Long-term treatment with testosterone alters ovary innervation in adult pigs

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Abstract

Background: Intraovarian distribution and density of nerve fibres immunoreactive (IR) to protein gene product 9.5 (PGP 9.5) and containing dopamine-β-hydroxylase (DBH), neuropeptide Y (NPY), somatostatin (SOM), galanin (GAL) were determined.

Methods: From day 4 of the first oestrous cycle to day 20 of the second studied cycle, experimental gilts (n = 3) were injected with testosterone (T), while control gilts (n = 3) received corn oil.

Results: After T administration the numbers of fibres IR to PGP 9.5 and fibres IR to DBH, NPY and SOM were decreased. Fewer PGP 9.5- and DBH-IR terminals were observed within the ground plexus and around arteries and medullar veins, and medium tertiary follicles, and DBH-IR terminals in the vicinity of small tertiary follicles. T decreased the density of NPY-IR fibres in the medullar part of the ground plexus, and SOM-IR in the cortical part of the ground plexus.

Conclusions: The obtained data show that long-term T treatment of gilts decreases the total number of intraovarian fibres, including sympathetic ones. These results suggest that elevated T levels that occur during pathological states may affect the innervation pattern of ovaries, and their function(s).

Keywords: Ovary, Follicles, Innervation, Hyperandrogenism, Gilts

Background

Hyperandrogenism is one of the most common and disturbing endocrine disorder of reproductive-aged women and can result from many pathological states. The ovary is an important source for androgen excess in women with polycystic ovary syndrome (PCOS) [1, 2] and androgen-secreting tumours [3]. Augmentation in the peripheral blood androgen levels is found in patients suffering from adrenal hyperplasia [4] and androgen-secreting tumours [5, 6]. Blood androgen levels are elevated in pigs with uterine inflammation [7] and polycystic ovaries [8], and in dogs with adrenal dysfunction [9].

The pig's ovary receives its nerve supply from sympathetic, parasympathetic and sensory components of the peripheral nervous system (PNS). The sympathetic ovarian innervation derives from the caudal mesenteric ganglion (CaMG), sympathetic chain ganglia (SChGs; Th10-L5 and S1), ovarian and aorticorenal ganglia, as well as from the cranial part of the paracervical ganglion (PCG). The last-mentioned ganglion is also the source of parasympathetic neuronal inputs to the ovary. The sensory ovary-projecting neurons (referred to further as ovarian perikarya or ovarian neurons) occur in the dorsal root ganglia (DRGs) neuromers from Th10 to L5 [10]. Intraovarian sympathetic nerve fibres, constituting the most numerous population, are localized around follicles in all stages of development, the corpora lutea (CL), blood vessels and the interstitial gland, as well as within the ground plexus. These fibres, besides catecholamines (mainly noradrenaline - NA), may also express and release other active substances, for example, neuropeptide Y (NPY), somatostatin (SOM) and galanin (GAL) [10, 11]. NA and the above-mentioned peptides were found to influence steroidogenesis in ovarian cells [12–16]. Moreover, NA, NPY and GAL, acting by specific receptors or modulating the release of co-localized substances from nerve fibres, affect the blood flow and steroidogenesis in ovaries [17–19]. NA also affects ovulation [20], while NA [[21] Curry], SOM [22] and GAL [23] may influence follicular development.
Previous studies mainly show the effects of oestrogens on ovarian innervation. An increase in the content of NA was found in the ovaries of adult rats after injection of oestradiol valerate (EV; long-acting oestrogen), which was accompanied by morphological changes in gonads [24–26]. In turn, prenatal exposure to diethylstilbestrol, long-acting oestrogen, in newborn rats resulted in a drop in the density of the intravaginal sympathetic nerve network and a disruption in follicular maturation [27], as well as a reduction in the number of sympathetic ovarian perikarya in the celiac ganglion (CG) [28]. Lakomy et al. [29, 30] revealed a rise in the content of NA and the activity of acetylcholinesterase in the ovaries of oestradiol-17β (E2)-treated prepubertal gilts, as well as a decrease in the values of these parameters after the application of E2 together with progesterone (P4). We reported that long-term treatment of gilts with E2 changes the morphology and chemical coding of ovarian perikarya in the SchGs [31], CaMG [32], DRGs [33] and PCG [34]. Moreover, E2 exposure of gilts increases the total number of intraovarian fibres, including sympathetic ones [35].

Our knowledge on androgen influence on the PNS neurons supplying the female reproductive tract, including ovaries, is limited. In rats, during the late pregnancy [36] and after parturition [37], androstenedione (A4), may mediate a luteotropic effect acting on the CG neurons. In fact, the pig, due to its embryological, anatomical and physiological similarity to humans, constitutes an especially valuable species for bio-medical research, including that of ovary functions [38, 39]. Our earlier studies revealed that long-term testosterone (T) treatment alters the morphological and chemical organization of the CaMG [40], PCG [41] and SchGs [42] ovarian perikarya in adult gilts. Taking into consideration the above-mentioned findings, we hypothesize that elevated levels of androgens during pathological states may also affect the innervation pattern, including sympathetic ones, in the ovaries, and finally the gonadal functions. Therefore, we examined the ovaries of sexually mature gilts to determine the effect of long-term T administration on: 1) the total number and distribution of nerve fibres (stained for a general pan-neuronal marker - protein gene product 9.5- PGP 9.5), 2) the distribution and density of nerve fibres containing DBH and/or NPY, SOM, GAL, and 3) the populations of DBH-, NPY-, SOM- and GAL-immunoreactive (IR) nerve fibres in relation to the total number of PGP 9.5-IR nerve fibres.

**Methods**

**Animals**
The study was carried out on 6 crossbred gilts (Large White x Landrace), aged 7–8 months and weighing 90–110 kg, having had two controlled consecutive oestrous cycles. Behavioural oestrus was detected using a boar. Three days before surgical operations the gilts were transported from a farm to a local animal house and kept in individual stalls under natural light and temperature (April, May). They were fed a commercial grain mixture and tap water ad libitum.

**Experimental procedures**

On day 3 of the first studied oestrous cycle (day 0 of the study), after induction of general anaesthesia by azaperone (2 mg/1 kg of body mass, Stresnil, Janssen Pharmaceutica N.V., Belgium) and sodium pentobarbital (30 mg/1 kg of body mass, Vetbutal, Biovet, Poland), a polyvinyl cannula (outer diameter 2.2 mm, inner diameter 1.8 mm, Tomel, Tomaszów Maz., Poland) was inserted into the jugular vein of each gilt in order to collect blood samples.

Next, the gilts were randomly assigned to one of two following groups: the control (group I, n = 3) and experimental (group II, n = 3). In the gilts of group I, from day 4 of the first studied oestrous cycle (day 1 of the study) to the expected day 20 of the second studied cycle, i.e., within 38 consecutive days, 2 ml of oil was injected i.m. every 12 h (h; at 07:00 and 19:00 h). In turn, in the gilts of group II, at the same time and in the same manner 1000 μg of T (catalog no. 35800, Serva Electrophoresis GmbH, Germany) in 2 ml of corn oil was injected. The applied dose of T was determined based on our preliminary experiment, showing that its application increases the peripheral blood T concentration about 3.5 fold. According to available reports, about a 3- and 5-fold increase in the total T and bioavailable T, respectively, in blood concentrations accompanies adrenal hyperplasia [4], while the free androgen index is about 5-fold higher in women with PCOS than in controls [1]. For estimation of T, A4, E2, oestrone (E1) and P4 levels blood samples were collected from gilts of both groups through the whole period of T/oil injection (twice a day - 09:00 and 21:00 h). The samples were then immediately placed in an ice bath, where they were kept until centrifugation (10 min, 1,500 × g, at 4 °C). The plasma was decanted and stored at −20 °C until further processing. The analysis of androgen and oestrogen concentrations in the peripheral blood of the gilts was described earlier [40]. After the last blood sample collection the gilts were slaughtered by electric shock (ENZ 300 Metalowiec, Bydgoszcz, Poland) and both ovaries from each gilt were immediately dissected out and weighed. Afterwards, the volume, length, width and height of the gonads, as well as the number of follicles were estimated. The follicles were divided into three size classes: 1–3, 4–6 and 7–10 mm in diameter. Following the inspection of the ovarian surfaces, for immunocytochemical studies, ovaries were cut into 3 parts (two lateral and the third
middle - containing the hilar region), and fixed by immersion in Zamboni's fixative for 30 min, washed with 0.1 M phosphate buffer (PB; pH 7.4) over two days, and finally transferred to and stored at 4 °C in 18 % buffered sucrose solution (pH 7.4) containing 0.01 % natriumzide (NaN₃) until further processing.

**Immunofluorescent procedures**

To investigate the distribution and density of PGP 9.5-, DβH-, NPY-, SOM- and GAL-IR intraovarian nerve fibres, from every third part of ovary (12-µm-thick) serial sections were cut in a cryostat (Frigocut, Reichert-Jung, Nussloch, Germany). The sections were mounted on chrome alum-coated slides and then subjected to a routine double-immunofluorescence technique described by Majewski and Heym [43]. Briefly, after air-drying at room temperature for 45 min and rinsing in 0.1 M phosphate-buffered saline (PBS; pH 7.4; 3 × 10 min), the sections were incubated in a blocking buffer containing 10 % normal goat serum (MP Biomedicals, Solon, OH, USA), 0.1 M PBS, 0.1 % donkey serum (Abcam, Cambridge, UK), 1 % Triton X-100 (Sigma-Aldrich, St. Louis, MO, USA), 0.05 % Thimerosal (Sigma-Aldrich, St. Louis, MO, USA), and 0.01 % NaN₃ for 1 h at room temperature to reduce non-specific background staining. Subsequently, after another wash in PBS (3 × 10 min), the sections were incubated overnight at room temperature with two different species-specific primary antisera raised against PGP 9.5 (mouse, 7863–2004, AbD Serotec, dilution 1:1000), as well as with DβH (rabbit, AB1585, Millipore, dilution 1:2000 and mouse, MAB308, Millipore, dilution 1:1000), NPY (rabbit, NA1233, Enzo Life Sciences International, Inc., dilution 1:4000), SOM (rabbit, 8330–0154, AbD Serotec, dilution 1:50), and GAL (rabbit, AB2233, Millipore, dilution 1:4000).

Following subsequent rinsing in PBS (3 × 10 min), the sections were incubated with secondary antisera Alexa Fluor 488 (donkey anti-mouse, A21202, Invitrogen, USA, dilution 1:1000), Alexa Fluor 546 (donkey anti-mouse, A10036, Invitrogen, USA, dilution 1:1000), Alexa Fluor 488 (donkey anti-rabbit, A21206, Invitrogen, USA, dilution 1:1000), Alexa Fluor 546 (donkey anti-rabbit, A11010, Invitrogen, USA, dilution 1:1000) for 2 h at room temperature to visualize the antibody combinations: PGP 9.5/DβH, PGP 9.5/NPY, PGP 9.5/SOM, PGP 9.5/GAL, DβH/NPY, DβH/SOM and DβH/GAL. Next, the washed sections were coveredslipped in carbonate-buffered glycerol (pH 8.6). Standard tests (preabsorption for the used antisera with the respective antigen at a concentration of 20–50 µg antigen/ml diluted antiserum, omission of primary or secondary antiserum and replacement by non-immune sera of all the primary antiserum used) were employed to control the specificity of immunofluorescence. Also, DβH, NPY, SOM and GAL staining in the porcine CaMG ovary supplying neurons were applied as positive controls (data not shown). The immunocytochemical staining procedure for one combination of examined substances was conducted on nine randomly chosen ovarian sections from every one-third part of the organ derived from each studied animal.

Double-immunolabeled nerve fibres were analyzed and photographed under an Olympus BX51 microscope equipped with epifluorescence and the appropriate filter sets for FITC (B1 module, excitation filter 450–480 nm, barrier filter 515 nm) and CY3 (G1 module excitation filter 510–550 nm, barrier filter 590 nm). Pictures were captured by a digital camera connected to a PC and analyzed with the AnalySIS software (version 3.02, Olympus Soft Imaging Solutions, Muenster, Germany). In our study the distribution and density of PGP 9.5-, DβH-, NPY-, SOM- and GAL-IR intraovarian nerve fibres were estimated within the ground plexus and around follicles, blood vessels and interstitial glands. All stained processes identified in the surrounding zone of the abovementioned structures were counted. Follicles, depending on the stage of development, were classified microscopically according to Wulff et al. [44] and Barboni et al. [45]: primordial - without granulosa cells, primary - surrounded by a single layer of cuboidal granulosa cells, secondary - with two or more granulosa cell layers without the antral cavity, tertiary - with antrum. Additionally, the tertiary follicles were divided into three size subclasses: small (to 3 mm in diameter), medium (4–6 mm in diameter) and large (7–10 mm in diameter). The diameter of follicles was measured using Microimage software (Olympus Polska sp. z o. o., Warsaw, Poland).

**Statistical analyses**

Data concerning the weight, volume and measurements of the ovaries, follicle numbers, as well as the density of innervation gained from two ovaries from each gilt were averaged per ovary. The mean (±SEM) weight, volume and size of the ovaries, the number of ovarian structures, as well as the total number of PGP 9.5-IR and the absolute numbers of DβH-, NPY-, SOM-and GAL-IR nerve fibres were compared between the groups using Student's t-test. To calculate the statistical significance of the mean (±SEM) numbers of PGP 9.5-, DβH-, NPY-, SOM- and GAL-IR nerve fibres, between the groups and within the same group, supplying the particular ovarian structures, one-way analysis of variance (ANOVA) followed by the Newman-Keuls test was performed. To indicate the differences in frequency of DβH-, NPY-, SOM-, GAL-IR nerve fibres occurrence in the total population of PGP 9.5-IR nerve fibres, the total number of the PGP-IR nerve population in each group was accepted as 100 %. The numbers of DβH-, NPY-, SOM-,
GAL-IR nerve fibres were expressed as a percentage (mean) of the total population of PGP 9.5-IR nerve fibres. Then, the Newman-Keuls test was applied for calculating the statistical significance of mean differences (ANOVA, InStat Graph Pad, San Diego, CA). Differences with a probability of $P < 0.05$ were considered significant.

Results

Macroscopic evaluation of the ovaries

Compared to the controls, treatment with T caused in the ovaries a reduction in the number of small (1–3 mm in diameter; 10.2 ± 0.33 vs. 20 ± 0.57, $P < 0.001$; respectively) and medium (4–6 mm in diameter; 3.16 ± 0.33 vs. 5.66 ± 0.6, $P < 0.05$; respectively) follicles, as well as a lack of large follicles (7–10 mm in diameter; 0 ± 0 vs. 4.4 ± 0.16, respectively). The weight and volume, as well as the length, width and height of the ovaries did not differ significantly in the control and experimental gilts (4.66 ± 0.29 vs. 3.51 ± 0.32 g; 4.79 ± 0.39 vs. 3.5 ± 0.29 ml; 1.4 ± 0.09 vs. 1.7 ± 0.08 cm; 2.91 ± 0.17 vs. 2.1 ± 0.29 cm; 2 ± 0.09 vs. 1.2 ± 0.28 cm, respectively).

The distribution and density of PGP 9.5-IR nerve fibres in the ovaries

In the ovaries of gilts treated with T, the total number of PGP 9.5-IR nerve fibres was lower ($P < 0.001$) than in the control ovaries (160.23 ± 4.2 vs. 213.29 ± 3.72, respectively).

Compared to the control group (Fig. 1a, c, e, g), the T treatment led to a drop in the number of PGP 9.5-IR intraovarian nerve fibres in the cortical ($P < 0.001$, Fig. 1b) and medullar ($P < 0.001$, Fig. 1f) parts of the ground plexus, the near cortical ($P < 0.001$) and medullar ($P < 0.001$, Fig. 1h) arteries and medullar veins ($P < 0.001$), as well as medium tertiary follicles ($P < 0.001$, Fig. 1d). The distribution and density of PGP 9.5-IR nerve fibres supplying the ground plexus, follicles, the interstitial gland and blood vessels in both examined groups are depicted in Table 1.

Ovarian DβH-, NPY-, SOM-, GAL-IR nerve fibres as a percentage of PGP 9.5-IR nerve fibres

Long-term T administration led to a decrease ($P < 0.01$) in the number of ovarian DβH-IR nerve fibres (normalised against the total population of PGP 9.5-IR nerve fibres) compared with that in the control group (46.6 % vs 55.3 %, respectively). However, there was no significant difference between the control and T-injected groups in terms of NPY-IR (21.6 % vs 20.9 %, respectively), SOM-IR (14.9 % vs 12.4 %, respectively) and GAL-IR (9.4 % vs 8.8 %, respectively) fibres normalised against the total population of PGP 9.5-IR nerve fibres.

The distribution and density of DβH-, NPY-, SOM- and GAL-IR nerve fibres in the ovaries

In the ovaries of T-treated gilts the absolute numbers of DβH-, NPY- and SOM-IR nerve fibres were lower than those calculated in the control group (74.71 ± 4.67 vs. 117.80 ± 2.1, $P < 0.01$; 31.89 ± 2.14 vs. 46.12 ± 1.78, $P < 0.01$; 19.9 ± 2.7 vs. 31.9 ± 0.91, $P < 0.05$; respectively). In turn, the absolute numbers of GAL-IR nerve terminals were similar in the control and T-treated gilts (20.25 ± 0.9 vs. 14.2 ± 2.69, respectively).

After T injections populations of the DβH-IR nerve fibres were lower within the cortical ($P < 0.001$, Fig. 2b)

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**Fig. 1** Ovarian PGP 9.5-IR nerve fibres (†) in the control and T-treated gilts. In the control ovaries numerous nerve fibres visible within the cortical (a) and medullar (e) parts of the ground plexus, and around the medullar artery (g) and medium tertiary follicle (c). Note a drop in the density of PGP 9.5-IR nerve fibres after T administration within cortical (b) and medullar (f) parts of the ground plexus, and in the vicinity of the medullar artery (h) and medium tertiary follicle (d)
and medullar (P < 0.001, Fig. 2j) parts of the ground plexus, around small (P < 0.001) and medium (P < 0.001, Fig. 2f) tertiary follicles, as well as cortical (P < 0.001) and medullar (P < 0.001, Fig. 2n) arteries and medullar veins (P < 0.001) than in the control group (Figs. 2a, e, i, m). In the ovaries of T-injected gilts a decrease (P < 0.01) in the number of NPY-IR nerve terminals was found in the area of the medullar ground plexus (Fig. 2l) compared to the control group (Fig. 2k). In turn, the population of SOM-positive nerve fibres was lower (P < 0.001) within the cortical part of the ground plexus following T administration (Fig. 2d) than in the control ovaries (Fig. 2c). The application of T did not significantly change the innervation pattern of particular ovarian structures by GAL-IR nerve fibres (Figs. 2o, p). The distribution and density of DβH-, NPY-, SOM- and GAL-IR nerve fibres supplying the ground plexus, follicles, the interstitial gland and blood vessels in both studied groups are given in Table 2.

The patterns of co-localization of DβH with/or NPY, SOM, GAL in nerve fibres in the ovaries
In the ovaries of the gilts receiving T, compared to the control ovaries, all DβH-IR nerve fibres in the vicinity of secondary follicles were simultaneously NPY-IR, while DβH with NPY was co-localized in the part of fibres supplying cortical blood vessels. Following T administration, in the cortical part of ovaries the appearance of SOM expression in the part of DβH-IR nerve fibres around arteries, and a lack of co-localization of these substances near veins were found. The co-expression of DβH with GAL did not differ in fibres around primordial and tertiary small and medium follicles and cortical blood vessels in both groups. Similarly, in the nerve fibres occurring within the medullar ground plexus and in the vicinity of blood vessels of the experimental and control ovaries the co-expression DβH with/or NPY, SOM, GAL was similar. The co-localization patterns of DβH with/or NPY, SOM, GAL in nerve fibres innervating ground the plexus, follicles, the interstitial gland and blood vessels in both studied groups are presented in Table 3.

Discussion
Our study shows that the long-term exposure of adult gilt ovaries to T resulted in disturbances in the development of follicles, a drop in the total population of nerve fibres (PGP 9.5-IR), including the total population of DβH-IR, as well as in the absolute numbers of NPY- and SOM-IR nerve fibres. Changes in the distribution and/or density of these fibres depending on the kind of chemical coding of the fibre and/or ovarian structure were also observed.

The peripheral blood androgen and oestrogen concentrations in gilts used in the current study have been reported earlier [40]. In T-injected gilts, in comparison to the control gilts, an increase in the T concentrations on all days of the study (approximately 3.5 fold), except for days 2, 12 and 35–37 was revealed. After T administration the mean daily concentrations of this steroid varied between 50 ± 13.6 and 141 ± 16.5 pg/ml. In T-treated gilts the levels of E2 were higher (approximately 1.6 fold) than in the control group on study days 4–14 and 21–29. In turn, mean daily E2 concentrations ranged from 3.2 to 16.8 ± 1.5 pg/ml. Following T injections the E1 levels decreased on days 8–11, 15, 16, 18, 19, 29–32, 37 and 38, while the A4 levels were significantly unchanged on all of the study days. Moreover, compared with the control group, mean P4 concentrations were lower in T-administered gilts on days 6–13 (ranged from 0.6 ± 0.1 to 3.4 ± 0.18 ng/ml), 22 (0.6 ± 0.07 ng/ml), 23 (0.4 ± 0.06 ng/ml) and 26–34 (ranged from 0.4 ± 0.04 to 0.8 ± 0.13 ng/ml) of the study [Jana et al. unpublished observations]. P4 concentrations allow the supposition that after T application the P4 synthesis in the corpora lutea was reduced in the first studied oestrous cycle, and the development of the corpora lutea did not occur in the second studied cycle. The disturbances in the course of the oestrous cycle were further confirmed by macroscopic examination of the ovaries, conducted on day 38 of the study (the expected day 20 of the second studied cycle), which revealed a drop in the number of small

### Table 1 Mean (±SEM) number of PGP 9.5-IR nerve fibres in the ovaries of gilts

| Ovarian structures   | C          | T          |
|----------------------|------------|------------|
| Cortex               | Ground plexus | 27.1 ± 0.29<sup>a</sup> | 15.23 ± 0.44<sup>x</sup> |
| Follicles:           | Primordial | 2.8 ± 0.2<sup>b</sup>  | 1.98 ± 0.39<sup>h</sup> |
|                      | Primary    | 2.69 ± 0.27<sup>b</sup> | 2.29 ± 0.31<sup>i</sup> |
|                      | Secondary  | 7.3 ± 0.37<sup>c</sup> | 8.27 ± 0.28<sup>g</sup> |
| Tertiary (mm):       | - up to 3  | 19.07 ± 0.73<sup>g</sup> | 20.32 ± 0.91<sup>x</sup> |
|                      | - 4–6      | 28.25 ± 1<sup>e</sup>  | 16.68 ± 0.44<sup>x</sup> |
|                      | - 7–10     | 40.14 ± 1.25<sup>e</sup> | l.s. |
| Arteries             | 14.27 ± 0.57<sup>e</sup> | 7.32 ± 0.39<sup>g</sup> |
| Veins                | 12.21 ± 0.06<sup>g</sup> | 5.46 ± 0.51<sup>c</sup> |
| Interstitial gland   | 5.2 ± 0.06<sup>c,f,g</sup> | 6.48 ± 0.9<sup>e</sup> |
| Medulla              | Ground plexus | 19.29 ± 0.73<sup>g</sup> | 10.55 ± 0.39<sup>c</sup> |
|                      | Arteries   | 19.64 ± 0.6<sup>s</sup> | 10.72 ± 0.6<sup>s</sup> |
|                      | Veins      | 15.36 ± 0.55<sup>b</sup> | 7.98 ± 0.38<sup>x</sup> |

<sup>C</sup> control gilts, T testosterone-treated gilts; means with different superscriptions (a, b, c, d, e, f, g) indicate differences (P < 0.05–0.001) in the same group among particular structures in ovarian cortex or medulla; x - indicates differences (P < 0.001) between both groups for the same structure; l.s. - lack of structure.
and medium tertiary follicles, and a lack of large tertiary follicles. These findings are congruent with previous studies performed on gilts receiving E\(_2\) for a long time [35]. The follicular development in the offspring of rats was also found to be suppressed after prenatal exposure to diethylstilbestrol [27]. The morphological changes in the ovaries of T-injected gilts (present study) were a consequence of long-term enhancement of T and E\(_2\) levels in the peripheral blood. It can be assumed that the effect of endogenous steroids on the ovaries of the T-injected gilts was in significant, which was confirmed by their decreased contents in gonadal tissue [Jana et al. unpublished observations]. We propose that changes in the morphology of ovaries and their lesser steroidogenic activity may result from a significant inhibition of the hypothalamic-pituitary axis function by higher E\(_2\) and T levels [46–48].

![Fig. 2 Ovarian D\(_\beta\)H- and/or NPY-, SOM-, GAL-IR nerve fibres (↑) in the control and T-treated gilts. Within the cortical ground plexus of the control gilts numerous D\(_\beta\)H+ (a) and not numerous SOM- (c) IR nerve fibres are visible, while in the gilts receiving T a noticeable decrease in the population of D\(_\beta\)H- (b) and SOM- (d) IR fibres is present. In the control ovary numerous D\(_\beta\)H-IR fibres near medium tertiary follicle (e). Note a drop in the number of these nerve fibres in the T-injected gilt (f). Around medium tertiary follicles of the control (g) and T-treated (h) gilts the population of NPY-IR nerve fibres are similar. Within the medullar ground plexus of the control gilts numerous D\(_\beta\)H- (i) or not numerous NPY- (k) IR nerve fibres are present. In turn, after T treatment, a decrease in the number of D\(_\beta\)H- (j) and NPY- (l) immunoreactivity is visible. A greater population of D\(_\beta\)H-IR fibres is present in the vicinity of the medullar artery in the control gilt (m) than after T treatment (n). The numbers of GAL-IR processes around medullar arteries of the control (o) and T-injected (p) gilts are similar. Negative controls for D\(_\beta\)H (r), SOM (s), NPY (t) and GAL (u)].
Table 2  Mean (±SEM) number of DβH-, NPY-, SOM- and GAL-IR nerve fibres in the ovaries of gilts

| Ovarian structures | DβH | NPY | SOM | GAL |
|--------------------|-----|-----|-----|-----|
|                    | C   | T   | C   | T   | C   | T   | C   | T   |
| Cortex             |     |     |     |     |     |     |     |     |
| Ground plexus      | 14.33 ± 1.93 ab | 5.91 ± 0.33 a | 4.75 ± 0.38 a | 1.61 ± 0.05 ab | 4.33 ± 0.17 a | 1.62 ± 0.02 b | 1.6 ± 0.48 a | 0 ± 0 a |
| Follicles:         |     |     |     |     |     |     |     |     |
| Primordial         | 1.16 ± 0.2 a | 0.9 ± 0.07 b | 0.66 ± 0.22 b | 0.25 ± 0.06 a | 0.33 ± 0.22 b | 0 ± 0 c | 0.5 ± 0.19 ab | 0.1 ± 0.03 b |
| Primary            | 2.44 ± 0.29 b | 1.8 ± 0.14 b | 0.33 ± 0.22 b | 0.35 ± 0.19 b | 0 ± 0 b | 0.91 ± 0.05 bc | 0 ± 0 b | 0 ± 0 b |
| Secondary          | 3.72 ± 0.27 c | 6.31 ± 0.15 a | 1.8 ± 0.6 c | 2.27 ± 0.05 ad | 1.4 ± 0.25 c | 2.03 ± 0.07 b | 0.4 ± 0.16 ab | 0 ± 0 b |
| Tertiary (mm):     |     |     |     |     |     |     |     |     |
| - up to 3          | 11.2 ± 1.2 a | 4.12 ± 0.33 ab | 2.5 ± 0.02 c | 1.12 ± 0.05 ad | 3.25 ± 0.31 ad | 2.16 ± 0.25 b | 1.33 ± 0.68 b | 2.82 ± 0.54 c |
| - 4-6              | 16.8 ± 0.17 b | 5.73 ± 0.33 a | 4.25 ± 0.51 b | 2.28 ± 0.35 ad | 3 ± 0.7 d | 1.58 ± 0.21 ab | 2.75 ± 0.37 b | 2.9 ± 0.29 c |
| - 7-10             | 24.75 ± 1.21 d | 7.25 ± 0.37 d | 7.0 ± 0.6 d | 3.3 ± 0.29 d | 4.33 ± 0.29 a | 3 ± 0.8 b | 0.8 ± 0.05 b | 1.6 ± 0.08 d |
| Arteries           | 7.33 ± 0.59 a | 2.1 ± 0.06 abx | 5 ± 0.68 a | 6 ± 0.05 a | 0.5 ± 0.19 bc | 0.1 ± 0.06 b | 0 ± 0 b | 0.6 ± 0.03 a |
| Veins              | 4.14 ± 0.34 ab | 1.35 ± 0.05 b | 5 ± 0.28 a | 2.1 ± 0.08 ab | 1.5 ± 0.32 c | 0.5 ± 0.05 a | 1.33 ± 0.22 b | 0.41 ± 0.04 ab |
| Intestinal gland   | 3.66 ± 0.04 c | 2.53 ± 0.33 bc | 0.6 ± 0.22 b | 1.59 ± 0.06 ad | 0 ± 0 b | 0 ± 0 c | 0 ± 0 b | 0 ± 0 b |
| Medulla            |     |     |     |     |     |     |     |     |
| Ground plexus      | 8 ± 0.33 a | 3.15 ± 0.08 b | 5.66 ± 0.17 ab | 1.92 ± 0.06 ab | 2.75 ± 0.31 ab | 0.79 ± 0.01 b | 1.8 ± 0.33 a | 1.6 ± 0.16 a |
| Arteries           | 2 ± 0.5 b | 4.3 ± 0.06 c | 3.25 ± 0.35 b | 1.12 ± 0.07 b | 3 ± 0.8 b | 0.8 ± 0.05 | 1.6 ± 0.34 a | 0.5 ± 0.08 b |
| Veins              | 8.6 ± 0.5 a | 3.17 ± 0.14 b | 4.5 ± 0.42 c | 3.36 ± 0.15 b | 1.6 ± 0.4 b | 1.9 ± 0.32 b | 0.4 ± 0.16 b | 0 ± 0 b |

C: control gilts, T: testosterone-treated gilts; means with different superscissions (a, b, c, d, e) indicate differences (P < 0.05–0.001) for particular substances in the same group among particular structures in ovarian cortex or medulla; x – indicates differences (P < 0.01, P < 0.001) for particular substances between both groups for the same structure; l.s. – lack of structure

We found that T injections in the gilts resulted in a decrease of the total population of ovarian nerve fibres, demonstrated by PGP 9.5 immunoreactivity. Similarly, both single and repeated (via 21 days) E2 administration led to a decrease in the total population of PGP 9.5-IR terminals in the mouse uterus [49] and rat vagina [50]. In contrast, the greater populations of these fibres were visible in the ovaries of the long-term E2-treated gilts [35]. A rise in the number of PGP 9.5-IR fibres was also observed in the mammary gland [51] and earlobe [52] in adult ovariectomized (OVX) rats after short-term (via 7 days) E2 exposure. Data exist showing that dehydroepiandrosterone (DHEA) exerted a stimulatory effect on nerve density in O VX rat vagina through an androgenic action [53]. These discrepancies in androgen and oestrogen effects on the population of nerve fibres are probably due to species differences, the kind of organ, as well as the time and doses of administered steroid hormones.

In the present study, in the ovaries of gilts treated with T, the population of DβH-IR nerve fibres, calculated in relation to the total population of PGP 9.5 fibres as well as the absolute number of DβH-IR fibres, markedly decreased. T treatment resulted in a drop in the absolute number of the NPY- and SOM-IR nerve fibres. In turn, populations of GAL nerve fibres (total and absolute) were similar in both studied groups. We also found that in the ovaries of T-treated gilts the distribution and/or density of fibres positive for DβH, NPY and SOM depended on the kind of chemical coding of the fibre and/or ovarian structure. However, changes in the innervation of the gonads by DβH-IR nerve fibres referred only to their density around/within the particular structures but not their distribution. Existing data show that in T-treated rats renal tyrosine hydroxylase (TH) activity [54] decreased, and that in rats exposed to E2 the level of this enzyme in the superior cervical ganglia was reduced [55]. After E2 injections some ovarian structures were found to be supplied by higher numbers of DβH-IR fibres [35]. Similarly, Anesetti et al. [56] reported an increase in the number of intraovarian TH-IR nerve endings in immature rats in response to cypionate estradiol. Moreover, in the ovaries of adult rats a greater number of NA-ergic nerve fibres was found after a single EV injection [24, 25]. Also, E2 treatment in adult OVX rats led to an increase in the density of TH-positive nerve terminals in the mammary gland [51]. A higher population of noradrenergic nerve fibres in porcine ovaries was also observed on 30 day of pregnancy, when E2 level is the highest [57]. However, this steroid did not exert any significant effect on the density of TH-IR nerve endings in rat urethra [58] or vagina [59]. Applied in our study, T injections resulted in a drop in the number of NPY- or SOM-IR fibres within the medullar and cortical parts of the ground plexus, respectively. After hormonal treatment the GAL-IR fibres were not visible within the cortical part of the ground plexus or around secondary follicles, while these fibres appeared around primary follicles. In our earlier study, E2 injections resulted in an increase in the number of NPY- or GAL-IR fibres within...
the cortical and medullar parts of the ground plexus, respectively. This hormonal treatment induced a parallel increase in the density of NPY-IR fibres around medullar arteries, while SOM- and GAL-IR fibres were not found in the vicinity of primordial follicles [35]. It was also reported that short-term (via 7 days) E2 application increased the number of calcitonin gene-related peptide (CGRP)-IR fibres in the earlobe [52] and mammary gland [51] in adult OVX rats. However, the population of these endings was not markedly changed in rat uterus after E2 treatment [52]. There is a lack of data concerning the effect of androgens on the innervation patterns of ovaries by NPY-, SOM- and GAL-positive nerve fibres. The abundance of peptidergic (NPY, VIP) fibres in rat prostate during postnatal development was regulated by androgens [60].

In the ovaries of T-treated gilts we also revealed changes in the co-localization patterns of DβH with/ or NPY, SOM in nerve fibres. There are no data concerning the co-localization of neurotransmitters in the intraovarian nerve fibres in response to androgen administration. However, it is important to add that the changes in co-localization patterns observed in the present study are similar to those revealed in the ovaries of E2-treated gilts [35].

It is difficult to indicate the mechanism(s) underlying the drop in the total number of nerve fibres, as well as the subsets of DβH-, NPY- and SOM-IR nerve fibres in the ovaries of gilts receiving T. We suppose that these changes may be associated with the reduction of perikarya within the ganglia innervating ovaries. We reported earlier that in the CaMG [40], SChGs [42] and PCG [41] of T-treated gilts, from which ovaries were obtained for the present study, down-regulation of the total number of ovarian perikarya was found. Moreover, fewer populations of DβH-, NPY- and SOM-IR fibres in the ovaries of T-injected gilts were coincident with decreased populations of ovarian perikarya expressing these substances in the above-mentioned sympathetic ganglia [40, 42]. The lower number of SOM-IR found after T treatment (present study) also corresponds with the reduced population of ovarian perikarya possessing SOM in the PCG [41]. In addition, changes in the innervation pattern of gonads in the gilts receiving T might result from T and E2 effects on other sympathetic peripheral ganglia supplying the ovaries [10]. Available data show that the reduction in the set of the intraovarian sympathetic nerve terminals in rats prenatally exposed to diethylstilbestrol [27] was accompanied by a decrease in the number of sympathetic ovarian perikarya in the celiac ganglion [28].

Table 3 The co-localization of DβH and/or NPY, SOM, GAL in the intraovarian nerve fibres of gilts

| Ovarian structures | Group | DβH | NPY | SOM | GAL |
|-------------------|-------|-----|-----|-----|-----|
| Cortex            | Ground plexus | C   | -   | +   | +   | -   |
|                   | T     | -   | +   | +   | l.f.|
| Follicles:        | Primordial | C   | -   | -   | -   | -   |
|                   | T     | -   | -   | l.f.| -   |
|                   | Primary | C   | -   | ++  | l.f.| l.f.|
|                   | T     | -   | ++  | +   | l.f.|
|                   | Secondary | C   | -   | +   | l.f.| l.f.|
|                   | T     | -   | ++  | +   | l.f.|
|                   | Tertiary (mm): | - up to 3 | C | - | + | + | - |
|                   |       | T | - | + | + | + |
|                   | - 4–6 | C   | -   | +   | +   | +   |
|                   | T     | -   | +   | +   | +   |
|                   | - 7–10 | C | - | + | + | + |
|                   | T     | l.s.| l.s.| l.s.| l.s.|
| Arteries          | C     | -   | ++  | -   | -   |
|                   | T     | -   | +   | +   | -   |
| Veins             | C     | -   | ++  | +   | +   |
|                   | T     | -   | +   | -   | -   |
| Interstitial gland| C     | -   | +   | l.f.| l.f.|
|                   | T     | -   | +   | l.f.| l.f.|
| Medulla           | Ground plexus | C | - | + | + | - |
|                   | T     | - | + | + | + |
| Arteries          | C     | -   | +   | +   | +   |
|                   | T     | -   | +   | +   | +   |
| Veins             | C     | -   | +   | -   | -   |
|                   | T     | -   | +   | -   | -   |

C control gilts, T testosterone-treated gilts; co-localization of DβH with NPY, co-localization of DβH with SOM, co-localization of DβH with GAL in the part (+) or all (++) of nerve fibres; (−) lack of co-localization; l.f. lack of fibres, l.s. lack of structure.
The changes in the populations of ovarian nerve fibres in T-injected gilts revealed in the present study may have an importance for gonadal functions. A drop in the numbers of fibres expressing DβH within the ground plexus, near blood vessels and follicles, and/or NPY (transmitter occurring in the greatest number of sympathetic nerve fibres) within the ground plexus may cause disturbances in a variety of ovarian sympathetic actions, including regulation of steroidogenesis and blood supply, and in relation to DβH-IR fibres also in the regulation of follicular development. This supposition is based on previous studies demonstrating that under physiological conditions NA and NPY were found to increase synthesis of P₄ and E₂ in granulosa and luteal cells in humans [12] and many animal species [13–15], as well as to affect the blood flow in ovaries [17–19]. Moreover, the participation of NA in follicular development is well-known [21]. In turn, a reduction in the population of fibres containing SOM (transmitter occurring in the greatest number of sympathetic and parasympathetic fibres) found within the ground plexus in the ovaries of T-treated gilts may be significant for gonadal steroidogenesis [16] and follicular development [22].

Conclusions

After long-term T administration in sexually mature gilts a decrease in the total population of intraovarian fibres, including fibres containing DβH, NPY and SOM is observed. Moreover, our study suggests that hyperandrogenism may affect the innervation pattern of the ovaries and consequently their function(s). However, further studies determining the mechanism(s) underlying changes in the ovarian innervation pattern during hyperandrogenic states are necessary. In sum, the present findings support the significant role of steroid hormones modulating neuronal plasticity.

Abbreviations

A4: Androstenedione; CaMG: Caudal mesenteric ganglion; CG: Celiac ganglion; CGRP: Calcitonin gene-related peptide; CL: Corpora lutea; DHEA: Dehydroepiandrosterone; DHT: Dihydrotestosterone; DRGs: Dorsal root ganglia; DβH: Dopamine-ß-hydroxylase; E1: Oestrone; E2: Oestradiol-17β; EV: Estradiol valerate; GAL: Galanin; IR: Immunoactive; NA: Noradrenaline; NaN₃: Natriumazide; NGF: Nerve growth factor; NPY: Neuropeptide Y; OVX: Ovariectomized; P4: Progesterone; PB: Phosphate buffer; PBS: Phosphate-buffered saline; PCG: Paracervical ganglion; PCOS: Polycystic ovary syndrome; PGG 9.5: Protein gene product 9.5; PNS: Peripheral nervous system; SCtGs: Sympathetic chain ganglia; SOM: Somatostatin; T: Testosterone; TH: Tyrosine hydroxylase

Acknowledgements

There are no acknowledgements.
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