Gonadotrophin-releasing hormone antagonist arrests premeiotic germ cell proliferation but does not inhibit meiosis in the male monkey: a quantitative analysis using 5-bromodeoxyuridine and dual parameter flow cytometry

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Abstract
Meiosis constitutes a crucial phase of spermatogenesis since the recombination of genetic information and production of haploid round spermatids need to be achieved. Although it is well established that gonadotrophic hormones are required for completion of the spermatogenic process, little is known about the dynamic and kinetic aspects of development of spermatocytes into spermatids and its endocrine control in the primate. In this study, S-phase germ cells were labelled using 5-bromodeoxyuridine (BrdU) incorporation and were then followed throughout meiosis under normal conditions and following GnRH antagonist (ANT)-induced gonadotrophin withdrawal in a nonhuman primate model, the cynomolgus monkey (Macaca fascicularis). Adult animals received either vehicle (VEH, n=4) or the ANT cetrorelix (n=5) throughout 25 days. On day 7 all animals received a bolus injection of BrdU. A biopsy was performed after 3 h, one testis was removed 9 days later (day 16 of treatment) and the other testis after 18 days (day 25 of treatment). Serum testosterone and inhibin levels, and testis weight were reduced (P<0·05) by ANT treatment. BrdU localized to pachytene spermatocytes 9 days after BrdU and to round spermatids 18 days after BrdU in both groups, demonstrating that BrdU-labelled pachytene spermatocytes had undergone meiosis. Flow cytometric analysis revealed that the relative number and number per testis of BrdU-tagged 2C and 4C cells were reduced significantly (P<0·05) within 16 days of ANT treatment. Numbers of 1C cells were lowered by day 25. The cell ratio for 1C:4C was similar with VEH and ANT (P>0·05). These findings indicate that ANT reduced the number of cells available for meiosis but did not alter the rate of transition into round spermatids. Unexpectedly, however, the stage-dependent progression of BrdU-tagged round spermatids was significantly (P<0·05) retarded under ANT as seen from the frequency of tubules containing BrdU-labelled round spermatids. The average duration of spermatogenic cycle was slightly prolonged (9·8 days in the VEH group and 10·8 days in the ANT group (P=0·09)). Since no atypical germ cell associations could be found, it remains unclear whether this slight prolongation is entirely due to altered spermatid progression or whether earlier phases are affected. We conclude for the nonhuman primate that (1) BrdU-labelling of premeiotic germ cells is suitable for tracing their meiotic transition into postmeiotic cells, (2) unlike in the rat, gonadotrophin suppression initially affects premeiotic cell proliferation and thus the number of cells available for meiosis, (3) the meiotic process continues quantitatively despite gonadotrophin deficiency and (4) prolonged gonadotrophin deficiency might alter the timing of germ cell development.

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Introduction
Mammalian spermatogenesis is a highly complex process comprising several entirely different events, i.e. multiple mitotic divisions of diploid spermatogonia, meiotic division of tetraploid spermatocytes giving rise to haploid spermatids and, finally, the morphological differentiation of round spermatids into elongated spermatids. These processes take place in a topographically and chronologically coordinated manner. Meiosis is a critical phase of gametogenesis for several reasons since recombination of the genetic information, reduction of chromosome numbers and formation of spermatids need to be accomplished. The overall spermatogenic process is under endocrine control and lack of gonadotrophic support to the testis provokes a progressive loss of germ cells (Weinbauer & Nieschlag 1996).
However, the testicular response to gonadotrophin deficiency seems to be different between rodents and primates. Detailed histological studies in the rat identified a specific loss of spermatocytes and spermatids as the earliest lesion following gonadotrophic hormone deprivation (Russell & Clermont 1977, Sinha-Hikim & Swerdlow 1993, O’Donnell et al. 1996). In the nonhuman primate, loss of spermatogonia (Weinbauer et al. 1991) and meiotic arrest (Aravindan et al. 1993) have been suggested. Prolonged withdrawal of gonadotrophic hormones eliminates spermatids, spermatocytes and even differentiated spermatogonia in the monkey (Marshall et al. 1986, Weinbauer et al. 1991). In contrast, some spermatocytes and round spermatids are retained despite continued elimination of luteinizing hormone (LH) and follicle-stimulating hormone (FSH) in the rat (Sun et al. 1990, Chandolia et al. 1991a,b, McLachlan et al. 1995).

Another difference between rodents and primates has been reported with respect to efficacy of spermatogenesis and daily sperm production. Daily sperm production (10⁶/g testis) was 10–24 in rats (Wing & Christensen 1982, Russell & Petersen 1984), 4–5 in nonhuman primates (Gopalkrishnan et al. 1987) and 3–7 in men (Johnson et al. 1980). In other species such as rabbit, bull, stallion, ram and boar, daily sperm production achieved values beyond 12 (10⁶/g testis) and it would appear that, in general terms, the primate testis produces comparatively low numbers of sperm per weight unit of testicular parenchyma (see for example Sharpe 1994). It has also been reported with respect to e

For the mechanism underlying the differences among rats and primates in the testicular response to gonadotrophin withdrawal and the comparatively lower germ cell production rate of the primate testis is not known, but such knowledge would have important implications for dealing with male infertility and designing male contraceptives. For primates, little is known about the dynamic and kinetic aspects of development of spermatocytes into spermatids and the role of gonadotrophins in this process. Germ cells in the S-phase of the cell cycle can be identified by the incorporation and localization of 5-bromodeoxyuridine (BrdU) (Thoolen 1990, van de Ven & Kant & de Rooij 1992) and can be traced during the spermatogenic cycle in the rat (Rosiepen et al. 1994).

Systematic investigation of these aspects of germ cell proliferation in man is not possible because of ethical constraints. We, therefore, have initiated studies on this topic in a nonhuman primate model, the cynomolgus monkey (Macaca fascicularis) since, in terms of the endocrine regulation of spermatogenesis, this primate species has proved to be a suitable animal model for the human (Weinbauer & Nieschlag 1988, 1996). S-phase germ cells were labelled using BrdU incorporation and were then followed throughout meiosis under normal conditions and following gonadotrophin-releasing hormone (GnRH) antagonist (ANT)-induced gonadotrophin withdrawal using quantitative histological and flow cytometrical techniques.

Materials and Methods

Animals

Ten adult cynomolgus monkeys (Macaca fascicularis), weighing between 4.4 and 6.3 kg, were caged individually and maintained under defined environmental conditions as previously described (Weinbauer et al. 1994). Animals were fed standard monkey pellet food supplemented with fresh fruit and had free access to tap water. The study was undertaken in accordance with the German Federal Law on the Care and Use of Laboratory Animals (license 72/92).

Experimental design

Animals were randomly allocated to the treatment groups (n=5/group) and either received daily s.c. injections of vehicle (5–25% glucose in saline, VEH) or the GnRH antagonist cetrorelix (ANT) at a dose of 450 µg/kg for 25 days. On study day 7 all animals received a single i.v. injection of BrdU under ketamine hydrochloride sedation (8 mg/kg, Ketamine, Parke-Davis, Munich, Germany). The dosage for BrdU was derived from clinical studies in which i.v. infusions of 150–700 mg BrdU/m² surface area had been given without myelotoxic effects (Hoshino et al. 1985). Following these studies a dose of 590 mg/m² BrdU (corresponding to 33 mg/kg) was administered as a single i.v. injection of a 2% solution BrdU in 0.9% NaCl for labelling only the generation of germ cells that are in the S-phase of the cell cycle at the time of BrdU application. Three hours after BrdU application a testicular biopsy was performed for the localization of BrdU. One testis was removed 9 ± 0.1 days after BrdU injection (day 16 of treatment) and the other testis after 18 ± 0.1 days (day 25 of treatment) for histological and flow cytometrical analyses. Blood was collected from the cubital vein for the analysis of serum T and inhibin concentrations prior to and on days 7, 16, 21 and 25 of the study under ketamine hydrochloride sedation.

Tissue processing and localization of BrdU

Testicular biopsy material was placed in Carnoy’s fluid. Testes were excised, weighed to the nearest 0.01 g, and cut into slabs that were fixed in Carnoy’s or Bouin’s fluid. Specimens were subsequently dehydrated and embedded.

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in Paraplast following routine procedures. Sections (3 μm) were cut from pre-cooled paraffin blocks for histological analysis and immunochemical localization of BrdU by immunogold–silver staining. For immunolocalization, sections were deparaffinized, dehydrated and incubated with periodic acid in a microwave oven (BIORAD 2500, München, Germany; 60% maximal power at 50 °C) for 30 min, rinsed in tap and distilled water and followed by Schiff’s reagent for 30 min. Periodic acid treatment served to hydrolyze the DNA allowing access of the BrdU antibody and to provide staining of the acrosome for staging of the spermatogenic process (van de Kant & de Rooij 1992). A monoclonal antibody against BrdU (Dako Diagnostica, Hamburg, Germany) was used at a dilution of 1:20 for 60 min and nonspecific binding was blocked using 5% normal goat serum. Gold-labelled goat anti-mouse IgG (Amersham Ltd, Amersham, Bucks, UK) at a dilution of 1:40 served as second antibody. Silver enhancement solution (Amersham Ltd) was added for 11 min. Sections were then counterstained with haematoxylin, dehydrated and mounted in Merckoglas.

**Dual parameter flow cytometry**

Snap-frozen testicular tissue, obtained on days 16 and 25 of the study, was stored at −80 °C until analysis. Tissue was blade-minced in C100 T solution (citrate and tween-20), homogenized (Ultra-Turrax T25, Janke & Kunkel, Heppenheim, Germany), further incubated for 30 min in C100 T and transferred into absolute ethanol for 24 h. Samples were then centrifuged, the pellet was resuspended in 2 M HCl/pepsin and incubated for 45 min at room temperature under constant agitation. Washing steps included phosphate-buffered saline. Monoclonal antibody against BrdU (Dako Diagnostica), diluted 1:10, was added and incubation lasted for 30 min in the dark. FITC-conjugated goat anti-mouse IgG (Sigma Chemicals, St Louis, MO, USA) was used as the secondary antibody diluted 1:10 with 5% normal goat serum. Gold-labelled goat anti-mouse IgG (Sigma Chemicals) was added to the pellet, gently vortexed and incubated for 60 min in the dark. A total of 16·3 ± 1·7 mg testicular tissue (ranging between 9 and 33 mg) were used for determination of the number of 1C, 2C and 4C cells (identified through labelling with propidium iodide) and of the number of 1C, 2C and 4C cells labelled with BrdU (identified by FITC emission).

Analysis was performed using the Coulter flow cytometer EPICS XL (Coulter, Krefeld, Germany) equipped with a 15 mW argon-laser at an excitation wavelength of 488 nm. In order to correct for non-specific FITC-signals, testicular tissue collected from an adult, intact cynomolgus monkey, not treated with BrdU, was processed as described above. Forward scatter signals were used as trigger signals and fluorescence emissions were measured by photomultiplier detectors with a band pass filter 525BP (504–540 nm) for light reflected by the dichroic 550 DL filter for FITC, and a 620BP filter after a 660 DL long pass filter for propidium iodide. Cell debris and aggregates were eliminated from analysis by gating. Each category of cells identified by the propidium iodide signal as 1C, 2C or 4C was analyzed for positive FITC signals. 2C cells are spermatagonia, secondary spermatocytes and somatic cells, 4C cells represent cells in mitotic or first meiotic division after DNA synthesis (primary spermatocytes, somatic cells and spermatogonia in G2) and 1C refers to haploid germ cells (Toppari et al. 1985).

**Histological analysis**

All sections were coded and evaluated in a randomized manner. The exposure to ANT was limited to 25 days because it is known from previous work that identification of spermatogenic stages is still possible whereas at later time points identification would become difficult or impossible owing to the progressive loss of round and elongated spermatids (Weinbauer et al. 1991). Germ cell-specific and stage-specific distribution of BrdU-labelled germ cells was performed. Spermatogenesis was divided into 12 stages according to Clermont & Leblond (1959). Frequencies were determined for spermatogenic stages I–IV (pooled), V, VI, VII, VIII and XI–XII (pooled). The number of seminiferous tubules containing BrdU-labelled germ cells in the various stages on days 9 and 18 after BrdU administration was determined and is referred to as the BrdU-staining frequency. Between 507 and 684 seminiferous tubule sections were evaluated per time point and animal for assessment of stage and staining frequency. The duration of one seminiferous cycle was calculated for each animal on the basis of individual stage frequencies, BrdU-staining frequencies at both time points after BrdU injection and the time interval between the two time points (Rosiepen et al. 1994, 1995, 1997).

The proportion of pachytene spermatocytes and round spermatids containing BrdU was also determined in stages V and VI by counting the total number of either pachytene spermatocytes (day 9 after BrdU) or round spermatids (day 18 after BrdU) and how many cells of these type contained BrdU. Randomly selected blocks were sectioned and analyzed. A total of 615–1466 pachytene spermatocytes (day 16) and 232–2491 round spermatids (day 25) in the respective stages were analyzed in each specimen. Since data are interpreted only with respect to relative numbers among the same cell type, corrections for tissue shrinkage and ANT-induced testicular involution were not necessary.

**Serum testosterone and inhibin**

Serum testosterone was measured by RIA using a commercial kit (DSL-4100, Diagnostic Systems Laboratories, Sinsheim, Germany). Intra- and interassay coefficients of
Figure 1 Immunolocalization of BrdU in testis of VEH-treated (a,b,f) and ANT-treated (c,d,e,) monkeys. (a,d) 3 h after BrdU injection: spermatogonia (open arrows) and early spermatocytes (closed arrows) are labelled; (b,e) 9 days after BrdU injection: pachytene spermatocytes are labelled; (c,f) 18 days after BrdU injection: round spermatids are labelled (blueish colour), note the lower number of spermatids following ANT treatment (c). Tubules in (b,c) are in stage VI; tubules in (e,f) are in stage VI and VII (where spermiation has occurred). Sections were counterstained with periodic acid–Schiff’s base and haematoxylin. Bars indicate 9 µm for (a, d) and 20 µm for (b,c,e,f).
variation were 6·5 and 13·4% respectively, and the detection limit was 0·17 nmol/l. Serum inhibin was measured by a RIA validated for cynomolgus monkey serum (Fingscheidt et al. 1989). This assay detects α subunit-containing forms including inhibin B and the pro α-C protein and was found to measure FSH-dependent inhibin in macaque monkeys (Weinbauer et al. 1991, Arslan et al. 1993). Hence, serum inhibin levels are used as an indication of circulating FSH activity. All samples were assayed in duplicate in a single assay. The detection limit was 1·2 U/ml serum and the within assay precision was 2·8%.

Statistical evaluation

Data are presented as means ± s.e.m. Germ cell ratios were tested by Student’s t-test. Other parameters were tested by two-way ANOVA for repeated measures followed by Student–Newman–Keuls test. Relative frequencies and percentage values were subjected to arcsin transformation prior to statistical analysis. Probability levels of $P<0·05$ were considered significant. Flow cytometric and histological data revealed abnormal spermatogenesis in one control animal and, hence, data from this animal were excluded from analysis.

Results

Histological analysis of BrdU-labelled cells

Three hours after BrdU bolus, spermatogonia and early spermatocytes displayed a BrdU signal (Fig. 1a and d). Sertoli cells containing BrdU were not seen whereas occasionally peritubular cells and interstitial cells displayed BrdU signal (not shown). The amount of biopsy material did not permit quantitative analysis of cellular staining proportions nor stage-related analysis but no qualitative differences of BrdU staining pattern could be seen between both groups. Nine days after BrdU administration, spermatocytes contained BrdU (Fig. 1b and e), and by day 18 after BrdU, spermatocytes and round spermatids were BrdU-positive (Fig. 1c and f) in both experimental groups. A schematic representation of the stage-dependent distribution of BrdU-labelled germ cell types and their progression in a VEH-treated animal and an ANT-treated animal is depicted in Fig. 2.

Flow cytometrical analysis of BrdU-labelled cells

Between 0·02 and 0·07% of testicular cells emanated a FITC-signal above background in the animal that had not been exposed to BrdU. These signals were regarded as non-specific and were corrected for each category (1C, 2C and 4C) during the analysis of all BrdU-exposed animals. Coefficients of variation for repeated dual parameter analysis of three to four preparations from the same animal ranged between 4 and 9%.

Relative numbers (percentages) of 2C and 4C cells labelled with BrdU (2C–BrdU, 4C–BrdU cells) were significantly lowered to 40 and 70% of VEH respectively, within 16 days of ANT treatment (Fig. 3). Proportions of 2C–BrdU cells were also reduced on day 25. When expressed as numbers per testis, 2C–BrdU and 4C–BrdU cells were reduced to 20 and 30% of VEH by day 16 and remained lower until day 25 (Fig. 3). For 1C cells, the proportion of BrdU-containing cells was negligible on day 16 (9 days after BrdU) in both groups ($<0·5\%$, Fig. 3). This observation corresponds to the immunocytochemical findings (Figs 1 and 2). By day 25 (18 days after BrdU), relative numbers and numbers per testis of 1C–BrdU cells were markedly increased in both groups (Fig. 3), reflecting the meiotic transition of spermatocytes into spermatids but these numbers were lower under ANT compared with VEH ($P<0·05$). The 1C–BrdU:4C–BrdU cell ratio was similar for VEH and ANT groups ($P>0·05$). Combined numbers of BrdU-labelled and unlabelled 2C, 4C and 1C cells were significantly reduced on day 25 when compared with VEH ($P<0·05$, data not shown).

Kinetic analysis of spermatocyte–spermatid transition

Stage frequencies did not differ significantly between VEH- and ANT-treated groups (Fig. 4). Differences were observed with regard to BrdU-staining frequencies, i.e. the number of tubules containing BrdUlabelled cells. By day 9 after BrdU injection, BrdU-staining frequencies (spermatocytes) for stages I–IV, V, VI, VII and VIII were similar for VEH and ANT groups ($P>0·05$). At the second time point (day 18 after BrdU), stage I–IV BrdU-staining frequencies were not significantly different ($P>0·05$) indicating that the duration of meiosis was not affected by ANT. However, the percentage of stage VI tubules containing BrdU-labelled germ cells (round spermatids) was significantly reduced ($P<0·05$) in the ANT group and no BrdU-tagged round spermatids could be discerned in stages VII and VIII. In contrast, 32% of stage VII and 14% of stage VIII tubules, on average, contained BrdU-labelled round spermatids in the VEH group. Thus, stage-dependent progression of round spermatids over time was altered by ANT (see also Fig. 2). Stage IX–XII tubules were not analyzed since BrdU-labelled round spermatids did not advance beyond stage VIII. Calculation of the duration of this process provided an estimate of the length of one spermatogenic cycle. Cycle length appeared somewhat longer in the ANT group but this effect did not attain statistical significance (VEH group: 9·80 ± 0·49 days, range: 9·10–11·03; ANT group: 10·88 ± 0·25 days, range: 10·36–11·45 days ($P=0·09$)). The ratio of BrdU-labelled:total number of germ cells in stages V and VI in VEH vs ANT group was 0·49 ± 0·07 vs 0·48 ± 0·08.
Stage of spermatogenesis

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for pachytene spermatocytes (day 9 after BrdU) and 0.33 ± 0.12 vs 0.22 ± 0.08 (P > 0.05) for round spermatids (day 18 after BrdU). A specific alteration of spermatid progression would imply the existence of atypical cellular associations which, however, could not be observed.

Endocrine parameters and testis weight

Mean testosterone levels were between 18.9 and 32.1 nmol/l in the VEH group throughout the study without significant change. Administration of ANT reduced testosterone concentrations from 20.7 nmol/l at...
Baseline to 6·1 ± 1·7 nmol/l within 7 days, 4·4 ± 1·1 nmol/l by day 16 and 2·8 ± 0·5 nmol/l by day 25. Baseline inhibin levels were 7·7 ± 2·4 µg/l and 9·4 ± 2·6 µg/l in VEH and ANT groups respectively. Inhibin levels declined over time in both groups but this effect was significantly (P<0·05) more pronounced with ANT: 3·4 ± 1·2 in VEH vs 1·5 ± 0·2 µg/l in ANT group at termination of the study. On day 16, testicular weights were 19 ± 1 g in the VEH group and were reduced, albeit not significantly, to 14 ± 2 g in the ANT-treated group (P>0·05). By day 25, testis weight was 19 ± 2 g in the VEH group and was further lowered with ANT to 10 ± 1 g, (P<0·05).

Discussion

Kinetic parameters of cell proliferation, meiosis and haploid germ cell development, and the gonadotrophin control of these processes, were studied in a nonhuman primate model (cynomolgus monkey) using qualitative and quantitative histological analysis of BrdU-labelled meiotic and postmeiotic germ cells as well as dual parameter flow cytometry. Cetrorelix, a highly potent antagonist of GnRH in this macaque species (Weinbauer & Nieschlag 1993) was used to inhibit gonadotrophic hormone secretion and the effectiveness of this treatment is evidenced by the reductions of serum testosterone.
and inhibin concentrations, and the decline of testis size.

By 3 h after BrdU injection, BrdU was localized to spermatogonia and early spermatocytes, by 9 days to spermatocytes and after 18 days to round spermatids. These observations demonstrate that early germ cells, having incorporated BrdU into their DNA, can progress beyond meiotic reduction division. Hence, a nonradioactive approach for tracing germ cells throughout the entire spermatogenic process is available for the nonhuman primate testis. An estimate of the duration of the spermatogenic cycle was derived from the 9-day and 18-day time points, i.e., the transition of spermatocytes into spermatids. Among control animals, the average duration of one cycle of spermatogenesis was 9.8 days. This value is similar to that obtained in the cynomolgus monkey by comparing BrdU-labelled germ cells 3 h and 9 days after BrdU (9.8 days, n=1 (G Rosiepen, GF Weinbauer & E Nieschlag, unpublished observation)). Based on thymidine labelling, the estimated length of the spermatogenic cycle was 9.4±0.3 days (n=4, Dang 1971) and 10.5 and 10.6 days (n=2, Fouquet & Dadoune 1986). Importantly, these comparative data indicate that the presence of BrdU had no adverse effect on the timing of the transitions from spermatocytes into spermatids in the present study.

The stage-dependent progression of BrdU-labelled germ cells was less advanced in the ANT-treated animals compared with VEH-treated animals. This is suggested by the observation that the frequency of stage V–VIII tubules containing BrdU-labelled round spermatids was reduced in gonadotrophin-deficient animals. Such a difference was unexpected since the timing of the spermatogenic cycle, including spermiogenesis, is believed to be independent of gonadotrophins (Clermont & Harvey 1965). A decreased labelling rate of germ cells at the time of injection in ANT-exposed animals could potentially explain this difference. Qualitative analysis of the biopsates taken 3 h after BrdU did not support that suspicion. The similarity of proportions of BrdU-labelled spermatocytes and of BrdU-staining frequencies on day 9 after BrdU would also argue against an adverse early effect of ANT on BrdU incorporation. It is of interest to note, however, that the BrdU-staining frequency for stage VIII tubules on day 9 was, on average, about 20% for VEH and about 10% for VEH (Fig. 5). At this time point pachytene spermatocytes are labelled in stage VIII. This effect was not statistically different, but raises the possibility that ANT might have slightly affected the progression of these cells. Estimation of the duration of one cycle based on the interval 3 h and 9 days after BrdU could clarify this aspect. However, the limited amounts of biopsy material prohibit quantitative assessment. Therefore, the possibility of ANT-induced alterations of BrdU uptake and of meiotic germ cell development cannot be ruled out with certainty.

The observation of altered spermiogenesis could also be explained by assuming that, during ANT exposure, only those germ cells having incorporated BrdU are affected specifically during meiotic transition into spermatids and/or during spermiogenesis. This possibility cannot be discarded with ultimate certainty but is unlikely to account entirely for the observations based on the following considerations. The percentage of BrdU-labelled round spermatids in stage V and VI differed by about 30% suggesting that, if BrdU-labelled spermatids are primarily affected under ANT, this effect is of substantial magnitude. The ratio of 1C:4C cells was similar in the present study (25 days of ANT) and in a previous study (28 days of ANT) in the absence of BrdU (Weinbauer et al. 1991). Biased assessment of the stages of spermatogenesis owing to ANT-induced morphological changes is ruled out by the consistency of stage frequencies over time and the lack of difference between ANT and VEH groups. These data suggest that BrdU itself had no major detrimental effect on the formation and progression of haploid germ cells.

If spermatids are specifically retarded during spermiogenesis under ANT administration, the occurrence of atypical cellular association is to be expected. However, we could not detect such atypical cellular associations. It is conceivable, that the magnitude of such an effect was too slow to be detected by our approach. However, in the light of the findings and considerations discussed above, the possibility must also be considered, that slight alterations of spermatocyte development or BrdU incorporation under ANT exposure might well have contributed to the reduced progression of spermatids. The duration of one spermatogenic cycle was found to be prolonged by about one day but this effect was not statistically significant (P=0.09). In an earlier investigation in the rat (Clermont & Harvey 1965) it was observed that the duration of the spermatogenic cycle is not under gonadotrophic control, although it is of interest to note from that study, that the spermatogenic cycle duration increased slightly in the hypophysectomized rat and was reversed by administration of testosterone and even more so with human chorionic gonadotrophin (hCG). Circumstantial evidence for man (Heller & Clermont 1964) does not support gonadotrophin control of the duration of the spermatogenic process: the progression of thymidine-labelled germ cells was compatible with the normal duration of spermatogenesis in one man with progestin-induced gonadotrophin deficiency and in another normal man treated with hCG. It remains unclear whether a hypogonadotrophic condition can indeed affect the duration of an entire cycle of spermatogenesis and further studies are necessary to clarify this issue.

Sertoli cells were consistently negative for BrdU, and peritubular and interstitial cells were rarely BrdU-positive. Hence, flow cytometric analysis of BrdU-tagged testicular cells essentially refers to germ cells. Since spermatogonia represent a main fraction of testicular 2C cells (Toppari et al. 1985), it is most likely that the majority of 2C–BrdU

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cells on day 9 after BrdU (day 16 of the study) are spermatogonia that were in the S-phase at the time of BrdU application. Within only 16 days of ANT treatment, the relative number and the number of 2C–BrdU cells per testis was significantly diminished. It would appear that ANT rapidly and pronouncedly inhibited spermatogonial proliferation in the nonhuman primate, possibly within a few days of initiation of treatment. The significant drop of 4C–BrdU cells (relative numbers and numbers per testis), representing either mitotic spermatogonia or meiotic spermatocytes, on day 16 further supports this view. Subsequent to this inhibitory effect on spermatogonial proliferation, spermatocytes and spermatids are progressively depleted over time as seen from the analysis of 4C–BrdU and 1C–BrdU cells on day 25. These kinetic data suggest that ANT-induced germ cell depletion results from an arrest of spermatogonial divisions in the nonhuman primate. Our observations extend previous histological reports on the significant reduction of numbers of renewing spermatogonia after 28 days of ANT treatment (Weinbauer et al. 1991) but do not confirm the assumption of a meiotic arrest as the cause of spermatogenic inhibition (Aravindan et al. 1993). The data on proportions of 4C and 1C cells suggest that the number of cells available for meiosis is reduced by ANT, whereas the development into round spermatids is not prohibited.

Figure 5 Relative frequency of seminiferous tubules containing BrdU-labelled germ cells (BrdU-staining frequency) on days 16 and 25 in VEH-treated (○) and ANT-treated (●) monkeys. This data represent the quantitative analysis of the cell-specific and stage-specific distribution of BrdU as indicated in Fig. 2. Stages are denoted by Roman numerals. Data represent means ± S.E.M. of four and five animals/group. *P<0.05 vs day 16 and **P<0.05 vs VEH.
Gonadotrophin deficiency in the nonhuman primate induces a marked increase in the number of 2C cells if expressed as percentage of the total number of cells analyzed (Weinbauer et al. 1991, 1994, Aravindan et al. 1993); this increase, however, is relative resulting from the loss of 1C and 4C cells. In fact, expression of 2C cell numbers per testis yields reduced numbers. Analysis of BrdU-tagged cells is more representative to describe the 2C population since the percentage of 2C–BrdU cells was also significantly decreased during ANT administration (Fig. 3, left panels). This issue becomes critical if ratios among the different cell populations, as indices of germ cell transformations, are based upon relative values; a 9-fold increase of the percentage of 2C cells along with unaltered percentage of 4C cells and a >2-fold decrease of the percentage of 1C cells in hypogonadotrophic monkeys was considered to indicate an arrest of meiosis (Aravindan et al. 1993). However, in the light of the findings from the present study using BrdU-tagged germ cells it is more likely that the loss of advanced germ cells per testis originates from an arrest of spermatogonial mitosis.

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