like behaviors are also reduced 24 h after stressor exposure. Metyrapone administration 3 h prior restraint exposure increases the time spent in open arms during an elevated-plus-maze test carried out 1 day after (Calvo et al., 1998; Calvo and Volosin, 2001).

Conversely, as stated by Rotllant et al. (2002), “metyrapone can act as a stressor”. Metyrapone increases c-fos mRNA expression in the hypothalamic paraventricular nucleus (PVN) (Herman et al., 1992) and fos-like immunostaining throughout the brain (Rotllant et al., 2002). Metyrapone also activates the hypothalamo-pituitary-adrenocortico (HPA) axis. The heteronuclear corticotropin-releasing factor (CRF) mRNA transcription is enhanced in the PVN (Herman et al., 1992) and vasopressin and CRF concentrations are increased in the pituitary portal blood (Conte-Devolx et al., 1992). Plasma ACTH concentration rises (Conte-Devolx et al., 1992; Herman et al., 1992; Rotllant and Armario, 2005) as well as that of 11-desoxycortisol (Conte-Devolx et al., 1992) and deoxycorticosterone (DOC, Krugers et al., 2000).

In order to examine the apparent controversial effects of metyrapone administration (anxiety concomitant to brain activation), a set of 6 investigations was carried out in freely moving rats. The first experimental step aimed at analyzing the immediate and delayed reactions to the injection of metyrapone through measuring two physiological variables. Abdominal temperature (\(T_{abdo}\)) was taken to approach anxiolytic properties of metyrapone through the stress-induced hyperthermia paradigm (Bouwknecht et al., 2007; Vinkers et al., 2009). Spontaneous locomotor activity (\(A_5\)) was recorded because it increases after social conflict (Sgoifo et al., 2002) and saline injection (Marinelli et al., 1997). Brain activation was assessed by recording the electroencephalogram (EEG). Arousal is reflected by fast β (19–30 Hz) and γ (γ1: 30–35 Hz and γ2: 35–50 Hz) frequency bands (Maloney et al., 1997). Variations in locomotor activity are associated with variations in the θ (4–8 Hz) band (Oddie and Bland, 1998). It increases with locomotion speed (Slawinska and Kasicki, 1998), but disappears when the animal is immobile (Whishaw and Vanderwolf, 1971). The second experimental step aimed at analyzing the effects of stress by applying a 60-min restraint after the extinction of the immediate response to metyrapone administration. The effects of metyrapone on stress-induced physiological activation were addressed using the time course of \(T_{abdo}\) and \(A_5\). The cerebral effects were assessed using brain c-fos mRNA expression (Chan et al., 1993). Metabolic effects of metyrapone administration were quantified in the blood using glycaemia, which increases after metyrapone (Werner, 1988; Rotllant et al., 2002) and stress (Armario et al., 1990), triglyceride concentration, which decreases after stress (Ricart-Juné et al., 2002), and lactate concentration, a marker of anaerobic metabolism. The role of the inhibition of glucocorticoid synthesis in the effects observed after metyrapone administration was evaluated through corticosterone supplementation. The place of the increased DOC production (Krugers et al., 2000) was analyzed by blocking the 5-α reductase using finasteride (Lephart et al., 1996). In fact, the transformation of DOC into tetrahydro-DOC (THDOC) by 3- and 5-α reductases (Raven et al., 1996; Rupprecht et al., 1998) acts as a positive modulator of GABA<sub>A</sub> receptor (Reddy, 2006).

2. Methods

2.1. Animals

The investigation was conducted in 250 male OFA Sprague–Dawley rats (Charles River Laboratories, L’arbresle, France) weighing 175–200 g upon arrival at the laboratory. Animals were housed at constant temperature (23 ± 2 °C) and relative humidity (50 ± 10%), and in a 12 h–12 h light-dark cycle (light on at 0800 h). The rats were accustomed to laboratory conditions during 10 days before surgery and were allowed 10 days to recover from the surgical operation. They were weighed 5 days a week to reduce handling stress (Briese and de Quijada, 1970). Experimental procedures were approved by the institutional ethics committee for animal care and performed in accordance with the principles of animal care (NIH publication no. 86-23, revised 1985) and the European Community Council Directive (86/609 EEC).

2.2. Drugs

Metyrapone was purchased from Interchim (Montluçon, France) for investigations no. 1, 3 and 6 (Fig. 1) and from Sigma–Aldrich (St-Quentin Fallavier, France) for investigations no. 2, 4 and 5 (Fig. 1). The substance was dissolved in 1 ml sterile saline (SAL) and injected IP. The chosen dosage was 150 mg kg<sup>−1</sup>, in order to block the stress-induced increase in blood corticosterone (Haleem et al., 1988). Corticosterone (CORT, Sigma–Aldrich) and finasteride (FIN, Interchim) were dissolved in 300 μL sesame oil (VEH) and injected SC. CORT was used either at a physiological (5 mg kg<sup>−1</sup>) or at a pharmacological dosage (40 mg kg<sup>−1</sup>). The latter dosages were chosen as they mimic the stress-induced blood corticosterone concentrations observed respectively in naive (Baez et al., 1996; Calvo and Volosin, 2001) and metyrapone-treated rats (Krugers et al., 1998). Finasteride was used at 50 mg kg<sup>−1</sup>, a dosage that blocks completely the 5-α reductase (Lephart et al., 1996).

2.3. Variables

2.3.1. \(T_{abdo}\) and \(A_5\)

\(T_{abdo}\) and \(A_5\) were assessed with a TA10TA-F40 implantable radiotransmitter and the telemetric signal was acquired using a DataQuest system running on ART-gold software 3.1
The TA10TA-F40 sensor was implanted into the abdominal cavity under deep anaesthesia (pentobarbital sodium, 60 mg kg\(^{-1}\), IP) according to a procedure described elsewhere (Michel et al., 2007a). As stated previously, the rats were allowed 10 days to recover from surgery. The \( T_{abd} \) and \( A_S \) values were measured every 2 min. \( A_S \) was calculated from variations in the transmitted power signal due to the rat position changes in its home cage. The \( A_S \) values \((x)\) were normalized using z-scores: \( z\)-score \((x) = (x - \mu)/\sigma\), standard deviation \((\sigma)\) and mean \((\mu)\) being calculated for each rat during the weekend preceding the experimental day (Chevrier et al., 2006). Weekends measures of \( A_S \) were thought to represent natural circadian variations, due to the absence of handling. Values in \( T_{abd} \) and \( A_S \) were averaged in 15-min and 1-h time periods. Baseline was assumed to be the mean value of the hour preceding injection or restraint exposure. Final data were presented in Table 1.

Figure 1  Questions to be answered by each experiment, means used to obtain an answer, related figure or table, and schematic view of experimental procedures. In Experiments 1 and 2, \( T_{abd} \), \( A_S \), EEG and EMG were measured before and after treatment injection. \( T_{abd} \) and \( A_S \) were both measured before and after drug injection and restraint in Experiments 3—6. Only data obtained after restraint are shown. Blood and brain samples were collected after stress and related data are presented in Table 1.
calculated as a difference to the relevant baseline (Δ values).

2.4. Electroencephalogram and electromyogram

To assess EEG and EMG, the TL10M3-F50-EEE transmitter (Data Sciences) was inserted into a pouch in the interscapular subcutaneous tissue under deep anaesthesia (pentobarbital sodium, 60 mg kg⁻¹, IP). The electrodes maintained by screws were placed in 5 holes drilled in the calvarium: one pair at 2 mm anterior and ±3 mm lateral to the bregma; and one pair at 4 mm posterior and ±2 mm lateral to the central suture. The reference electrode was placed 10 mm posterior to the bregma. Electrodes were anchored with dental cement (Dentalon Plus, Heraeus Kulzer, Dormagen, Germany). The EMG electrodes were placed inside the dorsal neck muscle. The rats received antibiotic (Extencilline, Sanofi-Aventis, 60,000 IU per rat, IP) and anti-inflammatory treatments (Ketofen 3 mg kg⁻¹, IM, Merial, Lyon, France) and were allowed 10 days to recover.

The signal was recorded at a sampling rate of 500 Hz (EEG) and 100 Hz (EMG). Data were imported to the Neuroscore software (v1.1.1, Data Science) for spectral analysis of the EEG by Fast-Fourier Transform (FFT) between 0.5 and 50 Hz with a 1 Hz resolution (hanning window, 10-s epochs). After artefact removal, data were analyzed according to two methods: (i) values were normalized by z-scores using parameters calculated for each rat on the signal assessed during the 2 h preceding injection. The calculated values were then averaged by 1-min epochs and plotted using SigmaPlot (Systat Software Inc., San Jose, CA, USA); (ii) values were divided into 8 frequency bands (δ: 0.5—4 Hz; θ: 4—8 Hz; α: 8—11 Hz; β1: 11—15 Hz; β2: 15—19 Hz; β2: 19—30 Hz; γ1: 30—35 Hz and γ2: 35—50 Hz) and expressed as percent of total spectral power (0—50 Hz). The EMG was averaged in 15-min epochs and expressed as root mean square of the activity (RMS).

2.5. Blood variables assessment

Plasma corticosterone concentration was analyzed using specific radioimmunoassay kits (¹²⁵I RIA kit, DPC France, La Garenne Colombes, France). Plasma glucose, lactate, triglycerides and cholesterol were assayed on a Hitachi 912 Analyser (Roche Diagnostics, Meylan, France) with the colorimetric method using Roche TM reagents (Roche Diagnostics, Meylan, France). Analyses were performed according to the manufacturer's instructions.

2.6. mRNA quantification by RT-PCR

The mRNA quantification of c-fos (NM_022197.2), CycA (NM_017101.1), β-actin (NM_031144.2) and ARBP (NM_022402.1) was done according to previously described methods (Michel et al., 2007a), using the same primers except for ARBP (Barbier et al., 2009). Briefly, the samples were conditioned in RNAlater (Qiagen, Courtaboeuf, France) until RNA extraction using MagNA Pure LC mRNA Isolation Kit II (Roche Applied Science, Mannheim, Germany). Reverse transcription was performed with oligo-dT (Reverse Transcription Core Kit, Eurogentec, Seraing, Belgium). Real-time PCR was carried out with the LightCycler Fast Start DNA Master SYBR Green kit (Roche Applied Science) with LightCycler (Roche Applied Science). Quantification cycles were assessed using the second derivative maximum method from a pool of cDNA samples as calibrator (Peinnequin et al., 2004), according to the comparative threshold cycle method (Livak and Schmittgen, 2001). Specificity of PCR amplification was verified with the LightCycler melting curve analysis (Peinnequin et al., 2004). The validity of internal control genes (CycA, β-actin and ARPB) was checked using geNorm (Vandesompele et al., 2002).

3. Experimental designs

Fig. 1 recapitulates the questions to be answered by each experiment, the experimental means that were used and the corresponding figures and tables presenting the results obtained.

3.1. Experiment 1

Experiment 1 analyzed (i) whether metyrapone induces stress using Tabd and Aδ time courses and (ii) whether metyrapone effects are related to glucocorticoid synthesis inhibition using a 40 mg kg⁻¹ corticosterone supplementation. The 43 rats were randomly distributed into 4 groups: saline—vehicle (SAL—VEH, n = 11), saline—corticosterone (SAL—CORT, n = 10), metyrapone—vehicle (MET—VEH, n = 11), and metyrapone—corticosterone (MET—CORT, n = 11). The rats, previously instrumented with TA10TA-F40 telemetric devices, received the assigned treatment and were left undisturbed for Tabd and Aδ recordings.

3.2. Experiment 2

Experiment 2 investigated (i) whether metyrapone elicits cerebral (central) effects and (ii) whether metyrapone-induced hypothermia triggers shivering through EEG and EMG recordings, respectively. The experiment was performed in 8 rats previously instrumented with TL10M3-F50-EEE telemetric devices. After recovery from surgery, each rat received an injection of saline and, 2 days later, another injection of metyrapone. Substances were given 120 min after lights were turned on and all recordings were made during the light period.

3.3. Experiment 3

Experiment 3 aimed at assessing the protective effect of metyrapone against stressor exposure by analyzing the time course of Tabd and Aδ during and after a 1-h restraint exposure carried out after normalization of metyrapone-induced Tabd and Aδ alterations. The 51 rats were randomly distributed into saline/vehicle (SAL—R, n = 12), saline/stress (SAL—S, n = 13), metyrapone/vehicle (MET—R, n = 13) and metyrapone/stress (MET—S, n = 13). The rats, previously instrumented with TA10TA-F40 telemetric devices, received the assigned pharmacological treatment and were then left undisturbed during 5 h. At this time, the rats belonging to stress groups were restrained during 1 h, then released and left again undisturbed. The rats belonging to rest groups remained undisturbed in their home cage throughout.
3.4. Experiment 4

Experiment 4 evaluated the involvement of glucocorticoid synthesis inhibition in the reduction of stress-induced physiological response after metyrapone. The time course of \( T_{abd} \) and \( A_\delta \) was studied in the same way as in Experiment 3. However, a 5 mg kg\(^{-1}\) corticosterone supplementation was administered concomitantly to metyrapone to obtain a corticosterone-clamp condition. The 43 rats were randomly affected to saline—vehicle (SAL—VEH, \( n = 11 \)), saline—corticosterone (SAL—CORT, \( n = 11 \)), metyrapone—vehicle (MET—VEH, \( n = 10 \)) and metyrapone—corticosterone (MET—CORT, \( n = 11 \)) treatment, the rats were exposed to a 1-h restraint (Fig. 2A; MET, \( p < 0.001 \); CORT, ns; interaction \( p < 0.01 \)) and A\(_\delta\) changes using finasteride co-treatment. The 45 rats were randomly assigned to saline—vehicle (SAL—VEH, \( n = 12 \)), saline—finasteride (SAL—FIN, \( n = 11 \)), metyrapone—vehicle (MET—VEH, \( n = 11 \)) and metyrapone—finasteride (MET—FIN, \( n = 11 \)). The experimental protocol was strictly similar to that of Experiment 4.

3.5. Experiment 5

Experiment 5 analyzed (i) whether metyrapone was capable of limiting stress-induced biological responses in the same experimental paradigm as in Experiment 4 and (ii) whether a 5 mg kg\(^{-1}\) corticosterone supplementation was sufficient to obtain stress-like levels of glucocorticoid blood concentration. The 60 rats were randomly distributed into saline/rest (SAL—R, \( n = 10 \)), metyrapone/rest (MET—R, \( n = 10 \)), saline/stress (SAL—S, \( n = 10 \)), metyrapone/stress (MET—S, \( n = 10 \)), metyrapone—corticosterone/stress (MET + CORT—S, \( n = 10 \)) and saline—corticosterone/stress (CORT—S, \( n = 10 \)). Five hours after treatment, the rats were exposed to a 1-h restraint (stress groups) or were left undisturbed in their home cage (rest groups). Immediately after restraint, the rats were quickly anaesthetized with 100% O\(_2\) ventilation containing 4% halothane (Laboratoire Belmont, Paris, France). Blood samples were placed in tubes containing lithium heparinate (Sarstedt, Marnay, France). Plasma was sampled after centrifugation and stored at \(-80^\circ C\) until assessment of blood variables. Prefrontal cortex and hippocampus were dissected and placed into RNAlater (Ambion, Courtaboeuf, France). These samples were maintained at 4 \(^\circ\) C during 24 h then stored at \(-20^\circ\) C until mRNA quantification by RT-PCR.

3.6. Experiment 6

Experiment 6 evaluated the role of THDOC in the metyrapone-induced \( T_{abd} \) and \( A_\delta \) changes using finasteride co-treatment. The 45 rats were randomly assigned to saline—vehicle (SAL—VEH, \( n = 12 \)), saline—finasteride (SAL—FIN, \( n = 11 \)), metyrapone—vehicle (MET—VEH, \( n = 11 \)) and metyrapone—finasteride (MET—FIN, \( n = 11 \)). The experimental protocol was strictly similar to that of Experiment 4.

3.7. Statistical analysis

Statistical analysis was performed with Statistica v7.1 (StatSoft-France, Maisons-Alfort, France). Comparisons were done by two-factor analysis of variance (ANOVA). The \( T_{abd} \) and \( A_\delta \) time courses were analyzed using ANOVA for repeated measures with factor effects. If necessary, post hoc tests were done using Bonferroni tests. Values are presented as means ± standard error of the mean (SEM).

4. Results

4.1. Experiment 1

During the baseline before injection, slight statistical differences (MET, ns; CORT, ns; interaction \( p < 0.01 \)) were observed for \( T_{abd} \) (SAL—VEH: 37.0 \pm 0.1 \(^\circ\)C; MET—VEH: 37.1 \pm 0.1 \(^\circ\)C; SAL—CORT: 37.2 \pm 0.1 \(^\circ\)C and MET—CORT: 36.9 \pm 0.1 \(^\circ\)C), but not for \( A_\delta \) (SAL—VEH: \(-0.2 \pm 0.1\); MET—VEH: \(-0.1 \pm 0.1\); SAL—CORT: \(-0.1 \pm 0.1\) and MET—CORT: \(-0.2 \pm 0.1\)).

Saline injection induced a strong hyperthermia (Fig. 2A). In rats receiving metyrapone, the hyperthermia was blocked and was followed by a 3-h hypothermia (Fig. 2A; MET, \( p < 0.001 \); and MET \times repetition, \( p < 0.001 \)). Corticosterone treatment did not modify \( T_{abd} \) time course with or without metyrapone co-treatment. Saline injection induced also an immediate increase in \( A_\delta \) (Fig. 2B), which was not observed after metyrapone. However, \( A_\delta \) was enhanced 2 h after metyrapone injection (MET, ns; MET \times repetition, \( p < 0.001 \)). Co-administration

![Figure 2](image-url)  
**Figure 2** Effect of acute administration of 150 mg kg\(^{-1}\) metyrapone IP and 40 mg kg\(^{-1}\) corticosterone SC on relative variations of abdominal temperature (A, \( T_{abd} \) expressed in \(^\circ\)C) and locomotor activity (B, \( A_\delta \), z-score). According to their experimental group, the rats received saline and sesame oil (SAL—VEH, \( n = 11 \)), metyrapone and sesame oil (MET—VEH, \( n = 11 \)), saline and corticosterone (SAL—CORT, \( n = 11 \)) or metyrapone and corticosterone (MET—CORT, \( n = 10 \)). Comparisons between groups used two ways factorial ANOVA and significant differences are indicated as follows: MET effect: \(^*\)\( p < 0.01\) and \(^{**}\)\( p < 0.001\); CORT effect: \(^*\)\( p < 0.01\) and interaction between MET and CORT effects: \(^{**}\)\( p < 0.05\). Data are expressed as means ± SEM.
of corticosterone did not affect $A_S$ time course after saline or metyrapone, although it amplified the increase in $A_S$ during the first 15 min after saline injection ($p < 0.05$). During the 5th hour post-injection, $T_{abd}$ and $A_S$ resumed pre-injection values and SAL—VEH and MET—VEH groups did not differ. This time period was thus selected as the reference period (stress—baseline) for the evaluation of metyrapone effects on stress reaction.

4.2. Experiment 2

Saline injection induced a 5-min decrease in the EEG spectral power in frequency bands between 0.5 and 25 Hz and a slight increase between 35 and 50 Hz (Fig. 3). No change was detected thereafter. Compared to saline, metyrapone modified the time course of EEG spectral power profoundly. It decreased $\alpha$ band relative power but did not affect the relative power of the $\delta$, $\theta$, $\alpha$ and $\beta$ bands. The main effect resided in the large increase that occurred in the $\gamma_1$, $\gamma_2$ bands and in the $\beta_2$ band. The effects of metyrapone lasted 2 h after injection for $\alpha$, $\beta_2$ and $\gamma_1$ frequency bands and 3 h for the $\gamma_2$ frequency band (data not shown). Concomitantly, metyrapone induced an increase in EMG activity, especially during the second hour following injection (Fig. 4; MET, ns and MET × repetition, $p < 0.05$).

4.3. Experiment 3

Before restraint stress, as in Experiment 1, saline injection triggered hyperthermia. Metyrapone administration blocked this stress-induced hyperthermia, and induced a subsequent hypothermia. The stress-induced increase in $A_S$ observed in saline rats was also blocked by metyrapone (data not shown).

During stress—baseline, no difference was observed between groups for $T_{abd}$ (SAL—R: 37.0 ± 0.1°C; MET—R: 36.7 ± 0.2°C; SAL—S: 37.0 ± 0.1°C and MET—S: 36.8 ± 0.2°C) and $A_S$ (SAL—R: −0.2 ± 0.1°C; MET—R: 0.0 ± 0.1°C; SAL—S: −0.3 ± 0.1°C and MET—S: −0.3 ± 0.1°C). During restraint, $T_{abd}$ increased sharply in saline rats (Fig. 5A; stress, ns; and stress × repetition, $p < 0.001$) and the stress-induced hyperthermia was blunted by metyrapone. In the resting condition, $T_{abd}$ levels remained higher in metyrapone than in saline rats (MET, ns; MET × repetition, $p < 0.001$ and MET × stress × repetition, $p < 0.001$).

After restraint release, saline rats exhibited a sharp increase in $A_S$ (Fig. 5B; stress, ns; and stress × repetition, $p < 0.001$). Metyrapone limited the stress-induced increase in $A_S$ (MET, ns; MET × repetition, $p < 0.001$ and MET × stress × repetition, $p < 0.05$).

4.4. Experiment 4

Similarly to observations made in Experiment 1, hyperthermia and increased $A_S$ were observed after saline injection, while hypothermia and a reduction in $A_S$ followed metyrapone injection. Corticosterone administration did not modify the concomitant $T_{abd}$ or $A_S$ time courses in saline and metyrapone rats (data not shown).

Figure 3 Change in EEG spectral profile after metyrapone administration. Eight animals received saline injection first (saline IP) and metyrapone 2 days later (150 mg kg$^{-1}$ metyrapone IP). Spectral powers between 0.5 and 50 Hz were analyzed with a 1-Hz resolution. The EEG spectral data were transformed into z-scores and averaged by 1-min epochs. Then, z-score data were averaged over 8 animals and color-coded.

Figure 4 Changes in EMG after metyrapone administration. Eight animals received saline injection first (saline IP) and metyrapone 2 days later (150 mg kg$^{-1}$ metyrapone IP). The EMG was expressed as the $10^5$ root mean square of the activity (RMS). Comparisons between groups were made through a two-way factorial ANOVA and significant differences are indicated as follows: MET effect: *$p < 0.05$; **$p < 0.01$ and ***$p < 0.001$. Data are expressed as means ± SEM.
During stress—baseline, compared to saline rats, metyrapone rats exhibited higher $T_{abed}$ (SAL−VEH: 37.0 ± 0.1 °C; MET−VEH: 37.2 ± 0.1 °C; SAL−CORT: 36.8 ± 0.1 °C and MET−CORT: 37.2 ± 0.1 °C, MET, $p < 0.01$) and A$_5$ (SAL−VEH: −0.3 ± 0.1; MET−VEH: −0.2 ± 0.1; SAL−CORT: −0.4 ± 0.1 and MET−CORT: 0.0 ± 0.1, MET, $p < 0.05$; MET × CORT, $p < 0.05$). Metyrapone reduced stress-induced hyperthermia (Fig. 6A; MET, $p < 0.01$; and MET × repetition, $p < 0.001$) and blunted the increase in A$_5$ (Fig. 6B; MET, $p < 0.01$; and MET × repetition, $p < 0.001$). Corticosterone supplementation did not change $T_{abed}$ and A$_5$ time courses in saline and metyrapone rats.

4.5. Experiment 5

Metyrapone did not modify blood corticosterone levels in resting condition (SAL−R: 125.9 ± 26.7 nmol l$^{-1}$, MET−R: 123.1 ± 16.0 nmol l$^{-1}$). Restraint induced an increase in corticosterone levels that was blunted by metyrapone (SAL−S: 476.2 ± 15.3 nmol l$^{-1}$, MET−S: 186.4 ± 8.0 nmol l$^{-1}$, MET, $p < 0.001$; stress, $p < 0.001$; MET × stress, $p < 0.001$).

In stressed rats, corticosterone supplementation limited the stress-induced increase in plasma corticosterone in saline rats (SAL−S: 476.2 ± 15.3 nmol l$^{-1}$, SAL + CORT−S: 358.3 ± 39.8 nmol l$^{-1}$; SAL−S vs. SAL + CORT−S: $p < 0.01$). Corticosterone addition had the opposite effect in metyrapone-treated rats (MET−S: 186.4 ± 8.0 nmol l$^{-1}$, MET + CORT−S: 284.6 ± 19.9 ng ml$^{-1}$; MET, $p < 0.001$; CORT, ns; CORT × MET, $p < 0.001$; MET−S vs. MET + CORT−S: $p < 0.05$).

Metyrapone did not alter resting levels of cortical and hippocampal c-fos mRNA expressions. c-fos mRNA expression increased after restraint in the frontal cortex (Table 1; stress, $p < 0.001$) and hippocampus (stress, $p < 0.01$). This reaction was not affected by prior metyrapone treatment (MET, ns; MET × stress, ns).

Table 1 Effect of metyrapone and stress on glucose (mMol L$^{-1}$), lactate (mMol L$^{-1}$) triglycerides (mMol L$^{-1}$) and cholesterol (mMol L$^{-1}$), and frontal cortex and hippocampus c-fos mRNA (a.u.). The animals received either saline and were stressed (SAL−S, n = 10) or rested (SAL−R, n = 10) or metyrapone and were stressed (MET−S, n = 10) or rested (MET−R, n = 10). The 1-h restraint was applied 5 h after treatment injection. Blood and brain samples were collected immediately at the end of restraint. The effects of metyrapone and stress were analyzed using a two-way ANOVA. Statistical differences are expressed in the first column with MET effect: * $p < 0.05$; *** $p < 0.001$ and stress effect: * $p < 0.01$; *** $p < 0.001$. Data are expressed as means ± SEM.

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<td>Frontal cortex c-fos</td>
<td>±±±</td>
<td>0.4 ± 0.1</td>
<td>0.4 ± 0.1</td>
<td>1.3 ± 0.3</td>
<td>1.0 ± 0.1</td>
</tr>
<tr>
<td>Hippocampus c-fos</td>
<td>±±</td>
<td>0.6 ± 0.1</td>
<td>0.6 ± 0.2</td>
<td>1.1 ± 0.1</td>
<td>0.9 ± 0.1</td>
</tr>
<tr>
<td>Glucose</td>
<td>***</td>
<td>10.9 ± 0.3</td>
<td>12.0 ± 0.3</td>
<td>12.1 ± 0.3</td>
<td>13.1 ± 0.3</td>
</tr>
<tr>
<td>Lactate</td>
<td>±±±</td>
<td>2.3 ± 0.2</td>
<td>2.3 ± 0.2</td>
<td>2.4 ± 0.4</td>
<td>2.4 ± 0.2</td>
</tr>
<tr>
<td>Triglycerides</td>
<td>*</td>
<td>1.2 ± 0.1</td>
<td>1.6 ± 0.2</td>
<td>1.1 ± 0.1</td>
<td>1.6 ± 0.2</td>
</tr>
<tr>
<td>Cholesterol</td>
<td>1.8 ± 0.1</td>
<td>1.6 ± 0.1</td>
<td>1.7 ± 0.1</td>
<td>1.6 ± 0.1</td>
<td></td>
</tr>
</tbody>
</table>
4.6. Experiment 6

Similarly to previous observations on immediate effects, metyrapone induced hypothermia and blocked the increase in $T_{\text{abd}}$ observed immediately after injection (data not shown). Finasteride co-treatment did not alter the concomitant $T_{\text{abd}}$ and $A_{\text{S}}$ time courses in saline- and metyrapone-treated rats (data not shown).

During stress—baseline, metyrapone rats showed higher $T_{\text{abd}}$ than saline rats (SAL—VEH: $37.0 \pm 0.1$°C; MET—VEH: $37.4 \pm 0.1$°C; SAL—FIN: $36.9 \pm 0.1$°C and MET—FIN: $37.3 \pm 0.1$°C, MET, $p < 0.001$). However, $A_{\text{S}}$ remained similar (SAL—VEH: $-0.4 \pm 0.1$; MET—VEH: $-0.3 \pm 0.1$; SAL—FIN: $-0.3 \pm 0.1$; MET—FIN: $-0.3 \pm 0.1$, ns). During restraint, metyrapone-treated rats exhibited a reduced hyperthermia compared to saline-treated rats (Fig. 6C; MET, $p < 0.001$; MET $\times$ repetition, $p < 0.001$). Finasteride did not change $T_{\text{abd}}$ time course in both saline and metyrapone rats. The increased locomotion observed after restraint in saline rats was reduced by metyrapone (Fig. 6D; MET, $p < 0.05$; MET $\times$ repetition, $p < 0.001$). Finasteride did not modify $A_{\text{S}}$ time course in both treatment groups.

5. Discussion

The present investigations demonstrated that metyrapone rapidly induced hypothermia and depressed locomotion...
while triggering cerebral arousal. After dissipation of these immediate effects, metyrapone also reduced the stress-induced hyperthermia and enhanced motor activity. The effects of metyrapone on body core temperature and locomotion were independent from glucocorticoid synthesis inhibition and THDOC synthesis.

5.1. Immediate effects of metyrapone

Immediately after saline injection, \( T_{abd} \) and \( A_s \) increased in relation to the injection-induced stress reaction. Hyperthermia occurs after handling (Briese and de Quijada, 1970; Vinkers et al., 2009), restraint (De Paula et al., 2000), and social (Sgoifo et al., 2002) and predator (Briese and de Quijada, 1970) confrontations. Increased motor activity is also observed after handling, saline injection (Vinkers et al., 2009) and social confrontation (Sgoifo et al., 2002). Both hyperthermia (Monda et al., 1998; Kiyatkin and Wise, 2001) and increased locomotion (Slawinska and Kasicki, 1998) are tightly linked to cerebral activation. These responses are evoked by cerebral injection of CRF (Morimoto et al., 1993) and reduced by anxiolytic substances such as GABA\(_A\) and 5-HT\(_{1A}\) receptor agonists (Bouwknecht et al., 2007; Conley and Bland, 1998; Slawinska and Kasicki, 1998). The decreased effect is congruent to other observations reporting that metyrapone treated rats were extinct. During stress—baseline, no differences were observed between saline- and metyrapone-treated rats regarding \( T_{abd} \), \( A_s \), EEG power spectra. Moreover, cerebral c-fos mRNA level and plasma corticosterone concentration were similar in both resting treatment groups 1 h after. As expected, \( T_{abd} \) increased in saline rats immediately after the beginning of restraint, and locomotion was enhanced just after restraint release. Concomitant cerebral activation was suggested by the increase in c-fos mRNA level in the hippocampus and frontal cortex (Senba and Ueyama, 1997; Chowdhury et al., 2000; Herman et al., 2003). The rats were also hyperglycaemic, such as usually observed in stressed fed rats (Selye, 1950; Nonogaki and Iuchi, 1997). Altogether, the above reactions suggest a strong stress response in saline-treated animals. Compared to saline, metyrapone administration induced a lower increase in \( T_{abd} \) and \( A_s \). Although metyrapone had increased basal blood glycemia, the latter was further increased after stressor exposure. Such an additive effect suggests that metyrapone and stress acted by different pathways and that metyrapone-treated rats were able to mobilize glucose when exposed to a stressor. Moreover, the c-fos mRNA level increase was statistically similar in both treatment groups after restraint. Metyrapone rats may have perceived their challenging environment, although the physiological consequences were different from those observed in saline-treated rats.

5.2. Subsequent effects of metyrapone

To evaluate whether metyrapone may have modified appraisal to stressor, the rats were submitted to restraint 5 h after injection. At this time, the immediate effects of metyrapone were extinct. During stress—baseline, no differences were observed between saline- and metyrapone-treated rats regarding \( T_{abd} \), \( A_s \), EEG power spectra. Moreover, cerebral c-fos mRNA level and plasma corticosterone concentration were similar in both resting treatment groups 1 h after. As expected, \( T_{abd} \) increased in saline rats immediately after the beginning of restraint, and locomotion was enhanced just after restraint release. Concomitant cerebral activation was suggested by the increase in c-fos mRNA level in the hippocampus and frontal cortex (Senba and Ueyama, 1997; Chowdhury et al., 2000; Herman et al., 2003). The rats were also hyperglycaemic, such as usually observed in stressed fed rats (Selye, 1950; Nonogaki and Iuchi, 1997). Altogether, the above reactions suggest a strong stress response in saline-treated animals. Compared to saline, metyrapone administration induced a lower increase in \( T_{abd} \) and \( A_s \). Although metyrapone had increased basal blood glycemia, the latter was further increased after stressor exposure. Such an additive effect suggests that metyrapone and stress acted by different pathways and that metyrapone-treated rats were able to mobilize glucose when exposed to a stressor. Moreover, the c-fos mRNA level increase was statistically similar in both treatment groups after restraint. Metyrapone rats may have perceived their challenging environment, although the physiological consequences were different from those observed in saline-treated rats.

5.3. Mechanisms of action of metyrapone

The biochemical mechanisms by which metyrapone may have produced these effects deserve discussion. Although metyrapone is known to block glucocorticoid synthesis (Jenkins et al., 1958), a major role of blood glucocorticoids in the blockade of the stress-induced hyperthermia and increased locomotion triggered by metyrapone is unlikely to occur. The following reasons may be proposed: (i) metyrapone did not induce glucocorticoid depletion because the resting blood corticosterone concentration was similar in metyrapone and saline rats 6 h after injec-
tion (Experiment 5), at a time at which stress reactivity was blunted (Experiments 3, 4, 6); (ii) corticosterone supplementation at a dosage that mimics stress levels of blood corticosterone failed to modify the time course of $T_{\text{abd}}$ and $A_S$ during both the early and the late phases; (iii) a high dose of corticosterone supplementation in metyrapone rats was also unable to modify the $T_{\text{abd}}$ and $A_S$ time course (Experiment 1), whereas it increases locomotion in saline rats suggesting treatment efficiency (Sandi et al., 1996). However, the fact that blood glucocorticoid concentration was not related to the previous effects does not rule out a role of glucocorticoid at the cellular level. Metyrapone can modify tissue glucocorticoid concentration by inhibiting specifically 11$\beta$-hydroxysteroid dehydrogenase type 1 (11$\beta$-HSD1) (Raven et al., 1995). This enzyme, which is widely expressed by neurons, converts inactive 11-dehydrocorticosterone into active corticosterone (Yau and Seckel, 2001).

Metyrapone may also act through the HPA axis activation since it increases brain CRF mRNA, arginine vasopressin and ACTH levels (Herman et al., 1992; Rotllant and Armario, 2005). Although CRF may have been implicated in arousal, it does not explain the blunt in $T_{\text{abd}}$ and $A_S$ since it produces the opposite effects (Morimoto et al., 1993). The accumulation of DOC (Conte-Devolx et al., 1992; Raven et al., 1996) further converted into the positive GABA$_A$ receptor modulator THDOC (Paul and Purdy, 1992) is also to be considered. Under GABA stimulation, metyrapone normalized the maximal chloride uptake previously reduced by restraint (Calvo et al., 1998). Such a positive GABA$_A$ modulation could account for metyrapone-induced hypothermia and diminished locomotion. However, the blockade of 5$\Delta$-reductase by finasteride, at a dose reducing brain concentration of neurosteroids (Verleye et al., 2005), failed to reverse the effects of metyrapone. Moreover, the metyrapone-induced brain activation suggested by EEG activation and cerebral c-fos induction (Herman et al., 1992) is not in accordance with such a mechanism. Therefore, metyrapone did not act through neurosteroid production.

Alternatively, metyrapone may act by modulating brain neurotransmission. Metyrapone is known to inhibit monoamine oxidase (MAO) and catechol-$\alpha$-methyl transferase (COMT) (Parvez and Parvez, 1973). It may therefore enhance noradrenaline and dopamine neurotransmission. However, COMT and MAO inhibitors are known to rather trigger hyperthermia and enhanced motor activity (Feldberg and Lang, 1970; Ashkenazi et al., 1983; Rivas et al., 1999). Metyrapone can also modify serotonin neurotransmission by upregulating serotonin 5-HT$_{1A}$ receptor through its effect on intracellular level of corticosteroid (Lopez et al., 1998). A 5-HT$_{1A}$-related mechanism may explain both its hypothemic and anxiolytic properties (De Vry et al., 2004; Hedlund et al., 2004).

Metyrapone may also simply act through metabolic modifications. Metyrapone may have reduced energetic metabolism as suggested by hypothermia. Actually, metyrapone induces metabolic depression in guinea pigs (Werner, 1988), although not in rats (Ohno and Kuroshima, 1986). The metabolic depression may be explained by an inhibition in glucose utilization (Bruno et al., 1972) and a shift towards lipid utilization, as supported by the decrease in respiratory quotient (Werner, 1988). This hypothesis is congruent with the development of hyperglycemia (Werner, 1988; Rotllant et al., 2002) and the observed increase in plasma triglyceride concentration.

6. Concluding remarks

Administration of metyrapone in rats triggered an immediate cerebral activation and a concomitant blockade of the injection-induced hyperthermia and enhanced locomotion. Subsequently, long after these early effects, metyrapone limited the consequences of restraint on body core temperature and locomotion. These effects were not mediated by blood corticosteroid concentration or neurosteroid production. These results shed light on the effect of P$_{450}$ inhibition on energy metabolism, brain activation and behavior, opening new potential pharmacological opportunities in the fields of stress-induce-anxiety disorders and brain ischemia.

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Conflicts of interest

The authors declare that over the past three years André Peinnequin has received compensation from ROCHE DIAGNOSTICS, the manufacturer of the qPCR reagents and devices used in this work. The authors have no other financial or competing interest to declare.

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