Postnatal testicular development in mouse species with different levels of sperm competition

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Abstract

Postcopulatory sexual selection leads to an increase in sperm numbers which is partly the result of an increase in relative testes mass and could also be the consequence of changes in testis architecture or function. Very little is known regarding developmental changes during the first spermatogenic wave that may lead to enhanced spermatogenic efficiency and increased sperm production. We examined testicular development after birth in four mouse species with different sperm competition levels to assess changes in testicular architecture and function. Differences in relative testes mass between species appeared soon after birth and were exacerbated thereafter. The volume of testes occupied by seminiferous tubules differed between species postnatally and were associated with sperm competition levels. Finally, changes over time in the proportions of tubules with different germ cell types were also associated with sperm competition levels, with the time taken for the transition between various cell stages being negatively associated with levels of sperm competition. We conclude that postnatal testis development differs between closely related species with different sperm competition levels influencing testis architecture and the rate of progression of spermatogenesis, leading to differences in testis function at reproductive maturity. Reproduction (2012) 143 333–346

Introduction

Postcopulatory sexual selection, in the form of sperm competition (Parker 1970), selects for a variety of male reproductive traits (Birkhead et al. 2009), such as ejaculates with increased sperm numbers and quality (Møller 1988, Birkhead et al. 1999, Hunter & Birkhead 2002, García-González & Simmons 2005, Firman & Simmons 2010, Gómez Montoto et al. 2011), as well as longer spermatozoa that perform better (Gomendio & Roldan 1991, 2008, Gomendio et al. 2007, 2011, Fitzpatrick et al. 2009, Lüpold et al. 2009a, Tourmente et al. 2011a, 2011b). Males that produce more spermatozoa are more likely to father offspring under conditions of sperm competition because the number of sperm is a key determinant of fertilization success (Birkhead & Møller 1998, and references therein). An increase in sperm output and more sperm cells per ejaculate are thus widespread responses to enhanced sperm competition levels (Gomendio et al. 1998).


Other mechanisms increasing sperm production and contributing to higher sperm output have been identified. Changes in testicular architecture may result in more sperm-producing machinery. Among primates, species with relatively larger testes have a greater proportion of seminiferous tubules than species with lower relative testes mass (Schultz 1938) and, in the great apes, the proportion of seminiferous tubules varies in agreement with relative testes mass being highest in the chimpanzee and lowest in the gorilla (Fujii-Hanamoto et al. 2011). Among rodents, the proportion of testicular volume occupied by seminiferous tubules was greater in Pseudomys australis (the Plains Rat), a species with high relative testes mass, than in Notomys alexis the Spinifex
Hopping Mouse, which has a low relative testes mass (Breed 1982, Peirce & Breed 1987, 1990). Furthermore, in comparative studies in birds, species with higher combined testes mass in relation to body mass exhibited an increase in the proportion of seminiferous tissue (Lüpold et al. 2009b, Rowe & Pruett-Jones 2011). Therefore, available evidence suggests that increases in the levels of sperm competition are associated with a higher proportion of seminiferous tubules which may, in turn, explain higher sperm output.

More spermatozoa may also be generated by modifications of testicular function resulting in a more efficient machinery. Among others, changes may include faster turnover of the sperm production cycle (Ramm & Stockley 2010), higher numbers of cell divisions during the proliferative, premeiotic phase (Ehmcke et al. 2006), or modifications in the relationship between germ cell proliferation, loss, and survival (Hess & França 2008).

Spermatozoa are produced in the seminiferous tubule compartment of the testes in a process known as spermatogenesis. The tubules are lined by the seminiferous epithelium, which consists of germ cells and Sertoli cells, with the latter supporting and nourishing the germinal epithelium. A lumen occupies the centre of the tubule to which the mature sperm cells are released. The space between the seminiferous tubules, referred to as the interstitial tissue, contains Leydig cells (which produce male sex hormones), blood and lymphatic vessels, and macrophages (Setchell & Breed 2006). The germ cells in the seminiferous epithelium go through three phases: proliferation, chromosome reduction (meiosis), and differentiation (spermiogenesis) (Clermont 1972, Russell et al. 1990, Kerr et al. 2006, Hess & França 2008, Hermo et al. 2010). The length of each of these phases is about one-third of the total process of spermatogenesis, and the total duration of spermatogenesis varies between species (Johnson 1995, 2000, Kerr et al. 2006); the length of spermatogenesis appears to be controlled by the genotype of the germ cells (Zeng et al. 2006, Arregui et al. 2008).

The cells that divide by mitosis during proliferation and constitute the group of cells from which meiosis and spermiogenesis proceed are termed spermatogonia; they thus constitute the stem cells for spermatogenesis (De Rooij & Russell 2000). Type A spermatogonia divide by mitosis to replace themselves and to provide a population of type B spermatogonia which, in turn, differentiate into type B spermatogonia from which primary spermatocytes arise (Kerr et al. 2006, Borg et al. 2010). Germ cells undergoing meiosis are the primary and secondary spermatocytes and completion of meiosis gives rise to haploid round spermatids that undergo a series of morphological changes with no cell division involved. Spermatids experience elongation during a series of developmental steps that include changes in cellular organelles such as acrosome formation, nuclear reshaping and condensation, development of the flagellum, reorganization and elimination of cytoplasm, and final release to the lumen of the seminiferous tubule during spermiation (Kerr et al. 2006, Tanaka et al. 2007).

The morphology and function of the testes that is observed in adult males develops during a protracted period of time before and during puberty (Kerr et al. 2006). Primordial germ cells first appear during foetal development. They proliferate, migrate, and establish the gonadal primordium. Nonmigrating germ cells in the gonad are called gonocytes and they arrest proliferation until around puberty. A series of cues trigger resumption of proliferation and the first spermatogenic wave begins. There are important differences between primates and most rodents because in the former there is a clear period of separation between postnatal and pubertal period, whereas this is not the case in the mouse and other rodents. In primates, spermatogenesis starts many months or years after birth, but in rodents it begins within a few days after birth. In the laboratory mouse, the first wave of spermatogenesis encompasses the proliferation and differentiation of germ cells and Sertoli cells over a period of 35 days, full fertility is reached by 6–7 weeks of age, and key time points for the appearance of particular types of germ cells are well defined (Bellvé et al. 1977, Bellvé 1993, Borg et al. 2010).

Studies of postnatal testis development have allowed a characterization of the first spermatogenic wave. A complex interaction between circulating follicle-stimulating hormone (FSH) and local testicular factors including activin and inhibin, and their effects on germ cell differentiation, was revealed during this first wave (Barakat et al. 2008, Mithraprabhu et al. 2010). Partial synchronization of spermatogenesis in some species has also been identified (Van Haaster & De Rooij 1994). Furthermore, germ cell apoptosis and underlying molecular mechanisms were also characterized in this first round of spermatogenesis (Jahnukainen et al. 2004). Expression of testicular germ cell genes during specific stages of spermatogenesis has also been assessed during postnatal development (Lahn et al. 2002, Anway et al. 2003, Wrobel & Primig 2005, Hansen et al. 2006, Cao et al. 2008, Itman & Loveland 2008, Albert et al. 2010). Finally, stages of postnatal development have been examined with the aim of identifying reproductive isolation in hybrid mice due to spermatogenic defects at several stages (Oka et al. 2010). Therefore, the first spermatogenic wave represents an important tool for the timing and characterization of cellular and molecular events during spermatogenesis, because it is possible to identify the progression through different cell types as they appear for the first time. Thus, the first spermatogenic wave is regarded as a useful model to study spermatogenesis.

In this study we have examined, using four closely related murid species that differ in sperm competition levels (Gomendio et al. 2006), if this selective force exerts an influence on the postnatal development of...
testes architecture and turnover of germinal epithelium which may explain differences in sperm production. In addition, the study of postnatal development allowed us to understand when and how differences in testes architecture and function originate.

Results

**Body mass and testes mass**

Changes in body mass during postnatal growth differed between the four mouse species ($F_{3,84} = 56.20, P<0.001$; Fig. 1A). Although they experienced a similar increase in body mass until 14 days after birth, from 20 days onwards the species started to differ. Significant differences in body mass were observed in adult mice (120 days old; $F_{3,20} = 45.36, P<0.001$).

Testes mass increased during postnatal growth and differed between species ($F_{3,84} = 240.36, P<0.001$; Fig. 1B) and Bonferroni’s *post-hoc* tests showed that there were significant differences between the four mouse species throughout postnatal development. From an early age (10 days), testes masses of *Mus spretus* and *Mus spicilegus* were higher than those of *Mus musculus* and *Mus pahari* and from 24 days onwards ($F_{3,8} = 25.87, P<0.001$) these two sets of species (*M. spretus–M. spicilegus* and *M. musculus–M. pahari*) differed significantly in testes mass. Adult testes mass of the four species were significantly different at 120 days postpartum ($F_{3,20} = 170.77, P<0.001$).

As a consequence of differences in body and testes masses, relative testes mass differed between species revealing ample variation in the levels of sperm competition among them (Fig. 1C). *M. spicilegus* showed the highest values of relative testes mass from 6 days to adulthood, followed by *M. spretus* (which had high-intermediate values), *M. musculus* (with low-intermediate values) and, finally, *M. pahari*, with the lowest relative testes mass throughout the period examined.

**Absolute measures and proportions of seminiferous tubules**

Different measures were obtained from seminiferous tubule cross-sections (Fig. 2). From birth to sexual maturity, all the components of the seminiferous tubules increased significantly in size. Most measures differed significantly between species throughout postnatal development. Thus, the seminiferous tubule cross-sectional area ($F_{3,84} = 29.26, P<0.001$; Fig. 3A), seminiferous epithelium height ($F_{3,84} = 43.29, P<0.001$; Fig. 3B), and seminiferous epithelium area ($F_{3,84} = 51.22, P<0.001$; Fig. 3D) were different between mouse species, whereas no significant differences were seen in lumen area (Fig. 3C). *Post-hoc* analyses revealed that, with the exception of the lumen area, there were clear differences between all four species in absolute measures.
of tubule size. In adult mice, the seminiferous tubule cross-sectional area (F_{3,20} = 9.63, P < 0.001) and the seminiferous epithelium area (F_{3,20} = 10.16, P < 0.001) were significantly higher in M. musculus than in M. spicilegus, whereas seminiferous epithelium height was higher in M. spretus than in M. spicilegus (F_{3,20} = 5.18, P = 0.008). Overall, the main differences between species in tubule cross-sectional area, epithelium height, and epithelium area were noted between 20 and 30 days postpartum and some remained thereafter. However, none of these differences exhibited a pattern in agreement with what was expected if they were associated with differences in the levels of sperm competition of these species.

The proportion of total seminiferous tubules and interstitial tissue was calculated at different times after birth (see Fig. 2). The percentage of total seminiferous tubules increased significantly during postnatal development (F_{8,84} = 132.60, P < 0.001), with an overall significant difference between species (F_{3,84} = 52.96, P < 0.001; Fig. 4A). In agreement with this, the percentage of interstitial tissue decreased during postnatal development (Fig. 4B; F_{8,84} = 130.77, P < 0.001). Differences between species showed a wide range of values shortly after birth (at 6 days, the range was 60–80% seminiferous tubules for the four species), but the range was narrower in adult animals (about 90–95% for all species). Thus, there was a marked increase in the percentage of seminiferous tubules in M. pahari (from ~60 to ~90%) and M. spicilegus (from ~65 to ~95%), with a less pronounced increase in M. musculus (from ~75 to ~90%) and M. spretus (from ~80 to ~95%). The differences in increase in percentage of seminiferous tubules between the species did not show any relationship with relative testes mass that would suggest that such increase is directly associated with levels of sperm competition. However, the percentage of seminiferous tubules in adult males showed significant differences (F_{3,20} = 26.43, P < 0.001), with two species (M. pahari, M. musculus) having lower values than the two other species (M. spretus, M. spicilegus; Fig. 4A), in agreement with their differences in relative testes mass.

Volumetric measures

The volume of testes increased from birth to adulthood (F_{8,84} = 269.98, P < 0.001) and such an increase varied significantly between species (F_{3,84} = 267.69, P < 0.001). The species also showed a considerable effect on the volume of testes occupied by seminiferous tubules (Fig. 5A; F_{8,84} = 266.47, P < 0.001), which increased in all species throughout postnatal development (F_{8,84} = 267.63, P < 0.001). The volume of testes occupied by seminiferous tubules was consistently lower in M. pahari and M. musculus than in M. spicilegus and M. spretus from 10 days onwards (F_{3,8} = 23.51, P < 0.001) with the values for M. spicilegus being significantly higher than those for the other species at 45 days postpartum (F_{3,8} = 95.88, P < 0.001). Adult males of M. pahari and M. musculus showed a similar volume of testes occupied by seminiferous tubules, which was significantly lower than that in M. spretus which, in turn, was significantly lower than that in M. spicilegus (F_{3,20} = 197.20, P < 0.001). These differences were associated with the levels of sperm competition in these species.

Relative to body mass, the volume of testes occupied by seminiferous tubules differed widely between species (Fig. 5B), M. spicilegus being the species with the highest values along all of postnatal developmental period, followed, in decreasing order, by M. spretus, M. musculus, and M. pahari, the species with the lowest relative volume of testes occupied by seminiferous tubules. These results were, again, associated with levels of sperm competition in the four mouse species.

Changes in cell types in seminiferous tubules during postnatal development

During postnatal testicular development, we quantified the proportion of seminiferous tubules according to the most advanced germ cell stage in each tubule, in an attempt to characterize the first spermatogenic wave (Fig. 6A-I). There were important changes in the...
predominant type of seminiferous tubules from birth to adulthood and clear differences were seen between species with regards to the pattern of predominant seminiferous tubule types.

A comparison between species revealed significant differences in the pattern of cells present in the testes at various time points during postnatal development (Fig. 7A–D). At the earliest time examined (14 days), *M. pahari* exhibited all seminiferous tubules with only Sertoli cells and spermatogonia, and this was also the predominant seminiferous tubule type of *M. musculus* testes, although some tubules contained spermatocytes. On the other hand, in *M. musculus* and *M. spicilegus*, spermatocytes were present in the majority of tubules. At 30 days postpartum, *M. pahari* exhibited about 30% of tubules with round spermatids as the most advanced stages, with the majority of tubules containing spermatocytes. In contrast, in *M. musculus*, the largest proportion of tubules contained round spermatids with about one-third of them already exhibiting spermatids (15% with elongating spermatids, and another 15% with elongated spermatids). Furthermore, in *M. spretus* and *M. spicilegus*, about half of the tubules contained elongated spermatids. By 45 days, *M. pahari* still had over 10% of tubules with spermatocytes as the most advanced stage and a high proportion of tubules (≈30%) with round spermatids, whereas in the other three species the vast majority of tubules contained elongating and, mainly, elongated spermatids. Finally, no major differences were seen between species in the pattern exhibited at 60 and 120 days postpartum, with the majority of seminiferous tubules containing elongated spermatids (Fig. 7).

A comparison of the progression over time in the proportion of tubules with different cell types showed differences between the species (Fig. 8A–D). Changes in the percentage of seminiferous tubules with Sertoli cells and spermatogonia, the earliest stage recognized, revealed that the decline over time was very fast in *M. musculus*, *M. spretus*, and *M. spicilegus*, whereas the decrease in *M. pahari* was much slower (Fig. 8A). Changes in the proportion of tubules with spermatocytes as the most advanced stage revealed a steep rise followed by a rapid decline in *M. musculus* and *M. spicilegus* while in *M. spretus* there was a continuous decline from the time of first sampling, suggesting that the first time point examined was already past the peak in the proportion of seminiferous tubules with spermatocytes; on the other hand, in *M. pahari*, the increase in percentage of tubules with spermatocytes developed slowly and the decrease was also slow (Fig. 8B). The timing of appearance of a majority of tubules with round spermatids varied between species and the same was observed with the decline in such proportion; however, the pattern (shape) of the curve was similar for most species (with similar time frames), but it was much faster for *M. spicilegus* with a steeper rise and decline (Fig. 8C).

Figure 3 Measures of seminiferous tubules from 6 to 120 days after birth in four mouse species (*M. pahari*, *M. musculus*, *M. spretus*, and *M. spicilegus*). (A) Seminiferous tubule cross-sectional area. (B) Height of the seminiferous epithelium. (C) Area of the lumen of the seminiferous tubule. (D) Area occupied by the seminiferous epithelium. For details of how measures were taken, see the ‘Materials and Methods’ section.
Finally, the rise in the proportion of tubules with differentiating spermatids (i.e. elongating and elongated) followed a sigmoid pattern that was similar for all species with the exception of *M. pahari* which had a much later start but which then exhibited a similar rate of increase as the other species (Fig. 8D). The proportion of tubules with differentiating spermatids reached a plateau at 45 days postpartum in three species (*M. musculus*, *M. spretus*, and *M. spicilegus*) and only at 60 days in the remaining one (*M. pahari*).

Based on the changes described above, we estimated the time taken for each species to undergo the transition between several stages, namely from spermatocyte (i.e. the beginning of meiosis) to round spermatids (the end of meiosis), and elongating and elongated spermatids (i.e. the completion of spermiogenesis), and the overall transition from spermatocytes to elongated spermatids. We found that *M. pahari* and *M. musculus* had a transition time from spermatocytes to elongated spermatids of 36 and 40 days respectively, whereas the estimated transition times of *M. spretus* and *M. spicilegus* were 31 and 25 days respectively (Fig. 9). Thus, species with higher levels of sperm competition have faster transition times from spermatocytes to elongated spermatids.

**Discussion**

The results of this study suggest that sperm competition selects for changes in testicular architecture and kinetics of germ cell development that are established early in postnatal development at the time of the first spermatogenic wave. The absolute and relative volumes of seminiferous tubules in the testis exhibit differences between species in agreement with their sperm competition levels and, similarly, the kinetics of germ cell development, as seen from the proportion of tubules with different germ cell types during postnatal development, are also associated with sperm competition levels.

Perhaps the most generally observed response to sperm competition is an increase in sperm numbers (Gomendio et al. 1998, Gómez Montoto et al. 2011) because more sperm enhance the chances of fertilization, as seen in the mouse (Robl & Dziuk 1984) and other species (Birkhead & Møller 1998, Gomendio et al. 1998). Thus, it is important to understand how males can increase sperm output under conditions of sperm competition. Adult males are known to invest in larger testes in relation to body mass when facing higher sperm competition levels (Harcourt et al. 1981, Hosken 1997, 1998, Stockley et al. 1997, Gomendio et al. 1998, Byrne et al. 2002). Recent studies have revealed that testis morphology and architecture may also be influenced by sperm competition (Lüpold et al. 2009b, 2011, Rowe & Pruett-Jones 2011). Furthermore, the duration of spermatogenesis may be modified to increase sperm production under sperm competition, with higher relative testes mass associated with changes in the kinetics of spermatogenesis that would presumably generate sperm at faster rates (Wistuba et al. 2003, Luetjens et al. 2005, Ramm & Stockley 2010).

We have used a group of closely related mouse species differing in sperm competition levels as revealed by testes mass relative to body mass. Values of body and testes masses of the males in this study, which are derived from natural populations but have been bred for a few generations under laboratory conditions, were similar to those recorded in natural populations (Frynta et al. 2009). Sperm competition in these species has favoured an increase in sperm numbers (quantity) in epididymal...
reserves, and in sperm quality (percentages of motile, normal, and acrosome-intact spermatozoa; Gómez Montoto et al. 2011). Differences in sperm competition levels in these mouse species were also associated with functional sperm traits such as the proportion of sperm that undergo changes in preparation for fertilization (capacitation) and the percentage of spermatozoa that undergo exocytosis of the acrosome in response to a physiological ligand (Gomendio et al. 2006). In addition, these species showed fertilization success in competitive contexts that mirrored their levels of sperm competition (Martín-Coello et al. 2009). As a consequence, these mouse species were regarded as a suitable model to examine the differences in testicular morphology and kinetics of germ cell transitions during the establishment of spermatogenesis.

We detected clear differences between these species in postnatal changes in body mass (i.e. growth) and testes mass and, as a consequence, in their relative testes mass at different times after birth. Testes mass in relation to body mass showed differences between species that were evident after 14 days, with two groups clearly noticeable, one of *M. pahari* and *M. musculus*, with low-intermediate sperm competition levels, and another one of *M. spretus* and *M. spicilegus*, with intermediate-high levels of sperm competition. These differences remained thereafter. Thus, differences in absolute testes mass, and in testes mass relative to body mass, appear very early in postnatal development.

We also examined changes in measures (size) of different components of seminiferous tubules. There was an increase in dimensions and area throughout the period of postnatal development. An increase in seminiferous tubule cross-sectional area, and seminiferous epithelium height and area, was observed in all species beginning at 20 days postpartum, but the differences observed did not follow the pattern predicted from the levels of sperm competition of these species. These results suggest that there is no direct association between sperm competition levels and absolute measures of seminiferous tubules and they agree with those reported by Lupold et al. (2009b) who also found differences between tubule dimensions in passerine birds, although such differences were not explained by relative testes mass or other indicators of female promiscuity such as a mating system.

Changes in the percentage of total seminiferous tubules in the testis, i.e. the proportion of testis area occupied by seminiferous tubules, could explain differences in sperm output. We reasoned that differential investments in sperm-producing machinery may become evident soon after spermatogenesis is switched on. We found that changes in the percentage of total seminiferous tubules varied considerably between species from the early stages after birth. It was lower for *M. pahari*, the species with the lowest relative testes mass, although values were not different for the other species throughout postnatal development, and therefore no clear association with sperm competition levels was seen. On the other hand, an association between the percentage of seminiferous tubules and sperm competition levels was seen among adult males and this agrees with recent studies in birds (Rowe & Pruett-Jones 2011). We also reasoned that, rather than the percentage of area occupied by tubules, the overall proportion of testes volume occupied by seminiferous tubules would be more informative. This was indeed the case, because volume of testes occupied by seminiferous tubules (*V* _TTb_; both absolute and relative to body mass) was different between species throughout postnatal development, and such differences were associated with the levels of sperm competition. It follows that testicular architecture is

![Figure 5](A) Volume of testes occupied by seminiferous tubules and (B) relative volume occupied by seminiferous tubules in the testes of four mouse species (*M. pahari*, *M. musculus*, *M. spretus*, and *M. spicilegus*) at different times after birth.

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**Figure 5** (A) Volume of testes occupied by seminiferous tubules and (B) relative volume occupied by seminiferous tubules in the testes of four mouse species (*M. pahari*, *M. musculus*, *M. spretus*, and *M. spicilegus*) at different times after birth.
established early on during the first spermatogenic wave, with the volume of machinery assigned to sperm production following the levels of sperm competition. To the best of our knowledge, this is the first evidence identifying such a developmental pattern.

Finally, we also assessed the progression in the proportion of seminiferous tubules categorized according to the most advanced cell stage in the sequence of germinal epithelium differentiation. Prior studies in the laboratory mouse have revealed a precise timing of appearance of ever more differentiated cell stages at different times after birth until continuous spermatogenesis is established (Bellvé et al. 1977, Bellvé 1993, Borg et al. 2010). This first spermatogenic wave may be rather synchronous in some species (e.g. Dzungarian hamster *Phodopus sungorus*), with the majority of the seminiferous tubules undergoing a parallel progression, or it may lack such synchrony (e.g. rat; Van Haaster & De Rooij 1994). Interestingly, it appears that this first spermatogenic wave may be slightly shorter than that seen in adult animals, and such shortening seems to take place perhaps in the early days of *postpartum* development, before testicular descent (with the shortening perhaps due to a higher
In all the species studied so far there is a clear association between the duration of the first spermatogenic wave and subsequent ones (Kluin et al. 1982, Van Haaster & De Rooij 1993). In addition, the first wave has the same pattern of cellular associations observed in the adult male, as seen in various species (Kluin et al. 1982, Van Haaster & De Rooij 1993). Furthermore, the morphology of the spermatagonia observed in immature mice during postnatal development is similar to that of adult spermatagonia (Drumond et al. 2011). The transit through the first spermatogonial phase seems to be the one exhibiting this shortening when compared to adults (Drumond et al. 2011), in agreement with the earlier studies (Kluin et al. 1982), whereas no shortening in the development of spermatocytes and/or spermatids was seen. Therefore, the use of this first spermatogenic wave to understand the kinetics of germ cell differentiation is reliable and well established.

Our study revealed important differences between species with regard to the onset and timing of the first spermatogenic wave. When the pattern of changes in this first spermatogenic wave of the different species was compared with their relative testes mass, a clear association between both was revealed, with the species showing the higher sperm competition levels exhibiting changes earlier than those in the other species. We assessed this further by estimating the duration of the transition from spermatocytes to elongated spermatids in the first spermatogenic wave. This analysis revealed that species with lower levels of sperm competition (M. pahari and M. musculus) had a much slower transition (i.e. more days taken to move along the cell stages, which was about 35–40 days in these two species). In comparison, species with higher sperm competition levels (M. spretus and M. spicilegus) had a much faster transition time, which was estimated to be about 31 and 25 days respectively. These results agree well with data from laboratory mice (of M. m. musculus and M. m. domesticus descent) in which the total duration of spermatogenesis is about 35 days (Oakberg 1956, Hess & França 2008). Our results also agree with the proposal by Ramm & Stockley (2010) that species with higher sperm competition levels, with higher relative testes mass, have shorter cycles of the seminiferous epithelium (indicative of a shorter duration of spermatogenesis), thus allowing for an increase in sperm production rate. Thus, the study of the first wave of spermatogenesis in closely related species will represent an outstanding model for the characterization of gene expression, hormone regulation and local factor control during germ cell proliferation, meiosis, and postmeiotic differentiation under different levels of sperm competition.

In conclusion, this study reveals how developmental patterns during the first wave of spermatogenesis vary...
between species with different levels of sperm competition. These early differences in testicular architecture and kinetics of sperm formation are likely to be responsible for the differences observed at sexual maturity, which influence sperm production rates.

Materials and Methods

Animals and sample collection

Males of *M. pahari*, *M. musculus*, *M. spretus*, and *M. spicilegus* were studied. Animals were born and maintained in our animal facilities where they were kept under standard laboratory conditions in environmentally controlled rooms at 20–24 °C, on a 1400 h light–1000 h darkness photoperiod, and provided with food and water *ad libitum*. Males aged 6, 10, 14, 20, 24, 30, 45, 60, and 120 days postpartum (day of birth was regarded as day 0) were sacrificed and were immediately weighed and dissected. For each species, sample size was three individuals for ages 6–60 days postpartum and six individuals for 120-day-old males. All animal handling was done following the Spanish Animal Protection Regulation RD1201/2005, which conforms to European Union Regulation 2003/65.

Testes were weighed and fixed in Bouin's solution during 24 h, transferred to 70% ethanol and maintained in the latter solution until processed. Testes were embedded in paraffin, sectioned transversally (5 μm thickness), and stained with haematoxylin–eosin. Images were taken using a digital camera (Digital Sight DS-5M, Nikon, Tokyo, Japan) and software for image capture (NIS-Elements F v.2.20, Nikon). Between five and 11 photographic images were taken from different and nonadjacent parts of one testis from each individual (*n* = 6.4 ± 0.7 images, mean ± S.E.M.). Each image included a group of transversal sections of seminiferous tubules. Image sizes were 2560 × 1920 pixels, corresponding to 0.03 mm² of the testis section, for males aged from 6 to 14 days, whereas the area was 0.1 mm² of the testis section for males aged from 20 to 120 days. Images were captured from testis areas where round cross-sections of the seminiferous tubules were prevalent. Measures of seminiferous tubules were taken using ImageJ v.1.41 software (National Institutes of Health, Bethesda, MD, USA).

Absolute measures and proportions of seminiferous tubules

We obtained the following measures from the seminiferous tubule cross-sections (Fig. 2); a) the ‘seminiferous tubule cross-sectional area’ (μm²) was obtained by tracing the tubule’s circumference. b) The ‘seminiferous epithelium height’ (μm) was obtained by measuring the height of the seminiferous epithelium in four different parts of each tubule and then calculating the mean value. c) The ‘seminiferous tubule lumen area’ (μm²) resulted from measuring the length and width of the lumen in each tubule, and calculating the lumen area as the area of an ellipse = π × lumen length/2 × lumen width/2 (semi-major axis) × lumen width/2 (semi-minor axis). Finally, d) the ‘seminiferous epithelium area in each tubule’ (μm²) was calculated by subtracting the lumen area from the seminiferous tubule cross-sectional area.

Figure 8 Comparison between four mouse species (*M. pahari*, *M. musculus*, *M. spretus*, and *M. spicilegus*) of postnatal changes in the seminiferous tubules. Each panel compares the timing of changes in different cell types for different species. Seminiferous tubules were classified according to the most advanced cell stage identified and the percentage of seminiferous tubules with (A) Sertoli cells–spermatagonias; (B) spermatocytes; (C) round spermatids and (D) elongating+elongated spermatids is shown.
To calculate the proportions of seminiferous tubules and interstitial tissue, each photographic image (corresponding to 0.03 or 0.1 mm² of the testis section) was measured to obtain the ‘total seminiferous tubule cross-sectional area’ (μm²), which was the total area of the image represented by seminiferous tubules. This was done by tracing the circumference of cross-sections of all seminiferous tubules in the image (Fig. 2). Then, the ‘percentage of total seminiferous tubules’ was calculated as the proportion of the testis area occupied by seminiferous tubules. The ‘percentage of interstitial tissue’ was calculated as the proportion of testis area not occupied by seminiferous tubules, i.e., the difference between the image area and percentage of area occupied by seminiferous tubules. Using the values of these proportions for each photographic image we then obtained the values of these proportions for each male. Finally, using the values of each proportion for all males of the same age, we calculated the mean value for each age and species.

**Volumetric measures**

The testes volume (V, mm³) of each male was calculated by dividing testes mass by testes density of each species (L Gómez Montoto, M Gomendio and E R S Roldan, unpublished results). Then, the volumetric value of the variable ‘percentage of total seminiferous tubules’ was calculated assuming a uniform distribution of this testis component in the entire organ. Thus, ‘volume of testes occupied by seminiferous tubules’ (VTb, mm³) = V × percentage of total seminiferous tubules/100. The value for this measure was calculated for each male and, subsequently, it was calculated for each age group for each species.

**Postnatal seminiferous tubule development**

The first wave of spermatogenesis was examined in sections from both testes of mice aged 14, 20, 24, 30, 45, and 60 days postpartum; testes sections of adult mice (120 days old) were also assessed. Earlier stages were not included in these analyses because at 6 and 10 days postpartum, cell types in one species (M. pahari) were not easily identified. Seminiferous tubules from the testes of each individual were examined to identify the germ cell types present. The earliest stage recognized comprised tubules containing only Sertoli cells–spermatogonia; spermatogonial types were not distinguished. Other cell types used to categorize seminiferous tubules were spermatocytes, round spermatids, elongating spermatids, and elongated spermatids. Each seminiferous tubule was categorized according to the most advanced cell type identified. Then, the proportion of tubules with each cell type (as the most advanced cell type) was calculated for each postnatal stage of development. The following categories were used to describe the seminiferous tubules: a) Sertoli cell–spermatogonia, b) spermatocytes (premeiosis), c) round spermatids (early spermiogenesis), and d) elongating spermatids and e) elongated spermatids (late spermiogenesis). Data were collected for each testis and means were calculated for each male (from both testes) and, subsequently, for each age group and species.

We estimated the time taken for the transition between some cell types in each species. This was done by taking into account the average time of appearance of the different cell types. We noted the earliest cell type that could be clearly identified in the tubules of all species (i.e. spermatocytes) and recorded the day when the maximum percentage of tubules with this cell type was found. Then, we identified the occurrence of the maximum percentage of tubules with round spermatids and, subsequently, of elongating spermatids. Finally, we noted the time point for each species when the percentage of elongated spermatids reached the highest point and remained elevated. Using these data we calculated the number of days for the different transitions.

**Statistical analyses**

A factorial ANOVA followed by Bonferroni’s post-hoc tests for multiple comparisons was used to test differences in variables of testicular morphology (absolute measures, proportions, and volumetric measures) between mouse species with different levels of sperm competition at different days postpartum. A one-way ANOVA by groups (ages), also followed by Bonferroni’s post-hoc tests for multiple comparisons, was used to detect differences in the previous variables within each age group between mouse species.

For data presentation in figures we calculated relative testes mass for each age group and species following Kenagy and Trombulak’s formula for rodents (Y = 0.031 × X0.77, where Y is predicted testes mass in grams for the observed body mass X; Kenagy & Trombulak 1986). Relative testes mass was calculated as the ratio of observed testes mass to the predicted testes mass Y. Similarly, the relative volume of testes occupied by seminiferous tubules (relative VTb) was calculated for each age group and species by dividing testes tubular volume (VTb) by body mass. Relative testes masses and relative VTb were not used in statistical analyses because they do not properly account for the allometric relationships between the variables (Tomkins & Simons 2002).
All statistical analyses were conducted with SPSS v.17 (SPSS Inc., Chicago, IL, USA) and P values were considered statistically significant at α<0.05.

Declaration of interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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