Derivation of sperm from xenografted testis cells and tissues of the peccary (*Tayassu tajacu*)

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Abstract

Because the collared peccary (*Tayassu tajacu*) has a peculiar Leydig cell cytoarchitecture, this species represents a unique mammalian model for investigating testis function. Taking advantage of the well-established and very useful testis xenograft technique, in the present study, testis tissue and testis cell suspensions from immature collared peccaries (*n* = 4; 3 months old) were xenografted in SCID mice (*n* = 48) and evaluated at 2, 4, 6, and 8 months after grafting. Complete spermatogenesis was observed at 6 and 8 months after testis tissue xenografting. However, probably due to de novo testis morphogenesis and low androgen secretion, functionally evaluated by the seminal vesicle weight, a delay in spermatogenesis progression was observed in the testis cell suspension xenografts, with the production of fertile sperm only at 8 months after grafting. Importantly, demonstrating that the peculiar testicular cytoarchitecture of the collared peccary is intrinsically programmed, the unique Leydig cell arrangement observed in this species was re-established after de novo testis morphogenesis. The sperm collected from the xenografts resulted in diploid embryos that expressed the paternally imprinted gene *NNAT* after ICSI. The present study is the first to demonstrate complete spermatogenesis with the production of fertile sperm from testis cell suspension xenografts in a wild mammalian species. Therefore, due to its unique testicular cytoarchitecture, xenograft techniques, particularly testis cell suspensions, may represent a new and very promising approach to evaluate testis morphogenesis and to investigate spermatogonial stem cell physiology and niche in the collared peccary.

Reproduction (2014) 147 291–299

Introduction

Testis tissue xenograft is a powerful approach developed in the last decade (Honaramooz et al. 2002). In this technique, small fragments of testis from different mammalian donor species (Schlatt et al. 2002, 2010, Honaramooz et al. 2004, 2008, Oatley et al. 2004, Snedaker et al. 2004, Rathi et al. 2006, Arregui et al. 2008, Abrishami et al. 2010, Ehmcke et al. 2011, Abbasi & Honaramooz 2012), are placed subcutaneously under the back skin of an immunodeficient mouse, where they respond to the rodent gonadotropins, initiating and leading to complete spermatogenesis (Honaramooz et al. 2002, Rodriguez-Sosa & Dobrinski 2009). As expected, this approach became an attractive strategy to recapitulate and study testis development in non-rodent species and, among several other important applications, could be used to preserve the germplasm of young individuals in which sperm collection is not an option (Rodriguez-Sosa & Dobrinski 2009). As a very important aspect, the graft-derived sperm fertility was confirmed using ICSI (Honaramooz et al. 2002, 2004, 2008), leading to the birth of healthy offspring (Ruddock et al. 2003, Nakai et al. 2010).

In another powerful approach, when dissociated testis cells were placed subcutaneously under the back skin of an immunodeficient mouse, the cells were able to organize and rearrange into seminiferous cords (de novo testis morphogenesis), which subsequently developed completely, including the production of viable sperm (Gassei et al. 2006, Honaramooz et al. 2007, Kita et al. 2007). Although it has been used for few donor species (Gassei et al. 2006, Honaramooz et al. 2007, Kita et al. 2007), this fascinating methodology became very important for studies involving cell interactions during testis morphogenesis and could be applied, for instance, to preserve the genome of valuable animals and endangered species.
Peccaries are suiforms that are found in the American continent, and among the three recognized species belonging to the Tayassuidae family, the collared peccary (*Tayassu tajacu*) is the most widely spread (*Groves & Grubb 1993*). Due to the high quality of its flesh and leather, this species is raised in captivity and is also one of the most hunted in neotropical forests (*Robinson & Redford 1991, Peres 1996*).

In recent studies carried out in our laboratory, we demonstrated that the collared peccary presents a unique testicular cytoarchitecture, in which Leydig cells are observed almost exclusively surrounding the seminiferous tubule lobes (*Costa et al. 2010, Campos-Junior et al. 2012*). Taking advantage of this peculiar feature, we found that, different from what is suggested in the literature (*de Rooij & Griswold 2012*), Leydig cells probably play a pro-differentiation role in spermatogonial stem cells (*Campos-Junior et al. 2012*). Therefore, the collared peccary may represent an excellent model for investigating the spermatogonial stem cell physiology and niche as well as for studies involving testis function, hormonal regulation of spermatogenesis and germ cell development. The aim of the present study was to use testis tissue and testis cell suspension xenografts from immature collared peccaries to morphofunctionally evaluate spermatogenesis progression and testis morphogenesis.

### Materials and methods

In the present study, eight collared peccaries (four males and four females) and 48 SCID mice were used, following approved guidelines for the ethical treatment of animals (Ethics Committee on Animal Experimentation – Federal University of Minas Gerais, CETEA – UFMG, protocol number 209/2013).

#### Donor testis

Four immature (~3 months old) collared peccaries (*T. tajacu*) weighing ~5 kg were used as testis donors. The animals raised in captivity were obtained from the Federal University of Pará and EMBRAPA/PA, located in the North region of Brazil (Amazon rainforest: 1°27’9”S, 48°29’9”W). Immediately after orchiectomy (*Guimarães et al. 2013*), the testes were transferred to the laboratory in ice-cold DMEM (Cat. No. 10-013-CM, Gibco) containing 2% antibiotic/antimycotic solution (Sigma) and supplemented with 10% of fetal bovine serum (FBS; Gibco).

#### Testis tissue xenograft

Under laboratory conditions, the collected peccary testes were washed three times with DMEM, the tunica albuginea was removed and the testicular parenchyma was then divided into small fragments (~3 mm³) and maintained in supplemented DMEM on ice until xenografting. To evaluate the donor testis histological status, before grafting, representative testis tissue fragments were fixed in Bouin’s solution and routinely prepared for histological analysis. Male immunodeficient SCID mice (6–8 weeks old; *n* = 24) were anesthetized (xylazine (7.9 mg/kg)/ketamine (69.8 mg/kg); *Campos-Junior et al. (2011)*) and castrated, and during the same procedure, four fragments of donor testis tissue were grafted into a small subcutaneous pocket under the back skin of the recipient mice. Two incisions were made on each side of the dorsal midline, just above the forelimbs and hindlimbs, and they were closed using wound clips (Michel Clips 7.5 mm, Miltex, York, PA, USA).

### Testis cell suspension xenograft

A cell suspension of testis cells was prepared by a two-step enzymatic digestion as described previously (*Honaramooz et al. 2002, 2007, Arregui et al. 2008*). Subsequently, aliquots of 2.5 × 10⁶ cells in DMEM were prepared and centrifuged at 1000 g for 5 min, and the pellets were maintained on ice until grafting. The surgical procedure was the same as that described previously for testis tissue xenograft, using, however, different male SCID mice of a similar age (6–8 weeks old; *n* = 24) as recipients.

To evaluate the cell content and viability of the testis cell suspension before xenografting, specific immunolabeling for Sertoli and germ cells and Annexin V/propidium iodide (Pl) apoptotic assay were carried out, in a flow cytometer, using four pellets from each donor. For immunolabeling, the cell samples were permeabilized with PBS with 1% of TWEEN 20 (Sigma) for 15 min. Non-specific binding was blocked with PBS with 1% of BSA (Sigma) for 30 min. Primary antibodies against VASA (DDX4, specific germ cell marker; Abcam, Ab13840; 1:200), CAMA-1 (1:200, Cambridge, Cambridgehire, England) and GATA4 (specific Sertoli cell marker; Santa Cruz Biotechnology, SC-1237; 1:200) were applied, and the cell suspensions were incubated for 60 min and antigens were detected by incubation with Alexa Fluor secondary antibodies (488 anti-rabbit (1:500) or 546 anti-goat (1:500)) for 60 min. All previously described steps were carried out at room temperature.

The Annexin V/PI assay was carried out using the ApopNexinAnnexin V FITC Apoptosis Kit (Millipore APT750, Billerica, Massachusetts, USA.) following the manufacturer’s instructions. The stained cells were acquired and analyzed using FACSscan (Becton & Dickinson, San Jose, California, USA,) and the CellQuest software.

### Biometrical and histological evaluations

Body and graft weights were assessed in mice that received both tissue and cell suspension xenografts at 2, 4, 6, and 8 months after grafting. As an indicator of the presence of bioactive testosterone (*Arregui et al. 2008*) originating from the collared peccary testsis xenograft, seminal vesicle weights were also recorded in mice that received the grafts at the same time points investigated (2, 4, 6, and 8 months). In the control (non-grafted mice), castrated (*n* = 4) and intact (*n* = 4) SCID mice, seminal vesicle weights were recorded only 8 months after grafting.

The grafts were fixed overnight in Bouin’s solution, routinely processed, embedded in paraplast, and sectioned (5 μm) for...
light microscopy histological evaluation, after staining with hematoxylin and eosin. To investigate the progression of spermatogenesis, the most advanced germ cell type present per seminiferous tubule cross section (150 per animal) was scored in the following five categories: i) Sertoli cell only; ii) gonocytes/spermatogonia; iii) spermatocytes; iv) round or elongated spermatids; and v) mature spermatids/sperm. These data are expressed in percentage. Furthermore, 3βHSD (SC-30820; Santa Cruz Biotechnology) immunofluorescence was also carried out as described previously (Campos-Junior et al. 2012).

**Blood perfusion analysis**

Blood perfusion analysis in the xenografted skin was carried out in a non-invasive way and without the use of tracer dyes in anesthetized mice using a laser Doppler perfusion image (LDPI) device (MoorLDPI-2, Moor Instruments, Axminster, Devon, UK). To direct the light beam (830 nm laser) and to avoid interference from surrounding cutaneous blood flow in the measurements, a dark plastic ring was positioned around the xenografts. Non-grafted skin was used as basal control. During the capture of images, the ambient light level was kept at a minimum to avoid any influence on the laser light and the recorded signals. To control for temperature variability, animals were kept at a constant temperature of 37 °C for 5 min before and during the measurements. The mean pixel value of each scanned image was calculated using the software MoorLDI V5.3, and the calculated mean flux is expressed as perfusion units (PU), representing the average blood flow in the region of interest.

**Sperm recovery from the xenografts**

To evaluate their fertility, sperm cells were recovered from both tissue (n=3) and cell suspension (n=3) xenografts at 8 months after grafting. For this purpose (Nakai et al. 2010), the grafts were minced and dispersed in supplemented DMEM. The obtained tissue suspension was centrifuged for 10 min at 600 g and the supernatant was discarded. The sperm cell pellets, mostly sperm, were resuspended in DMEM and maintained at room temperature.

**Oocyte sampling and in vitro maturation and ICSI**

Using standard procedures, ovaries from sexually mature female collared peccaries (n=4) were obtained by ovariectomy (Guimarães et al. 2013). Thirty-two cumulus–oocyte complexes (COCs) were collected from follicles in TCM 199 (Sigma) supplemented with 10% FBS (Gibco, Life Technologies), 20 mM HEPES (Dojindo Laboratories, Kumamoto, Japan), 100 IU/ml penicillin G potassium (Sigma), and 0.1 mg/ml streptomycin sulfate (Sigma). Briefly, about 20 COCs were cultured in 100 μl of maturation medium for 48 h in TCM. After culture, cumulus cells were removed from the oocytes by treatment with 150 IU/ml hyaluronidase (Sigma) and gentle pipetting. Denuded oocytes with the first polar body were harvested under a microscope.

**Figure 1** Histological and cell suspension evaluations of immature collared peccary testis parenchyma before xenografting. (A) At 3 months of age, the donor testis parenchyma exhibited the peculiar Leydig cell (LC) cytoarchitecture already described for sexually mature peccaries. In the seminiferous cords (SC), gonocytes (G) were the only germ cell type present at this age and, in the cell suspension, these germ cells (B) were characterized as VASA positive (DDX4 – green). Flow cytometric analysis of the testicular parenchyma cell suspensions revealed that gonocytes (C) and Sertoli cells (D) represented approximately one-sixth (~19%) and one-third (~35%) respectively of the cell population. In the Annexin V-FITC/PI staining assay (E, F, and G), the cell suspension was found to contain mostly viable cells (~92%; A~/PI~/~) and few early-phase (A+/PI−) and late-phase (A+/PI+) apoptotic cells, as well as necrotic cells (A~/PI+/). Bar: A=200 μm; inset in A=30 μm and that in B=30 μm.
stereomicroscope, and 18 oocytes were considered as in vitro matured.

As the number of obtained viable oocytes was small, ICSI procedure was carried out using spermatozoa only from testis cell suspension xenografts (eight oocytes), and parthenogenetic embryos were also produced as a control (Pereyra-Bonnet et al. 2008). Embryo rate was assessed 24 and 48 h after ICSI, and the DNA content of the parthenogenetic and ICSI embryos was evaluated using NanoDrop (ND1000, Thermo Scientific). To investigate whether the embryos contained sperm DNA, the expression of the paternally imprinted gene NNAT was also assessed by RT-PCR as follows. Total RNA was isolated from individual embryos using TRIzol Reagent (Life Technologies). Samples were quantified using NanoDrop (ND1000, Thermo Scientific), and their integrity was examined by gel electrophoresis in 1% agarose. Amplification was carried out in the Mastercycler (Eppendorf, Hauppauge, New York, USA.), with initial incubation at 94 °C for 2 min and 94 °C for 2 min, followed by 35 cycles of 94 °C for 15 s, 55 °C for 15 s, and 72 °C for 1 min. PCR fragments were visualized by gel electrophoresis in 1% agarose.

**Statistical analyses**

All the data were tested for normality and homoscedasticity of the variances. Parametric data were analyzed by ANOVA, and the differences were compared by Tukey’s test, whereas Student’s t-test was used for two-parameter analysis. All the data are expressed as means ± S.D., and the significance level considered was P < 0.05. All the analyses were carried out using the GraphPad Prism (version 5, La Jolla, California, USA). The flow cytometer data were analyzed using the FlowJo software (treeStar Software, Ashland, OR, USA).

**Results**

**Collared peccary testis sample evaluations before xenografting**

As has been observed in the testis tissue fragments obtained from 3-month-old peccaries (Fig. 1A), gonocytes were the most advanced germ cell type present in the seminiferous cords and the Sertoli cells were still immature. This figure also clearly shows that the peculiar peccary testis structure observed in sexually matured animals obtained from 3-month-old peccaries (Fig. 1A), gonocytes were the most advanced germ cell type present in the seminiferous cords and the Sertoli cells were still immature. This figure also clearly shows that the peculiar peccary testis structure observed in sexually matured

![Biometric data obtained at different time points after collared peccary testis tissue and cell suspension xenografting.](image-url)
Among the four experimental groups at any time point evaluated (Fig. 2A). It can be observed in this figure that the weights of both graft types increased gradually and that, at all the four time points investigated, the xenografts derived from the testis tissue were bigger (P<0.05) than those derived from the cell suspensions (Fig. 2B). The graft recovery rates obtained in the present study were 100% (96/96) and 78% (75/96) respectively for the testis tissue and cell suspension xenografts. Except at 8 months, at the other three time points studied the seminal vesicle weights of mice that received testis tissue xenografts were greater (P<0.05) than that observed for the mice grafted with cell suspensions (Fig. 2C and D). However, in contrast to the castrated mice that did not receive grafts, at 8 months after xenografting, the seminal vesicle weights of both the grafted mouse groups reached that of the intact mice (P>0.05; Fig. 2D). At the four time points investigated, testis tissue and cell suspension xenograft weights were significantly and positively correlated with the seminal vesicle weights (r=0.65 and r=0.97 respectively).

**Biometric data of recipient mice**

Body, graft, and seminal vesicle weights at different time points after testis tissue and cell suspension xenografting are shown in Fig. 2. Although the body weights of mice increased continuously during the experimental period, no significant differences (P>0.05) were observed among the four experimental groups at any time point evaluated (Fig. 2A). It can be observed in this figure that the weights of both graft types increased gradually and that, at all the four time points investigated, the xenografts derived from the testis tissue were bigger (P<0.05) than those derived from the cell suspensions (Fig. 2B). The graft recovery rates obtained in the present study were 100% (96/96) and 78% (75/96) respectively for the testis tissue and cell suspension xenografts. Except at 8 months, at the other three time points studied the seminal vesicle weights of mice that received testis tissue xenografts were greater (P<0.05) than that observed for the mice grafted with cell suspensions (Fig. 2C and D). However, in contrast to the castrated mice that did not receive grafts, at 8 months after xenografting, the seminal vesicle weights of both the grafted mouse groups reached that of the intact mice (P>0.05; Fig. 2D). At the four time points investigated, testis tissue and cell suspension xenograft weights were significantly and positively correlated with the seminal vesicle weights (r=0.65 and r=0.97 respectively).

**Figure 3** Histological status of collared peccary testis tissue evaluated at different time points after xenografting. As can be observed, overall, the characteristic testicular cytoarchitecture of the collared peccary was maintained and spermatogenesis progressed until the formation of sperm. Therefore, the most advanced germ cell types observed at 2, 4, and 6–8 months were respectively spermatogonia (arrowhead in A′), pachytene spermatocytes (arrowhead in B′), and mature spermatids/sperm (arrowheads in C′ and D′). LC, Leydig cells; SC, seminiferous cords; ST, seminiferous tubules; *, tubular lumen; and arrow, Sertoli cell. Bar: A, B, C, and D = 200 μm; A′, B′, C′, and D′ = 70 μm; and A″, B″, C″, and D″ = 40 μm.

**Figure 4** Histological status of collared peccary testis cell suspensions evaluated at different time points after xenografting. As can be observed, after de novo testis morphogenesis, the peculiar testicular cytoarchitecture of the collared peccary was maintained and, although at a slower pace, complete spermatogenesis was observed at 8 months of age. Spermatogonia (inset in A″ and arrowhead in B″), early primary spermatocytes (arrowhead in C″), and mature spermatids/spERM (arrowhead in D″) were respectively the most advanced germ cell types observed at 2, 4, 6, and 8 months. LC, Leydig cells; SC, seminiferous cords; ST, seminiferous tubules; *, tubular lumen; arrow, Sertoli cell. Bar: A, B, C, and D = 200 μm; A′, B′, C′, and D′ = 70 μm; and A″, B″, C″, and D″ = 40 μm; inset in A‴ = 30 μm.
Spermatogenesis progression and de novo testis morphogenesis in the xenografts

Histological evaluations of both testis tissue and cell suspension xenografts were first carried out at 2 months after grafting. In comparison with the testis histological status observed before grafting (Fig. 1A), the only difference observed was that gonocytes advanced up to spermatogonia. Moreover, testis structure appeared healthy and presented the peculiar characteristic of the collared peccary testis, Sertoli cells were still immature, and no evidence of lumen was observed in the seminiferous cords (Fig. 3A). At 4 months after grafting, the germ cells progressed until the meiotic phase of spermatogenesis and the presence of tubular lumen indicated that the Sertoli cells were more mature (Fig. 3B). Complete spermatogenesis was established at 6 and 8 months after grafting (Fig. 3C and D) and therefore Sertoli cells presented characteristics of fully mature cells and the tubular lumen was prominent, particularly at 8 months after grafting.

At 2 months after testis cell suspension xenografting, the testis cells interacted and de novo testis morphogenesis had already occurred. As has been observed for the testis tissue grafts, the peculiar collared peccary testis parenchyma cytoarchitecture was re-established, and very few spermatogonia were observed (Fig. 4A). At the other three time points investigated, in comparison with the testis tissue xenografts, spermatogenesis progression was delayed (Fig. 4B and C). Therefore, complete spermatogenesis was observed for the first time only at 8 months after grafting (Fig. 4D). Figure 5 shows, through DAPI and the Leydig cell marker 3βHSD, that in both graft approaches used, the unique testicular cytoarchitecture of the collared peccary was preserved.

Although carried out only at 8 months, the quantitative histological evaluations of the xenografts confirmed that spermatogenesis progression was delayed in the cell suspension xenografts. For instance, whereas most of the seminiferous cross sections in the testis tissue grafts presented spermatids, the cell suspension xenograft tubular profiles evaluated presented mainly spermatocytes (Fig. 6A). At the same time point, in comparison with the non-grafted SCID mice, the blood perfusion analysis indicated that the testis tissue and cell suspension xenografts were ~3.5-fold more vascularized in the grafted mice (P<0.05; Fig. 6B and C).
Sperm fertility evaluation

As can be observed in Fig. 7A and B, the collared peccary sperm produced in the xenografts were successfully harvested. These graft-derived sperm, evaluated by ICSI, were able to produce two-cell and four-cell embryos (Table 1 and Fig. 7C and D) and, suggesting that these embryos were diploid, the DNA content of these graft-derived ICSI embryos was approximately twofold higher than that found in the parthenogenetic control embryos (Fig. 7E). Corroborating this finding, the expression of the paternally imprinted gene NNAT was observed exclusively in the four-cell (two of three) ICSI embryos, whereas that of GAPDH (constitutive gene) was observed in the injected and parthenogenetic embryos (Fig. 7F).

Discussion

Using testis tissue and testis cell suspension xenografts from immature collared peccaries, we observed the occurrence of complete spermatogenesis. Particularly, our data suggest, for the first time in a wild mammalian species, that sperm produced from the testis cell suspension grafts are able to generate diploid embryos expressing NNAT, a paternally imprinted gene. The peculiar testicular cytoarchitecture of the collared peccary (Costa et al. 2010, Campos-Junior et al. 2012) was also re-established when the testis cell suspensions were xenografted in immunodeficient mice. Therefore, among other potential applications, collared peccary testis xenografts can be used to investigate spermatogenic stem cell physiology and niche, as well as mechanisms involved in testis morphogenesis in this species presenting a unique Leydig cell arrangement.

Similar to several other mammalian species (Schlatt et al. 2002, Honaramooz et al. 2004, 2008, Oatley et al. 2004, Snedaker et al. 2004, Rathi et al. 2006, Arregui et al. 2008, Abrishami et al. 2010, Abbasi & Honaramooz 2011, Gourdon & Travis 2011), in the present study, we demonstrated that testis fragments from pre-pubertal collared peccaries were successfully preserved under the back skin of SCID mice, with complete spermatogenesis being observed at 6 months after grafting. Additionally, when testis cells isolated from the same peccaries were xenografts, they were able to interact and de novo testis morphogenesis occurred, with complete spermatogenesis being observed only at 8 months after xenografting. Except for a few other species investigated in this aspect (mice, Kita et al. (2007), domestic pig, Honaramooz et al. (2007), and sheep, Arregui et al. (2008)), to our knowledge, this is the first study to demonstrate the morphogenetic capacity of testis cells in a wild species.

Under captivity conditions, sperm release (puberty) in the seminiferous tubule lumen in the collared peccary is first observed at 11–12 months of age (Guimaraes et al. 2013). Therefore, probably due to the more mature hypothalamic–pituitary–gonadal axis in the recipient mice (Rodriguez-Sosa & Dobrinski 2009), the first spermiation in peccary testis tissue xenograft is clearly advanced (Schlatt et al. 2002, Honaramooz et al. 2004, 2008, Oatley et al. 2004, Rathi et al. 2006, Arregui et al. 2008, Abrishami et al. 2010, Abbasi & Honaramooz 2011, Gourdon & Travis 2011). However, because the initial cellular interactions and testis reorganization in peccaries investigated in the present

Table 1 In vitro development of collared peccary oocytes injected with sperm derived from the testis cell suspension xenograft.

<table>
<thead>
<tr>
<th>Experimental group</th>
<th>Number of mature oocytes (%)</th>
<th>Two-cell embryos (%)</th>
<th>Four-cell embryos (%)</th>
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<tr>
<td>Graft-ICSI</td>
<td>8</td>
<td>75 (6/8)</td>
<td>37.5 (3/8)</td>
</tr>
<tr>
<td>Parthenogenetic</td>
<td>10</td>
<td>40 (4/10)</td>
<td>10 (1/10)</td>
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study take ~ 6 weeks to occur (PHA Campos-Junior, GMJ Costa & LR França 2013, unpublished data), this advancement was less evident in the testis cell suspension xenografts. Furthermore, the unique testicular cytoarchitecture of the collared peccary was re-established after de novo testis morphogenesis, which strongly suggests that this peculiar feature is intrinsically programmed by the donor cells, and this distinct characteristic provides a new and very useful tool to investigate testis development and function.

In agreement with the literature (Schlatt et al. 2002, Honaramooz et al. 2004, 2008, Oatley et al. 2004, Arregui et al. 2008), the increase in the seminal vesicle weight of recipient mice demonstrated that the collared peccary grafts were responsive to mouse gonadotropins, producing enough bioactive androgens to support complete spermatogenesis. The significant and positive correlation between the seminal vesicle and graft weights, as well as the graft histological status (i.e., Sertoli cell nuclear morphology, lumen formation, and spermatogenesis progression), over the four time points investigated is a good indication of the importance of androgen secretion for graft development and maturation. Therefore, in comparison with the testis tissue xenografts, the delay in spermatogenesis progression observed in the testis cell suspension grafts is a functional consequence of low androgen levels.

Although some studies have shown that sperm produced in testes xenografts are able to produce diploid embryos and healthy progeny (Honaramooz et al. 2004, 2008, Nakai et al. 2010), the fertility of cell suspension graft-derived sperm has never been evaluated in a wild species. In the present study, we were able to graft-derived sperm has never been evaluated in a

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.

Funding

Support by FAPEMIG, CNPq and FAPESPA.

Acknowledgements

The authors thank Mara Lívia Santos for providing technical help. They also thank the FAPEMIG and CNPq for awarding scholarships to PHACJ, GMJC, LSB, and SMSNL. LSB additionally thanks INCT-Nanobiofar (CNPq/MCT/FAPEMIG).

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Received 11 November 2013
First decision 25 November 2013
Revised manuscript received 4 December 2013
Accepted 9 December 2013