Impairment of Spermatogonial Development and Spermiation after Testosterone-Induced Gonadotropin Suppression in Adult Monkeys (Macaca fascicularis)*

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ABSTRACT

Human male hormonal contraceptive regimens do not consistently induce azoospermia, and the basis of this variable response is unclear. This study used nine adult macaque monkeys (Macaca fascicularis) given testosterone (T) implants for 20 weeks to study changes in germ cell populations in relation to sperm output. Germ cell numbers were determined using the optical dissector stereological method. Four animals achieved consistent azoospermia (azoospermic group), whereas five animals did not (nonazoo group). T-induced gonadotropin suppression in all animals decreased A pale (Ap) spermatogonia to 45% of baseline within 2 weeks, leading to decreased B spermatogonia (32–38%) and later germ cells (20–30%) after 14 and 20 weeks. Though the reduction in later germ cell types could be primarily attributed to the loss of spermatogonia, the data suggested that some cells were lost during the spermatocyte and spermatid phase of development. B spermatogonial number was more markedly suppressed in azoospermic animals, compared with the nonazoo group, as was the conversion ratio between Ap and B spermatogonia. Abnormal retention of elongated spermatids (failed spermiation) was also prominent in some animals and spermiation failure. We conclude that: 1) the variable suppression of sperm output is attributed to the degree of inhibition of germ cell development from type B spermatogonia onwards, and this is related to the degree of FSH suppression; and 2) inhibition of Ap and B spermatogonial development and failure of spermiation are the major defects caused by long-term T administration to monkeys. (J Clin Endocrinol Metab 86: 1814–1822, 2001)

The administration of exogenous testosterone (T), either alone or in combination with a progestin, reduces the secretion of the pituitary gonadotropins, LH and FSH, and thereby sperm production, and is a promising approach to male contraception (1). A feature of many such contraceptive formulations is the variable induction of azoospermia, ranging from 70–95%, depending on the regimen and ethnic group under study (1–4). Though sperm counts of less than 3 million per mL may provide adequate contraception (4), there is a general consensus that the reliable induction of azoospermia is important to ensure contraceptive efficacy and the widespread acceptance of male hormonal contraception. An understanding of the biological basis for the variable response is essential to this goal.

Although many studies have investigated the effects of hormone suppression on spermatogenesis in the rat (5–7), fewer studies have investigated the effects of FSH and LH/testicular T suppression on germ cell development in monkeys and man. In monkeys rendered gonadotropin-deficient by hypophysectomy or GnRH antagonist treatment, spermatogenesis are the most sensitive germ cell type (8–11); however, which spermatogonial subtypes are regulated by hormones is controversial (9, 11–14). A recent study in men receiving weekly injections of T enanate suggested that the inhibition of spermatogonial development was also the primary lesion in response to gonadotropin withdrawal and further suggested that the failure of sperm release may also be important (15). In accordance with this latter observation, our recent study in rats showed that spermatiation failure contributes to the suppression of spermatogenesis after hormone withdrawal (16).

Potential T-based contraceptive regimens in monkeys and humans clearly inhibit various phases of germ cell development; however, the relationship between these changes and the extent of suppression of sperm output is unclear. Our recent results in monkeys given T implants indicated that animals who achieve stable azoospermia in response to 20 weeks of T administration have significantly lower levels of circulating FSH, compared with those animals who did not achieve consistent azoospermia, whereas no differences were seen in LH or T (serum or testicular) levels (17).

The purpose of this study is to describe the time course of the effects of exogenous T administration on germ cell development in macaque monkeys and to relate the degree of...
suppression of germ cell development to sperm output. The optical disector stereological technique was used for the unbiased determination of Sertoli cell and germ cell number in testicular biopsies taken before, during, and after T administration. The data show that reductions in type A pale (Ap) and B spermatogonia are the key lesions in spermatogenesis and that failure of spermiation is also important.

Materials and Methods

Animals and study design

Nine normal adult male crab-eating macaque monkeys, Macaca fascicularis (age, 5–12 yr; BW, 5–9 kg) were housed in the Regional Primate Research Center at the University of Washington, under defined environmental conditions. All procedures were performed in accordance with the Endocrine Society Guidelines for the Care and Use of Experimental Animals (18). All procedures were carried out under Ketamine (Fort Dodge, IA) HCl (10 mg/kg, im) anesthesia. Valium (1 mg/kg; im, Elkins-Sinn, Inc., Cherry Hill, NJ) was given, before Ketamine HCl, to all monkeys, for the electroejaculation procedure.

The study involved a pretreatment (control) phase (4 weeks), treatment phase (20 weeks) terminating with the removal of the implants, and recovery phase (16 weeks). T was administered via sterilized sc SILASTIC implants (5.5-cm long; inside diameter, 0.33 cm; outside diameter, 0.46 cm) for 20 weeks. Two implants were inserted sc in the midscapular region of each animal.

Testicular biopsies

Open testicular biopsies were performed alternatively on the right and left testis, in aseptic conditions, under ketamine/xylazine anesthesia. All animals were biopsied 4 weeks before the treatment (control) and then at weeks 2, 14, and 20 during the treatment and at 12 weeks post treatment. Approximately 300–500 mg tissue was taken per biopsy and was divided in half; half was frozen in liquid nitrogen for the measurement of testicular androgens (Narula et al., submitted). The other half was immersion-fixed in Bouin’s fluid for approximately 5 h, then stored in 70% ethanol. Biopsies were observed under a dissecting microscope and cut along a plane lying at right angles to the dominant tubule orientation, to maximize the number of circular tubule profiles.

Testicular function and hormone data

Body weights, testis volumes, and sperm counts were assessed every 2 weeks during control, treatment, and recovery periods. Testis volumes were measured using an orchidometer. Seminal fluid was collected after rectal probe electroejaculation, and total sperms counts (millions/ejaculate) were determined using an orchidometer. Blood samples were collected from the femoral vein every 2 weeks during the study (Narula et al., submitted). Bioassays were used to measure the bioactivity of LH and FSH as detailed elsewhere (Narula et al., submitted). The detection limits of the assays were as follows: bioLH, 2.4 IU/L; bioFSH, 0.08 IU/L.

Testicular T was measured using previously published methods (19) as detailed elsewhere (Narula et al., submitted). The other half was immersion-fixed in Bouin’s fluid for approximately 5 h, then stored in 70% ethanol. Biopsies were observed under a dissecting microscope and cut along a plane lying at right angles to the dominant tubule orientation, to maximize the number of circular tubule profiles.

Cell identification and groupings

Each seminiferous tubule was staged into one of the following stage groupings based on the cellular associations present in stages I-VI, VII-VIII-XI, and XII (see (Ref. 20), and the cells were counted in the following groups: Sertoli cells, type A dark (Ad) spermatogonia (present in all stages); type Ap spermatogonia (present in all stages); type B (B) spermatogonia (present in stages X-XII and I-VI); preleptotene + leptotene + zygotene (PI-Z) primary spermatocytes (present in stages VII-XI); pachytene primary spermatocytes (PS) (present in all stages); round spermatids (rSt 1–7) (stages I-VII); elongating spermatids (eSt 8–12) in steps 8–12 (stages VIII-XII); and elongated spermatids (eSt 13–14) in steps 13–14 (stages I-VI).

Sertoli cells and germ cells, in 25-μm thick PAS-stained methacrylate sections, were classified on the basis of their nuclear morphology, as previously described by Zhengwei (20), based on the criteria of Clermont (21, 22). Spermatogonia were subdivided into type Ad and type Ap spermatogonia, and type B spermatogonia as follows (see Fig. 1 for morphology of the various subclasses of spermatogonia):

Ad spermatogonia. Ad spermatogonia were variable in size, with a nucleus that showed darker staining, compared with Ap spermatogonia. The nuclei were rounded and often broadly applied to the basement membrane. Many nuclei contained a vacuole and/or lightly stained regions; and in most cases, the cytoplasm contained strong PAS-positive material (Fig. 1).

Ap spermatogonia. Ap spermatogonia had nuclei that were generally larger than Ad spermatogonia and were paler in color. The nuclei were almost always round and showed one or two clumps of chromatin associated with the nuclear membrane. The cytoplasm did not show PAS positive material (Fig. 1).

FIG. 1. Morphology of spermatogonia and Sertoli cell nuclei in 25-μm-thick sections of PAS stained, methacrylate-embedded monkey testicular biopsy material. A, Sertoli cell nucleus, showing characteristic irregular shape and central nucleolus. Bar, 5 μm. B, Ad spermatogonium, showing PAS-positive material in the cytoplasm. This morphology was seen in the majority of Ad spermatogonia. C, Ad spermatogonium with a darkly stained nucleus containing small vacuolated areas and a visible halo around the nucleus. D, Ap spermatogonium with a large, rounded, lightly stained nucleus containing two clumps of chromatin. E, Ap spermatogonium with a slightly lighter stained nucleus and two clumps of chromatin. F, B spermatogonium with a rounded, darkly-stained nucleus, showing the characteristic peripherally located chromatin and a central nucleolus. All micrographs were taken at the same magnification.
**B spermatogonia.** B spermatogonia had nuclei that were rounded, darker in color, with peripherally located clumps of chromatin and a central nucleus. The nuclei were situated further away from the basement membrane than Ad spermatogonia (Fig. 1).

Sertoli cells were clearly distinguishable from spermatogonia by their irregularly shaped pale-colored nucleus and the presence of a single prominent nucleus (see Fig. 1). Criteria for classification of the later germ cell types have been described in detail in our previous study (20).

**Stereological analysis**

The optical dissector method was used to determine the numbers of germ cells and Sertoli cells in each biopsy as previously described (20, 23). This method is an unbiased and efficient technique for estimating cell number, including nonspherical objects such as elongated spermatids, and assumes only that cell number equates to nuclear number. The 25-μm methacrylate sections were optically sectioned by means of a high numerical aperture lens (100× objective lens, numerical aperture 1.4) on an Olympus Corp. (Tokyo, Japan) BX-50 microscope equipped with a microcater (Heidelberg D83301; Traunreut, Germany) attached to the microscope stage to monitor the depth scanned. The image was captured by a Pulnix TMC-6 video camera coupled to a Pentium personal computer using a Screen machine H fast multimedia video adapter (FAST, Hamburg, Germany). A software package, Image 1.10 (Olympus Corp., Munich, Germany) was used to superimpose a set of unbiased counting frames on the video image. Fields to be counted were selected by a systematic uniform random sampling scheme, with the use of unbiased counting frames on the video image. Fields to be counted were selected by a systematic uniform random sampling scheme, with the use of unbiased counting frames on the video image.

**Assessment of spermiation failure**

The release of mature step 14 spermatids via spermiation occurs at the end of stage VI/early stage VII; and thus, mature step 14 spermatid heads located in the basal portion of the epithelium from stage VII and beyond represent retained spermatids that do not undergo spermiation. The number of retained spermatids in stages VII-XII were counted, as per other germ cells, and expressed per Sertoli cell.

To compare the number of spermatids retained within the epithelium, with the number of normal spermatids available for spermiation, the number of retained spermatids in stages VII-XII and steps 13–14 spermatids in stages I-VI (N/SC) were compared for the different duration of stages in which they are present, and were thus multiplied by the relative stage frequency for that stage grouping. The number of retained spermatids per Sertoli cell in stages VII-XII was then divided by the number of steps 13–14 spermatids (in stages I-VI) per Sertoli cell and multiplied by 100, as an estimation of the percentage failing to spermiate. It should be noted that the comparison of retained spermatids to steps 13–14 spermatids is only an approximation of the percentage of spermatids failing to spermiate; because retained spermatids will undergo phagocytosis and disappear over the 136-h period from stage VII to XII, and thus the number of retained spermatids counted will be an underestimation of the actual number retained. However, because of the severity of spermatogenic suppression in most animals, it was not possible to subdivide between these stages; and thus, all retained spermatid data were pooled.

Spermiation failure was only assessed in animals that had spermiogenic cells present in the biopsy during the T-administration period. Because three animals had severe spermatogenic arrest and no elongating or elongated spermatids present in the tests at either the 14- or 20-week treatment periods, these animals were excluded from this analysis.

**Statistics**

For comparison of one endpoint across the study period, data were log-transformed and tested for homogeneity of variance before repeated-measures one-way ANOVA. When a P value of less than 0.05 was obtained by ANOVA, post hoc analysis was performed using Dunnett’s test for differences compared with the control group, and Newman-Keuls post hoc comparison was performed for other differences between the various time points. When comparing between different germ cell populations, expressed as percent control, the loss of germ cells at different points in the spermatogenic cycle was associated with significant heterogeneity in variances; and thus, nonparametric analyses for correlated samples (Friedman ANOVA and Wilcoxon signed-rank test) were used. When comparing data between groups of animals achieving either consistent azoospermia (azoos group) and those that did not (non-azoos group), significant heterogeneity of variance was observed; and thus, the nonparametric Mann-Whitney U test was used. Differences at the level of P < 0.05 were considered significant. Data are expressed as mean ± SEM (n = 9 for all groups except the 14-week treatment group, where n = 8).

**Results**

**Germ cell populations before, during, and after T administration.**

The effect of T administration on the number of each germ cell type (N/SC), in comparison with control values for all nine animals, is shown in Table 1. After 2 weeks of T administration, there were no significant changes in germ cell populations as grouped in the categories shown in Table 1. After 14 and 20 weeks of T administration, significant decreases were seen in all germ cell populations from B spermatogonia through to steps 13–14 elongated spermatids. The number of A spermatogonia (Ad + Ap spermatogonia) was...
and 20 weeks of T administration, and 12 weeks after the removal of implants (recovery). All data is mean ± SEM (n = 9 in all groups except 14 weeks, where n = 8). A, Data for spermatogonial and spermatocyte populations, type A spermatogonia (A, ■), type B spermatogonia (B, ○), preleptotene-zygote spermatocytes (PI-Z, ◇), pachytene spermatocytes (PS, □). B, Data for spermatid populations, steps 1–7 rSt (rSt 1–7, ▼), steps 8–12 elongating spermatids (8–12, ●), steps 13–14 elongated spermatids (13–14, ○).

Table 1. Effect of T implants on germ cell populations (in numbers per Sertoli cell), sperm counts, and hormone parameters in monkeys

<table>
<thead>
<tr>
<th>Germ cells N/SC</th>
<th>Control n = 9</th>
<th>T administration</th>
<th>Recovery n = 9</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2 wks n = 9</td>
<td>14 wks n = 8</td>
<td>20 wks n = 9</td>
</tr>
<tr>
<td>A spermatogonia</td>
<td>0.59 ± 0.04</td>
<td>0.60 ± 0.04</td>
<td>0.38 ± 0.05</td>
</tr>
<tr>
<td>B spermatogonia</td>
<td>0.34 ± 0.04</td>
<td>0.29 ± 0.06</td>
<td>0.09 ± 0.03</td>
</tr>
<tr>
<td>Preleptotene-zygote spermatocytes</td>
<td>1.57 ± 0.22</td>
<td>0.98 ± 0.20</td>
<td>0.40 ± 0.17</td>
</tr>
<tr>
<td>Pachytene spermatocytes</td>
<td>5.86 ± 0.95</td>
<td>5.34 ± 0.60</td>
<td>1.52 ± 0.77</td>
</tr>
<tr>
<td>Round spermatids (steps 1–7)</td>
<td>10.8 ± 1.89</td>
<td>13.0 ± 1.61</td>
<td>2.75 ± 1.50</td>
</tr>
<tr>
<td>Elongating spermatids (steps 8–12)</td>
<td>8.53 ± 1.51</td>
<td>8.87 ± 0.73</td>
<td>1.72 ± 0.76</td>
</tr>
<tr>
<td>Elongated spermatids (steps 13–14)</td>
<td>7.15 ± 0.84</td>
<td>9.33 ± 1.11</td>
<td>2.00 ± 0.92</td>
</tr>
</tbody>
</table>

Sperm counts & hormones

<table>
<thead>
<tr>
<th></th>
<th>2 wks n = 9</th>
<th>14 wks n = 8</th>
<th>20 wks n = 9</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sperm count × 10⁶ per ejaculate</td>
<td>57.7 ± 14.2</td>
<td>48.7 ± 20.4</td>
<td>9.2 ± 4.6</td>
</tr>
<tr>
<td>bio FSH IU/L</td>
<td>4.4 ± 0.5</td>
<td>2.3 ± 0.4</td>
<td>2.0 ± 0.9</td>
</tr>
<tr>
<td>intra-testicular T nmol/L</td>
<td>860.8 ± 309.4</td>
<td>134.0 ± 22.6</td>
<td>56.3 ± 10.8</td>
</tr>
</tbody>
</table>

Repeated-measures ANOVA, followed by Dunnett’s test for comparison to the control group. Data is shown from the control phase, at 2, 14, and 20 weeks of T administration, and 12 weeks after the removal of implants (recovery). All data is mean ± SEM. The number of animals (n) included in each time point is indicated.

a P < 0.05, compared with control.
b P < 0.01, compared with control.

Significantly below control only at the 20-week time point. All germ cell types recovered within 12 weeks after removal of T implants.

To appreciate the relative changes in each germ cell population during T administration, germ cell numbers were expressed as percent of control values (Fig. 2). The total number of type A spermatogonia remained between 73–99% of control, across the treatment period. After 14 and 20 weeks of T administration, the number of type B spermatogonia was suppressed to 20–38% of control values. All later germ cell types were suppressed to a similar degree.

To further investigate the effects of T-induced hormonal changes on the progression of germ cell development, germ cell conversion ratios (CR) were examined after 14 and 20 weeks, when spermatogenic suppression had reached steady state. The primary defect was a significant decrease (P < 0.01) in the CR of A-to-B spermatogonia at the 14- and 20-week treatment time points (CR, 0.57 ± 0.05 in control biopsies vs. 0.21 ± 0.04 after 14 weeks, and 0.25 ± 0.06 after 20 weeks, both P < 0.01). The only other significant changes in CR were seen after 20 weeks, when there was a decrease in the transition of pachytene spermatocytes through to round spermatids (1.88 ± 0.12 control vs. 1.07 ± 0.37 20 weeks, P < 0.05) and a decrease in the transition through the later phases of spermiogenesis, from steps 8–12 to steps 13–14 spermatids [1.04 ± 0.14 control vs. 0.73 ± 0.18 (20 weeks), P < 0.05].

Effect of T administration on sperm counts and hormone levels

T administration caused significant decreases in sperm counts, intratesticular T concentrations, and the levels of circulating bioactive FSH and LH (Table 1). All parameters returned to control levels within 12 weeks after the removal of T implants. All animals achieved maximal suppression of hormone levels and spermatogenic parameters after 12 weeks of T implants (Fig. 3). In addition, there was a decrease in the transition of pachytene spermatocytes through to round spermatids (1.88 ± 0.12 control vs. 1.07 ± 0.37 20 weeks, P < 0.05) and a decrease in the transition through the later phases of spermiogenesis, from steps 8–12 to steps 13–14 spermatids [1.04 ± 0.14 control vs. 0.73 ± 0.18 (20 weeks), P < 0.05].

Effect of T administration on spermatogonial subtypes

In the monkey, the type A spermatogonia population is comprised broadly of two morphologically distinct classes of spermatogonia. Ad spermatogonia (see Fig. 1) are considered to be the nonproliferative reserve spermatogonial population (21, 22, 24), whereas the Ap spermatogonia (see Fig. 1) are considered to provide the differentiating cells committed to spermatogenesis (21, 22, 24, 25). The effects of T adminis-
As shown in Fig. 3, type Ap spermatogonia were suppressed when the CR between the spermatogonial subtypes was considered separately. Changes were seen when the total A spermatogonial number was considered, whereas Ap spermatogonia were significantly below control values. When the number of type A spermatogonia (i.e., Ad + Ap spermatogonia) was unaltered (Table 1, Fig. 3), changes were seen when the type A spermatogonial subtypes were considered separately. As shown in Fig. 3, type Ap spermatogonia were suppressed to 45% of control ($P < 0.05$), and Ad spermatogonia increased to 165% of control ($P < 0.05$). Despite these changes, no change was seen in B spermatogonia at this time.

After 14 weeks of T administration, Ap spermatogonia remained suppressed, whereas Ad spermatogonia had declined to control levels (Fig. 3). Type B spermatogonia were now suppressed to an extent similar to that of the Ap spermatogonia (38% and 32% of control for Ap and B, respectively). After 20 weeks of T administration, the number of Ad spermatogonia had risen slightly, whereas the number of Ap spermatogonia decreased further (compared with 14 weeks). When the total A spermatogonial number was considered, there was a significant decrease at 20 weeks. Type B spermatogonia remained suppressed after 20 weeks of T administration. When the CR between the spermatogonial subtypes were analyzed, the only significant decrease was seen in the transition of Ap to B spermatogonia after 20 weeks (not shown).

Twelve weeks after removal of the T implants, the total number of A spermatogonia returned to control values (Fig. 3). Ad spermatogonia were significantly elevated above control, whereas Ap spermatogonia were significantly below control values.

### Failure of spermiation during T administration

There was a significant linear relationship between sperm count and the number of steps 13–14 elongated spermatids per Sertoli cell, after both 14 ($r = 0.93$, $P < 0.001$, $n = 8$) and 20 weeks ($r = 0.81$, $P < 0.01$, $n = 9$) of T administration. However, in some animals, the sperm count did not reflect the number of mature spermatids (steps 13–14) in the testis. For example, animal no. 97114 was azoospermic at 18 and 20 weeks but had 35% the control number of steps 13–14 elongated spermatids in the testis, suggesting a failure of spermiation. Therefore, the failure of spermiation was assessed during T administration. Because three animals had severe spermatogenic suppression after long-term T administration, and no elongating or elongated spermatids were present in their testes, these animals were excluded from the analysis of spermiation.

In control testicular biopsies, few mature retained spermatids were seen in the basal portion of the epithelium ($0.023 \pm 0.016$ N/SC, $n = 6$), and similar numbers were seen after 2 weeks of T administration ($0.046 \pm 0.016$, $n = 6$). After 14 and 20 weeks of T administration, the numbers of retained spermatids per Sertoli cell increased to $0.071 \pm 0.018$ (n = 5) and $0.235 \pm 0.179$ (n = 6), respectively; however, there was marked between-animal variability. To make an estimate of the degree of spermiation failure, the number of spermatids retained in the epithelium was compared with the number of spermatids available to spermiate (Table 2). Animals that had higher sperm counts had less retained spermatids and a lower percentage of spermatids retained (Table 2). A significant negative correlation was seen between sperm count and the percentage of spermatids retained after 14 ($r = -0.83$, $n = 5$, $P < 0.02$) and 20 weeks ($r = -0.94$, $n = 6$, $P < 0.001$).

### Heterogeneity of response to T administration

Animals showed heterogeneous suppression of sperm counts, with some animals achieving consistent azoosperma, whereas others achieved variable degrees of spermatogenic suppression (17). As described elsewhere (17; and Narula et al., submitted), animals were assigned into two groups: those showing sustained azoosperma up to 20 weeks of T administration (azoos group, n = 4), and those that did not show consistent suppression, with consistently detectable sperm counts ranging between 1 and 40% of their control values (nonazoos group, n = 5).

Figure 4 shows germ cell development (N/SC expressed as a percent of control) at 14 weeks of T administration in the azoo and nonazoo groups. There was a general trend for germ cell numbers to be lower in the azoo, compared with the nonazoo group, from B spermatogonia onwards. When the data were expressed as percent of control, significant decreases in the azoo group were seen in pachytene sper-
TABLE 2. Spermatid retention in the seminiferous epithelium, compared with sperm counts, after 14 and 20 weeks of T administration

<table>
<thead>
<tr>
<th>Animal ID</th>
<th>14 weeks T administration</th>
<th>20 weeks T administration</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Retained spermatids N/SC</td>
<td>Retained spermatids N/SC</td>
</tr>
<tr>
<td>94029</td>
<td>7.20</td>
<td>0.058</td>
</tr>
<tr>
<td>94028</td>
<td>3.43</td>
<td>0.030</td>
</tr>
<tr>
<td>97105</td>
<td>3.59</td>
<td>0.121</td>
</tr>
<tr>
<td>86185</td>
<td>1.56</td>
<td>0.103</td>
</tr>
<tr>
<td>93042</td>
<td>na</td>
<td>na</td>
</tr>
<tr>
<td>97114</td>
<td>0.70</td>
<td>0.041</td>
</tr>
</tbody>
</table>

na, Biopsy not available. Data from individual animals is shown.  
<sup>a</sup> Animals that did not have any elongated spermatids present in the testis were not included in the analysis (n = 3).  
<sup>b</sup> The number per Sertoli cell of steps 13–14 elongated spermatids.  
<sup>c</sup> The number per Sertoli cell of step 14 elongated spermatids retained in the basal portion of the epithelium in stages VII–XII.  
<sup>d</sup> Calculated by dividing the N/SC of retained spermatids by the N/SC of steps 13–14 spermatids, correcting for the different stage durations in which they are present.  
<sup>e</sup> Millions per ejaculate.

matocytes, round spermatids, and steps 13–14 elongated spermatids. When the data were expressed as N/SC, germ cell numbers were significantly lower in the azoo, compared with the nonazoo, group. There were no significant differences in the numbers of germ cells between azoo and nonazoo groups in any other germ cell CR.  

Fig. 5 shows the changes in the spermatogonial subtypes in the azoo vs. nonazoo groups after 14 weeks of T administration. Animals in the azoo group had significantly less Ad and B spermatogonia per Sertoli cell but similar numbers of Ap. After 20 weeks, a similar pattern was observed, except that there was no difference in the number of Ad spermatogonia between the two groups (not shown).

When the CR of germ cells were considered, to assess the efficiency of the transition of germ cells through each developmental step, there was a significant decrease in the transition of Ap to B spermatogonia in the azoo (vs. nonazoo) group at both 14 weeks (0.46 ± 0.10 azoo vs. 1.27 ± 0.05 nonazoo, P < 0.05) and 20 weeks (0.23 ± 0.06 azoo vs. 0.82 ± 0.21 nonazoo, P < 0.05) of T administration, but there were no differences in any other germ cell CR.

Although there was no difference in the levels of FSH between the azoo and nonazoo groups during the control phase, animals in the azoo group had significantly lower immuno-FSH and bioFSH across the 20-week T-administration phase, as detailed elsewhere (17; and Narula et al., submitted). After 14 weeks of T administration, bioFSH was 8% of control in the azoo group vs. 77% of control in the nonazoo group (P < 0.05). Similarly, after 20 weeks, bioFSH was 22% of control in the azoo group vs. 65% of control in the nonazoo group (P < 0.05). No differences were seen between the azoo vs. nonazoo groups, during the treatment period, in any other endocrine parameter, including serum T, bioLH, and intratesticular androgens (17; and Narula et al., submitted).

**Discussion**

In this study, we have used stereological techniques to measure the changes in germ cell development in adult macaque monkeys administered exogenous T for 20 weeks. This animal provides an excellent model for the study of human hormonal contraception, showing profound suppression of serum gonadotropin levels, but variable patterns of sper-
matogenic suppression, in response to steroid treatment (17; Narula et al., submitted). We show that the primary lesion in all animals relates to type Ap spermatogonial development. Moreover, in animals achieving consistent azoospermia, compared with animals with variable spermatogenic suppression, the key germ cell determinant of the response of sperm counts seemed to be the more profound suppression of the conversion of type Ap to B spermatogonia, which was related to a more profound suppression of circulating FSH. We also show that during T administration, other sites in spermatogenesis are affected, including spermatocytes, spermatids, and the spermiogenesis process.

The arrest of spermatogenesis at the level of type A spermatogonia, in response to gonadotropin withdrawal, is supported by other studies in monkeys, after GnRH antagonist administration (9–11) and hypophysectomy (8), and in humans given weekly injections of T enanthate (15). In the current study, we showed that, whereas total A spermatogonial numbers were unchanged after 2 weeks of T administration, a decline in Ap spermatogonial number was apparent; and thus, Ap spermatogonia were the first germ cell type to show a significant decrease after gonadotropin withdrawal. In all animals, a major reduction in Ap spermatogonial number was seen at 14 and 20 weeks of T administration, and CR data showed that the progression of germ cells through the pachytene spermatocyte stage and spermiogenesis was also impaired, albeit to a lesser extent. We speculate that losses of pachytene spermatocytes and spermatids result from apoptosis, as has been previously described in gonadotropin-deprived rats (26, 27).

We recently reported that monkeys achieving consistent azoospermia had significantly lower levels of circulating FSH (both immuno- and bioactive FSH) (17; Narula et al., submitted), whereas no differences were seen in serum bi- or T, or testicular androgen levels. These data suggest that residual FSH plays a key role in supporting spermatogenesis during T-induced gonadotropin suppression. The heterogenous sperm count response seen in these monkeys also allowed us to compare germ cell numbers, sperm counts, and FSH levels in animals achieving either consistent azoospermia or variable spermatogenic suppression. The 14-week time point during T administration was investigated to understand changes in the spermatogenic process that would promote subsequent changes in sperm output during the remaining 6 weeks of the T-administration phase. Analysis of the germ cell data at 14 weeks showed that, whereas Ap spermatogonia were similarly suppressed in both the azo and nonazo groups, all other germ cell types were lower in the azospermic animals. Analysis of CR between germ cell types showed that the reduction in the transition of Ap to B spermatogonia was a major subsequent determinant of sperm output. Therefore, induction of consistent azoospermia is associated with lower circulating FSH levels (17; and Narula et al., submitted), a greater impairment of either B spermatogonial survival or in the transition of Ap to B spermatogonia, leading to fewer germ cells being available to proceed through spermatogenesis.

Type A spermatogonia in the primate and the human can be divided into at least two morphologically distinct subtypes: Ad and Ap (22). Type Ap spermatogonia divide in the later stages of the spermatogenic cycle, to produce type B spermatogonia, as well as to renew their own population (21, 22, 24, 25, 28, 29). Ad spermatogonia, however, rarely divide and are thus considered to be resting or reserve stem cells (21, 22, 24, 25, 28, 29). Ap spermatogonia have been suggested to be the true stem cells of the testis, because Ap (and not Ad) spermatogonia are seen in humans after radiation therapy (30), after long-term estrogen therapy, and in the postpubertal cryptorchid testes (31). Thus, Ap spermatogonia are probably the stem cells of the testis that renew their own population as well as produce cells committed to differentiation. Various studies have suggested that Ap spermatogonia can undergo a transition, probably without division, into Ad spermatogonia (24, 28, 29). Because Ad spermatogonia are not proliferative, the transition of Ap into Ad may be a means of inactivating a certain proportion of the spermatogonial population (28, 29). Moreover, studies in irradiated monkeys suggest that Ad spermatogonia may be able to undergo a transition back to Ap spermatogonia, thus allowing repopulation of the testis (28, 29), further suggesting that Ad spermatogonia are reserve stem cells.

T implants caused a decrease in Ap spermatogonia within 2 weeks, and this decrease in Ap spermatogonia eventually led to subsequent decreases in germ cells from type B spermatogonia onwards. Other investigators have shown reductions in Ap spermatogonia after long-term gonadotropin suppression (9) or hypophysectomy (8) in macaques. This reduction may involve the inhibition of the mitosis of Ap spermatogonia, because GnRH antagonist treatment of macaque monkeys decreased Ap spermatogonia in the stages in which mitosis occurs (11), and this led to a disappearance of proliferating cell nuclear antigen-labeled Ap spermatogonia from the testis (25). The relative role of androgens and FSH in this loss of Ap spermatogonia is unclear. FSH treatment stimulated Ap spermatogonial number in either GnRH antagonist-treated (9) or normal (12) macaques; however, their number can also be maintained by androgens (8), suggesting a role for each hormone. In the current study, Ap spermatogonia were maximally suppressed within 2 weeks, at a time when there were more substantial reductions in testicular T than in serum FSH. Our study paradigm does not allow us to identify the relative contributions of the suppression of T and FSH to the impairment of Ap spermatogonial number. Clearly, the fall in intratesticular T levels could have contributed to the reduction in Ap spermatogonia; however, we cannot exclude the possibility that the reduction in serum FSH bioactivity is the major regulator of this impairment.

Species differences between primates in the regulation of spermatogonial number may exist. In rhesus monkeys, B spermatogonia, rather than Ap spermatogonia, seem to be hormone-sensitive, because only B spermatogonial numbers increased after administration of FSH to hypophysectomized animals (13). Also, unilaterally orchidectomized rhesus monkeys, which had significant increases in circulating FSH levels, showed increases in B spermatogonia but not Ap populations (14). However, although a change in Ap numbers was not seen, it is possible that the mitotic activity of Ap spermatogonia may still be stimulated by FSH in these monkeys, leading to the increased numbers of B spermatogonia. In juvenile rhesus monkeys, FSH alone (32), or both FSH and
Thus, Ad spermatogonia are increased during gonadotropin withdrawal, as has been demonstrated leading to few B spermatogonia being produced. Therefore, all of the effects of FSH suppression may be at the level of Ap spermatogonia rather than B. A third explanation for the data is that Ap spermatogonia may be regulated by FSH and/or T (8, 9, 12), whereas B spermatogonia may be primarily regulated by FSH, as has been postulated (13).

The suggestion that Ap and B spermatogonia may be differentially regulated may explain the discrepancy between our current and previous studies (11). Though the current study shows that 2 weeks of T administration caused a decrease in Ap spermatogonia (yet no change in type B), previous studies in monkeys given a GnRH antagonist for 16 or 25 days demonstrated decreases in B spermatogonia in the absence of significant changes in Ap spermatogonia (11). Although that study showed Ap spermatogonial mitosis was impaired, as evidenced by a decrease in the proportion of Ap spermatogonia in stages VII-XII, wherein Ap spermatogonial mitosis occurs, the primary defect was in type B spermatogonia (11). FSH levels were not measured in the GnRH antagonist-treated animals (11), but it is possible that GnRH antagonist treatment caused a more rapid suppression of circulating FSH than T administration, leading to more marked and rapid effects on B spermatogonia.

The increase in Ad spermatogonia after 2 weeks of T administration, and a second increase after removal of the T implants, was unexpected. It is believed that Ad spermatogonia do not proliferate (22, 25), and thus they are unlikely to proliferate in response to T administration. We suggest that Ap spermatogonia cease to proliferate after gonadotrophin withdrawal, as has been demonstrated (25), and instead undergo a transition into resting Ad spermatogonia. Ap spermatogonia have been suggested to form Ad spermatogonia in normal monkeys via transitional type A (At) spermatogonia (24). In our study, At spermatogonia were classified as Ad spermatogonia. Studies in irradiated monkeys suggest that Ap are inactivated into Ad spermatogonia soon after irradiation (28, 29). Thus, it is possible that gonadotropins stimulate Ap proliferation, whereas gonadotropin suppression causes a transition into nonproliferative Ad forms. Studies in rhesus monkeys have shown decreases in Ad spermatogonia after either administration of FSH to juvenile monkeys (34) or a unilateral-orchidectomy-induced rise in FSH (14). Thus, Ad spermatogonia are increased during gonadotro-
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