Peripubertal orchidectomy transitorily affects age-associated thymic involution in rats

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Abstract

The role of gonadal hormones in induction and, particularly, maintenance/progression of rat thymic involution, which normally starts around puberty, was reassessed by examining the effects of peripubertal orchidectomy on thymic weight and morphometric parameters at different times up to the age of 10 months. Up to 6 months post-castration both thymic weight and cellularity in orchidectomized (Cx) rats were greater than in age-matched control rats, sham Cx (Sx). The increase in thymic cellularity reflected an increase in thymocyte proliferation rate (the proportion of proliferating cells was 18.6 ± 0.7% in 2-month-old Cx (N = 5) vs 13.4 ± 0.3% (N = 5) in age-matched Sx rats) followed by reduced sensitivity to apoptotic signals (apoptotic thymocytes were 9.8 ± 0.9% in 2-month-old Cx (N = 5) vs 15.5 ± 0.3% (N = 5) age-matched Sx rats). However, 9 months post-orchidectomy, neither thymic weight and cellularity nor any of the morphometric parameters analyzed differed between Cx and control rats. The reduction of thymic cellularity in Cx rats to control values may be related to increased sensitivity of their thymocytes to apoptotic signals in culture (72.6 ± 1.2% in 10-month-old vs 9.8 ± 0.9% in 2-month-old Cx rats) followed by reduced responsiveness to proliferative stimuli (14.1 ± 0.2% in 10-month-old vs 18.6 ± 0.7% in 2-month-old Cx rats). Thus, the study indicates that the effects of peripubertal orchidectomy on thymic weight and cellularity, as well as on the main morphometric indices, are long-lasting but not permanent, i.e., that removal of the testes can only postpone but not prevent age-related organ atrophy and consequently functional deterioration of the immune system.

Key words
• Rat thymus
• Aging
• Orchidectomy
• Stereological analysis
• Thymocyte proliferation
• Thymocyte apoptosis

Introduction

Thymic involution is one of the most clear-cut effects of aging on the immune system of both laboratory animals and humans. In rodents, there is a sharp increase in immunological activity after birth, reaching a peak at sexual maturity (1). Thereafter, a gradual decline in immune function occurs. Changes in thymus weight and structure, as well as in its capacity to produce T lymphocytes, also become evident around puberty.
Thus, as the thymus involutes, the output of newly generated T cells declines, with a reduction in the level of naive T cells in the periphery. These changes have been related to the age-associated decline in function of the immune system, leading to an increased frequency of specific infections, malignant diseases and autoimmune disorders in old individuals (4). The fact that thymic involution is associated with immunosenescence and its various associated diseases has prompted many studies aimed at understanding the causes and mechanisms of thymic degeneration which may ultimately lead to the possibility of manipulation. This might not only prolong life, but also improve its quality, due to the absence of severe associated pathologies.

Since thymic involution in rodents becomes obvious at the same time as the increase in production of gonadal steroids, a causal link between increasing sex steroid level and age-related thymic atrophy has been suggested (1,2). This notion is supported by data indicating that androgens in males modulate both thymic weight and T-cell maturation by acting either directly on the thymocytes (5) or indirectly on thymic epithelial cells (TEC) (6) supporting T-cell differentiation/maturation. Namely, substantial alterations in the composition of thymocyte subsets at different stages of maturation (delineated by expression of the major differentiation antigens CD4, CD8, T-cell receptor), which are localized in clearly defined thymic compartments, have been observed in animals subjected to gonadectomy (7,8). A putative role for sex steroids in the induction of thymic involution is also supported by numerous experiments showing reversal of orchidectomy-induced thymic hypertrophy by androgen replacement (7). However, the long-term commonly held view that gonadal hormones have a pivotal role in the induction and maintenance of thymic involution has recently been disputed. Namely, thymic involution in humans is proposed to be a sex steroid-independent event (9), while lack of gonadal hormones was shown not to produce a delay in age-associated thymic involution in hypogonadal HPG/ Bm-hpg/hpg mice (10). The obvious discrepancy between these and numerous earlier studies may be reconciled by the fact that the role of sex steroids in thymic involution was estimated by exploring the effects of gonadal hormone deprivation/administration for not more than a few months. Therefore, the present study was undertaken to reassess the putative role of gonadal hormones in the induction and, particularly, in the maintenance/progression of age-associated thymic atrophy in rats. To this end, we followed the effects of peripubertal orchidectomy on the rat thymus for up to 9 months after surgery. Since it has been shown that the functional capacity of the aged residual thymus correlates with anatomical measurements of so-called true thymic tissue (4), we subjected this thymic component (i.e., cortical and medullary lymphoid tissue) from both orchidectomized (Cx) and control rats to a tentative morphometric analysis. Additionally, since changes in thymic cellularity mainly reflect alterations in the main homeostatic processes, i.e., apoptosis and proliferation, thymocytes from Cx and control rats were examined for their sensitivity to both apoptotic and proliferative signaling.

**Material and Methods**

**Animals**

Male inbred AO rats were maintained under a standard 12-h photoperiod, at 21 ± 2°C, with free access to food and water. At the age of 30 days, the rats were submitted to bilateral Cx or sham Cx (Sx) as previously described (8). All animal experimentation was conducted according to local ethical guidelines and was approved by our Institutional Animal Care and Use Committee.
One, 3, 6, or 9 months after orchidectomy, the rats were killed under ether anesthesia by exsanguination. Thirty-day-old intact rats, as well as 2-, 4-, 7-, and 10-month-old Sx and non-operated rats were included in the study as controls. All groups consisted of 5 animals. Each thymus was aseptically isolated, trimmed of all excess body fat and gently blotted on gauze to remove excess blood. The two lobes were divided and weighed individually. The right lobe was used for preparation of thymocyte single cell suspensions and the left lobe was fixed in Bouin’s solution and further processed for morphometric analysis.

**Computer-assisted morphometric measurements**

Thymic specimens fixed in Bouin’s solution and embedded in paraffin were serially cut into 5-µm thick sections. Every 40th section (approximately 20-30 sections per organ) was stained with hematoxylin and eosin. Morphometric measurements were made by a point counting method as previously described (11) using an Olympus BX50 microscope and image analysis software (Micro Image, Version 4.0, Olympus Optical Co. GmbH, Hamburg, Germany). The test areas were randomly chosen and each image, acquired using a digital camera, was saved, overlaid with the corresponding grid and analyzed.

Absolute volumes of the main thymic compartments, i.e., cortex, medulla and subcapsular/interlobular connective/adipose tissue, were estimated from the volume of the processed and embedded organ and volume density (Vv) of the corresponding compartment. The relative amount of thymic tissue shrinking during processing and embedding (approximately 34%) in all groups was determined stereologically, as described earlier (11). Thus, all stereological data refer to fixed thymic tissue. Each thymic compartment Vv was determined at 40X magnification using an orthogonal test grid with 130 points, and by dividing the number of test points hitting the analyzed structure by the total number of test points falling on the organ. The overall number of test areas was 100 per animal.

The total number of thymocytes in the thymic compartment was calculated from the numerical density (Nv) of thymocytes, as the number of cells per volume unit, and the absolute volume of that compartment. When calculating the overall number of cortical thymocytes it was taken into consideration that: i) two cortical subcompartments, i.e., outer and deep cortex (12) can be delineated morphologically and functionally, ii) three quarters of cortical thymocytes are situated in the deep cortex (13). The Nv of thymocytes was estimated at immersion magnification using a grid that corresponds to the multipurpose M42 test-system. The test grid was placed randomly, but positioned parallel to and just touching the capsule for the outer cortex and the cortico-medullary junction for the deep cortex analysis, respectively. To estimate the Nv of medullary thymocytes, the grid was placed randomly throughout the medulla. For each thymic compartment 60 test areas per animal were measured.

**Preparation of thymocyte suspensions**

Thymocyte suspensions were prepared by gently grinding the thymic tissue on a sterile 60-µm sieve screen in complete RPMI-1640 medium. These cell suspensions were washed in ice-cold complete RPMI-1640 medium, and the number of cells in each of them was enumerated using an improved Neubauer hemocytometer. As expected (14), the total number of thymocytes/thymus thus estimated did not significantly differ from the corresponding value measured morphometrically. The viability of cell preparations (as determined by Trypan blue exclusion) was routinely greater than 95%.
Analysis of cycling cells

Propidium iodide binding to DNA was utilized to identify cells in active phases of the cell cycle. As previously described (14), 100 µL RPMI 1640 with 5 µg/mL concanavalin A (ConA, Sigma-Aldrich Chemie, Taufkirchen, Germany) was added to 100 µL of a cell suspension containing 2.5 x 10^5 thymocytes dispersed into plastic 96-well plates (Nunc A/S, Roskilde, Denmark) to attain a final ConA concentration of 2.5 µg/mL. All cultures were run in triplicate. Cells were harvested after 48 h of culture. A total of 0.5 x 10^6 cells was fixed by the addition of 400 µL ice-cold absolute ethanol, on ice, for 30 min. The cells were then centrifuged at 300 g for 10 min and the supernatant was decanted. After the addition of 1 mg/mL RNAse 1A (Sigma-Aldrich Chemie) in 250 µL phosphate-buffered saline, pH 7.4, to the pellet and vortexing, the samples were incubated in a water bath at 37ºC for 20 min. Propidium iodide (Sigma-Aldrich Chemie) at 20 µg/mL in 250 µL was then added and the samples were gently mixed and incubated for at least 10 min at room temperature. Next, the cells were passed through a fine nylon mesh and analyzed with a FACScan flow cytometer (Becton Dickinson, Mountain View, CA, USA). The gate was set for singlet populations so that doublets and higher order cell aggregates would be excluded from DNA analysis.

Detection of apoptotic thymocytes

Since apoptotic thymocytes are normally rapidly eliminated by phagocytes in vivo, the relative number of apoptotic cells was quantified after 18 h of cultivation, as previously suggested (15). A 100-µL aliquot of the cell suspension (at a concentration of 2.5 x 10^6/mL in RPMI 1640 complete medium) was added to each well of a 96-well flat-bottom plate (Nunc A/S). The cells were incubated for 18 h at 37ºC in a 5% CO₂ humidified atmosphere. Apoptotic thymocytes were detected using merocyanine 540 (MC540). Similarly to annexin-V, this lipophilic dye stains an altered packing order of phospholipids in the outer leaflet of the apoptotic cell plasma membrane. Thus, the percentage of apoptotic cells labeled with MC540 is shown to be equivalent to that obtained by annexin-V staining (16).

The procedure described by Mower and collaborators (17) was used for MC540 staining. All samples were analyzed on the same day with the FACScan flow cytometer (Becton-Dickinson) using CellQuest Software (Becton-Dickinson).

Statistical analysis

Data are reported as means ± SEM. Thymic weight and stereological parameters from Sx and non-operated rats of the same age were compared by the Mann-Whitney U-test. Since none of the values of the parameters analyzed differed significantly between Sx and non-operated rats of the same age, the data for these two groups were pooled and presented as one group (Sx) in all graphs. The differences among groups of different ages were tested by non-parametric Kruskal-Wallis one-way analysis of variance followed by the Mann-Whitney U-test whenever differences were found. The SPSS 10.0 software for Windows was used for statistical analysis.

Results

Histology

Thymuses from 2- and 4-month-old control rats displayed normal morphology: well-defined cortical regions, densely packed with thymocytes, distinct medullary regions, thin interlobular septa, and small amounts of subcapsular adipose tissue (Figure 1, Panel A). However, in 7- and particularly in 10-month-old rats (Figure 1, Panel B), striking struc-
structural changes were observed. The thickness of the thymic capsule and septa increased and a substantial quantity of adipose tissue accumulated under the capsule and in the interlobular spaces. The borderline between the cortex and medulla appeared less distinct.

No prominent differences in thymic structure were observed between Cx and age-matched controls.

**Computer-assisted morphometric analysis**

Age-associated changes in control rats. The results showed that the relative thymic weight (ratio to 100 g body weight) of control rats progressively decreased from the age of 30 days to 7 months, and then remained at that level until the age of 10 months (Figure 2, Panel A, b). Furthermore, they revealed that: i) the absolute thymic weight of the same animals was maximal around 2 months of age; ii) between the ages of 2 and 4 months regressive changes took place, so that at 4 months of age thymic weight was significantly reduced by an average of 27% (Figure 2, Panel A, a). This decrease reflected a significant reduction in the size of the thymic cortex (Figure 2, Panel B, a). Absolute thymic weight did not decrease further with age (Figure 2, Panel A, a). However, the volume of thymic cortex continued to shrink until the age of 7 months, and remained at that level for the next 3 months. In spite of the decrease in cortical volume between the ages of 4 and 7 months, the absolute thymic weight remained unaltered due to an increase in the volume of thymic connective/adipose tissue (Figure 2, Panel B, c). Although further enlargement in the volume of this tissue was recorded between the ages of 7 and 10 months, overall thymic weight remained unaffected.

The relative thymic cellularity (ratio to 100 g body weight) showed a progressive decrease from 30 days up to 10 months of age (Figure 3, Panel A, b). The absolute thymic cellularity exhibited a similar pattern of changes to that shown by absolute organ weight except for an additional significant decrease.
decrease between the ages of 7 and 10 months (the average thymic cellularity diminished by 66% from 2 to 10 months of age; Figure 3, Panel A, a). The age-associated changes in thymic cellularity reflected alterations in cortical thymocyte numbers at all time points except for the age of 10 months, when the reduction in organ cellularity was related to a significant decrease in the number of medullary thymocytes due to a decline in their numerical density (Figure 3, Panels B, C).

Effects of orchidectomy. Orchidectomy at the age of 30 days produced a significant increase in relative and absolute thymic weight (Figure 2, Panel A, a,b), as well as in relative and absolute thymocyte number at all time points except for the age of 10 months (Figure 3, Panel A, a,b). At that age neither the relative and absolute thymic weight nor any of the morphometric parameters significantly differed between Cx and age-matched control rats. Regardless of age (2, 4, or 7 months), the increase in the thymic weight of Cx animals reflected an increase in thymic cortical volume (Figure 2, Panel B, a), whereas at the ages of 2 and 4 months increases in medullary volume also contributed to enlargement of the organ (Figure 2, Panel B, b). The significant increase in cortical volume in all groups of Cx rats correlated with the rise in cellularity in this compartment (Figure 3, Panel B, a). However, only in 4-month-old rats did the increase in medullary volume reflect changes in the number of thymocytes (Figure 3, Panel C, a). In 2-month-old animals the greater volume of this compartment was most likely related to an increase in volume of its non-lymphoid component. Namely, in the medulla of 2-month-old rats a significant reduction in thymocyte Nv was found (Figure 3, Panel C, b) and accordingly, in spite of the increased

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**Figure 2.** Age-related changes in A, absolute thymic weight (a) and relative thymic weight (b), as well as B, volume of different thymic compartments: (a) cortex, (b) medulla, and (c) connective/adipose tissue, in rats orchidectomized at the age of 30 days (Cx) and control animals (Sx). Data are reported as means ± SEM for N = 5. A2-month-old Sx vs 30-day-old Sx; B4-month-old Sx vs 2-month-old Sx; C7-month-old Sx vs 4-month-old Sx; D10-month-old Sx vs 7-month-old Sx; #Cx vs age-matched controls; 4-month-old Cx vs 2-month-old Cx; 7-month-old Cx vs 4-month-old Cx; 10-month-old Cx vs 7-month-old Cx. *P < 0.05 and **P < 0.01; #P < 0.05 and ##P < 0.01 (Mann-Whitney U-test).
Figure 3. A, Age-related changes in the (a) total and (b) relative number of thymocytes, as well as in the number of thymocytes (B, a and C, a) and in thymocyte numerical density in the (B, b,c) thymic cortex and (C, b) medulla in rats orchidectomized at the age of 30 days (Cx) and control animals (Sx). Data are reported as means ± SEM for N = 5. A2-month-old Sx vs 30-day-old Sx; B4-month-old Sx vs 2-month-old Sx; C7-month-old Sx vs 4-month-old Sx; D10-month-old Sx vs 7-month-old Sx; *Cx vs age-matched controls; B4-month-old Cx vs 2-month-old Cx; C7-month-old Cx vs 4-month-old Cx; D10-month-old Cx vs 7-month-old Cx. *P < 0.05 and **P < 0.01; #P < 0.05 (Mann-Whitney U-test).
volume of the whole compartment, the overall number of lymphoid cells in this compartment remained unaltered (Figure 3, Panel C, a).

**Age-associated changes in Cx rats.** Analysis of age-associated changes in Cx rats demonstrated a progressive decrease in relative and absolute thymic weight between the ages of 2 and 10 months, so that at 10 months neither relative nor absolute thymic weight differed significantly between Cx and Sx rats (the average thymic weight was reduced by 57%; Figure 2, Panel A, a,b). This decrease was particularly pronounced between the ages of 4 and 10 months (the average weight was reduced by 45%). The reduction of absolute thymic weight mainly reflected the decrease in cortical volume (Figure 2, Panel B, a). A significant decrease in medullary volume (Figure 2, Panel B, b) was found to contribute to the reduction of thymic weight only in 4-month-old rats. In 10-month-old rats a significant enlargement of connective-adipose tissue volume was also recorded (Figure 2, Panel B, c).

In Cx rats both absolute and relative thymic cellularity also exhibited a progressive decrease between the ages of 2 and 10 months (the average number was reduced by 82%; Figure 3, Panel A, a,b). The changes in thymic cellularity primarily reflected the reduction in number of cortical thymocytes (Figure 3, Panel B, a) due to the progressive loss of these cells in the deep cortex (at all time points) and in the outer cortex (at 10 months of age), which was evident from the reduced Nv of thymocytes in the deep and outer cortical compartments, respectively (Figure 3, Panel B, b,c). Between the ages of 7 and 10 months a significant loss in medullary thymocytes leading to a decrease in Nv of thymocytes was also recorded (Figure 3, Panel C, a,b).

**Thymocyte proliferation**

To clarify the mechanisms underlying age-dependent alterations in thymic cellularity in ConA+ thymocyte cultures from 2- and 10-month-old rats (i.e., at time points when the most striking changes in thymic weight and cellularity were recorded) the percentage of cells in the S/G2M active phases of the cell cycle was examined.

The results showed that, irrespective of age, in thymocyte cultures from Cx rats there was a greater percentage of cells in the S/G2M phases of the cell cycle than in the corresponding cultures from age-matched
control rats (Figure 4, Panel B). Furthermore, irrespective of gonadal presence, the percentage of cells in the S/G2M phases of the cell cycle was greater in thymocyte cultures from 2-month-old rats than in corresponding cultures from 10-month-old rats.

It should also be pointed out that the percentage of apoptotic cells forming a flow cytometric peak to the left of the G1 peak (Figure 4, Panel A), which is often referred to as the sub-G1 peak (18), was significantly (P < 0.01) greater in thymocyte cultures from 10-month-old rats (68.16 ± 1.4% in Sx rats and 75.28 ± 0.8% in Cx rats) than in the corresponding cultures from 2-month-old rats (25.71 ± 1.7% in Sx rats and 22.15 ± 2.34% in Cx rats).

Thymocyte apoptosis

Since not only alterations in thymocyte proliferation, but also in thymocyte apoptosis may affect thymic cellularity, thymocyte sensitivity to apoptotic signaling was estimated at the same time points at which thymocyte proliferative capacity was examined. In thymocyte cultures from 2-month-old Cx rats the percentage of apoptotic cells was significantly lower than in the age-matched controls. However, on the contrary, in thymocyte cultures from 10-month-old rats the percentage of apoptotic cells was increased in Cx compared to Sx rats (Figure 5, Panel B). Nevertheless, the percentage of apoptotic cells was significantly lower in thymocyte cultures from both Cx and control 2-month-old rats than in the corresponding cultures from 10-month-old rats (Figure 5, Panel B).

According to the intensity of MC540 fluorescence, on the one hand, and forward scatter, on the other (19), two subsets of apoptotic cells can be distinguished: 1) cells in early apoptosis, and 2) cells in advanced/late apoptosis (Figure 5, Panel A). Cells in early apoptosis exhibit a high level of MC540 staining, while those in advanced/late apoptosis show lower levels of MC540 staining.
and lower forward scatter. In thymocyte cultures from 2-month-old Cx rats the reduction in the percentage of apoptotic cells reflected decreases in the relative numbers of cells in both the early and late phases of apoptosis (Figure 5, Panel B). On the other hand, the higher percentage of apoptotic cells in 10-month-old Cx rats compared to age-matched control rats was mainly due to a rise in the proportion of cells in late apoptosis (Figure 5, Panel B).

Discussion

This study describes the age-related changes in weight, cellularity and morphometric indices of the thymus in peripubertally Cx and control AO rats. We have confirmed that gonadal hormone removal in the peripubertal period leads to a significant increase in thymic weight and cellularity in young adult rats (8), and have shown that this organ reaches its maximum weight and cellularity one month after surgery. Finally, to the best of our knowledge, for the first time it has been demonstrated that in rats the effects of peripubertal orchidectomy on the thymus are long-lasting but not permanent, i.e., that removal of the testes can only postpone but not prevent the age-related organ atrophy.

The present findings confirm that, after an initial decline in early adulthood, thymic weight in rats (3), as in humans (9) and some other mammals (20), but not in mice (21-23), remains unaltered during a relatively long period of time (from 4 to 10 months of age). Although thymic weight in 10-month-old rats did not differ from values for 4- and 7-month-old animals, the total number of thymocytes in both thymic compartments, and hence the overall organ cellularity, were significantly reduced reflecting, most likely, an age-associated increase in thymocyte sensitivity to apoptotic stimuli followed by decreased responsiveness to proliferative signaling. This assumption emerged from our findings obtained with thymocyte cultures, which are fully consistent with previously reported data on cell sensitivity to apoptotic and proliferative signaling in thymocyte cultures from aged Wistar rats (14). The age-associated expansion of connective-adipose tissue described here, which compensates for the loss of lymphoid tissue, has also been observed in aged male Wistar rats (24). Therefore, these results support our previous findings that the overall thymic weight of aging rats does not correlate with the amount of functional lymphoid thymic tissue (24).

Orchidectomy at the age of 30 days produced a pronounced increase in both thymic weight and cellularity at all examined points from 1 to 6 months post-castration, but at 9 months post-castration, none of these parameters significantly differed between Cx and age-matched controls. At all time points examined between 1 and 6 months post-castration both the size and the cellularity of the cortical compartment were increased, while these parameters of the thymic medulla were augmented only at 3 months after castration. The increases in thymic weight and cellularity evoked by orchidectomy may be associated with: i) improved precursor generation in the bone marrow (25), ii) alterations in the thymic microenvironment resulting in more efficient attraction of thymocyte progenitors (26), iii) increased thymocyte sensitivity to proliferative stimuli (27), and iv) reduced thymocyte sensitivity to apoptotic stimuli (28). Our results showed that 1 month after surgery (when thymic weight and cellularity were at their maximum), the increase in the overall thymocyte number most probably reflected an increased thymocyte sensitivity to proliferative stimuli on the one hand, and reduced sensitivity to apoptotic stimuli on the other. In agreement, there are data showing that androgens inhibit thymocyte proliferation (29) and accelerate thymocyte elimination via apoptosis (28,30). The present findings also demonstrate that 9 months after orchidectomy nei-
ther thymic weight nor any estimated morphometric parameter significantly differed from those in age-matched controls. This is in keeping with the very recent conclusion of Min and colleagues (10) that the orchidectomy-induced effect on thymic weight in mice is of limited duration. In other words, our results indicate that removal of the testes in rats, as in mice (10), postpones age-associated thymic involution, but does not prevent it completely. Furthermore, our results also indicate that a new balance between thymocyte apoptosis and proliferation was established in these animals. Differently from thymocyte cultures from 2-month-old rats, the percentage of apoptotic cells was significantly greater in thymocyte cultures from Cx rats compared with age-matched controls. Moreover, although the proportion of cells in the S/G2M phases of the cell cycle was significantly greater in ConA+ thymocyte cultures from 10-month-old Cx rats than in those from age-matched Sx rats, it was significantly less than in thymocyte cultures from 2-month-old Cx rats. A compensatory intrathymic increase in androgen production sufficient to cause accelerated thymocyte apoptosis, but still not sufficient to prevent proliferation of these cells may be assumed to explain these findings. In favor of this hypothesis are data suggesting that both lymphocytes (31) and TEC may produce steroid hormones (32). Furthermore, analysis of thymocyte differentiation/maturation under the same experimental conditions showed that alterations in the relative proportion of thymocytes (delineated by expression of CD4, CD8 and T-cell receptor) become quantitatively more pronounced with duration of gonadal deprivation (8). Thus, the present findings, taken together with those previously published (8), indicate that, although 9 months after orchidectomy a new balance between the main thymic homeostatic processes (i.e., apoptosis and proliferation) is established, so that thymic cellularity does not significantly differ between Cx and control rats, the thymocyte differentiation kinetics remains substantially altered, producing an accumulation of cells at one maturational step and depletion at the subsequent downstream steps. Moreover, the age-associated decline in thymic weight and cellularity in Cx rats was even more pronounced than that in non-operated animals and, in contrast to control rats, both thymic weight and cellularity exhibited a progressive reduction between the ages of 2 and 10 months. This suggests that: i) age-associated thymic atrophy, which is initially sensitive to circulating androgen levels, becomes independent of them with progression of the process and ii) the kinetics of age-associated thymic involution depends on the presence of circulating gonadal hormones.

To explain the limited duration of effects induced by gonadal hormone ablation on age-related thymic atrophy several possibilities should be taken into consideration. First, as mentioned above, lymphocytes themselves have been shown to produce androgens (31), so that a post-castrational compensatory increase in intrathymic androgen production from re-entrant and/or developing lymphocytes cannot be ruled out. Furthermore, since TEC possess a complete intracellular machinery for the synthesis of steroid hormones, and since they are supposed to produce these hormones (32), a compensatory increase in TEC androgen production should also be taken into account. In support of the previous assumption are data indicating that: i) the ratio of non-lymphoid to lymphoid component volume was increased in the thymic medulla of rats subjected to gonadal deprivation for 9 months and ii) androgens influence TEC proliferation in a biphasic mode, reducing their proliferation at greater concentrations than optimal in a dose-dependent manner, and increasing it at less than optimal concentrations (33). Second, compensatory production of adrenal androgens should be taken into account. However, it has been shown...
that the adrenals of rats and mice do not synthesize androgens (34), so that in these animals orchidectomy is not followed by increased serum levels of either dehydroepiandrosterone or testosterone (35) and this hypothesis should be rejected. Third, it may be assumed that thymic involution is mediated by distinct mechanisms acting during different periods of ontogenesis or that some other, non-androgen-mediated mechanism (e.g., growth hormone-dependent) (36), may take over the androgen role in long-term androgen-deprived rats. At present there are no data available to strongly support this hypothesis. Finally, it may also be hypothesized that gonadal hormones are important in the initial phases of thymic regression, while later accumulating age-related functional or structural defects (37) are responsible for the maintenance/progression of thymic involution. In agreement with this hypothesis are data showing that: i) normal, age-related androgen depletion (38) exerts no beneficial effect on the aged thymus (39), and ii) castration of old rats does not restore thymic weight to the level found in young animals (40).

In conclusion, this study has clearly shown that gonadal ablation cannot prevent thymic involution, suggesting that even if circulating gonadal hormones are predominantly responsible for the initiation of age-associated thymic involution, maintenance of their circulating levels is not necessary to secure maintenance/progression of the age-associated regressive thymic changes. To further clarify the role of androgens in the initiation, and particularly in the maintenance/progression of age-related thymic atrophy, experiments addressing not only thymic effects evoked by long-lasting withdrawal of circulating androgens, but also those induced by blockade of intrathyMICally synthesized/released androgen action, should be undertaken.

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