Differential expression of mesotocin receptors in the uterus and ovary of the pregnant tammar wallaby

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Abstract

Mesotocin, an oxytocin-like peptide, is released in highest concentrations during parturition in macropodid marsupials. In late pregnant wallabies, uterine sensitivity to mesotocin increases markedly in the myometrium of the gravid uterus. This coincides with a significant increase in myometrial mesotocin receptor concentrations 3–4 days before term. To date, there is no information on mesotocin receptor gene expression in female wallaby reproductive tissues. This study aimed to examine mesotocin receptor gene expression in the uterus and ovaries of pregnant tammar wallabies, and to localise mesotocin receptors within the uterus. An RT-PCR strategy produced a consensus nucleotide sequence of 834 bp, which encoded 278 amino acids of transmembrane domains I to VI. This protein sequence has approximately 80% homology with the bovine and rat oxytocin receptor exon 2 region. Only one mesotocin receptor was detected in the tammar genome. The myometrium and mammary gland both expressed a 4.1 kb mesotocin receptor gene transcript. Myometrial mesotocin receptor gene expression increased on day 22 of the 26-day gestation and was significantly higher in the gravid than the non-gravid uterus in late pregnancy. This pattern of mesotocin receptor gene expression paralleled mesotocin receptor concentrations. Mesotocin binding sites were localised only to the myometrium, the highest densities being observed in the gravid uterus. Finally, this study showed high expression of mesotocin receptors in the corpus luteum. The pattern of luteal mesotocin receptor expression differed from the myometrium, with a decrease in mesotocin receptors occurring on the day of expected births.


Introduction

The oxytocin (OT) receptor belongs to the class 1 G-protein-coupled, seven-transmembrane domain receptor superfamily and primarily signals via Gαq proteins through the phospholipase C-beta isoform (PLC-β). Kimura et al. (1992) first isolated and identified a cDNA encoding the human OT receptor. To date, the OT receptor-encoding sequences have been identified in the pig (Gorbulev et al. 1993), rat (Rozen et al. 1995), sheep (Riley et al. 1995), cow (Bathgate et al. 1995), mouse (Kubota et al. 1996) and rhesus monkey (Salvatore et al. 1998). Unlike eutherians, which secrete OT, marsupials secrete mesotocin (MT), which differs from OT by one amino acid, an isoleucine for leucine at position 8. The presence of OT-like receptors in marsupials was first demonstrated in the brushtail possum uterus, mammary gland and median vaginal sacs (Sernia et al. 1990, 1991). These studies reported that the OT-binding site was not specific for MT over OT with similar high affinities, whereas AVP, lysine [Lys-8]-vasopressin (LVP) and [Phe 2]-vasopressin (PP) had much lower binding affinities. This is consistent with binding to an OT-like receptor, and not an AVP receptor.

The tammar wallaby (Macropus eugenii) is a well-established model to differentiate between the role of maternal systemic and fetal-specific factors in the regulation of myometrial MT receptors during pregnancy. Female tammar wallabies give birth to small, altricial young after a short gestation period of only 26 days after reactivation of the diapausing blastocyst (Renfree et al. 1996). They have two anatomically separate uteri, which open into the anterior vaginal expansion via separate cervices (Tyndale-Biscoe & Renfree 1987). Due to the unique reproductive tract anatomy of marsupials, the influence of the conceptus on the regulation of myometrial MT receptors can be isolated from maternal factors, with direct comparison between the gravid and non-gravid uterus. From day 23 of gestation, MT receptor concentrations in the myometrium of the gravid uterus increase markedly, whereas in the
non-gravid myometrium they remain unchanged (Parry et al. 1997). In fact, MT receptors are significantly decreased in the non-gravid myometrium on the last 2 days of gestation. The changes in MT receptor density in the gravid uterus are matched by large increases in uterine sensitivity to exogenous MT (Parry et al. 1997). A recent study in unmated tammars showed no increase in MT receptors in either uterus despite the similar lengths of the luteal phase in non-pregnant and pregnant tammars (Siebel et al. 2002a). These data confirm that the increase in MT receptors is not only unique to the gravid uterus, but also pregnancy specific.

To date, there is no information on MT receptor gene expression in relation to MT receptor concentrations during pregnancy in female wallaby reproductive tissues. Moreover, MT receptors have not been localised within the wallaby uterus. Earlier work in the tammar prostate gland elucidated a partial sequence of the tammar MT receptor gene, which encoded a protein of 196 amino acids (Parry & Bathgate 1998). The derived amino-acid sequence had relatively high homology (74–77%) with the region extending from putative transmembrane domains II to VI in all eutherian OT receptors and relatively low homology (38–52%) with AVP receptors. Although MT receptor gene transcripts were demonstrated in the prostate gland (Parry & Bathgate 1998), this study did not include female reproductive tissues. Therefore, the overall aim of this study was to examine myometrial MT receptor gene expression in the reproductive tract and ovaries of the pregnant tammar wallaby. This study also included autoradiography to localise MT receptors within the uterus.

Materials and Methods

All animal experiments were conducted with approval from the La Trobe University Animal Ethics Committee (no. AEC 00/3 L) and the Department of Natural Resources and Environment (permit no. 10002261).

Animals

Female tammar wallaby (*M. eugeni*) tissues were collected on Kangaroo Island (SA, Australia) from wild-shot animals with approval from the South Australian Department for Environment and Heritage (permit no. Q24436) and the Victorian Department of Natural Resources and Environment (permit no.10000449). Females were housed with males in open, grassed enclosures at the Wildlife Reserve (La Trobe University, Bundoora, VIC, Australia) with water and kangaroo pellets readily available.

A pilot study demonstrated no significant differences in either MT receptor mRNA or receptor concentrations between wild-shot and colony animals at the same stage of gestation. Therefore, data from wild and colony-housed pregnant tammars were pooled at each stage of gestation. Gestational age was estimated from fetal crown rump length and head-length measurements, and verified by specific developmental characteristics validated from a large sample size (>300 animals) obtained from time-mated animals in a breeding colony (L J Parry, unpublished observation). In all cases, these measurements are accurate to within half a day. To examine MT receptor mRNA expression in a range of tissues at a specific stage of gestation (day 23), pregnancies were synchronised in a group of colony-housed adult wallabies by removing the pouch young (R PY) from females assumed to be carrying a blastocyst in embryonic diapause.

Tissue collection

All tissues used for RNA analysis were collected as quickly as possible after the animal had stopped breathing. Colony-derived animals were killed by an overdose injection (3–5 ml) of pentobarbitone sodium (Lethabarb; Virbac Australia Pty Ltd, Parkhurst, NSW, Australia) (325 mg/ml) into the heart. The nipples and mammary glands were collected before the reproductive tract was removed via laparotomy. The ovaries and oviducts were removed, and the two uteri dissected from the extraterine tissue and cut longitudinally to open the uterine cavity. The fetus, placenta, cervix and median vagina were all collected, as was the endometrium when separated from the myometrium. All tissues were placed in liquid nitrogen and stored at −80°C until further processing. The gravid and non-gravid uteri were collected from day-23 pregnant females and cut in transverse section so that autoradiography binding studies could be performed on the intact myometrium, endometrium and placenta (gravid uterus only). These tissues were fixed in Tissue-Tek O.C.T. embedding medium (ProSciTech, Thuringowa, QLD, Australia) and frozen at −80°C. Serial sections (8–10 μm) were cut with a cryostat at −20°C and thaw mounted on gelatin-coated slides. Sections were then stored at 4°C overnight in the presence of silica gel.

Mesotocin (MT) receptor gene

The nucleotide and derived amino-acid sequence of the 5′-coding region of the tammar MT receptor were obtained with the First-Choice RLM-RACE kit (Ambion, Geneworks, Adelaide, Australia), according to the manufacturer’s instructions. Total RNA was extracted from the gravid myometrium (day 23), using 2 ml RNAWiz (Ambion) per 100 mg tissue. After isopropanol precipitation, the RNA pellet was resuspended in H2O treated with RNA Secure (Ambion). First-strand cDNA was synthesised from 5 μg total RNA, using 0.5 μg/μl random hexamers or a tammar MT receptor-specific primer (MTR 490, 5′-CCT CAT CTC TTA CCT GTG AGG-3′; Geneworks) and 100 U Superscript II RNase H− reverse transcriptase (Invitrogen, Mulgrave, Australia) in a total volume of 20 μl. A volume of 2 μl of the cDNA was used as a template for the PCR reaction to amplify the 5′-end of the MT receptor transcript, with two nested tammar MT
receptor-specific primers (MTR 5’F1, 5’-GCG ACT TCT AGT AGT GTA GCC-3’ and MTR 5’R5, 5’-CAG ATC TAG TGG TGG CCTG TGT-3’). The PCR conditions were as follows: 40 cycles of 1-min denaturation at 95°C, 1 min at the annealing temperature of 55°C and a 2-min extension at 72°C. A further 15-min extension step was added at 72°C. This yielded a DNA product of approximately 350 bp, which was eluted, ligated into the pGEM-T vector (Promega), transformed into JM109 competent cells and sequenced as described previously (Parry et al. 1997).

**Southern blot analysis of genomic DNA**

Genomic DNA (15 µg) extracted from male liver tissue was digested with a single restriction enzyme (HindIII) and electrophoresed on a 0.5% agarose gel in 0.5 X TBE. After depurination in 0.2 M HCl, alkali denaturation and neutralisation, DNA was capillary transferred to nylon membranes (Hybond N+; Amersham Life Sciences, Castle Hill, Australia) in 20 X saline-sodium citrate (SSC) and UV cross-linked. Membranes were prehybridised for 3 h at 65°C in a solution containing 0.25 M NaHPO$_4$ (pH 7.2), 1 mM EDTA, 20% (w/v) sodium dodecyl sulfate (SDS) and 0.5% (w/v) blocking reagent (Roche), and hybridised overnight at 65°C with a 580 bp tammar MT receptor cDNA probe labelled with digoxigenin (0.35 mM; Roche) by PCR, according to the manufacturer’s instructions. After hybridisation, membranes were washed three times in 0.2 M NaHPO$_4$, 1 mM EDTA and 1% (w/v) SDS at 62°C. Hybridisation signals were subsequently detected by chemiluminescence with CPSD (disodium 3-(4-methoxy spiro[1,2-dioxetane-3,2'-5'-chloro] tricyclo[3.3.1.1$^{3,7}$]decan)-4-y1) phenyl phosphate) as substrate. Membranes were subsequently exposed to film at −80°C for 48 h.

**Northern hybridisation**

Total RNA was extracted from myometrial samples obtained on days 17, 23 and 25 of gestation, and also from mammary gland tissue on days 4–5 post-partum and liver on day 23 of gestation to use as positive and negative controls respectively. Approximately 20 µg total RNA was denatured in 20 X 3-morpholino-propanesulfonic acid (MOPS), 50% (w/v) formamide (Asia Pacific Specialty Chemicals, Seven Hills, Australia) and 2.2 M formaldehyde (Asia Pacific Specialty Chemicals) for 15 min at 65°C. Samples were then subjected to electrophoresis on a 1.3% (w/v) agarose/2.2 M formaldehyde/1 X MOPS gel and transferred to an optimised Hybond-NX membrane (Amersham Life Sciences) by overnight capillary transfer. Samples were then covalently attached to the membrane by UV cross-linking at 65°C for 2 h. An RNA ladder (0.24 9.5 kb) (Promega) was included on the gel as a molecular size marker. Specific MT receptor transcripts were identified with a 480 bp tammar MT receptor cDNA probe, derived by PCR and labelled with $[^{13}P]$ dCTP to a specific activity of 6.4 X 10$^8$ c.p.m./µg DNA by random primer extension. The membranes were prehybridised in buffer (0.25 M sodium phosphate, 1 mM EDTA and SSC: 0.3 M sodium citrate and 3 M NaCl, pH 7.0) at 65°C for 1 h. Denatured radiolabelled probe (45 000 c.p.m./µl) was then added to fresh buffer and incubated at 65°C overnight. After hybridisation, membranes were washed twice in 2 X SSC/0.1% (w/v) SDS at room temperature for 10 min. The final wash used 0.2 X SSC/0.1% (w/v) SDS at 65°C for 30 min. Membranes were then exposed to film (BIOMAX MS – $^{32}$P; Kodak Australia), with a single intensifying screen at −80°C for 48 h.

**Quantitative (Q-)PCR analysis**

For each sample, 600 ng total RNA was reverse transcribed in a 30 µl reaction containing 1 X TaqMan buffer, 5.5 mM MgCl$_2$, 500 µM dNTPs, 2.5 µM oligo d(T), 0.4 µM RNase inhibitor and 1.25 U/µl MultiScribe reverse transcriptase (Applied Biosystems, Scoresby, VIC, Australia). A second reaction mix using 30 ng total RNA from each sample and a series of myometrium RNA dilutions (100–0.001 ng) was prepared for the endogenous reference 18S ribosomal RNA PCR reactions and to generate the 18S standard curves respectively. First-strand cDNA synthesis for all samples was carried out simultaneously at 25°C for 10 min, 42°C for 45 min and 95°C for 10 min with a final cooling temperature at 4°C, before storage at −20°C. The 18S and tammar-specific MT receptor primers and FAM (6-carboxy fluorescein)-labelled probes were designed with Primer Express (Applied Biosystems) and provided by Keystone Division (Biosource International, Foster City, CA, USA), as described by Siebel et al. (2002b). All reactions were carried out in triplicate 25 µl volumes consisting of 1 X TaqMan Universal PCR Master Mix, 0.8 µM forward and 0.8 µM reverse MT receptor primers, 0.4 µM MT receptor probe and 2.5 µl cDNA template. Each plate included a sample in triplicate with water to replace the cDNA template (NTC) and a series of 18S standards with known RNA concentrations. The relative CT standard curve method was used in this study; therefore, MT receptor and 18S gene expression was assessed in separate PCR reactions with C$_T$ values for both genes related to those of the 18S standards. The amount of MT receptor mRNA expressed in each sample was calculated from regression lines generated from the standard curves and presented as MTR/18S mRNA (Siebel et al. 2002b).

**Radioreceptor assay**

Radioreceptor assays were carried out as described by Siebel et al. (2002b), using the labelled ligand [125I]-129-vasotocin (125I-OTA; ProSearch International Australia Pty Ltd, Malvern, VIC, Australia) with a specific activity of 2200 Ci/mmol. The receptor assay mixture consisted of duplicate aliquots of 100 µl diluted tissue suspension, 100 µl 129-OTA (15 000–20 000 c.p.m./tube) and 100 µl assay buffer (50 mM Tris–HCl, 5 mM MgCl$_2$ and 0.2% bovine serum albumin) and incubated at 37°C for 90 min. The receptor assay mixture was then added to a mixture of [125I]-vasotocin (125I-OTA) or 125I-vasopressin (125I-OP) and competed with increasing concentrations of unlabeled vasotocin (125I-vasotocin or 125I-vasopressin). The resulting binding was determined by filtration through glass microfibre filters (Whatman GF/B) and counting in a gamma counter (Beckman LKB Wallac, Australia). The apparent dissociation constant (Kd) was determined by Scatchard analysis.

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alumnum (BSA), pH 7.6) containing a range of 0.01–1 pmol/tube unlabelled OTA standards. Protein concentrations in the membrane fractions were measured with a DC Protein Assay Kit (Bio-Rad) with BSA as the protein standard. Protein concentrations were 60–120 μg/ml, which is within the range where specific binding is linearly correlated with protein concentration. Data were analysed by non-linear regression, using the Ligand computer program (Munson & Rodbard 1980), to obtain the binding affinity (Kd) and the receptor content (Rn) for radiolabelled ligand binding.

**Competitive binding studies**

Competitive binding assays determined the ligand specificity of the 125I-OTA binding site in the ovary. A 100 μl aliquot of membrane preparation obtained from pooled corpora lutea collected on day 25 of gestation was incubated with 100 μl 125I-OTA in competition with a series of ligand solutions of varying molar concentrations prepared in assay buffer. A 100 μl aliquot of the following ligands was used: OT (0.1, 0.5, 1, 5, 10 and 100 pmol/tube), MT (0.1, 0.5, 1, 5, 10 and 100 pmol/tube), AVT (0.1, 0.5, 1, 5, 10 and 100 pmol/tube), LVP (1, 5, 10, 50, 100 and 500 pmol/tube) and PP (10, 50, 100, 500, 1000 and 2000 pmol/tube) (all from Sigma Pharmaceuticals), as well as OTA (0, 0.01, 0.02, 0.1, 0.2, 1 and 100 pmol/tube) (kindly provided by Dr Maurice Manning). The interaction of each peptide with 125I-OTA was expressed as a relative displacement curve (B/B0) versus log molar concentration and fitted with sigmoidal curves (GraphPad Prism, San Diego, CA, USA). Two independent experiments were conducted with duplicates of each ligand to determine the respective IC50 values.

**Localisation of MT binding sites**

Tissue sections were first brought to room temperature before incubation for 30 min in washing buffer (50 mM Tris–HCl (pH 7.4)). Immediately after washing, sections were incubated with 100 μl 125I-OTA (50–100000 c.p.m.) in incubation buffer (50 mM Tris–HCl, 5 mM MgCl2 and 0.1% BSA) for 1 h at room temperature. Non-specific binding was determined by the addition of 2 μM unlabelled OTA peptide under the same conditions as for 125I-OTA binding. The incubation step was terminated by washing sections in ice-cold washing buffer. Sections were then air-dried and placed in the Fujix BAS IP magazine holder 2040 TR (Fujifilm, Mulgrave, VIC, Australia) apposed to a phosphoimager plate for 24 h. This was visualised with Analytical Imaging Station software on the Bio-imaging analyser (Fujix BAS 2000). For a clearer image, the sections were exposed to a film sensitive to 125I-radioactivity (MS film: Amersham Life Science) for 24–48 h at −80°C. The film was developed in an AGFA CP 1000 developer (AGFA-Gavaert Ltd, Nunawading, Australia) for localisation of the radioactivity. For a direct comparison between radiolabelling and tissue histology, serial sections were briefly fixed in 4% paraformaldehyde (PFA) and stained with haematoxylin and eosin by well-established procedures.

**Statistical analysis**

Data for both MT receptor mRNA and receptor concentrations did not show homogeneity of variance and were therefore log-transformed. To test for significant differences between stages, one-way ANOVA was used with a least squares difference post-hoc test (SPSS, Inc., Chicago, IL, USA) at the 95% confidence interval. Paired t-tests were performed on the log-transformed data to test for significant differences between the gravid and non-gravid uteri. All data are reported as mean±S.E.M.

**Results**

**MT receptor gene**

Sequencing analysis of the various PCR fragments obtained with 5’RACE with tammar MT receptor-specific primers produced a consensus nucleotide sequence of 834 bp (GenBank accession no. AY206419), which encodes a protein of 278 amino acids. This sequence accounts for the equivalent of the bovine/rat OT receptor exon 2, or mouse/human OT receptor exon 3 region. There was 100% sequence homology compared with the previous tammar MT receptor nucleotide sequence reported by Parry and Bathgate (1998). An additional 180 nucleotides were identified at the 3’-end, which starts approximately 33 amino acid residues into the amino-terminus. This sequence does not include the methionine start site, but comprises the first six transmembrane domains. 3’-RACE was attempted but was unsuccessful probably due to the large 3’-UTR of the MT receptor, as predicted from the size of the transcript in the Northern blot.

The derived 278-amino-acid sequence of the putative tammar MT receptor exon 2 region has approximately 80% homology compared with the human (Kimura et al. 1992), ovine (Riley et al. 1995) and bovine (Bathgate et al. 1995) OT receptor sequences. Highest similarity (96%) is seen in the transmembrane domains II and III and extracellular loop 1 (Fig. 1). In contrast, there is only 82% homology in the region containing the amino-terminus, the first transmembrane domain and the intracellular loop. The wallaby MT receptor also shows 72% and 78% homology to human vasopressin receptors, V1A and V1B respectively, but there is no significant similarity between the tammar MT receptor and the only complete vasotocin receptor sequence, that of the bullfrog (Rana catesbeiana). Hybridisation of HindIII-digested genomic DNA with a tammar MT receptor-specific probe indicated a specific band at approximately 4.1 kb (Fig. 2), demonstrating a single MT receptor gene in the wallaby genome.

**MT receptor gene expression**

Initial RT-PCR analysis revealed a single PCR product strongly expressed in the myometrium, corpus luteum,
cerebellum and hypothalamus of the brain. MT receptor gene transcripts were also weakly expressed in the heart, follicle and oviduct (data not shown). Northern hybridisation of total RNA demonstrated the presence of a MT receptor transcript in both gravid and non-gravid myometrium across a range of pregnancy stages and in the mammary gland (Fig. 3). The size of the transcript was approximately 4.1 kb, estimated from the size of the 28S (4.7 kb) and 18S (1.9 kb) ribosomal RNA bands and an RNA marker. Maximal expression was observed in the gravid myometrium during late pregnancy (Fig. 3, lanes 3 and 5), with a distinctive decrease in MT receptor transcript expression in the non-gravid myometrium (lanes 4 and 6). The signal intensity of the MT receptor transcript in the mammary gland (lane 7) from early lactation was relatively low and equivalent to that observed in non-gravid myometrium from day 23 of pregnancy. No MT receptor transcript was detected in the liver (lane 8), which was used as a negative control.

A comparison of MT receptor gene expression between different tissues by Q-PCR revealed that MT receptor mRNA levels were significantly higher in the gravid myometrium than the non-gravid myometrium (paired t-test, \( P = 0.017 \)), placenta (\( P = 0.023 \)), gravid endometrium (\( P = 0.011 \)) and median vagina (\( P = 0.007 \)) (Fig. 4). However, the highest expression of MT receptor mRNA was detected in the corpus luteum collected on day 23 of gestation, which was significantly higher than that of all other tissues obtained from animals on day 23 of gestation, including the gravid myometrium (\( P = 0.034 \)).

There was a significant (one-way ANOVA, \( P < 0.001 \)) increase in uterine MT receptor mRNA expression in the later stages of gestation (Fig. 5a). Myometrial MT receptor mRNA concentrations were significantly (\( P = 0.016 \)) upregulated in the gravid uterus on day 20 of gestation, and remained relatively high until the day of expected birth. Differences in MT receptor gene expression between the gravid and non-gravid uteri were observed on days 23–26 of gestation, with higher MT receptor mRNA concentrations detected in the gravid uterus. However, these differences were not significant until days 25 and 26 (\( P < 0.009 \)) of gestation. A significant (\( P = 0.003 \)) downregulation in MT receptor mRNA expression occurred in the non-gravid uterus at term compared with the non-gravid uterus at day 22 of gestation. Only six stages of gestation were available for the Q-PCR analysis of MT receptor mRNA expression in the corpus luteum. Overall, there was no significant (one-way ANOVA, \( P = 0.516 \)) difference in luteal MT receptor mRNA expression across pregnancy stages (Fig. 5b). However, a direct comparison between days 23 and 26 of gestation showed a significant (independent t-test, \( P = 0.006 \)) decrease in MT receptor mRNA expression on the day of expected birth.

**MT receptor concentrations**

A comprehensive study examined myometrial MT receptor concentrations in gravid and non-gravid uteri collected from day 11 of gestation, through to the expected day of birth (day 26), and then in post-partum tissues. Protein concentrations (\( \mu g/ml \) membrane preparation) did not change significantly during pregnancy, and the differences between uteri were negligible. Therefore, MT receptor concentrations were presented as fmol/mg protein to determine receptor changes in myometrial tissues. In mid-pregnancy, days 11–20, MT receptor concentrations were
The downregulation of MT receptors in the non-gravid myometrium was significant \( (P < 0.05) \) on day 23 of gestation, compared with the non-gravid myometrium on day 22. It was not until day 22 of gestation that myometrial MT receptor concentrations were significantly (paired \( t \)-test, \( P = 0.029 \)) higher in the gravid than in the non-gravid myometrium. There was also a significant difference in MT receptors between uteri on both days 23 and 26 \( (P < 0.01) \) of gestation (Fig. 6). On the day after birth (days 0–1 post-partum), MT receptor concentrations were significantly \( (P = 0.01) \) decreased in the post-partum myometrium when compared with the gravid uterus on day 26 of gestation. Receptor concentrations also remained low in the non-post-partum myometrium. There was no difference in MT receptor expression between uteri up to 1 week after birth.

Characterisation of corpus luteum MT receptors

The specificity of 125I-OTA binding in the corpus luteum of pregnant animals was assessed in competitive displacement experiments using different OT and AVP receptor agonists and antagonists (Fig. 7). These data show that AVT and MT bound with relatively high affinity to the 125I-OTA binding site, as did OT. In contrast, the two AVP receptor agonists LVP and PP had much lower binding affinities, as higher molar concentrations were needed to displace the 125I-OTA from the binding site. In general, the ligand affinities for the 125I-OTA binding site in the corpus luteum were in the following order of decreasing affinity: AVT = MT > OT > LVP ≫ PP and indicate binding to an OT-like receptor, as in previous results in the myometrium (Siebel et al. 2002b). At the one stage of gestation examined (day 25), luteal MT receptor concentrations were 140–435 fmol/mg protein \( (n = 2) \). Insufficient tissues were available to assess ovarian MT receptor concentrations throughout gestation.

Localisation of MT binding sites in the tammar uterus

MT binding sites were localised specifically to the myometrium in both the gravid (Fig. 8A) and non-gravid (Fig. 8D) uterus, with intense 125I-OTA labelling in the myometrial smooth muscle layer. No labelling was present in the endometrium, as confirmed by histological analysis (Fig. 8C and F). In the negative control, 2 \( \mu \)M OTA displaced 125I-OTA binding in the tammar uterus (Fig. 8B and E).

Discussion

This study reported a partial 834-bp sequence of the tammar MT receptor cDNA. The derived sequence of 278 amino acids comprises six of the seven transmembrane domains and starts approximately 33 amino acid residues into the amino-terminus. Further comparisons with OT receptor sequences from eutherian species have highlighted several regions of highly conserved amino acids.
in the OT/MT receptor that may be important for its biological action. Of particular interest are the highly conserved regions in transmembrane domains II, III and VI. This class of G-protein-coupled receptors typically binds ligand within the transmembrane domains, confirming their importance for ligand binding and selectivity (Kimura & Ivell 1999). The transmembrane domains may also influence the ligand selectivity of signal transduction of the OT receptor (Chini et al. 1996). Experiments introducing point mutations into selected transmembrane domains have confirmed that there are different contact sites within the receptor molecule for the ligand and ligand-specific antagonists (Yarwood et al. 1997). Many of the key amino acids shown by mutagenesis to be important in ligand binding appear to be present in the tammar MT receptor sequence.

Figure 3 Northern hybridisation using a specific tammar MT receptor cDNA probe and total RNA (20 μg) from gravid (GR) and non-gravid (NG) myometrium (myo) on days 17, 23 and 25 of gestation, with mammary gland (m. gland) and liver as a negative control. The approximate sizes of the 28S and 18S RNA, and the tammar MT receptor transcript (4.1 kb) are indicated.

Figure 4 Mean ± S.E.M. MT receptor mRNA concentrations in reproductive tissues from day-23 pregnant females (n = 4). a, significantly (P < 0.05) higher than all tissues; b, significantly (P < 0.05) higher than all tissues, except corpora lutea (CL). gr myo: gravid myometrium; ng myo: non-gravid myometrium; gr endo: gravid endometrium; m vag: median vagina; YSM: yolk sac membrane.

Figure 5 Mean ± S.E.M. MT receptor mRNA concentrations in the (a) myometrium of the gravid and non-gravid uterus and (b) corpora lutea from day 17 of gestation to the day of birth (day 26; n = 4 at each stage). a, P < 0.05 higher than the gravid myometrium on days 17–18; b, P < 0.05 higher than the non-gravid uterus; c, P < 0.05 lower than day-22 non-gravid myometrium; d, P < 0.05 lower than day-23 corpora lutea.
Northern analysis confirmed the expression of a single MT receptor transcript of approximately 4.1 kb in both the myometrium and mammary gland of the tammar wallaby. In the human, there are two distinct OT receptor transcripts, approximately 3.6 kb in the mammary gland and 4.4 kb in ovary, uterine endometrium and myometrium (Kimura et al. 1992). In the rat, the OT receptor gene is highly expressed in the uterus and gives rise to three separate polyadenylation variants, with the predominant transcript detected at 6.5 kb (Rozen et al. 1995). Two major transcripts were identified in the bovine myometrium and endometrium at 2.0 kb and 6.5 kb, with a third minor band at 3.5 kb (Bathgate et al. 1995). Usually, these variants differ in the length of the 3'-UTR, due to different polyadenylation sites. However in the tammar, there were no apparent polyadenylation variants of the MT receptor gene after exposure of the blot to film for up to 3 days.

Q-PCR demonstrated high expression of MT receptors in the uterus and ovary of the tammar wallaby. As in previous RT-PCR results, MT receptor mRNA expression in the median vagina and placenta of the tammar was almost undetectable. Supposedly, then, the median vagina does not respond to MT in the tammar. This is in contrast to the brushtail possum, where MT receptors are present in both the uterus and median vaginal sacs, implying that these tissues respond to MT as a single contractile unit (Sernia et al. 1991). The highest expression of the MT receptor transcript was detected in the tammar myometrium 4–5 days before birth and, for the first time, in the corpus luteum of pregnancy. In the cow, OT receptor gene transcripts are detected in most uterine tissues, with the highest levels of expression in the endometrium and myometrium at term. However, no OT receptor gene transcripts have been detected in the corpus luteum at any stage of pregnancy (Ivell et al. 1995). Similarly, the rodent myometrium is abundant in OT receptor mRNA during pregnancy (Larcher et al. 1995), but there are no OT receptors in the corpus luteum.

The radioreceptor assay data in this study demonstrated significant increases in MT receptor concentrations in the gravid myometrium as early as day 21 of gestation. Differences between the uteri occurred on days 22 and 23 of gestation, as well as on the day of birth. However, receptor concentrations do not precisely parallel MT receptor mRNA expression. Although there was a marked gravid uterus-specific increase in MT receptor mRNA concentrations on day 20 of gestation, differences in MT receptor mRNA expression between uteri were not significant until day 25 of gestation. This appeared to be due to a large variation in MT receptor mRNA concentrations in the gravid myometrium. In support of previous findings, there is a significant downregulation in both MT receptor mRNA and receptor concentrations in the non-gravid myometrium on day 25 of gestation (Parry et al. 1997). The upregulation of MT receptors in the gravid myometrium relates to the differential uterine responsiveness to exogenous MT between the gravid and non-gravid myometrium in late pregnancy (Parry et al. 1997). An increase in MT receptors in the gravid myometrium at the end of pregnancy is consistent with the idea that MT stimulates uterine contractions at birth. This stimulatory effect appears to be essential for normal parturition, as infusion

Figure 7 Competitive displacement of 125I-OTA binding in the corpus luteum on day 25 of gestation in the tammar wallaby by d(CH2)5 [Tyr(Me)2, Tyr4, Orn8, Tyr-NH2]-vasotocin (OTA), mesotocin (MT), oxytocin (OT), arginine [Arg-8]-vasotocin (AVT), lysine [Lys-8]-vasopressin (LVP) and [Phe2]-vasopressin (PP).

Figure 6 Mesotocin receptor concentrations in the myometrium of the gravid and non-gravid uterus from day 11 of gestation to 2 days post-partum (pp). Data are mean ± S.E.M. (fmol/mg protein, n = 2–4 at each stage). a, significantly (P < 0.05) higher than gravid myometrium on days 16–18 of gestation; b, significantly (P < 0.05) higher than non-gravid uterus; c, significantly (P < 0.05) lower than day-22 non-gravid myometrium; d, significantly (P < 0.05) lower than day-26 gravid myometrium.
of an OT receptor antagonist in late-pregnant tammars delays birth (Renfree et al. 1996), and an increase in plasma MT is observed only during delivery (Parry et al. 1996). Therefore, evidence suggests that MT is an important part of the hormonal cascade associated with delivery in this species.

For further characterisation of uterine MT receptor expression in the tammar, specific MT binding sites were localised to the myometrium of both the gravid and non-gravid uterus. Previous studies in sheep also reported specific OT binding to smooth muscle cells of the myometrium, as well as endometrial tissue and the oviduct with $^{125}$I-OTA (Ayad et al. 1991). Overall, the autoradiography studies in the tammar support the MT receptor assay and gene expression data, which demonstrated higher MT receptor concentrations in the gravid myometrium than in the non-gravid myometrium and endometrium during late pregnancy.

In the present study, relative MT receptor mRNA expression was highest in the corpus luteum, even when compared with the gravid myometrium. However, it is not clear whether this is due to an increase in the number of cells expressing MT receptor mRNA or to an upregulation of MT receptor mRNA expression within each cell (Ivell et al. 2001). Pharmacological studies demonstrated the presence of a single high-affinity OT-like receptor in the corpus luteum of the pregnant tammar wallaby. The affinity ($K_a$) of this receptor for $^{125}$I-OTA is similar to that of the MT receptor in the myometrium.

Functional OT receptors have been detected in bovine granulosa cells, suggesting that OT may play an autocrine role in follicular growth (Okuda et al. 1997). In contrast, there is significant OT gene expression in the bovine corpus luteum, but no OT receptors, following the onset of labour. A suggested paracrine role for OT has been investigated within the primate ovary with the localisation of OT and OT receptors in luteal tissue and ovarian remnants of the marmoset monkey (Einspanier et al. 1994, 1997). This was the first evidence of the potential involvement of OT in the induction of luteinisation in any eutherian species.

Early and mid-luteal phase corpora lutea of the brushtail possum contain MT peptide concentrations similar to that of OT in the non-ruminant corpus luteum (Sernia et al. 1994). The presence of immunoreactive MT in a substantial number of an OT receptor antagonist in late-pregnant tammars delays birth (Renfree et al. 1996), and an increase in plasma MT is observed only during delivery (Parry et al. 1996). Therefore, evidence suggests that MT is an important part of the hormonal cascade associated with delivery in this species.

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population of cells is reminiscent of the situation in the sheep, where OT expression is restricted to the large luteal cells (Rodgers et al. 1983). Early work found no MT or OT protein in extracts of corpora lutea from pregnant tammar wallabies (Curlewis et al. 1988). However, evidence of an MT gene transcript has been shown in the prevulatory follicle and corpus luteum (Parry et al. 2000). These studies confirm that the tammar ovary has the ability to synthesise MT and suggest that an ovarian OT physiology was present early in the evolution of mammals.

The mechanism of luteal regression in the tammar wallaby is unknown. Q-PCR analysis of corpora lutea collected from late-pregnant tammars confirmed that MT receptor mRNA expression remains relatively high throughout most of gestation. However, at term, there was a decrease in MT receptor mRNA expression coinciding with luteolysis. These data support the hypothesis that a fall in ovarian MT receptor expression may be required for luteal regression.

In conclusion, MT binding sites were localised to the myometrium of the uterus in the pregnant tammar wallaby. Uterine MT receptor mRNA and receptor concentrations increased significantly in the gravid myometrium throughout most of gestation. However, at term, there was a significant decrease observed on the expected day of birth. MT receptor mRNA expression in the uterus and ovary of the tammar wallaby is under significant decrease observed on the expected day of birth. Luteal MT receptor mRNA expression was relatively high throughout pregnancy, with a significant decrease observed on the expected day of birth. It therefore appears that MT receptor expression in the uterus and ovary of the tammar wallaby is under differential regulation, with the feto-placental unit having a major influence on myometrial MT receptors.

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