Aromatase is abundantly expressed by neonatal rat penis but downregulated in adulthood

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Abstract

Although synthesis of estrogen by male gonads has been well documented for over half a century, it is only recently that the role of estrogen in male reproductive events has gained appreciation. We recently reported abundant expression of estrogen receptor (ER)-α and -β in different cell types of the rat penis, whose levels diminished with advancing age. The present study, which builds on data from the ER study, was designed to determine whether the penis is capable of generating its own local estrogen by examining evidence of the expression of aromatase, a microsomal enzymatic complex which irreversibly converts androgens to estrogens, using immunohistochemistry, Western blotting, in situ hybridization and real-time PCR analyses. Secondly, the effects of sex steroid hormones on penile aromatase were examined. Discrete aromatase immunoreactive cells were localized in primordial corpus cavernosum, corpus spongiosus and os penis, blood vessels and sensory corpuscle of glans penis. In situ hybridization signals corresponded with immunohistochemical findings. Western blot, enzyme immunoassay and real-time PCR analyses of rat penile samples revealed an age-dependent expression of aromatase and estrogen, with levels at week 1 almost resembling those of the ovary, but they decreased sharply by week 8, and decreased further by week 35. This expression pattern was strikingly similar to that of ER-α reported previously. Testosterone and diethylstilbesterol administered prenatally upregulate levels of aromatase mRNA and protein, and estrogen postnattally. Dihydrotestosterone upregulated aromatase mRNA and protein, but not estrogen. We conclude that estrogen acts via ER in a paracrine and/or autocrine manner to regulate penile events, particularly during development, and that estrogen synthesis is regulated by estrogen and androgens.

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Introduction

The first report documenting synthesis of estrogen by male gonads was published over half a century ago (Zondek 1934, Goldzieher & Roberts 1952, Leach et al. 1956, Morse et al. 1962, Attal 1969, Raeside & Renaud 1983, Carreau et al. 1999). However, it is only in the past decade that interest in the role of estrogen in male fertility has increased, largely due to reports that exposure to environmental estrogens may have detrimental effects on male reproductive development and health, and that deletion of the estrogen receptor (ER)-α gene leads to impairment of male fertility (Greene et al. 1939, Meistrich et al. 1977, Stillman 1982, Carreau et al. 1999, McKinell et al. 2001). The role of estrogen in male fertility could include regulation of germ cell development or spermatogenesis, as reviewed by recent studies using mice lacking aromatase (Robertson et al. 1999).

In the male, estrogen is produced mainly by the testis, and to a lesser extent, by the brain and adrenal gland, in quantities overall far less than in the female (Carreau et al. 1999). However, although the ovaries, overall, produce more estrogen than the testes, it is interesting to note that local tissue concentrations of estrogen in male reproductive tissues, such as the efferent ducts, exceed the levels
of circulating estrogen in females (Free & Jaffe 1979). Equally interesting are recent revelations that circulating estrogen levels in adult males resemble those found in females during the early follicular phase and that levels in postmenopausal women are less than in age-matched males (Longcope et al. 1990, de Ronde et al. 2003).

Clearly, estrogen exerts strong effects on male reproductive events. These effects are principally mediated by the classical ER, ER-α and the recently discovered ER, ER-β – members of the steroid/thyroid nuclear receptor superfamily which regulate gene transcription through estrogen-responsive elements (Koike et al. 1987, Kuiper et al. 1996). ER has been localized in the gonads and tracts of both males and females (Hess et al. 1997a, Sar & Welsch 1999, Mowa & Iwanaga 2000, 2001a,b, Jesmin et al. 2002). In male reproductive tissues other than the testis, the most intense signals of ER-α mRNA in adult rodents were localized in the epithelia of the efferent duct and initial segment of the epididymis, sites responsible for sperm concentration (Hess et al. 1997, Mowa & Iwanaga 2001a). ER-α mRNA expression is also intense in the muscle layer of the vas deferens of the rat, while ER-β mRNA is weakly expressed in the efferent duct, epididymis and vas deferens (Hess et al. 1997b, Mowa & Iwanaga 2001a). We recently described the existence of ER-expressing cells in the penis, whose levels were expressed age-dependently, with the most intense signals observed during the perinatal period, but declining thereafter with age (Jesmin et al. 2002). Initial signals of ER-α were localized to the mesenchyme and subepithelial stroma, and later (postnatal day 2) to the corpus spongiosus, corpus cavernosum and urethral epithelia (Jesmin et al. 2002). ER-β was initially detected by postnatal day 2 and was localized diffusely in corpus spongiosus and cavernosum in immature rats (Jesmin et al. 2002).

Because circulating levels of estrogen are low in the male during the perinatal period, we sought to determine whether the penis was the major source of estrogen acting on ER-expressing penile cells by examining the expression of aromatase. The cytochrome P450 aromatase (P450 arom) is a microsomal enzymatic complex present in the endoplasmic reticulum of various vertebrate tissues and it irreversibly converts the aromatizable androgens, such as testosterone, to estrogens (Lephart 1996, Pereyra-Martinez et al. 2001, Carreau et al. 2002, Wiszniewska 2002). Aromatase is expressed in a tissue-specific manner and is composed of two proteins - a ubiquitous NADPH-cytochrome P450 reductase and a cytochrome P450 aromatase - which contain the heme and the steroid pocket (Lephart 1996, Carreau et al. 1999, 2002).

The present study localized aromatase in the penis; like ER, aromatase was expressed in an age-dependent manner. Sex steroids administered prenatally upregulate levels of aromatase mRNA and protein and estrogen postnatally.

Materials and methods

Animals and treatments

Male Wistar rats at postnatal days 1 and 3 (n=30 for each) were anesthetized by gaseous diethyl ether, killed and had their penises carefully harvested for Western blot analyses, immunohistochemistry, in situ hybridization and real-time PCR experiments. In addition, Wistar rats aged 1, 8 and 35 weeks were also anesthetized, killed and the penises were carefully harvested for Western blot analyses, ELISA and real-time PCR (n=10 for each). To determine the effects of varying estrogen:androgen ratios on aromatase expression, rats at day 10 of pregnancy were injected subcutaneously every alternate day until parturition with one of the following sex steroid hormones, dissolved in 0.1 ml sesame oil: (1) diethylstilbestrol (DES; Sigma, St Louis, MO, USA) 10 µg/rat (high dose), 0.1 µg/rat (low dose); (2) dihydrotestosterone propionate (DHT; Sigma) 12.5 mg/rat; and (3) testosterone propionate (Sigma) 1 mg/rat (n=3 mothers for each treatment). Pregnant rats, of equivalent gestational age (day 10), served as controls and were treated with an equivalent volume (0.1 ml) of sesame oil. After birth, the neonatal rats (pups) were treated at postnatal day 0 and day 2 with the sex steroid hormones (DES: 0.005 µg/pup (low dose), and 0.5 µg/pup (high dose); DHT: 750 µg/pup; testosterone: 50 µg/pup) dissolved in 50 µl sesame oil. Age-matched control pups were treated with an equivalent volume (50 µl) of sesame oil and processed as the treated groups. All the pups were killed on the third day after anesthetizing them with gaseous diethyl ether.
Penile tissues were carefully harvested and processed as described below. Ovaries from pregnant or adult rats were used as positive control tissue for aromatase. All procedures were carried out in accordance with the regulations laid down by the Hokkaido University School of Medicine Animal Care and Use Committee.

Characterization of antisera

A number of antisera of aromatase were initially tested in rat penile tissues for immunohistochemistry. In our hands, CYP19 C-16 (Santa Cruz Biotechnology, Santa Cruz, CA, USA), an affinity purified goat polyclonal antibody raised against a peptide at the carboxyl terminus of human CYP19, and OBMMCA2077 (Oxford Biomarketing, Oxford, UK), a mouse anti-human cytochrome P450 synthetic peptide corresponding to amino acids 376–390, produced the most consistent robust signals. Thus, these antisera were used in the present study. Controls included omission of the primary antiserum, and omission of the secondary antibody.

Immunofluorescence staining

For immunohistochemical studies of aromatase, tissue specimens were fixed in 4% buffered formalin solution, dehydrated and then embedded in paraffin. The tissues were sliced in 4 µm sections transversely, deparaffinized and treated for 10 min with citrate buffer (10 mM citric acid, pH 6·0) in a microwave oven (750 W) before immunostaining. In another set of experiments, frozen tissue samples were cut by cryostat in 8 µm sections, fixed in acetone and air-dried. To prevent non-specific binding by secondary antibody, the sections were blocked by non-immune serum (1% bovine albumin in Tris) for 30 min at room temperature. After overnight incubation at 4 °C with primary antibodies, the sections were rinsed in phosphate buffer solution and then exposed to the fluorescence secondary antibody, Cy3-conjugated AffiniPure anti-mouse IgG or fluorescein-conjugated AffiniPure rabbit anti-goat IgG (Jackson ImmunoResearch Laboratories, Westgrove, PA, USA) for 2 h, according to the manufacturer’s instructions. The samples processed without primary antibodies served as negative controls. Immunofluorescence images were observed under a Laser Scanning Confocal Imaging System (MRC-1024, Bio-Rad Laboratories, Hemel Hempstead, Herts, UK). In our hands, the intensity of aromatase immunoreactivity in frozen tissue sections was stronger compared with paraffin-embedded tissues.

Western blot analysis

After careful sampling of the penises and ovaries, the tissues (penises and ovaries) were rinsed in phosphate-buffered saline (PBS) on ice, minced with scissors, homogenized, and then centrifuged at 1000 g for 15 min to pellet any insoluble material. The total protein concentration of the supernatant was determined by the method of Lowry et al. (1951), with BSA as standard. Samples (10 µg) were run on SDS-PAGE, using 8% polyacrylamide gel, and electrotransferred to a polyvinylidene difluoride filter (PVDF) membrane. To reduce non-specific binding, the PVDF was blocked for 60 min at room temperature in Tris-buffered saline (TBS: 20 mM Tris-HCl, 500 mM NaCl, pH 7·5) containing 1% albumin. Thereafter, the PVDF was washed three times for 5 min in TBS-Tween buffer (TTBS: 20 mM Tris-HCl, 500 mM NaCl, 0·05% Tween 20, pH 7·5) and incubated overnight at 4 °C with specific antibody (1:200 dilution for aromatase) in TTBS containing 1% albumin. After extensive washing with TTBS, the PVDF was incubated with horseradish peroxidase-conjugated anti-mouse or anti-goat antibody (Amersham, UK) diluted at 1:2000 dilution for aromatase) in TTBS containing 1% albumin. After extensive washing with TTBS, the blots were visualized using the enhanced chemiluminescence detection system (Amersham), exposed to X-ray film for 5 min, and analyzed by image analysis software from NIH produced by Wayne Rasband (National Institute of Health, Bethesda, MD, USA; http://rsb.info.nih.gov/nih-image/download.html). The results are expressed as a percentage of the band obtained from the ovary in each experiment. To check for protein loading/transfer variations, all blots were stained with Ponceau Red (washable, before incubation with antibodies) and with Coomassie Brilliant Blue. The intensity of total protein bands per lane was evaluated by densitometry. Negligible loading/transfer variation was observed between samples.
Generation of the aromatase polyclonal antibody and tissue ELISA

The ELISA technique was used to test for penile aromatase and estrogen concentrations to supplement Western blot data. Because there is no aromatase antibody suitable for determining local aromatase concentration in tissues using ELISA, we developed a new antibody specific for aromatase. Mice were immunized with the peptide CALEDDVIDGYPKKG, representing amino acid residues 376–390 of human aromatase, and serum was collected and titrated. Antibodies were affinity-purified over Amino-Link columns (Pierce Biotechnology, Rockford, IL, USA), conjugated with immunogen and eluted with 0·1 M glycine, pH 3. The new aromatase antibody generated was validated by Western blot. Observation of a single band at the expected molecular mass allowed the use of the antibody in ELISA. Moreover, the specificity of the aromatase antibody was further verified by isolation of an aromatase peptide (Immunobion, Sapporo, Japan), which was then used for the pre-adsorption test. The newly generated antibody was purified by peptide affinity column and then used for ELISA. Elutes were dialyzed against PBS and kept frozen in aliquots at −80 °C. Biotinylated primary antibodies were prepared using Biotin-(AC5)2-Osu (Dojin, Kumamoto, Japan). Samples, including standards of known target protein concentration and unknowns, were pipetted into wells coated with specific antibody. A second biotinylated antibody IgG was added. Samples were incubated for 4 h at room temperature and then aspirated to remove any excess or unbound biotinylated antibody. The enzyme streptavidin-peroxidase, which binds to the target protein-bound biotinylated antibody, was added. After a second incubation and washing to remove any unbound enzyme, substrate solution was added. This solution acts on bound enzyme to produce color. Absorption of the colored product was measured by spectrophotometry.

In situ hybridization

Specific 60-basepair (bp) long oligonucleotides complementary to either a 5′-region, i.e. 221–281 bp (Aro 5′ ologo, 5′-CCA-CAA-GAA-TCT-GCC-ATG-GGA-AAT-GAG-AGG-CGC-GAT-TCC-CAG-ACA-GTA-GCC-AGG-ACC-TGG-3′), or the heme-binding region, i.e. 1351–1411 bp (Aro 3′ ologo, 5′-CAC-CAT-GGC-GAT-GTA-CTT-CCC-AGC-ACA-GGC-TCC-GGC-CCA-AAG-CCA-AAT-GGC-TGA-AAA-TA-3′), of the rat cytochrome P450 aromatase cDNA (Lauber & Lichtensteiger 1994) were purchased. Both probes produced almost the same signal, but the latter probe produced the most robust and consistent signal. The oligonucleotides were labeled with 35S-dATP, using terminal deoxyribonucleotidyl transferase (Promega, Madison, WI, USA) at a specific activity of 0·5 × 10^9 d.p.m./µg DNA.

Tissues (penis or ovary) were rapidly removed and frozen in liquid nitrogen. Cryostat sections, 15–20 µm in thickness, were prepared and mounted on glass slides precoated with 3-aminopropyltriethoxysilane. The in situ hybridization protocol used has been described in detail previously (Mowa & Iwanaga 2000). Briefly, tissue sections were fixed in 4% paraformaldehyde for 10 min and acetylated with 0·25% acetic anhydride in 0·1 M triethanolamine-HCl (pH 8·0) for 10 min. Slide-mounted sections were prehybridized for 2 h in a buffer containing 50% formamide, 0·1 M Tris-HCl (pH 7·5), 4 × SSC (1 × SSC; 150 mM NaCl and 15 mM sodium citrate), 0·02% Ficoll, 0·02% polyvinylpyrrolidone, 0·02% bovine serum albumin, 0·6 M NaCl, 0·25% SDS, 200 µg/ml tRNA, 1 mM EDTA and 10% dextran sodium sulfate. Hybridization was performed at 42 °C for 10 h in the prehybridization buffer supplemented with 10 000 c.p.m./µl 35S-labeled oligonucleotide probes. The slides were washed at room temperature for 20 min in 2 × SSC containing 0·1% sarkosyl and twice at 55 °C for 40 min in 0·1 × SSC containing 0·1% sarkosyl. The sections were either exposed to Hyperfilm-βmax (Amersham) for 4 weeks or dipped in Kodak NTB2 nuclear track emulsion and exposed for 4–8 weeks.

Specificity of oligonucleotide probe

The specificity of the oligonucleotide probes for in situ hybridization was determined by labeling 15-µm cryostat sections of ovary from pregnant rats as positive control tissue. Corpora lutea of pregnant rats possess abundant levels of aromatase mRNA (Lauber et al. 1994). Furthermore, the specificity of the probes was confirmed by the disappearance of signals when excess doses of the corresponding non-labeled (35S-dATP) anti-sense oligonucleotides
(cold) were added to the labeled anti-sense oligonucleotides (hot) hybridization fluid. Consistent aromatase mRNA signals above background levels were considered positive and were subjectively scored as weak, moderate or strong.

**RNA preparation and real-time quantitative PCR**

Total RNA samples were prepared from tissues by the guanidinium thiocyanate-phenol-chloroform single-step extraction method with Isogen (Nippon Gene, Toyama, Japan), used routinely in our laboratory (Matsuda et al. 1999). After isolation, treatment with DNase I and quantification, RNA was reverse-transcribed to cDNA by the use of a ReverTra Ace (TOYOBO, Osaka, Japan). The single-stranded cDNA was then used in real-time quantitative PCR to evaluate relative expression levels of aromatase mRNA. DNA amplification was performed in the Applied Biosystems (ABI 7900HT) real-time PCR machine using the GeneAmp 7900HT Sequence detection system software (Perkin-Elmer Corp., Foster City, CA, USA) and the detection was determined by measuring the binding of the fluorescence dye SYBR Green I to double-stranded DNA. The PCR reactions were set up in microtubes in a volume of 20 µl. The reaction components were 2 µl cDNA synthesized in 10 µl of 2 × SYBR Green master mix (Perkin-Elmer Corp.) and 0·4 µM of each pair of oligonucleotide primers, as described above. The program was performed as follows: an initial step at 95°C for 10 min, and then 40 cycles of 95°C for 15 s and 60°C for 60 s. Regression curves were drawn for each sample and the relative amount of aromatase mRNA was calculated from the threshold cycles with the instrument’s software (SDS 2·0), according to the manufacturer’s instructions. The PCR products were analyzed by gel electrophoresis, to confirm the specificity of the generated products. Relative expression levels of aromatase were normalized to the geometric mean of the two internal control genes, α-fib and glyceraldehyde-3-phosphate dehydrogenase (G3PDH). The primer sequences for real-time PCR were obtained from the Genebank, accession number NM-017085 (forward, CCT GGA GA TGA CGT GAT TG, position 1189; reverse, CGA TGT ACT TCC CAG CAC AG, position 1385), yielding a product of 197 bp.

**Measurement of 17β-estradiol in penile tissues by enzyme immunoassay (EIA)**

For determining tissue estrogen levels, the penile and ovarian tissues were washed in cold PBS just after tissue harvest and snap frozen on dry ice. The tissues were homogenized in 0·1 molar (M) Tris-HCl (pH 7·2) buffer by a high speed homogenizer, at a consistent ice-cold temperature, centrifuged at 1000 g for 15 min to pellet any insoluble material, and thereafter the protein concentrations were measured. Finally, the local tissue concentrations of 17β-estradiol were quantified using an enzyme immunoassay kit (ESTRADIOL EIA, PANTEX, Santa Monica, CA, USA), according to the manufacturer’s instructions. The specificity of this kit has previously been confirmed in our laboratory using various types of rat tissue including serum, plasma, ovary, brain and penis. The values obtained were compared with those of the Rodent Estradiol ELISA Test Kit (Endocrine Technologies, Inc., Newark, CA, USA). The lowest level of 17β-estradiol detectable by the kit used in this study is 4·6 pg/ml. Intra-assay and interassay coefficients of variation (%CV) were 5·9 and 5·2 respectively.

**Statistical analysis**

Data are presented as means ± S.D. Statistical assessment of the data was performed by ANOVA with multiple comparisons by Fisher’s protected least significance t-test. P values less than 0·05 were considered significant.

**Results**

**Characterization of aromatase antisera and immunofluorescence**

Immunofluorescence staining for aromatase protein showed that its expression was localized to the penis spongiosus, penis cavernosus, urethra, sensory corpuscle of glans penis, blood vessel, dorsal nerve and primordial os penis at postnatal day 1 (Fig. 1A–K). In penile spongiosus, aromatase was expressed in both stromal and muscle cells. The observed immunoreactivity of aromatase was considered specific because primary antibodies (data not shown) or secondary antibodies alone showed no staining in the penile spongiosus, cavernosus or urethra. A representative photograph
Figure 1 (A) to (D).
Figure 1 (E) to (K).
of urethra (u) at postnatal day 1 obtained by omission of the primary antibody served as a negative control (Fig. 1I). Furthermore, sections of ovaries from adult female pregnant rats stained with aromatase showed discrete immunoreactivity in the corpus lutea and granulosa. It is noteworthy that, in both ovary and penis, aromatase immunolabeling was localized in cytoplasm, as expected, and not in the nucleus (Fig. 1A–K).

**In situ hybridization**

**In situ** hybridization studies showed intense to moderate and diffused signals of aromatase mRNA in spongiosus, cavernosus, urethra, glans penis, sensory corpuscle of glans penis and neurovascular bundles of postnatal day 3 penis (Fig. 2). Both basal and luminal epithelium of the urethra showed aromatase mRNA expression. The most intense signals were observed in penile spongiosus, penile cavernosus, glans penis and sensory corpuscles of glans penis. Only moderate amounts of aromatase mRNA were detected in neurovascular bundles and urethra.

**Western blot analysis**

Immunoblot analysis of the ovary and penises showed a single band with a molecular mass of ~54 kDa, which matched the molecular mass of aromatase (Fig. 3). Quantification of the immunoblot bands by densitometry revealed a decrease in the levels of penile aromatase protein with age. When levels of penile aromatase protein were expressed as a percentage of aromatase protein of the ovary, they were 85 ± 11% at 1 week ($P<0.001$), 37 ± 7% at 8 weeks ($P<0.001$) and 13 ± 4% at 35 weeks of postnatal age ($n=5$; $P<0.001$) (Fig. 3). These findings were also confirmed by immunofluorescence (data not shown). Moreover, no band was observed when aromatase antiserum was pre-adsorbed with the immunizing peptide (data not shown).

**Real-time PCR**

Consistent with the results of immunoblot, aromatase gene expression in penile tissues was decreased in a time-dependent manner (Fig. 4B). Ovarian tissue was used as a positive control. **In situ** hybridization experiments also showed an age-related decrease of aromatase mRNA expression in penile tissues (data not shown).

**Quantitation of aromatase and 17β-estradiol in penile tissues by ELISA and EIA**

The ELISA technique, using antibody generated in our laboratory, also showed a time-dependent decrease in levels of penile aromatase protein, consistent with the data obtained with immunoblot described above. As with Western blots, the pregnant ovary was used as a positive tissue (Fig. 4A). Furthermore, we measured the tissue concentration of estrogen in the penis to determine whether penile aromatase was functional or was capable of irreversibly converting androgens to estradiol. The estrogen concentration in the 1-week-old penis was 23 ± 6 pg/mg compared with 46 ± 8 pg/mg in the positive control tissue (ovary). By week 8 of age, the penile estrogen concentration dropped by about a half, to 13 ± 3 pg/mg, and by threefold of week 1 levels at 35 weeks of age (9 ± 3 pg/mg) (Fig. 4C), a trend which interestingly resembles that of aromatase concentrations.

**Effects of sex steroid hormone treatment on levels of penile aromatase**

Treatment of pregnant mothers from day 10 (10 µg, on alternate days) and neonates from birth...
Figure 2 (A).
Figure 2 (B) to (D).
Figure 2 (E) to (J).
to day 3 (0.5 µg, on alternate days) with high doses of DES caused an increase of 30 ± 3% in aromatase protein expression \((n=5; \ P < 0.001)\) compared with vehicle-treated rats (Fig. 5). However, low doses of DES (1 µg for mother, 0.005 µg for neonate) had insignificant effects on penile aromatase expression. Similarly, treatment with androgens (DHT and testosterone) up-regulated aromatase expression in the penis by 70 ± 7% (DHT) and 40 ± 6% (testosterone) (3-day-old penis) (Fig. 5).

The effects of treatment with the potent synthetic estrogen, DES, and with the androgens, DHT and testosterone, on aromatase gene expression in the 3-day-old penis corresponded with data obtained by immunoblot analysis, as revealed by real-time PCR above – i.e. high doses of DES increased the levels of the aromatase gene, whereas low doses had no effect; both testosterone and DHT upregulated aromatase gene expression (Fig. 6A).

High doses of DES markedly increased estrogen levels \((39.3 ± 7 \text{ pg/mg})\) in penile tissues compared with control \((17 ± 4 \text{ pg/mg})\), whereas low DES upregulated estrogen only slightly \((20.6 ± 4 \text{ pg/mg})\), consistent with data shown above. Surprisingly, although DHT upregulated aromatase protein and mRNA levels, it had no effect on tissue levels of penile estrogen \((17.4 ± 3.6 \text{ pg/mg})\) (Fig. 6), whereas testosterone upregulated estrogen concentrations in the penis \((26.7 ± 5 \text{ pg/mg})\), consistent with its effects on aromatase expression.

Discussion

The present study provides evidence of local estrogen production in the penis expressed age-dependently using Western blots, enzyme immunoassay, real-time PCR, immunohistochemistry and in situ hybridization. Aromatase protein and mRNA are (1) localized in a variety of cells found in different penile compartments, (2) expressed age-dependently in the penis, declining with advancement of age, in parallel with tissue estrogen levels, and (3) upregulated by exogenous sex hormone steroids (testosterone, DHT, DES). Interestingly, these findings correlate with our recent report on the penile ER and confirm our earlier speculation that a local autocrine/paracrine estrogen–ER system exists in the penis, which may play a role in development in association with the classic sex steroid hormone, androgen.

Real-time PCR analysis showed significant expression of aromatase gene in the penis. This was further confirmed by in situ hybridization analysis, which revealed localization of aromatase mRNA in various penile compartments. Immunoblots, enzyme immunoassay and immunofluorescence showed that aromatase mRNA was translated into protein. On Western blots, we detected a single and distinct band migrating at 54 kDa. This molecular mass is very close to the values reported for seminiferous tubules and crude germ cells in mature rats \((55 \text{ kDa})\) (Levallet \textit{et al.} 1998) and in both infant and adult testicular cell lines in humans \((53 \text{ kDa})\) (Brodie \textit{et al.} 2001). Levels of penile aromatase protein and estrogen concentrations were strikingly similar across ages, as revealed by enzyme immunoassay, implying that penile aromatase is not only transcribed and translated locally, but is functionally active in the penis where it irreversibly converts androgens into estrogens. Two types of promoters for aromatase have been identified in the rat testis, namely proximal promoter II and distal promoter I, which are also present in the ovary and adipose/bone tissues respectively (Bourguiba \textit{et al.} 2003, Simpson 2003). However, at this point, the specific promoters of penile aromatase are not known.

Earlier studies have shown that testosterone downregulates rat corpora cavernosa androgen receptors via aromatization to estrogen, an effect blocked by aromatase inhibitor (Lin \textit{et al.} 1993). Other studies, using human penile smooth muscle cell lines, reported that estrogen attenuates

Figure 2 In situ hybridization analysis showing expression of aromatase mRNA in the ovary and penis of pregnant and postnatal day 3 rats respectively. Abundant expression was observed in (A) the corpus luteum of ovary (cl) (inset: high magnification), (B) penile spongiosus (ps) and urethra (u), (E) cavernosum (pc), (G, H) sensory corpuses (sc), (I) blood vessels (bv), and (J) dorsal nerve (dn). When the cold probe was added in excess of the hot probe, the signal disappeared in penile spongiosus (ps) and urethra (u) (C) and in penile cavernosum (pc) (F). Magnification ×200. At high magnification (×400), clear aromatase signal was seen in penile spongiosus (ps), and urethral luminal and basal epithelium (D).
proliferation and induces expression of oxytocin receptors in these cells (Crescioli et al. 2003, Vignozzi et al. 2004). These findings are consistent with the present data that demonstrate aromatase in various penile compartments. One of the key novel findings of the present study is that aromatase and estrogen in the penis are abundant and are regulated age-dependently, with levels in the first week of life almost resembling those of the ovary, the classical tissue. It is noteworthy that this pattern of aromatase expression resembles those of penile ER reported recently (Jesmin et al. 2002), confirming our earlier speculation that estrogen’s role in the penis may be functional predominantly during development, particularly during the neonatal period (Jesmin et al. 2002). It is possible that expression of penile aromatase, like testicular aromatase, appears prenatally to influence the initial stages of penile differentiation (Greco & Payne 1994).

The functional significance of local extra-gonadal estrogen in cell physiology and pathophysiology is just beginning to be appreciated in tissues such as the nervous, adipose and adrenal gland (Simpson 2003). Although the role of estrogen in the penis is currently unclear, evidence supporting a role for local estrogen in penile events is beginning to emerge (Greene et al. 1939, Stillman 1982, Iguchi et al. 1990, North & Golding 2000, Jesmin et al. 2002, Crescioli et al. 2003, Dietrich et al. 2004, Vignozzi et al. 2004). It is interesting to note that aromatase, like ER, is broadly expressed in similar penile compartments (urethra, spongiosus, cavernous) and cell types (neuronal, vascular, stromal and epithelial) (Jesmin et al. 2002), suggesting the existence of a local autocrine and/or paracrine system of estrogen action in penile cells. The diversity of cell types expressing aromatase and ER indicates the complexity of estrogen’s potential functions in the penis (Jesmin et al. 2002). Such functions may, in general, include regulation of development, differentiation and, specifically, may involve function of penile blood vessels, nerves, stromal and epithelial cells (Patrone et al. 1999, 2000, Calles-Escandon & Cipolla 2001, Papka et al. 2001, Jesmin et al. 2002). Indeed, we have previously shown that caspase 3 and Bcl-2, respectively, co-localize with ER-α in the sensory corpuscles of the glans penis, suggesting that estrogen acting in an autocrine fashion via ER-α may promote survival and/or viability of sensory neurons in the glans penis (Jesmin et al. 2002). The use of aromatase knockout mice should provide more insight into the role of aromatase in penile functions.
events, even though findings of earlier studies show that these mice are fertile and capable of copulating (Robertson et al. 1999). However, it is important to note that transgenic mice expressing human aromatase, with high concentrations of circulating estrogen and prolactin and low testosterone, are infertile and display multiple structural and functional male reproductive abnormalities (Li et al. 2003). These findings may emphasize the
importance of the local estrogen:androgen balance in male reproduction.

In the local cellular physiology of sex steroid-sensitive cells, the androgen:estrogen ratio may be more important than single hormone action (Carreau et al. 1999, Simpson 2003). This ratio is mainly controlled by aromatase and its expression is regulated by many factors, including estrogen, testosterone, DHT, follicle-stimulating hormone, luteinizing hormone, cyclic AMP, growth factors, tumor necrosis factor-α and interleukin-1 (Carreau et al. 2002, Chen et al. 2002, Simpson 2003). The present study begins to address the regulatory effects of estrogens and androgens on penile aromatase mRNA and protein. We demonstrate that testosterone, DHT and DES, when administered prenatally, upregulate levels of penile aromatase mRNA and protein postnatally. However, unlike DES and testosterone, DHT had no effect on local penile estrogen concentration, suggesting that the influence DHT on aromatase is limited to transcription and translation, but not to aromatase activity. The regulation of aromatase expression is complex and is tissue specific, with possibly one specific promoter for each tissue (Carreau et al. 2002, Simpson 2003). For instance, proximal promoter II regulates expression of ovarian aromatase and distal promoter I.1 modulates placental aromatase, whereas distal promoter I.4 regulates aromatase expression in the adipose and bone tissues (Simpson 2003). The effects of the sex steroid hormones on penile aromatase, as demonstrated here, are not surprising, in that aromatase promoters express response elements of these hormones and have similar effects on aromatase expression in other tissues (Lephart 1996, Callard et al. 2001, Carreau et al. 2002, Simpson 2003). For example, estrogen increased expression of aromatase in zebrafish brain, whereas DHT and testosterone were ineffective (Kishida et al. 2001). In another study, androgens in the brain region regulated aromatase in a region- and gender-specific manner (MacLusky et al. 1994). More studies are needed in future to identify the specific promoter and transcripts of penile aromatase and to determine in more detail the specific regulatory pathways of testosterone, DHT and estrogen. Equally important will be the effects of environmental estrogens.

The interrelationship between testosterone and estrogen biosynthesis has made it hard to differentiate their precise effects. Indeed, much of the conventional knowledge on the biological effects attributed to androgen in penile events may represent effects of estrogen or androgen plus estrogen; for example, previously androgen was thought to restrict penile growth via down-regulation of its receptor (Takane et al. 1990). However, a later study, which utilized specific
agonist and antagonist/inhibitors for sex steroids and enzymes, demonstrated that estrogen synthesized from testosterone down-regulates the androgen receptor, and not testosterone, as previously thought (Lin et al. 1993). Thus, there is a need to re-visit this subject since we are now equipped with a variety of research tools, such as animal models lacking ER, androgen receptor, estrogen, DHT and their specific inhibitors or antagonists, to better understand this subject.

In summary, the novel findings of this study, demonstrating the presence, age- and sex steroid-dependent regulation of aromatase in the penis, provides further evidence of a role for local estrogen in penile events and the interaction of the estrogen and androgen systems. It is essential that future studies determine the specific promoter of penile aromatase and investigate the exact mechanisms and factors regulating it.

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