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Endogenous Opioids and Attenuated Hypothalamic–Pituitary–Adrenal Axis Responses to Immune Challenge in Pregnant Rats

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In late pregnant rats, the hypothalamic–pituitary–adrenal (HPA) axis is hyporesponsive to psychogenic stressors. Here, we investigated attenuated HPA responses to an immune challenge and a role for endogenous opioids. ACTH and corticosterone were assayed in blood samples from virgin and 21 d pregnant rats before and after endotoxin [lipopolysaccharide (LPS); 1 μg/kg, i.v.], interleukin-1β (IL-1β; 500 ng/kg, i.v.), or vehicle. In virgins, plasma ACTH concentrations increased 1 h after LPS and 15 min after IL-1β, as did corticosterone, with no responses in pregnant rats. In situ hybridization revealed increased corticotrophin releasing hormone (CRH) mRNA expression in the dorsomedial parvocellular paraventricular nucleus (pPVN) and increased anterior pituitary pro-opiomelanocortin mRNA expression 4 h after IL-1β in virgins; these responses were absent in pregnant rats. In contrast, immunocytochemistry showed that Fos expression was similarly increased in the nucleus tractus solitarius (NTS) A2 region in virgin and pregnant rats 90 min and 4 h after IL-1β. Naloxone pretreatment (5 mg/kg, i.v.) restored ACTH and pPVN CRH mRNA responses after IL-1β in pregnant rats but reduced the CRH mRNA response in virgins without affecting ACTH. Proenkephalin-A and μ-opioid receptor mRNA expression in the NTS was significantly increased in the pregnant rats, indicating upregulated brainstem opioid mechanisms. IL-1β increased noradrenaline release in the pPVN of virgin, but not pregnant, rats. However, naloxone infused directly into the pPVN increased noradrenaline release after IL-1β in pregnant rats. Thus, the HPA axis responses to immune signals are suppressed in pregnancy at the level of pPVN CRH neurons through an opioid mechanism, possibly acting by preterminal autoinhibition of NTS projections to the pPVN.

Keywords: ACTH; CRH mRNA; IL-1β; μ-opioid receptor mRNA; noradrenaline; nucleus tractus solitarius; proenkephalin-A mRNA

Introduction

Responsiveness of the hypothalamic–pituitary–adrenal (HPA) axis to psychogenic stressors is strongly attenuated during late pregnancy (Neumann et al., 1998; Brunton et al., 2000; Johnstone et al., 2000), but HPA axis responses in pregnancy to immune challenge, a physical stressor, have not been reported. In non-pregnant rats, HPA axis activation by immune challenge reflects the bidirectional relationship between the immune and stress axes (Besedovsky and Del Rey, 1992; Turnbull and Rivier, 1995; Mulla and Buckingham, 1999), with a vital role of the HPA axis in modulating immune responses (Turnbull and Rivier, 1995). Systemic administration of endotoxin [lipopolysaccharide (LPS)] increases corticotrophin releasing hormone (CRH) release (Givalois et al., 1995), CRH mRNA expression in the hypothalamic paraventricular nucleus (PVN) (Kakucsa et al., 1993), and adrenocorticotrophic hormone (ACTH) and corticosterone secretion (Turnbull et al., 1998). LPS stimulates cytokine production by macrophages, hence increasing circulating cytokine levels (Andersson et al., 1992), including interleukin-1β (IL-1β), which can mediate HPA responses to endotoxin (Besedovsky and Del Rey, 1992). Accordingly, systemic administration of IL-1β potently activates the HPA axis (Besedovsky and Del Rey, 1992; Ericsson et al., 1994). The key role of PVN CRH neurons in this response (Berkenbosch et al., 1987; Sapolsky et al., 1987; Rivest and Rivier, 1991) involves their activation by direct noradrenergic projections from the brainstem A1 and A2 neurons in the nucleus tractus solitarius (NTS) and ventrolateral medulla (VLM) (Ericsson et al., 1994; Buller et al., 2001b), with the A2 neurons of primary importance (Ericsson et al., 1994; Buller et al., 2001b). Systemic IL-1β stimulates these neurons (Ericsson et al., 1994) by first acting on IL-1 type I receptors on the endothelium of brain blood vessels to activate local production of prostaglandins (Ericsson et al., 1995; Rivest et al., 2000), which can then excite the NTS neurons via EP3 and EP4 receptors (Zhang and Rivest, 1999).

The attenuation of HPA axis responses to psychogenic stres-
sors in pregnancy involves inhibition by a central endogenous opioid mechanism that emerges in pregnancy (Douglas et al., 1998). Possible sources of endogenous opioid that could regulate PVN CRH neurons include noradrenergic NTS neurons, which coexpress proenkephalin-A (pENK-A) mRNA (Ceccatelli et al., 1989). μ-Opioid receptors mediate presynaptic inhibition of noradrenaline release in the hypothalamus (Onaka et al., 1995); thus, enkephalins may autoinhibit NTS neurons.

Here, we tested whether HPA axis responses to immune challenge are attenuated in late pregnancy and whether NTS neurons respond to IL-1β in pregnant rats. We also investigated whether differences in noradrenaline release in the PVN could account for attenuated HPA axis responses and investigated a role for endogenous opioids. Finally, we established whether pENK-A and μ-opioid receptor mRNA expression in NTS neurons is upregulated in pregnancy.

Materials and Methods

Animals

Female Sprague Dawley rats (initial body weight, 260–290 g; Bantin and Kingman, Hull, UK) were maintained under standard conditions of temperature (20–22°C), humidity, and lighting (12 h light/dark cycle; lights on at 7:00 A.M. with ad libitum access to food and water. Rats were initially housed in groups of five to six and then caged individually after surgery. Pregnant rats were obtained by mating overnight with a sexually experienced male; pregnancy was confirmed by finding a vaginal plug of semen on the cage floor the following morning (day 1 of pregnancy; day 22 of pregnancy is the expected day of parturition). Virgin rats were used as controls. All procedures were performed in accordance with current United Kingdom Home Office regulations.

Surgery: jugular vein cannulation

Five days before the experiment, virgin and day 16 pregnant rats were fitted with a SILASTIC jugular vein cannula (wall, 0.25 mm; internal diameter, 0.5 mm; filled with sterile heparinized 0.9% saline; 50 U/ml) under inhalational anesthesia (2–3% halothane in 600 ml/min each of oxygen and nitrous oxide) for subsequent blood sampling and drug administration.

Blood sampling

On the morning of the experiment (between 7:30 and 9:00 A.M.), jugular cannulas were connected to polyvinylchloride (PVC) extension tubing (wall, 1 mm; internal diameter, 0.5 mm) filled with heparinized saline (50 U/ml 0.9% saline) and attached to a 1 ml syringe for remote sampling. Rats were then left undisturbed for 90 min. Blood samples for radioimmunoassay (RIA) of ACTH (0.5 ml) or ACTH and corticosterone (0.65 ml) were collected into 1 ml syringes containing 20 μl of 5% EDTA chilled on ice and centrifuged to separate plasma, which was frozen and stored at −20°C until assay. Each blood sample was replaced immediately with an equivalent volume of sterile 0.9% saline. Pregnancy status was checked postmortem, and only pregnant rats with at least six viable fetuses were included in the study (actual range, 9–18 pups). The timings of blood samples for the different experiments are given below.

Experiment 1: ACTH responses to LPS

After two basal blood samples (0.5 ml for ACTH) taken 15 min apart, rats were given either 1 μg/kg LPS endotoxin (Serotype 055:B5; 2 μg/kg; i.v.; Sigma, Poole, Dorset, UK) or vehicle (0.9% saline, i.v.). Additional blood samples were taken 60, 120, 180, and 240 min after treatment. These time points were deemed to be the most appropriate after reviewing the literature (Turnbull et al., 1998).

Experiment 2: HPA axis and NTS responses to IL-1β

After two basal blood samples taken 30 min apart, rats were given either 500 ng/kg human recombinant (rh) IL-1β (1 μg/ml; i.v.; R & D Systems, Oxon, UK) or vehicle (0.2% bovine serum albumin (BSA) in PBS). Additional blood samples were withdrawn 15, 30, 60, 90, and 120 min after treatment. Rats were killed by conscious decapitation 4 h after treatment (optimal time to measure PVN CRH mRNA expression after stimulation) (Harbuz and Lightman, 1989). Trunk blood was collected into tubes containing chilled 5% EDTA (0.2 ml/100 g body weight), and plasma was separated and stored as above. The forebrains, brainstems, and pituitary glands were rapidly removed, frozen on dry ice, and stored at −70°C for CRH mRNA in situ hybridization (ISH), Fos immunocytochemistry, and POMC mRNA ISH (to assess corticotroph exposure to secretagogue), respectively.

To evaluate Fos expression in the NTS at 90 min after treatment (the time for peak neuronal Fos expression after stimulation) (Sharp et al., 1991), additional groups of pregnant and virgin rats were fitted with a jugular cannula as above. Five days later (on day 21 of pregnancy), the rats were given either 500 ng/kg rhIL-1β (1 μg/ml, i.v.) or vehicle (0.2% BSA in PBS) and killed 90 min later by conscious decapitation. Brainstems were rapidly removed, frozen on dry ice, and stored at −70°C for Fos immunocytochemistry.

Experiment 3: effects of naloxone on ACTH responses to IL-1β

Pregnant (day 21) and virgin rats were implanted with a jugular vein cannula as above. Immediately after the first basal blood sample was taken (0.5 ml for ACTH RIA), the rats were given 5 mg/kg naloxone (10 mg/ml) or 0.9% saline intravenously. After a second blood sample 15 min later, all rats were given 500 ng/kg rhIL-1β (1 μg/ml, i.v.). Additional blood samples (0.5 ml for ACTH RIA) were taken 15, 30, 60, and 90 min after IL-1β injection.

Experiment 4: effects of naloxone on PVN CRH mRNA responses to IL-1β

Pregnant (day 21) and virgin rats were used and were implanted with a jugular vein cannula. The rats were given, as above, naloxone or vehicle followed by IL-1β intravenously and killed by decapitation 4 h later. Brains were rapidly removed, frozen on dry ice, and stored at −70°C as before for CRH mRNA ISH. These rats were blood sampled as above for other measurements (not reported here).

Experiment 5: pregnancy and pENK-A mRNA and μ-opioid receptor mRNA expression

Virgin, midpregnant (day 10), late-pregnant (day 21), parturient (90 min after the birth of the first pup), and lactating (days 7–10) rats were killed by conscious decapitation. Brainstems were rapidly removed, frozen on dry ice, and stored at −70°C until processing for pENK-A and μ-opioid receptor mRNA ISH. Brains were also collected from untreated virgin and pregnant (d21) rats. Because enkephalin is coexpressed with CRH in the parvocellular PVN (pPVN) (Hokfelt et al., 1983) and is also expressed in the periforgical region (Khachaturian et al., 1983) (a region that may influence PVN neuron responses), sections cut through the pPVN and periforgical region were also processed for pENK-A mRNA ISH.

Experiment 6: effects of naloxone on IL-1β-evoked noradrenaline release in the PVN in virgin and pregnant rats

Surgery: implantation of microdialysis probe. Virgin and pregnant (day 18/19 of pregnancy) rats were anesthetized with halothane (as above), and a U-shaped microdialysis probe (tip length, 1.4 mm; dialysis surface, 2.4 mm²) (Landgraf and Ludwing, 1991) was implanted into the right PVN [coordinates: caudal, 1.7 mm; lateral, 1.6 mm; 7.4 mm deep from the brain surface; according to Paxinos and Watson (1996) at an angle of 10° to the vertical to avoid puncturing the sagittal sinus]. Probes were secured in place with two jeweler’s screws fixed to the skull and dental acrylic. Probes were flushed with sterile artificial CSF (aCSF) (in mm: 138 NaCl, 3.36 KCl, 9.52 NaHCO₃, 0.49 Na₂HPO₄, 2.16 urea, 0.49 NaH₂PO₄, 1.26 CaCl₂, 1.18 MgCl₂, pH 7.2). Lengths of fluorinated ethylene propylene (FEP) tubing (CMA Microdialysis, Solna, Sweden) were connected to the probe inlet and outlet. Rats were also fitted with a jugular vein cannula (as before) for administration of IL-1β. After surgery, rats were housed in commercially available purpose-designed Perspex bowls (CMA Microdialysis) with food and water available ad libitum. The rats were connected to the microdialysis tubing (CMA Microdialysis Probe, CMA/Microdialysis Sweden) and attached to a 1 ml syringe for remote sampling. Additional blood samples were obtained 15, 30, 60, and 90 min after IL-1β injection.
nous cannulas were connected to PVC extension tubing attached to a 1 ml syringe filled with heparinized saline. A length of FEP tubing con-
nected to a 1 ml Hamilton syringe prefillled with aCSF was attached to the inlet port of the probe. Hamilton syringes (1 ml) were secured in a microinfusion pump. Probes were flushed with aCSF (flow rate, 1.5 μl/ min) for 90 min before the experiment commenced. After flushing, mi-
crodialysis samples were collected at 15 min intervals. Three dialysis
samples were collected under basal conditions and then probes were
either perfused with aCSF or with aCSF containing naloxone (1 mg/ml)
for the remainder of the experiment. An additional three dialysate
samples were collected and then all rats were administered IL-1β (500 ng/kg, i.v.). Five additional dialysate samples were collected after IL-1β admin-
istration. In each case, dialysates were collected in chilled microdialysis
buffer [1% BSA (w/v), 5% dextran sulfate (w/v), 15 mM dietho-
tetriol]. The detection sensitivity for a 10\(^{-11}\) sample injection was 5–25 pg/ml.

Radioimmunoassays. Plasma ACTH and corticosterone concentra-
tions were determined using commercially available radioimmunoassay
kits (Immunodiagnostic Systems, Tyne and Wear, UK). ACTH was mea-
sured in unextracted plasma samples using a two-site immunoradiomet-
metric assay (Hodgkinson et al., 1984). Corticosterone was measured in
unextracted plasma (diluted 1:10 in assay buffer) using a double antibody
radioimmunoassay with\(^{125}\)I-corticosterone as the tracer (Keith et al., 1978).
For the ACTH and corticosterone assays, sensitivity was 1 pg/ml
and 0.4 ng/ml, respectively, and the intra-assay variation was <11
and <6%. All samples from any one experiment were assayed together.

In situ hybridization. Tissue was sectioned coronally on a cryostat at 15
μm and thaw-mounted onto gelatinized slides. Sections were fixed with
4% (w/v) paraformaldehyde in 0.1M PBS, washed in PBS, acetylated in
triethanolamine/acetic anhydride solution, dehydrated in an ascending
ethanol series (70 –100%), delipidated in chloroform, and partially rehy-
drated in 95% (v/v) ethanol. All sections from a particular experiment
were processed in the same hybridization reaction, and in each case,
synthetic oligonucleotide probes were used (MWG-Biotech, Ebersberg,
Germany). To detect CRH mRNA expression in the PVN, a 42-mer probe
(5'-CCT GTT GCT GTG AGC TTC AGC TAA CTC TCG TGC CCT GCC-3'),
complementary to bases 496–537, which encode amino acids 22–35 of the rat CRH peptide (Jingami et al., 1985), was used.
The POMC probe was a 48-mer oligonucleotide (5'-GGT CAT GAA GCC GTA GGG GGG GGG GTT GCC CCA GGC-3')
complementary to rat POMC mRNA (Nakanishi et al., 1979). The pENK-A probe was a 39-mer probe (5'-TGC ATC CTT CTT CCA TAA CTC GCC CCT GCC-3'),
complementary to bases 721–759 of pENK-A mRNA (Rosen et al., 1984), and the μ-opioid receptor
probe was 39-mer (5'-TTC AGA CCG CAG CTT GGA TCG GAC TGG TTG CCA TCA ACG TGG-3') and complimentary to bases 282–320 of
rat μ-opioid receptor mRNA (Chen et al., 1993).

Probes were 3'-labeled with \(^{32}\)P-dATP using terminal deoxynucleoti-
dyl transferase and purified using spin columns (QUlAquick nucleotide
removal kit; Qiagen, Crawley, West Sussex, UK). Sections were hybrid-
ized with radiolabeled probe (10\(^{-5}\) cpm labeled probe/section) in hybridiz-
ing buffer [1% BSA (w/v), 5% dextran sulfate (w/v), 15 μM dietho-
tetritol, 2 mM EDTA, 1% ficoll (w/v), 50% formamide (v/v), 0.1 mg/ml
PolyA, 1% polyvinylpyrrolidone (w/v), 0.2 mg/ml salmon testes DNA,
1.2 mM sodium chloride, 2.5% sodium pyrophosphate (w/v), 20 mM Tris,
PH 7.6, 0.1 mg/ml yeast RNA, 0.1 mg/ml yeast total RNA] overnight at
37°C in humidified chambers. Sections were rinsed three times in 1×
SSC at room temperature, washed four times for 15 min each in 1× SSC
at 45–63°C (depending on hybridized probe melting temperature), and
twice for 30 min each at room temperature. Sections were air-dried and
exposed to autoradiographic film (Hyperfilm-β-max; Amersham Bio-
sciences, Buckinghamshire, UK) at room temperature, with \(^{14}\)C polymer
strip standards to check optimal exposure (Amersham Biosciences).
Exposure times were 21 and 14 d for CRH and POMC mRNA, respectively.
Film was then developed (Kodak D-19; Sigma) and fixed (Hypam rapid
fixer; Ilford, Cheshire, UK). Slides were dipped in liquid autoradiog-
graphic emulsion (Kodak type NTB-3; Anachem, Bedfordshire, UK) and
exposed (at 4°C) for 12 weeks for CRH mRNA and 21 d for pENK-A
mRNA and μ-opioid receptor mRNA. Slides were developed and fixed as
above and counterstained with hematoxylin and eosin. The film autora-
diograms were quantified using a computerized image analysis system
(objective magnification, 2.5×; NIH Image version 1.62). Grain area
was measured over a minimum of six PVN profiles (in three consecutive
sections) and six anterior pituitary sections from each rat for quantifica-
tion of CRH and POMC mRNA expression, respectively. The area of each
profile was measured to calculate grain area/PVN or anterior pituitary
(square millimeter/square millimeter). Background measurements were
made over areas adjacent to the region of interest and subtracted.
Measurements over a \(^{14}\)C polymer strip were used to confirm that tissue
measurements were on the dynamic part of the radioactivity/silver grain
density curve.

The number of CRH mRNA-expressing cells in the PVN was counted in
emulsion-dipped sections. A positive cell was defined as one with more
overlying silver grains than the mean over 10 cells lateral to the PVN
(background) plus three SDs. Data are presented as the number of pos-
tive cells per square millimeter PVN profile. For quantitative analysis of
pENK-A and μ-opioid receptor mRNAs in the NTS, silver grain area
was measured in >150–200 cells in the NTS per rat. Background
measurements were made over the equivalent of 20 cells in neuropil adjacent
to the NTS and subtracted. pENK-A mRNA expression in the pPVN was
determined by counting the number of positively expressing cells
(emulsion-dipped sections) and by measuring grain density over the
pPVN (film autoradiographs). Both were measured in a minimum of six
PVN profiles per rat.

![Figure 1](image-url)
For all ISH measurements, average values for each rat were used to calculate group means ± SEM.

**Fos immunocytochemistry.** Coronal brain-stem cryostat sections (15 μm) cut through the A2 cell region of the NTS (at the level of the area postrema) were mounted on gelatinized slides. Sections were fixed in 4% paraformaldehyde in 0.1 M phosphate buffer (PB), pH 7.3, for 30 min at room temperature, followed by three 5 min washes in 0.1 M PB, pH 7.4. Endogenous peroxidases were deactivated with 1.5% hydrogen peroxide in 0.1 M PB for 15 min and then sections were washed in PB containing 0.3% Triton X-100 (PB-T). Sections were incubated in normal sheep serum (NSS) for 60 min before incubation with rabbit anti-Fos (Ab-2) antibody (Calbiochem, La Jolla, CA) (diluted 1:1000 in PB-T containing 1% NSS and 0.02% sodium azide; 750 μl/slide) at 4°C for 48 h. Sections were washed in PB-T (three times for 15 min each), incubated in excess (750 μl/slide) goat anti-rabbit peroxidase complex (Vector Laboratories, Burlingame, CA), and diluted 1:500 in PB-T containing 1% NSS for 24 h at 4°C. Next, sections were washed three times for 15 min each in 0.1 M PB, followed by a 10 min rinse in acetate buffer (0.33 M, pH 6.0). Sections were incubated in acetate buffer (0.33 M, pH 6.0) containing 2.5% nickel ammonium sulfate and 0.5% diaminobenzidine for 7–10 min until Fos-immunoreactive nuclei were visualized. The reaction was terminated with a 5 min wash in 0.165 M acetate buffer. Fos-positive cells were manually counted bilaterally in 8–10 NTS sections per rat (VLM A1 neurons are not activated by IL-1β at the dose used) (Ericsson et al., 1994). The area of the NTS profiles was measured using an AppleMac computer (Apple Computers, Cupertino, CA) and NIH Image (version 1.62) software.

**Statistical analysis.** Two-way repeated-measures (RM) ANOVA, followed by Student–Newman–Keuls multiple-comparison tests were used to analyze ACTH, corticosterone, and noradrenaline data. CRH mRNA, POMC mRNA, and Fos data were analyzed using a two-way ANOVA. pENK-A and μ-opioid receptor mRNA data were analyzed using a one-way ANOVA or Student’s t-test. Values shown are group means ± SEM. p < 0.05 was considered statistically significant.

**Results**

**Experiment 1: ACTH responses to LPS**

Basal plasma ACTH concentrations (1.53 ± 0.19 pg/ml in the virgin rats, n = 22 vs 0.87 ± 0.21 pg/ml in the pregnant rats, n = 17) were not significantly different between any of the groups (Fig. 1) (two-way RM ANOVA). Intravenous LPS significantly increased plasma ACTH concentration in the virgin group (p < 0.001; two-way RM ANOVA) within 60 min of the injection, which peaked at 120 min, and remained significantly elevated 240 min after LPS administration. However, LPS evoked only a small, nonsignificant increase in plasma ACTH concentration in the pregnant group (Fig. 1). Vehicle injection had no significant effect on plasma ACTH in either the virgin or the pregnant rats (Fig. 1).

**Experiment 2: HPA axis and NTS responses to intravenous IL-1β**

To circumvent the possibility that changes in endogenous cytokine generation may underlie the attenuated response to LPS in pregnancy, we measured HPA responses to IL-1β in subsequent experiments. Basal plasma ACTH concentrations were not significantly different between any of the groups (Fig. 2a) (p > 0.05; two-way RM ANOVA). Intravenous IL-1β rapidly increased ACTH secretion within 15 min in the virgin group (Fig. 2a) (p < 0.001; two-way RM ANOVA), which remained significantly elevated for at least 120 min after IL-1β, returning to baseline levels within 240 min. In contrast, in the pregnant group, IL-1β induced only a nonsignificant increase (1.7-fold; p = 0.093 and p = 0.084; two-way RM ANOVA vs basal and pregnant vehicle-treated groups, respectively) in plasma ACTH concentration 30 min after the injection (Fig. 2a). Plasma ACTH concentrations in the pregnant group were significantly less than those in the virgin group at all time points measured after IL-1β (p < 0.001; two-way RM ANOVA). Vehicle injection did not affect plasma ACTH concentrations in either the virgin group or the pregnant group (Fig. 2a).

Basal plasma corticosterone concentration was not significantly different between groups (Fig. 2b) (p > 0.05; two-way RM ANOVA). After IL-1β administration, plasma corticosterone concentration increased significantly within 30 min in the virgin group and was still significantly greater than basal at 240 min (p < 0.001; two-way RM ANOVA). Intravenous IL-1β had no significant effect on plasma corticosterone concentration in the pregnant group (Fig. 2b). Vehicle injection had no significant effect on plasma corticosterone concentration in either group.

POMC mRNA expression in the anterior pituitary was similar in the vehicle-treated virgin and pregnant rats. IL-1β treatment significantly increased POMC mRNA expression in virgin rats (Fig. 3a) (p < 0.001; two-way ANOVA) but had no such effect in pregnant rats (Fig. 3a).

Four hours after IL-1β treatment, CRH mRNA expression (measured as grain density and number of positive cells) in the dorsomedial PVN was significantly increased in the virgin group (Fig. 3) (p < 0.001; two-way ANOVA), but there was no significant change in CRH mRNA expression in the pregnant group (Fig. 3). CRH mRNA expression in the PVN was not significantly different between the virgin and pregnant control rats (Fig. 3) (p = 0.233; two-way ANOVA).

IL-1β significantly increased Fos expression in the A2 cell region of the NTS of both virgin and pregnant rats killed 90 min (Fig. 4a,b) or 240 min (Fig. 4c) after treatment; however, there were no significant differences in the number of Fos-positive neurons between the virgin and pregnant IL-1β-treated groups.
There were no significant differences in the number of Fos-immunopositive NTS neurons between the virgin and pregnant groups after vehicle (Fig. 4).

Experiment 3: effects of naloxone on ACTH responses to IL-1β
Basal plasma ACTH concentrations were not significantly different between any of the groups, and neither naloxone nor vehicle injection significantly affected basal plasma ACTH levels in any of the groups (Fig. 5a) (two-way RM ANOVA). IL-1β rapidly increased plasma ACTH concentration in the vehicle-treated virgin group, which peaked within 15 min (63.5 ± 2.8 pg/ml); however, there was no significant change in the vehicle-treated pregnant group (17.9 ± 3.5 pg/ml). Pretreatment with naloxone had no significant effect on the ACTH secretory response to systemic IL-1β in the virgin group (Fig. 5a). In contrast, in the pregnant group given naloxone, there was a significant ACTH response to IL-1β (p < 0.001 vs basal levels; p < 0.002 vs pregnant vehicle-treated group at 15 min; two-way RM ANOVA), although ACTH concentrations were significantly less than those in the virgin groups (Fig. 5a) at all sample times after IL-1β.

Experiment 4: effects of naloxone on PVN CRH mRNA responses to IL-1β
CRH mRNA expression in the pPVN 4 h after IL-1β treatment was 7.4-fold greater in the virgin vehicle-pretreated rats than in the pregnant vehicle-pretreated rats (Fig. 5b,c) (p < 0.001; two-way ANOVA). Naloxone pretreatment significantly increased CRH mRNA expression in response to IL-1β in the pregnant group (p < 0.001 vs pregnant/vehicle group; two-way ANOVA) but significantly reduced CRH mRNA expression in response to IL-1β in the virgin group (p < 0.001 vs virgin/vehicle group; two-way ANOVA). There were no differences in CRH mRNA expression between the virgin and pregnant IL-1β-treated groups pretreated with naloxone (p = 0.457; two-way ANOVA).

Experiment 5: pENK-A mRNA expression in the pPVN, perifornical region and NTS, and μ-opioid receptor mRNA expression in the NTS in pregnancy
pENK-A mRNA expression in the NTS was significantly greater on day 21 of pregnancy compared with virgin and lactating rats (Fig. 6a,b) (p < 0.01; one-way ANOVA). There was no significant difference in the level of pENK-A mRNA expression in either the pPVN or the perifornical region between virgin and late-pregnant rats (Table 1) (p = 0.50 and p = 0.58, respectively; Student’s t test). μ-Opioid receptor mRNA expression in the NTS was significantly increased in pregnant, parturient, and lactating rats versus virgin controls; peak expression was at parturition (Fig. 6c,d) (p < 0.001; one-way ANOVA).

Experiment 6: effects of naloxone on IL-1β-evoked noradrenaline release in the PVN in virgin and pregnant rats
Effect of IL-1β on noradrenaline release in the PVN of virgin and pregnant rats
Basal noradrenaline levels were not significantly different between the virgin group (513 ± 149.8 pg/ml) and the pregnant group (618.5 ± 206.3 pg/ml; Student’s t test; p = 0.466). Statistical analysis of log transformed basal noradrenaline data (to reduce variance) also showed no significant difference between virgin and pregnant rats (Student’s t test; p = 0.20). Systemic IL-1β administration evoked a significant increase in noradrenaline release in the PVN of virgin rats but not pregnant rats (Fig. 7a) (two-way RM ANOVA; p < 0.001); to control for interanimal variability, data were normalized as a percentage of within-rat mean basal values calculated from the six samples before IL-1β.
Effect of naloxone treatment on IL-1β-evoked noradrenaline release in the PVN of virgin and pregnant rats

In the pregnant rats, naloxone treatment alone had no significant effect on noradrenaline release in the PVN (Fig. 7b) (for analysis of effects of naloxone, data before naloxone and after IL-1β were normalized to the mean of the three post-naloxone microdialysate noradrenaline contents). After stimulation with IL-1β, noradrenaline release in the PVN was significantly increased in the pregnant rats pretreated with naloxone (two-way RM ANOVA; p < 0.02 at peak), contrasting with the lack of a significant effect of IL-1β on noradrenaline release in the PVN of the pregnant rats pretreated with vehicle (Fig. 7b).

Naloxone treatment alone had no significant effect on noradrenaline release in the virgin rats. In virgin rats, naloxone had no significant effect on noradrenaline release in the PVN after IL-1β administration, which was indistinguishable from that observed in the virgin rats treated with IL-1β only (data not shown).

Discussion

Our results demonstrate that in late pregnancy, the HPA axis is almost nonresponsive to immune challenge with systemic endotoxin or IL-1β as a result of endogenous opioid inhibition of pPVN CRH neurons. This inhibition may be exerted by enkephalins produced by NTS neurons and acting via μ-opioid receptors on noradrenergic terminals in the PVN. The results complement findings of reduced HPA responses in pregnancy to emotional stressors (Douglas et al., 1998; Neumann et al., 1998; Johnstone et al., 2000) and to orexin (Brunton and Russell, 2003), indicating global suppression of CRH neuron responses to stressors in late pregnancy.

We showed HPA axis activation by intravenous LPS in virgin females as in male rats (Turnbull et al., 1998). In late pregnancy, the ACTH secretory response to LPS was strongly attenuated. It is not known whether direct LPS actions on the brain (Chakravarty and Herkenham, 2005) are involved in HPA axis responses or altered in pregnancy. The reduced ACTH response to LPS in late pregnancy was not a result of reduced cytokine generation by immune cells, because the ACTH and corticosterone responses to IL-1β were also suppressed. The attenuated ACTH response to LPS or IL-1β is not attributable to failure of corticotrophs to respond to secretagogues in pregnancy (Ma et al., 2005), nor is metabolic clearance of corticosterone enhanced (Waddell and Atkinson, 1994). The attenuated ACTH and corticosterone responses to immune challenge result from reduced CRH drive (and/or vasopressin, although CRH predominates) (Turnbull et al., 1998) by the pPVN, because CRH mRNA expression in the pPVN was not stimulated by intravenous IL-1β in late pregnancy in contrast with virgin rats. Increased pPVN CRH mRNA expression after IL-1β indicates stimulation of the CRH neurons, which releases CRH from the axon terminals in the median eminence (Herman and Morrison, 1996). As CRH increases anterior pituitary POMC mRNA expression (Bruhn et al., 1984), the increase after IL-1β only in virgin rats indicates greater corticotroph exposure to CRH in virgin rats than in late-pregnant rats. Moreover, intracerebroventricular IL-1β depletes median eminence CRH in virgin rats but not in pregnant rats (Nakamura et al., 1998).

Reduced activation of CRH mRNA expression in late pregnancy after IL-1β is probably a consequence of altered input from brainstem noradrenergic neurons. Noradrenaline excites CRH neurons (Cole and Sawchenko, 2002), and CRH mRNA expression is tonically stimulated by noradrenergic and adrenergic pathways (Kiss et al., 1996). Here, increased noradrenaline release in the PVN of virgin rats, but not late-pregnant rats, after
IL-1β may explain the differences in CRH neuron activation. A putative source of this noradrenaline is the brainstem NTS neurons, which project directly to the PVN, relaying peripheral input to the hypothalamic CRH neurons. Lesioning the noradrenergic projection from the NTS may also mediate HPA responses to systemic IL-1β. NTS and VLM catecholamine cells may also mediate HPA responses to systemic IL-1β indirectly via the central amygdala and bed nucleus of stria terminalis (Buller et al., 2001a), and signaling through this pathway might be suppressed in pregnancy.

Naloxone reinstated an ACTH response to IL-1β in late-pregnant rats, indicating intact IL-1β signaling to the NTS in late pregnancy. Thus, signaling from brainstem blood vessels after systemic IL-1β [involving prostaglandin generation via cyclooxygenase (COX)] (Rivest et al., 2000) is unlikely to be suppressed in late pregnancy. We propose that reduced signaling by systemic IL-1β is at the level of the pPVN CRH neurons, including the terminals of the NTS projection. Altered intrahypothalamic COX signaling in pregnancy, underlying reduced LPS-induced fever responses (Mouihate et al., 2002), might contribute to reduced pPVN CRH neuron activation after systemic IL-1β. NTS and VLM catecholamine cells may also mediate HPA responses to systemic IL-1β indirectly via the central amygdala and bed nucleus of stria terminalis (Buller et al., 2001a), and signaling through this pathway might be suppressed in pregnancy.
pregnant rats, which strongly indicates a role for endogenous opioids in the attenuated response to systemic IL-1β. Naloxone also increases ACTH secretion after forced swimming in late pregnancy (Douglas et al., 1998) and during parturition (Wigger et al., 1999). Reduced corticotroph responses to CRH may explain the still-reduced ACTH secretory response to IL-1β in late-pregnant rats after naloxone (Neumann et al., 1998; Ma et al., 2005). However, a central inhibitory action of opioids in late pregnancy was clearly indicated, because naloxone triggered a pPVN CRH mRNA response to IL-1β and, infused directly into the PVN, naloxone restored IL-1β-evoked noradrenaline release in late-pregnant rats. In contrast, naloxone reduced the pPVN CRH mRNA response to IL-1β in virgin rats but not the ACTH response; naloxone reduces the ACTH response to swim stress in virgin rats (Douglas et al., 1998). The inferred excitation by endogenous opioids of HPA axis responses in virgins is consistent with potentiation of HPA stress responses by morphine (Buckingham, 1982) and stimulation of CRH release by enkephalins (Buckingham, 1982). Hence, we deduce that endogenous opioids promote pPVN CRH neuron stress responses in virgin rats; however, in late pregnancy, a dominant inhibitory opioid mechanism is activated, attenuating pPVN CRH neuron and, hence, ACTH secretory responses.

Naloxone does not distinguish the opioid receptor type(s) mediating the opposing effects on HPA activity in virgin and pregnant rats. However, the hypothalamus is almost devoid of δ-receptors (Mansour et al., 1987), and selective μ-receptor antagonists abolish ACTH stress responses in male rats (Cover and Buckingham, 1989), whereas blockade of κ-receptors exaggerates stress responses (Cover and Buckingham, 1989). Thus, in the present study, naloxone may have attenuated pPVN CRH mRNA responses to IL-1β in virgins via μ-receptors. However, the central endogenous opioid inhibition of oxytocin secretion that emerges in pregnancy is exerted via μ- not κ-receptors (Douglas et al., 1995). Furthermore, μ-receptors mediate attenuated HPA responses to IL-1β in male rats treated chronically with morphine (Chang et al., 1996), indicating the potential for inhibitory actions via μ-receptors.

The present finding of increased μ-opioid receptor mRNA expression in the NTS of pregnant rats is consistent with μ-receptor mediation of endogenous opioid inhibition of pPVN CRH neuron activation by IL-1β. Upregulated μ-opioid receptor production in NTS neurons could lead to increased transport to their terminals in the hypothalamus, where μ-opioids inhibit noradrenaline release in the supraoptic nucleus (Onaka et al., 1995). HPA axis hyporesponsiveness to immunological stressors in lactation (Shanks et al., 1999) and late pregnancy is likely a result of different mechanisms, because in the present study, proenkephalin-A mRNA expression in the brainstem was similar in virgin and lactating rats.

There are several possible sources of central endogenous opioid restraint that CRH neuron responses in pregnancy, including the arcuate nucleus, pPVN CRH neurons, perifornical region, and the NTS neurons themselves. Arcuate nucleus β-endorphin cells project directly to the PVN (Sawchenko et al., 1982), and in late pregnancy, POMC mRNA and β-endorphin expression in the arcuate nucleus are increased (Douglas et al., 2002). However, arcuate neurons are unlikely to mediate the immediate opioid actions on responses to systemic IL-1β, because they respond later than NTS and pPVN neurons (Brady et al., 1994). Enkephalin is expressed in the perifornical region and is coexpressed with CRH in pPVN cells, but here we showed that pENK-A mRNA expression is not increased in either the pPVN or perifornical region in late pregnancy. Furthermore, because these pPVN cells are not activated in pregnant rats after IL-1β, autoinhibition of pPVN neurons by enkephalin seems implausible.

The NTS is a more likely source of opioids that inhibit pPVN CRH neuron responses to IL-1β. Enkephalins and dynorphins are synthetized by NTS neurons (Ceccatelli et al., 1989, 1992). Thus, if the NTS neurons showing upregulated pENK-A mRNA expression in pregnancy include noradrenergic neurons projecting to the PVN (Ericsson et al., 1994), their activation by IL-1β is expected to release more enkephalin from their terminals to act on the upregulated μ-opioid receptors. Hence, the proposed site of naloxone action in pregnancy is the brainstem noradrenergic nerve terminals in the PVN, as supported by increased noradren-
aline release after systemic IL-1β during local nalofoxine application in pregnant rats.

In conclusion, in late pregnancy, HPA axis responsiveness to LPS and IL-1β is markedly attenuated, resulting from reduced drive by pPVN CRH neurons. This seems not to be a consequence of interrupted signaling to NTS neurons in pregnancy but is parsimoniously proposed to be through upregulated presynaptic autonomic endogenous opioid action on the brainstem noradrenergic projection to the CRH neurons, suppressing noradrenaline release. Reduced HPA responses to immune challenge in pregnancy will protect the fetuses from excessive glucocorticoid exposure, which can cause adverse programming and may also underlie the altered incidence of autoimmune disorders in pregnancy (Dunna and Finlay, 1989; Runmarker and Andersen, 1995; Ostensen and Villiger, 2002).

References


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