DNA flow-cytometric analysis of testicular germ cell populations of the bonnet monkey (*Macaca radiata*) as a function of sexual maturity*

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**Summary.** Testicular germ cell populations of biopsies from 32 male bonnet monkeys in 5 different age groups were quantitated in a flow cytometer after labelling of germ cell DNA with the specific fluorochrome, 4,6-diamidino phenyl indole. The 5 quantifiable populations were spermatogonia (2C), preleptotene spermatocytes (S phase), primary spermatocytes (4C), round spermatids (1C) and elongate spermatids (HC). The seminiferous tubules of immature 3–4-year-old monkey had only Sertoli cells and spermatogonia (2C).

At 5–6 years, germ cells in S-phase (9-5%), 4C (11-1%), 1C (41-8%) and HC (17-1%) stages of maturation appeared for the first time but at 7–8 years of age and beyond all cell types except HC decreased while 1C remained relatively constant. Histometric analysis correlated well with the flow-cytometric data. The decrease in cells of 2C, S-phase and 4C stages was associated with an increase in mitotic index, signifying acceleration in the kinetics of germ cell transformation into subsequent cell types. The total turnover in cell transformation (1C:2C) was significantly (P < 0:01) increased at and beyond 7–8 years. Maximum transition from 2C to 4C occurred at 5–6 years (4C: 2C ratio 0:8 at 5–6 years and 0:6 at 7–8 years). The ratio HC:1C (kinetics of cell transformation during spermiogenesis) attained near total efficiency only by 10 years of age (1:08 at 10–14 years; 0:9 at 18–20 years). Also, the cell associations within the seminiferous tubules of monkeys ≥10 years of age were better defined than those of younger animals.

The changes in germ cell ratios correlated well with alterations in testicular volume, sperm numbers in the ejaculate and surges of testosterone and increments in FSH in the serum, characteristic of development of sexual maturity. It is apparent from this study that DNA flow cytometry of testicular germ cell populations reveals subtle changes in spermatogenic status of bonnet monkeys with a high degree of sensitivity.

**Keywords:** spermatogenesis; DNA flowcytometry; germ cells; testosterone; FSH; puberty; primate

**Introduction**

Spermatogenesis is conventionally evaluated by carrying out time consuming and painstaking histological examination of testicular biopsies (Clausen et al., 1978a). The use of DNA flow cytometry, in contrast, facilitates rapid and accurate quantitation of large numbers of testicular cells;
germ cells and hence is gaining acceptance as an effective method of assessing spermatogenesis. Although flow-cytometric analyses of human testicular germ cell populations have been reported, they are for tissues obtained at autopsy (Thorud et al., 1980) or to investigate infertility (Clausen & Abyholm, 1980; Evenson & Melamed, 1983), and systematic studies have been confined to the mouse (Toppari et al., 1988) and the rat (Clausen et al., 1978a, b, 1979, 1982; Toppari et al., 1986; Van Kroonenburgh et al., 1986). Testicular germ cell populations of monkeys have not so far been measured by DNA flow cytometry.

Much basic information is available concerning the reproductive cycle of the female non-human primate (Prahalada et al., 1975; Maneckjee et al., 1976; Knobil, 1980; diZerega & Hodgen, 1980; Ravindranath & Moudgal, 1987), but relatively less is known about the regulation of the spermatogenic cycle in the male non-human primate (Bercu et al., 1983). The present studies were carried out to obtain baseline values for percentages of testicular germ cell types of bonnet monkeys as a function of age and to establish their correlation with histometric analysis of testicular biopsies, physiological attributes and endocrine profiles characteristic of sexual maturity.

**Materials and Methods**

Most of the monkeys used in this study were born in the colony and whenever wild-caught monkeys were added to a specific group to make up the required numbers, their age was determined by comparing their dentition with that of colony-born monkeys of similar age. The 32 males used were divided into 5 groups based on their age: pre-pubertal (3–4 years), pubertal (5–6 years), young adults (7–8 years), normal adults (10–14 years) and older adults (18–20 years). All monkeys were maintained in well ventilated rooms with a regulated light:dark schedule (lights on for 12:00 h). Blood was taken from unanaesthetized monkeys by femoral venepuncture into Vacutainer tubes at 10:00 h and 22:00 h to determine the occurrence of nocturnal testosterone surges. The standardized animal management practices followed in the colony have been published elsewhere (Ravindranath & Moudgal, 1987).

**Hormones and chemicals.** Pepsin, Tween-20 and polyethylene glycol (PEG) were obtained from Sigma Chemical Co. (St Louis, MO, USA); 4,6-diamidinophenylindole (DAPI) from Serrafine Biochemica (Heidelberg, FRG); Dulbecco's phosphate-buffered saline (PBS) from Himedia (Bombay, India); the nylon filter, Nybolt P-25 from Swiss Silk (Zurich, Switzerland); potassium hydrochloride from Parke-Davis (Detroit, MI, USA); crystalline testosterone from Steraloids, Inc. (NH, USA); and tritiated testosterone from Amersham Laboratories (Amersham, Bucks, UK).

**Testicular biopsies and cell preparation.** Approximately 100 mg testicular tissue were removed from one testis of each monkey, anaesthetized with ketamine hydrochloride, and transferred to Petri dishes containing PBS. While a portion of the tissue was fixed in Karnovsky's fluid for eventual histometric analysis, the remainder was finely minced (Evenson & Melamed, 1983) in a small volume of PBS and the preparation was vortexed to maximize the release of germ cells from seminiferous tubule minces. This procedure revealed germ cell ratios similar to those obtained from other attempted procedures using collagenase or trypsin. Subsequent fractions of cells recovered from the minced tubule debris, after the collection of the first fraction, exhibited decreased total cell yield but unchanged ratios, validating the fact that this procedure did not affect specific cell type losses. The cell suspension was aspirated, washed with PBS, filtered using nylon filter (50 µm) and fixed in 70% chilled ethanol and stored at 4°C until analysis.

**DNA staining and flow cytometry.** The staining procedure followed was as described by Otto et al. (1981). Briefly, the fixed cells were washed with 0.9% (w/v) NaCl and incubated with 0.5% pepsin in isotonic PBS (pH 2.0) for 1 h at 37°C. After centrifugation, the cells, by now rendered permeable to the stain, were resuspended in 1 volume of 0.2 M-citric acid (pH 1.8) containing 0.5% Tween-20 and 1% PEG and allowed to stand for 20 min at room temperature. Nine volumes of 5 µM 4,6-diamidino phenyl indole (DAPI) in 0.4 M-disodium-hydrogen-phosphate (pH 9.0) were added (final pH 7.4) and incubated at room temperature in the dark for 30 min. The stained cells were filtered through a nylon filter (50 µm) and analysed in a Phye ICP-22 (Phye A.G., Gottingen, West Germany) flow cytometer, equipped with a high-pressure mercury lamp, at 700 V, using filters of 360 nm wavelength for excitation and 450 nm for emission. Although the germ cell ratios measured by flow cytometry did not alter irrespective of whether 10 or 50 000 cells were studied, for the purpose of this study, at least 50 000 cells were analysed. The different cell populations thus obtained based on their DNA content (as opposed to ploidy expression, N) are expressed as 'C' values. A sample of human peripheral blood leucocytes was used as the diploid DNA standard (2C) to ascertain peak positions of the flow-cytogram distributions of populations and the cell in each peak were measured by planimetry (Otto et al., 1984).

**Histometric analysis.** Testicular biopsies were obtained from monkeys and fixed in modified Karnovsky's fluid (David et al., 1973), washed in 0.1 M-cacodylate buffer and post-fixed in 1% osmium tetroxide in 0.1 M-cacodylate buffer, dehydrated through graded series of acetone and embedded in araldite. Semithin sections of 0.5-µm thickness were cut at frequent intervals and stained with toluidine blue and observed under bright-field optics. The tubular
sections of the testicular biopsy were evaluated systematically and scored according to the method of Johnsen (1970). The score was given according to the presence or absence of the main cell types observed. The counting was performed with a × 10 objective exposing several tubules in one field. Damaged tubules were not included. The testicular biopsy scoring was done by the same observer. The mean score was calculated by multiplying the number of tubules recorded at each score with the score and the sum of all multiplications was divided by total number of tubules recorded. The total number of tubules recorded per biopsy was 50.

Seminiferous tubules were examined by optical microscopy using a × 100 oil-immersion objective and about 200 randomly distributed germ cells were counted; the number of dividing germ cells was divided by number of non-dividing germ cells to give the mitotic index.

**Hormone assays and physiological attributes.** Serum testosterone was estimated using an RIA system standardized and routinely used in the laboratory (Murty et al., 1979). This assay had a sensitivity of 10 pg and the inter- and intra-assay variations were of the order of 12.3% and 5.6% respectively. Serum FSH was estimated using a solid-phase RIA system developed in the laboratory. This system used cynomolgus monkey FSH as standard (kindly provided by Dr G. Bialy, NICHD, Bethesda, MD, USA), with 3.0 ng FSH/ml sensitivity and inter- and intra-assay variations of the order of 12% and 9%, respectively.

Collection of semen by electroejaculation and determination of sperm counts were done according to methods described by Mastroianni & Manson (1963). Testicular volumes were determined using Prader beads.

**Statistical analysis of data.** All analyses of hormonal and physiological attributes for individual monkeys were performed at least 3 times for each group and the mean ± s.e.m. values and the means of flow-cytometric data on the percentages of germ cell populations and ratios (Thorud et al., 1980; Toppari et al., 1986) were statistically analysed using Student's *t* test.

**Results**

**Physiological and endocrine attributes of sexual maturity**

As shown in Table 1, the body weight and testicular volumes of monkeys aged 3–4 years and 5–6 years were significantly different from each other (*P* < 0.001) and lower (*P* < 0.01) than those of the older age groups. The onset of adulthood, in physical terms, is therefore apparent from 7–8 years onwards. However, the nocturnal testosterone concentration in the 5–6-year-old monkeys was comparable to that seen in adults (Table 1). The 3–4-year-old animals did not exhibit the nocturnal testosterone surge and the values in the evening were significantly (*P* < 0.001) lower than those of all the other groups. The sperm counts of monkeys, first seen in the 5–6-year-old group, were also significantly different from those of the 7–8-year-old (*P* < 0.05) and older (*P* < 0.01) monkeys. The immature monkeys (3–4 years) did not ejaculate. Serum FSH values increased with age, concentrations being significantly lower (*P* < 0.05) in 3–4-year-old than in 5–6- and 7–8-year-old monkeys. However, after 10 years, there was an apparent decline in serum FSH values to attain a steady state.

**Histometric analysis**

The 3–4-year-old monkeys did not show any active mitosis but only Sertoli cells and basal spermatogonia with no other cell types (Fig. 1a). Some mitotic activity, greater in some tubules, and occasional spermatozoae were seen in the seminiferous tubules of the 5–6-year-old pubertal group (Fig. 1b). Advanced germ cell types were seen in the 7–8-year-old young adults (Fig. 1c). Monkeys aged 10–14 and 18–20 years showed histological profiles that indicated normal active spermatogenesis (Fig. 1d). Mean testicular biopsy scores and mitotic indices are shown in Table 2.

**DNA flow cytometric analysis of germ cells**

Using DNA flow cytometry to distinguish germ cell DNA contents, 5 different populations of germ cells could be identified in the sexually mature adult testis. Their positioning in the flow cytomgram was based on the fluorochrome binding to germ cell DNA resulting in fluorescence emission intensities proportional to the DNA contents of cell populations. Since C-values refer to DNA content and ploidies to amount of chromosomes, after locating the 2C (diploid spermatogonia)
Table 1. Physiological and hormonal correlates of monkeys at different ages

<table>
<thead>
<tr>
<th>Groups</th>
<th>Age (years)</th>
<th>No. of monkeys</th>
<th>Body weight (kg)</th>
<th>Testicular volume (cc)</th>
<th>Testosterone (ng/ml)</th>
<th>FSH (ng/ml)*</th>
<th>Sperm counts (× 10⁶/jaculate)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>3–4</td>
<td>3</td>
<td>2.7 ± 0.2bcde</td>
<td>1.0bcde</td>
<td>4.8 ± 0.7</td>
<td>5.0 ± 1.2bc</td>
<td>55 ± 18ab</td>
</tr>
<tr>
<td>b</td>
<td>5–6</td>
<td>15</td>
<td>5.4 ± 0.2cde</td>
<td>7.4 ± 0.7cde</td>
<td>4.6 ± 0.4</td>
<td>10.9 ± 1.2a</td>
<td>225 ± 63ab</td>
</tr>
<tr>
<td>c</td>
<td>7–8</td>
<td>5</td>
<td>7.2 ± 0.5abc</td>
<td>17.9 ± 1.6b</td>
<td>6.3 ± 1.5</td>
<td>11.8 ± 2.5a</td>
<td>283 ± 28ab</td>
</tr>
<tr>
<td>d</td>
<td>10–14</td>
<td>5</td>
<td>7.8 ± 0.2b</td>
<td>16.8 ± 0.9b</td>
<td>4.9 ± 0.3</td>
<td>8.7 ± 4.1</td>
<td>199 ± 34ab</td>
</tr>
<tr>
<td>e</td>
<td>18–20</td>
<td>4</td>
<td>9.6 ± 0.8abc</td>
<td>17.3 ± 3.1b</td>
<td>5.9 ± 1.0</td>
<td>8.5 ± 1.9</td>
<td></td>
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</tbody>
</table>

Within columns, values with different superscript letters indicate groups with significantly (P < 0.005–0.01) different values.

*Cynomolgus FSH equivalent.

Table 2. Testicular biopsy score and mitotic activity in bonnet monkeys of different ages

<table>
<thead>
<tr>
<th>Age (years)</th>
<th>Testicular biopsy score</th>
<th>Mitotic index</th>
</tr>
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<tbody>
<tr>
<td>3–4</td>
<td>2.0 ± 0.1</td>
<td>0</td>
</tr>
<tr>
<td>5–6</td>
<td>5.0 ± 0.2</td>
<td>24.0 ± 2.4</td>
</tr>
<tr>
<td>7–8</td>
<td>6.5 ± 0.2</td>
<td>32.2 ± 2.8</td>
</tr>
<tr>
<td>10–14</td>
<td>6.6 ± 0.7</td>
<td>38.8 ± 5.9</td>
</tr>
<tr>
<td>18–20</td>
<td>7.1 ± 0.6</td>
<td>38.0 ± 4.1</td>
</tr>
</tbody>
</table>
peak using human peripheral blood leucocytes as the 2C standard, the other major populations were identified according to their relative fluorescence intensities and classified as 4C (diploid primary spermatocytes), 1C (haploid round spermatids) and apparently hypohaploid HC (haploid elongate spermatids; see 'Discussion'). The HC peak was present only in adult monkeys with mature spermatooza in their ejaculates.

Representative DNA flow cytograms of testicular germ cells of monkeys from different age groups are presented in Fig. 2 and the percentages of individual populations in Fig. 3. The different cell percentages provided, however, do not add up to 100 as all non-specific fluorescent pulses in the flowcytograms were also quantitated to determine percentages but are not indicated in the histogram (Fig. 3). The 3–4-year-old monkeys exhibited 86.7% of 2C cells (Fig. 3), indicating the presence of mostly spermatogonia in the testicular seminiferous tubules (Fig. 2). The onset of active spermatogenesis was apparent in the 5–6-year-old animals with cells in S-phase (9.5%), 4C (11.1%), 1C (41.8%) and HC (17.1%). Compared to these monkeys, significant changes ($P < 0.01$) were observed in the percentages of all the different testicular germ cell types (Fig. 3) in each of the 3 age groups of older monkeys. However, the 1C values (45.7 ± 3.0; 37.3 ± 5.8 and 39.3 ± 2.8%, for the 3 groups respectively) did not vary from that of 5–6-year-old animals. Beyond 5–6 years of age the percentage of cells in 2C, S-phase, 4C and HC populations did not show any significant changes and the flow cytograms were very similar (Fig. 2).

The ratios of the different populations are presented in Table 3. The ratios could not be calculated for the 3–4-year-old monkeys because their testicular tissue contained only one type of cells, i.e. 2C. The 1C:2C ratios indicate the overall turnover of cells from spermatogonia up to the round spermatid stage. This ratio, however, can be subdivided into 3 components, 4C:2C; 4C:S-phase and 1C:4C. The 1C:2C values for the 7–8 and 18–20-year-old animals were significantly ($P < 0.01$) different from those of 5–6-year-old monkeys. However, the 10–14-year group did not exhibit significant variation, possibly because of the high s.e.m. values. The 4C:2C and 1C:4C ratios of monkeys belonging to all the older groups were also significantly ($P < 0.05–0.01$) different from those of the 5–6-year-old monkeys. Although the 4C:S-phase ratios of 10–14- and 18–20-year-old animals were significantly ($P < 0.05$) different from those of the 5–6-year-old group, these ratios for the 7–8 and 18–20 year old groups were similar. The 1C:4C ratios of all the three oldest groups were significantly ($P < 0.01$) different from those of 5–6-year-old monkeys.

**Discussion**

In the current study DNA flow cytometry has been used in quantitating the relative testicular germ cell populations in the monkey as a function of sexual maturity. The differentiated germ cell populations with cells in all stages of spermatogenesis is seen for the first time in 5–6-year-old monkeys and this correlates well with physiological characteristics such as testicular volume, hormone profile and the appearance of spermatooza in the ejaculate that signify onset of sexual maturity. Histological observations showed the presence of primary spermatocytes, round and some elongate spermatids at this age group and there was increased mitotic activity, emergence of S-phase, 4C, 1C and some HC cells in the flow-cytometric profiles. A marked increase in testicular volume after 30 months of age has also been reported for the rhesus monkey (M. mulatta: Bercu et al., 1983) and the crab eating monkey (M. fascicularis: Steiner & Bremner, 1981). Although testosterone rhythms have been described for most primates including the bonnet monkey (Mukku et al., 1976) the current study suggests that these surges may be involved in promoting spermatogenesis in terms of initiation of transformation of spermatogonia into more advanced cell types as seen at 5–6 years. That this group is still adolescent is evident from the fact that the testicular volumes and testicular biopsy scores are significantly lower than and the germ cell ratios markedly different from those exhibited by older monkeys.

In man (Clausen & Abyholm, 1980) the flow-cytometrically quantifiable germ cells released from testicular minces can be classified into: (1) 2C—primarily spermatogonial cells in the G1
Fig. 2. Characteristic flow cytograms of testicular germ cell populations of monkeys of different age groups.

Fig. 1. Semithin (0.5 μm) sections of testis from bonnet monkeys of different age groups × 400. (a) 3–4 years: only Sertoli cells and spermatogonia. (b) 5–6 years: Sertoli cells, spermatogonia, spermatocytes and occasional elongate spermatids are observed. Tubules appear disorganized and characteristic cell associations are ill defined. (c) 7–8 years: normal spermatogenesis with all cell types still with ill defined associations. (d) 10–20 years: well defined germ cell associations and normal spermatogenesis.
Fig. 3. Histogram depicting the percentages of monkey testicular germ cells quantitated by DNA flow cytometry. Only beyond 10 years of age, these percentages were maintained constant, indicating achievement of steady state spermatogenic kinetics in terms of total cell turnovers. Values are mean ± s.e.m.

Table 3. Ratios of germ cell populations from monkey testicular biopsies studied using DNA flow cytometry

<table>
<thead>
<tr>
<th>Age (years)</th>
<th>1C:2C</th>
<th>4C:2C</th>
<th>4C:S-phase</th>
<th>1C:4C</th>
<th>HC:1C</th>
</tr>
</thead>
<tbody>
<tr>
<td>3–4†</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>5–6</td>
<td>2.9 ± 0.2</td>
<td>0.8 ± 0.1</td>
<td>1.2 ± 0.1</td>
<td>3.8 ± 0.4</td>
<td>0.4 ± 0.06</td>
</tr>
<tr>
<td>7–8</td>
<td>4.0 ± 0.3**</td>
<td>0.5 ± 0.1*</td>
<td>1.4 ± 0.7</td>
<td>8.3 ± 12**</td>
<td>0.6 ± 0.1</td>
</tr>
<tr>
<td>10–14</td>
<td>4.2 ± 1.1</td>
<td>0.5 ± 0.1*</td>
<td>0.7 ± 0.02*</td>
<td>8.4 ± 10**</td>
<td>1.08 ± 0.3*</td>
</tr>
<tr>
<td>18–20</td>
<td>4.6 ± 0.5**</td>
<td>0.6 ± 0.1*</td>
<td>0.9 ± 0.04*</td>
<td>7.8 ± 10**</td>
<td>0.9 ± 0.2*</td>
</tr>
</tbody>
</table>

*Significantly (P < 0.05) different from values at 5–6 years.
**Significantly (P < 0.01) different from values at 5–6 years.
†The germ cell preparations from the biopsies of monkeys from this group had only a 2C peak rendering calculations of ratios impracticable.

phase; (2) S-phase—preleptotene spermatocytes and some proliferative spermatogonia undergoing DNA synthesis; (3) 4C—leptotene, zygotene, pachytene and diploten epithelial spermatocytes and some spermatogonial cells in the G2 phase; (4) 1C—round spermatids; and (5) hypofluorescent (due to hypercondensation of nuclear chromatin) apparently hypohaploid cells referred to here as HC-elongate spermatids. In the present study, the HC peaks observed were positioned exactly coinciding with flow-cytometrically analysed DAPI-stained ejaculated spermatozoa (data not shown), supporting the notion that the HC peak consists of elongate and mature spermatozoa in which the nuclear chromatin has undergone greater compaction. Cell sorting from this peak (Clausen et al., 1982) has revealed it to consist of mature as well as maturing spermatozoa. Furthermore, chemical decondensation of the nuclear chromatin of the compacted HC population caused it to shift to the 1C position (Zante et al., 1977).
The testicular germ cell distribution in adult bonnet monkeys closely resembles that reported for man with the 1C population approximating 40% of the total cells, unlike the 75% reported for the rat (Clausen et al., 1978b). In contrast to immature (3–4-year-old) monkeys, in which 87% of the pulses were found in the diploid region (spermatogonial cells), in the immature rat (Clausen et al., 1978a, b; Vaishnav et al., 1988), a flow-cytometrically quantifiable 4C population as well as cells in S-phase in percentages almost similar to the adult have been observed.

By 5–6 years there is a significant reduction ($P < 0.001$) in 2C values in bonnet monkeys, suggesting that spermatogonia begin active DNA synthesis at this time and are transformed sequentially to pre-leptotene (S-phase, 9–5%) and primary spermatocytes (4C, 11–1%), round spermatids (1C, 41–8%) and elongate and mature spermatozoa (HC, 17–1%). Histologically, the seminiferous tubules of monkeys of this age revealed disorganization and scattering of the germ cell types with the expected characteristic cell associations being ill defined (Fig. 1b). However, the kinetics of this process appears accelerated (increase in mitotic activity) by the time they attain 7–8 years of age. In this group, therefore, the percentage of cells in 2C, S-phase and 4C are further reduced compared with those in 5–6-year-old monkeys ($P < 0.01$). Although the percentage of round spermatids appears not to be altered, the HC population is significantly increased, suggesting an accelerated formation of mature spermatozoa as well as an active replenishment of 1C cell type. Overall cell transformation from spermatogonia to round spermatids (1C:2C, 4:0) as well as conversion of 4C to 1C (1C:4C, 8:3) is significantly ($P < 0.01$) increased at 7–8 years.

Histological examination (Figs 1b & c) and morphometric quantitations (Table 2) also revealed higher biopsy scores and mitotic indices for the 7–8-year-old animals. The characteristic germ cell associations are also better defined in this group. As a final end point, significantly ($P < 0.01$) increased sperm output (from $55 \times 10^6$ to $225 \times 10^6$) is observed in the ejaculate (Table 1). Although the HC:1C ratio of 5–6-year-old monkeys (0.4) was not significantly different from that of 7–8-year-old animals (0.6), both were significantly lower ($P < 0.05$) than those of the older age groups, suggesting that the rate of transformation of round spermatids into elongate spermatids during spermiogenesis attains maximal efficiency only by 10 years of age.

The endocrine control of sexual development of the male monkey is not clearly understood. In immature rats FSH is obligatorily required for both spermatogonial proliferation and spermiogenesis (Huckins et al., 1973). FSH deprivation in immature and maturing rats results in a spermatogenic block at the level of primary spermatocytes leading to reduced formation of spermatids (Vaishnav et al., 1988). Bercu et al. (1983) observed no significant increases in FSH concentrations with age (studied between 3 and 60 months of age) in rhesus and crab-eating monkeys but did report the presence of micro- and macropulses of FSH in post-natal and pre-pubertal monkeys, respectively. Although the need for FSH in sperm production in the primate has been proposed (Murty et al., 1980; Matsumoto et al., 1986; Moudgal et al., 1988; Van Alphen et al., 1988), in the present study, except for a 2-fold increase in serum FSH values between 3–4 and 7–8 years of age, the FSH concentrations did not show marked changes. Whether initiation of spermiogenesis is dependent upon serum FSH or nocturnal testosterone levels or both remains to be established. The reduced HC:1C ratios observed in the present study in monkeys of younger age groups could be a reflection of the relatively lower serum FSH:testosterone ratios (0.6 at 3–4 years, 1.45 at 5–8 years, 1.85 at 10–20 years) observed. Monkeys desensitized with GnRH or immunized with FSH show a marked change in this germ cell ratio (unpublished observations).

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