The importance of oxytocin mechanisms in the control of mouse parturition*

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The role of oxytocin in parturition in mice was investigated. Pup birth profiles, blood samples and brains were collected from parturient mice observed under red light conditions in a reversed light:dark photoperiod. Peripheral administration of an oxytocin antagonist in a dose-dependent manner delayed the birth of subsequent pups, indicating that oxytocin is required for a normal pup birth profile. Oxytocin neurones were activated during birth as shown by both increased immediate early gene (Fos) expression in oxytocin neurones in the supraoptic nucleus and increased plasma oxytocin concentrations during birth. In addition, the nucleus of the tractus solitarius and the olfactory bulbs, sites that process inputs to oxytocin neurones, become activated during parturition. Exposure to stress during parturition halted subsequent deliveries; at this stage plasma oxytocin concentrations were not higher than those of virgin mice, and birth was restored by administration of oxytocin. Administration of β-adrenergic antagonist (propranolol) also restored stress-delayed birth, whereas administration of ritodrine (β-agonist) delayed birth in non-stressed mice, indicating that adrenergic mechanisms contribute to stress-delayed births in mice. Administration of morphine (µ-opioid agonist) delayed births transiently, but naloxone (opioid antagonist) did not prevent stress-delayed birth, indicating that endogenous opioids do not appear to contribute to neuroendocrine or uterine mechanisms that promote birth in mice. Therefore, despite evidence in oxytocin knockout mice that oxytocin is not essential for parturition in this species, the results of the present study indicate that oxytocin neurone activity and secretion contribute to the birth process in normal mice.

Introduction

Oxytocin has strong uterotonic actions and is considered to drive parturition. Thus, administration of oxytocin antagonist delays the onset and progress of birth (Antonijevic et al., 1995a) by preventing uterine contractions and is used to prevent preterm labour in women (Atosiban, Ferring: Goodwin et al., 1994). The magnocellular oxytocin system is activated in rats during birth: oxytocin secretion increases (see for example, Higuchi et al., 1986), oxytocin neurones increase their firing rate (Summerlee, 1981), expression of Fos (Luckman et al., 1993; Antonijevic et al., 1995b) and oxytocin hnRNA (Douglas et al., 1998) and dendritic oxytocin release (Neumann et al., 1996). Oxytocin neurones are activated at parturition by neural signals from the contractile uterus, relayed by the brainstem, and from olfactory bulb neurones (Meddle et al., 2000; Douglas et al., 2001).

Oxytocin also plays a role in parturition in several other species, including pigs (Lawrence et al., 1997) and humans (Fuchs et al., 1991). However, in mice, deletion of DNA sequences in the gene encoding oxytocin has no apparent effect on birth (Nishimori et al., 1996; Young et al., 1996), although it does abolish the milk ejection reflex during suckling. These first transgenic animal models lacking the oxytocin peptide indicate that oxytocin is not essential for birth.

Stress exposure before and during parturition disrupts birth in many species, including humans (Laatikainen, 1991), rats (Leng et al., 1987), pigs (Lawrence et al., 1995) and mice (Newton et al., 1968), and this may, at least in part, relate to the interruption of the usual pattern of oxytocin secretion or oxytocin action in some species (Leng et al., 1987; Lawrence et al., 1995). Activation of peripheral or central β-adrenergic mechanisms in response to stress (Sanchez-Ramos et al., 1996) may also lead to the observed delays in births. These and other inhibitory factors, such as opioids, appear to modulate oxytocin secretion and mechanisms in parturition and actions on the uterus (Acevedo and Conrreras, 1987; Leng et al., 1987; Laatikainen, 1991; Petherick et al., 1993; Sanchez-Ramos et al., 1996; Wigger et al., 1999). Thus, in rats, exogenous opioids delay births in established parturition (Russell et al., 1989, 1991; Douglas et al., 1993), and the delaying effect of stress during parturition is prevented by the opioid antagonist naloxone (Leng et al., 1987).

In the present study, the role of oxytocin in parturition in the normal outbred mouse was investigated using an oxy-
tocin antagonist to block the action of oxytocin at oxytocin receptors, by determining the activity of oxytocin neurones in terms of their gene expression and secretion during parturition, and activation of their potential inputs from the brainstem and olfactory bulb. The effect of stress on oxytocin secretion and the efficacy of oxytocin in restoring births after interruption by stress exposure in parturient mice were also studied. In addition, the contribution of β-adrenergic mechanisms and endogenous opioids to stress-delayed birth was investigated.

Materials and Methods

Outbred MF1 female and male mice (6–8 weeks of age on arrival; Bantin and Kingman, Hull) were housed in groups of five in a temperature- and humidity-controlled ventilated room on a 12 h light:12 h dark reversed photoperiod, with lights on at 19:00 h (because mice predominantly give birth during the dark phase). Two to three weeks after adjustment to the reversed light cycle, mice were mated (five females to one male). From the next day (designated day 0 of pregnancy), the mice were housed in groups until 5–7 days before expected delivery, when they were transferred to individual cages. On the day of expected birth (days 18.5–19.5), the mice were observed continuously from 08:00 h in red light conditions and the times of birth of all pups were recorded. For all experiments, mice were selected randomly for treatments. All experiments were performed with ethical approval from the UK Home Office.

Temporal profile of pup births, and the effect of oxytocin antagonist or morphine on the progress of parturition

After the birth of the second pup, parturient mice were either left undisturbed in their cages (n = 4), or were injected s.c. with vehicle (isotonic saline, 100 μl per 50 g body weight; n = 6), oxytocin antagonist (F382; Ferring Pharmaceuticals, Malmö; at a dose of 30 or 300 μg per kg body weight, n = 5, 9, respectively) or with morphine (a μ-opioid agonist, 5 mg kg⁻¹ body weight; n = 5). The mice were then returned to their home cages and the births of subsequent pups were recorded.

Effect of parturition on plasma oxytocin concentration

After the birth of the second pup, parturient mice (n = 5) were decapitated when conscious and trunk blood (approximately 0.8–0.9 ml) was collected into ice-cold tubes containing EDTA (anti-coagulant, 5%, 150 μl per sample) and aprotinin (a protease inhibitor; Sigma, Poole; 0.039 trypsin inhibitor units in 10 μl per sample). Controls were pregnant mice on the day of expected parturition (n = 7) and (iii) postpartum mice (20–24 h after birth, n = 4). Pregnant and postpartum mice were killed time-matched with parturient mice. Forebrains, brainstems and olfactory bulbs were removed, frozen in crushed dry ice and stored at −70°C until processing for Fos immunocytochemistry.

Oxytocin concentration during parturition after exposure to stress. On the same day as the collection of the blood samples above, parturient mice were placed in a clean glass jar (8.0 cm × 5.5 cm × 12.0 cm, width by depth by height; n = 3) for 15 min after the birth of the second pup and then killed by decapitation when conscious. Trunk blood was collected and processed at the same time as above.

Effect of parturition on Fos expression in the mouse brain

Single labelling. Mice were killed by decapitation after brief sodium pentobarbitone anaesthesia (i.p. injection of 0.1 ml (6 mg) Sagatal; Rhone Merieux, Harlow). Groups included: (i) parturient mice (90 min after the birth of the second pup, n = 4), (ii) pregnant mice on the day of expected parturition (n = 7) and (iii) postpartum mice (20–24 h after birth, n = 4). Pregnant and postpartum mice were killed time-matched with parturient mice. Forebrains, brainstems and olfactory bulbs were removed, frozen in crushed dry ice and stored at −70°C until processing for Fos immunocytochemistry.

Double labelling. Additional groups of mice were perfusion-fixed at 90 min after the birth of the second pup to examine whether Fos expression in the supraoptic nucleus occurred in oxytocin neurones. Parturient (n = 3) and time-matched virgin mice (n = 5), pregnant mice (day of expected parturition, n = 5) and mice at 20–24 h post partum (n = 4) were deeply anaesthetized with sodium pentobarbitone (i.p. 0.2–0.3 ml, Sagatal; diluted 50:50 in sterile water to give 6–9 mg) and perfusion-fixed transcendially with heparinized saline (5 μl ml⁻¹, for approximately 60 s) and then with 4:100 (w/v) paraformaldehyde (in 0.1 mol PBS l⁻¹, for 25–30 min, at a flow rate of one drop per s). The brains were removed, post-fixed (in 15:100 (w/v) succrose in paraformaldehyde overnight and then in 30% (w/v) sucrose in PBS overnight) and frozen before double immunocytochemistry for Fos and oxytocin.

Effect of stress on the progress of parturition

On the birth of the second pup, mice were either left in their home cages (unstressed) or were stressed by transferring them into a clean glass jar, as described above. Subsequent pup births were recorded.

Stress interruption of pup births and effect of oxytocin. Mice were either untreated and left in their home cage (n = 8), or injected s.c. with vehicle and returned to their cage (n = 6), placed in a glass jar for 60 min (n = 6) or injected with oxytocin (Syntocinon; Alliance Pharmaceuticals Ltd, Chippenham; 100 mU in 100 μl, n = 6) and placed in a jar for 60 min.

Effect of β-adrenergic mechanisms in the responses to stress. Propranolol (β-adrenergic receptor antagonist; 5 mg kg⁻¹, 100 μl per 50 g body weight; Sigma) or vehicle was injected s.c. before the mice were transferred to a jar.
Effect of opioid antagonist on the responses to stress. Naloxone (general opioid receptor antagonist, 5 mg kg\(^{-1}\), i.p. in 100 \(\mu\)l per 50 g bodyweight, Sigma) or vehicle was injected s.c. immediately before mice were transferred to a jar (for 30 min, \(n = 3\), 3, respectively, or for 60 min, \(n = 4\), 5, respectively) or returned to the home cage (naloxone: 0.5 mg kg\(^{-1}\), \(n = 5\) or 5 mg kg\(^{-1}\), \(n = 5\); vehicle, \(n = 8\)).

Hormone analysis

Plasma oxytocin concentration was measured by an oxytocin radioimmunoassay using the Higuchi antibody (Higuchi et al., 1986) and modified according to Douglas et al. (1995). The assay sensitivity was 5 pg ml\(^{-1}\) and the intra-assay coefficient of variation was 19%.

Immunocytochemistry procedures

Immunocytochemistry for Fos was performed on coronal cryostat-cut brain sections (15 \(\mu\)m in thickness) that contained the supraoptic nucleus (at 6060–6280 \(\mu\)m anterior to the interaural line), olfactory bulb (through main and accessory parts) and the nucleus of the tractus solitarius (NTS; A1/C1 region, also containing area postrema, brainstem), according to published protocols (Douglas et al., 1995; Meddle et al., 2000). Briefly, sections were fixed and then incubated in rabbit anti-rat polyclonal anti-Fos antibody (1:1000 (v/v); Oncogene Sciences, Cambridge) for 24 h at 4\(^{\circ}\)C. After incubation with the secondary antibody (1:500 (v/v); Cambridge Bioscience, Cambridge), Fos labelling was visualized using the nickel-enhanced (glucose amino oxidase) 3',3'-diaminobenzidine (DAB) method, resulting in black cell nuclei. Slides were dehydrated and a coverslip was added. The number of Fos-positive neurones in the supraoptic nucleus (at 6060–6280 \(\mu\)m anterior to the interaural line), olfactory bulb (through main and accessory parts) and the nucleus of the tractus solitarius (NTS; A1/C1 region, also containing area postrema, brainstem) was counted on coded slides in eight sections per mouse, under a microscope (\(\times\) 25 objective).

Statistical analysis

Data are presented as group mean \(\pm\) SEM. Statistical analysis was performed using SigmaStat Software (Jandel Scientific, Worthing). Two-way ANOVA for repeated measures was used to compare parturition profiles between groups, taking into account the birth times for all pups after treatment (with the Student–Newman–Keuls’ post hoc test), and one-way ANOVA or Student’s \(t\) test for comparison of Fos data or plasma oxytocin data. \(P < 0.05\) was considered to be significant.

Results

Temporal profile of pup births and the effect of oxytocin antagonist or morphine on the progress of parturition

The cumulative time between the birth of pup two and pup ten in untreated mice in their home cage was 60.3 \(\pm\) 4.9 min and there was no significant effect on the profile of pup births for mice that were treated with vehicle s.c. (Fig. 1). Administration of the oxytocin antagonist significantly delayed the births of subsequent pups (two-way ANOVA for repeated measures, across the group: \(P < 0.01\); interaction between pup number and group: \(P < 0.01\); Student–Newman–Keuls’ post hoc test revealed that after the low dose of oxytocin antagonist, overall pup births were slower than the untreated and vehicle-treated groups, and
after administration of the high dose, pup births were slower than all other groups (Fig. 1)). Further statistical analysis at the level of each pup revealed that after administration of the high dose of the antagonist, births of pups four, five, six, seven and eight were significantly slower than those in the untreated and vehicle groups, and were slower at the births of pup nine and pup ten than all other groups (one-way ANOVA across groups, P < 0.01 for all, Student–Newman–Keuls’ post hoc tests, each P < 0.05). One-way ANOVA did not reveal any differences after administration of the low dose antagonist at any individual pup birth.

Compared with the same untreated and vehicle-treated mice, overall, morphine had no significant effect on pup births (two-way ANOVA for repeated measures, across groups: not significant). However, administration of morphine transiently delayed the birth of pup four (cumulative time from the birth of pup two to pup four, 16.0 ± 5.1, 26.2 ± 4.0 and 38.8 ± 5.4 min, respectively; one-way ANOVA, P < 0.05). There was no significant difference at the birth of the other pups (for example, cumulative time from the birth of pup two to pup eight was 40.5 ± 8.0, 53.5 ± 4.2 and 61.0 ± 5.6 min, respectively, one-way ANOVA, not significant).

**Effect of parturition on plasma oxytocin concentration and effect of stress exposure**

Plasma oxytocin concentrations were significantly higher in parturient mice at the birth of the second pup compared with both time-matched late pregnant and virgin mice (P < 0.05, one-way ANOVA, Fig. 2a, post hoc test). The plasma oxytocin concentration of mice placed in the jar for 15 min after the birth of their second pup was 5.0 ± 0.1 pg ml⁻¹, which was significantly lower than that of parturient mice killed after the birth of their second pup (one-way ANOVA, P < 0.01, post hoc test P < 0.05; Fig. 2b).

**Effect of parturition on Fos expression in the mouse brain**

In sections processed for Fos immunocytochemistry alone, parturient mice showed significantly more Fos expression.
in the supraoptic nucleus, olfactory bulb and NTS compared with both late pregnant mice and mice at 20–24 h post partum (one-way ANOVA, \( P < 0.01 \) for all brain regions measured; post hoc tests indicated that Fos in the parurient group in all cases was significantly greater than the other groups, \( P < 0.05 \); Fig. 3a–c). In the main olfactory bulb, there was no Fos expression in five of the 15 mice, one of which was parurient, and in the accessory olfactory bulb no Fos expression was observed in seven of the 15 mice, none of which were parurient. Therefore, further non-parametric statistical analysis was performed and confirmed a significant difference between the parurient and other groups of mice in the accessory olfactory bulb (Kruskal–Wallis, \( P < 0.01 \)) but not in the main olfactory bulb. The cumulative time from the birth of pup two to pup ten was 48.0 ± 11.0 min.

Fos expression was observed in many oxytocin neourones of the supraoptic nucleus during parturition but expression of Fos in oxytocin cells was scarce in the supraoptic nucleus of all other groups (Fig. 4a–d). The number of cells in the supraoptic nucleus expressing Fos significantly increased in parturition compared with all other groups (\( P < 0.05 \), one-way ANOVA, Student–Newman–Keul’s post hoc test; Fig. 4e), confirming the above observations (Fig. 3a). There was an increase in the number of double-labelled supraoptic nucleus cells positive for both Fos and oxytocin during parturition compared with all other groups (\( P < 0.0001 \), one-way ANOVA; Student–Newman–Keul’s post hoc test, \( P < 0.05 \) parturient versus all other groups; Fig. 4f). In comparison with virgin mice, the number of oxytocin neurones did not change with pregnancy, parturition or lactation (data not shown). The cumulative time for the birth of pup two to pup ten was 68.0 ± 10.0 min in parturient mice in this experiment.

Effect of stress on the progress of parturition

Stress interruption of pup births and the effect of oxytocin. Further births were completely halted in mice that were transferred to the glass jar for 60 min on the birth of the second pup (in six of six mice) and resumption of birth occurred only after the mice were returned to their home cage (Fig. 5a). The cumulative time between the birth of pup two and pup five in mice that were injected with vehicle, placed in the jar for 60 min and then returned to their home cages was 101.8 ± 6.3 min compared with 31.5 ± 7.5 min in vehicle-treated mice that remained in their home cages (untreated mice: cumulative time between the births of pup two and pup five 19.8 ± 2.8 min; one-way ANOVA, \( P < 0.001 \); post hoc test \( P < 0.05 \) pup group versus all other groups). Once birth resumed, the speed of delivery was similar to that in control mice (the cumulative time between the birth of pup five and pup eight was: vehicle plus jar 15.3 ± 3.3 min; vehicle plus cage 17.7 ± 1.5 min; and untreated 13.5 ± 2.1 min; one-way ANOVA, not significant). Administration of oxytocin s.c. to mice before they were placed in the jar completely prevented the delay in pup births and there was no significant difference between the oxytocin-treated, stressed group and the unstressed controls (oxytocin group: time between the birth of pup two and pup five, 23.7 ± 4.5 min; one-way ANOVA as described above. Taking into account all data: two-way ANOVA for repeated measures, across group: \( P < 0.001 \); interaction between group and pup number: \( P < 0.001 \), Fig. 5a).

Effect of \( \beta \)-adrenergic mechanisms in the responses to stress. Administration of propranolol prevented the stress-induced delay in pup births (Fig. 5b, cumulative time between the birth of pup two and pup five was 30.5 ± 6.1 min, compared with 101.8 ± 6.3 min in vehicle-treated stressed mice; one-way ANOVA, \( P < 0.001 \); \( P < 0.05 \) post hoc test vehicle-treated mice in jar versus all other groups except ritrodrine-treated mice in cage) and there was no significant difference between the propranolol-treated stressed mice and the non-stressed control mice. Propranolol-treated non-stressed mice showed a similar birth profile to vehicle-treated non-stressed mice, whereas ritrodrine-treated non-stressed mice showed a greatly prolonged delivery time: the mean time to the birth of pup three was 127.8 ± 23.6 min (one-way ANOVA at pup three, \( P < 0.001 \); \( P < 0.05 \) post hoc test ritrodrine-treated mice in cage versus all groups except vehicle-treated mice in the jar. Taking into account all data except the ritrodrine group: two-way ANOVA for repeated measures, across group: \( P < 0.0001 \); interaction between group and pup number: \( P < 0.0001 \), Fig. 5b).

Effect of opioid antagonist on the responses to stress. Administration of naloxone had no significant effect on pup births when mice were stressed in the jar (for 30 or 60 min) in comparison with those given vehicle and placed in the jar, but births in all stressed mice were significantly delayed compared with unstressed, vehicle-treated controls (two-way ANOVA for repeated measures, across group \( P < 0.05 \), Fig. 6). Pup births in unstressed rats given naloxone were not different from those of vehicle-treated, unstressed controls (0.5 mg naloxone kg⁻¹: from pup two to pup five, 44.8 ± 7.2 min; 5 mg naloxone kg⁻¹: from pup two to pup five, 50.2 ± 12.2 min; vehicle: from pup two to pup five, 39.3 ± 7.3 min; one-way ANOVA, not significant).

Discussion

The results of the present study show that not only is parturition delayed in mice by administration of an oxytocin antagonist, indicating a key role for oxytocin receptors and, hence, oxytocin in the timing of births, but also that oxytocin neurones are activated as Fos expression in magnocellular oxytocin neurones and oxytocin secretion into the blood increase. From this finding, and reports that posterior pituitary oxytocin content decreases during parturition (Fuchs, 1985), it is evident that hypothalamic neurones secrete oxytocin from their nerve terminals into the systemic circulation at birth in mice. Therefore, in the present study it was shown that, despite reports indicating that lack of
oxytocin in mice has no obvious effect on parturition (Nishimori et al., 1996; Young et al., 1996), evidently oxytocin does play a role in the birth process in mice, as it does in rats (Summerlee, 1981; Higuchi et al., 1986; Luckman et al., 1993; Antonijevic et al., 1995a) and broadly as in other species. However, there are to date no published studies of the temporal profile of parturition in the oxytocin knockout mice to determine whether parturition in the above models has a normal pattern and duration.

The findings from the present study are supported by reports that oxytocin receptor expression reaches a peak in the mouse uterus (Kubota et al., 1996; Mahendroo et al., 1996) and uterine sensitivity to oxytocin increases just before parturition (Suzuki and Kuriyama, 1975; Stepke et al.,

**Fig. 4.** Activation of oxytocin neurones in the supraoptic nucleus (SON) during parturition. Mice were perfused-fixed 90 min after the birth of the second pup (Part.), on the day of expected parturition (pregnant), 20–24 h post partum or were virgins. The brains were removed and double immunocytochemistry for Fos and oxytocin was performed on sections containing SON. Photomicrographs of the SON from (a) virgin, (b) pregnant, (c) parturient and (d) postpartum mice. OC: optic chiasm; long arrow: double labelled cell; short arrow: single labelled Fos-positive cell, including cells at the boundary of the SON. Scale bars represent 20 μm. (e–f) Quantification of Fos: (e) number of Fos-positive cells per SON profile, one-way ANOVA, $P < 0.05$; $^{*}P < 0.05$ versus all other groups, post hoc test; (f) number of cells containing both Fos and oxytocin per SON profile, one-way ANOVA, $P < 0.0001$; $^{*}P < 0.05$ versus all other groups, post hoc test.
1994). This is key to the onset of birth, as prevention of oxytocin receptor upregulation delays onset of birth (Sugimoto et al., 1997). In other species it has been proposed that oxytocin secreted from the uterus and placenta (Mitchell and Chibbar, 1995; Zingg et al., 1995) could act locally in the uterus to drive labour, supplementing or replacing neurohypophysial oxytocin. However, the mouse uterus does not express oxytocin mRNA (Murphy and Ho, 1995).

In rats, prostaglandins are uterotonic, although they are less potent than oxytocin (Fuchs, 1972). Deletion of the COX-1 (cyclo-oxygenase responsible for the synthesis of PGF$_{2\alpha}$) gene or the PGF$_{2\alpha}$ receptor gene in mice reveals that prostaglandins are essential for parturition in mice (Sugimoto et al., 1997; Gross et al., 1998). The prevention of parturition in these mice is related to the lack of oxytocin receptors in the uterus, as oxytocin receptor expression does not increase at term (Sugimoto et al., 1997). Ovariectomy in these mice, or a combination of deletion of the oxytocin and COX-1 genes (Sugimoto et al., 1997; Gross et al., 1998) restores parturition, indicating that prostaglandins may be essential only to induce luteolysis at term. Oxytocin receptor expression in the corpora lutea decreases as they enter luteolysis, preventing the luteotrophic effect of oxytocin, but oxytocin receptor expression in the uterus is simultaneously increased, enhancing oxytocin-induced uterine contraction (Gross et al., 1998; Imamura et al., 2000).

Thus, the mechanisms of initiation and progress of parturition in mice involve both prostaglandins and oxytocin; evidently there is redundancy that enables birth to proceed in oxytocin-deficient mice (Russell and Leng, 1998). Vasopressin, the other nonapeptide hormone synthesized in the supraoptic nucleus and secreted from the posterior pituitary, could also be involved. Vasopressin acts partly via the oxytocin receptor in the mouse uterus, and sensitivity to vasopressin increases at the end of pregnancy (Stepke et al., 1994). In the present study, a few Fos-positive cells that were not co-labelled with oxytocin were found in the supraoptic nucleus, indicating activation of vasopressin cells during birth, and supporting previous reports in rats (Kumaresan et al., 1979). Thus, vasopressin can contribute to the generation of uterine contractions during parturition, and may compensate for oxytocin in the case of oxytocin deficiency.

**Fig. 5.** Effect of oxytocin and β-adrenergic mechanisms on the temporal profile of pup births during stress exposure. Mice were transferred to a glass jar at the birth of the second pup (stressed) or they remained in their home cages (non-stressed). Data are mean ± SEM cumulative times to the birth of subsequent pups. (a) Mice were untreated (unstressed, ○), or injected s.c. with vehicle (unstressed, □; stressed, ■), or oxytocin (100 mU, stressed, ●) after the birth of the second pup. Two-way ANOVA for repeated measures $P < 0.0001$ across groups; $^* P < 0.05$ all data from vehicle-treated stressed group versus all other groups, post hoc test. (b) Mice were given vehicle s.c. (same unstressed [□] and stressed [■] mice as in (a)), or propranolol (5 mg kg$^{-1}$, unstressed, △; stressed, ▲) or ritodrine (0.5 mg, unstressed, ▽) after the birth of the second pup. Two-way ANOVA for repeated measures across group $P < 0.0001$; $^* P < 0.05$ all data from vehicle-treated stressed or ritodrine-treated unstressed groups versus unstressed vehicle- and propranolol-treated groups, and the stressed, propranolol-treated group, post hoc test.

**Fig. 6.** Effect of naloxone on the profile of pup births during exposure to stress. Mice were injected s.c. with vehicle (unstressed, □; stressed for 30 min, ◊; stressed for 60 min, ▼) or naloxone (5 mg kg$^{-1}$, stressed for 30 min, □; stressed for 60 min, ▼) at the birth of the second pup. Data are mean ± SEM cumulative times to the birth of subsequent pups. Two-way ANOVA for repeated measures, $P < 0.05$ across groups; $^{**} P < 0.05$ all data from stressed groups versus unstressed vehicle-treated group, post hoc test.
The NTS and the olfactory bulb, as well as the supraoptic nucleus, neurones strongly express Fos during, but not before or after, birth. Activated NTS neurones include neurones that project directly to the supraoptic nucleus (Meddle et al., 2000) and mediate positive feedback from the contracting uterus to the supraoptic nucleus, as reported in rats (Douglas et al., 2001). Olfactory bulb neurones also project to the supraoptic nucleus in rats and are activated during parturition (Meddle et al., 2000). Activation of these inputs provides further evidence of an important role for supraoptic nucleus neurones in parturition. As accessory olfactory bulb neurones expressed Fos, indicating input from the vomeronasal organ, activation may reflect a response to pup odours that may be involved in evoking maternal behaviour.

The present study has also provided further evidence that reduced plasma oxytocin concentrations accompany interruption of pup births, and furthermore, that oxytocin administration prevents the stress-induced delay in pup births, extending and confirming reports in mice (Newton et al., 1968), rats (Bosc and Nicolle, 1979; Leng et al., 1987), dogs (Bleicher, 1962) and pigs (Lawrence et al., 1995). In addition, the β-adrenergic antagonist, propranolol, was able to restore birth in stressed mice, whereas the β-agonist, ritodrine, delayed birth in non-stressed mice. Adrenergic actions in the uterus alter during the oestrous cycle and in pregnancy (Cruz and Rudolph, 1986; Cruz et al., 1990), and β-adrenergic antagonists are used clinically to promote birth (Sanchez-Ramos et al., 1996), whereas agonists act as tocolytics on the human uterus (Segal et al., 1998), also inhibiting uterine contractions and delaying birth in rats and mice (Woodward and Cheng, 1982; Yoshizawa et al., 1998). Thus, it seems likely that β-adrenergic mechanisms are activated during stress-delayed birth in mice. Whether stress induces peripheral sympathetic activation is unclear. Enhanced sympathetic tone may reduce uterine blood supply and inhibit smooth muscle contractions during birth. Reports that oxytocin opposes catecholamine-induced tocolysis (Segal et al., 1998) indicate that oxytocin can overcome β-adrenergic block of parturition, and propranolol and oxytocin administered together promote birth (Sanchez-Ramos et al., 1996). Oxytocin alone was highly effective at overcoming the interruption of births by stress indicating that the uterus remains well-perfused and that the inhibition of oxytocin secretion contributes to the delay in births. The inhibition of uterine contractions by β-adrenergic mechanisms would reduce afferent neural stimuli to oxytocin neurones and hence reduce positive feedback stimulation of oxytocin secretion.

The inhibitory effect of stress on pup births and the restorative effect of propranolol could also act via central actions on oxytocin neurones. In lactation, propranolol facilitates the milk ejection reflex (Poulain and Dyer, 1984), indicating β-adrenergic receptor-mediated inhibition of oxytocin neurones. Conversely, catecholamines act in the supraoptic nucleus to increase basal and suckling-induced secretion of oxytocin through α-receptors (Clarke et al., 1979; Parker and Crowley, 1993). In addition, presumed excitatory noradrenergic pathways to oxytocin neurones, mediated by α1 receptors, are activated during parturition in rats (Douglas et al., 2001). Any role of a central inhibitory β receptor-mediated mechanism on oxytocin neurones in stress responses during birth in either rats or mice is not known.

Morphine transiently slows births in mice, as shown in rats (Russell et al., 1989, 1991; Douglas et al., 1993), but in rats morphine strongly inhibits oxytocin neurone activity and secretion (Douglas et al., 1993; Luckman et al., 1993). Opioids also inhibit labour in humans (Lindow et al., 1992). However, naloxone did not restore birth in stressed mice, indicating that endogenous opioids are not involved in the delay of pup delivery. This is in contrast to findings in rats and pigs in which naloxone prevents the slowing of births by stress (Leng et al., 1987; Lawrence et al., 1992), and increases oxytocin secretion, indicating that endogenous opioid inhibition of oxytocin neurones accounts for the disruption in parturition by stress in these species. Morphine evidently does not act directly on the uterus in rats (Russell et al., 1989), but may act on opioid receptors in the spinal cord to mediate analgesia, which might be expected to promote birth. However, the site of morphine action in delaying birth in mice is not known.

Thus, in conclusion, during parturition in mice, oxytocin plays an important role in promoting birth, as in other mammals (Russell and Leng, 1998), despite evidence that parturition proceeds in oxytocin-deficient mice (Nishimori et al., 1996, Young et al., 1996). However, the mechanisms controlling oxytocin secretion and action during the stress-delayed birth differ from other species.

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