Spermatogenesis does not require the local production of follistatin

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Abstract

It has been proposed that follistatin can modulate the actions of activins and/or other members of the transforming growth factor-β superfamily of proteins on testicular function, since mice overexpressing follistatin showed spermatogenic disruption. However, since mice with targeted disruption of the follistatin gene die soon after birth, it is not feasible to determine the effect of the absence of follistatin on testicular function using this model. To further understand the role of follistatin on the development and maintenance of spermatogenesis, fetal testes, collected by Caesarean section at day 18 of gestation from follistatin null mice, were transplanted to the external ear of castrated recombination activating gene 1 immunocompromised male mice. The testicular grafts were then analysed 7–8 weeks after transplantation and showed that full spermatogenesis developed in both the testes of wild-type and follistatin null mice. This study indicates that, if follistatin is required to modulate spermatogenic development, it is not supplied by local testicular production but by circulating follistatin from the host mouse.


Introduction

The inhibins and activins were isolated from the gonads on the basis of their capacity to specifically modulate follicle-stimulating hormone (FSH) secretion. The activins stimulate, whereas the inhibins suppress FSH secretion by pituitary cells (Robertson et al. 1985, Ling et al. 1986, Vale et al. 1986). Follistatin, a glycosylated single-chain protein with no structural similarity to inhibin, can suppress FSH secretion by pituitary cells (Robertson et al. 1987, Ueno et al. 1987), an action subsequently shown to be due to its capacity to bind and neutralize the actions of the activins (Nakamura et al. 1990). Follistatin is produced in two forms through an alternative splicing event to produce follistatin 288, which binds to heparin sulphate proteoglycans and is considered locally active (Esch et al. 1987). A larger form, follistatin 315, does not bind to heparin sulphate proteoglycans and is considered to be a circulating form. Recent data also indicate that follistatin can bind other members of the transforming growth factor-β superfamily of proteins, namely bone morphogenetic proteins (BMP) 2, 4, and 7, albeit with less than 10% of its affinity for activin (Iemura et al. 1998, Glister et al. 2004), thereby expanding the potential physiological functions of this protein.

The activins and inhibins have been shown to have local actions within the testis. Activin A enhanced spermatogonial proliferation in vitro (Mather et al. 1990, Hakovirta et al. 1993) and promoted the reaggregation of Sertoli and germ cells in the absence of basement membranes and peritubular cells (van Dissel-Emiliani et al. 1989, Mather et al. 1993). In contrast, the inhibins suppressed spermatogonial proliferation when injected locally into the adult hamster testis (van Dissel-Emiliani et al. 1989). Activin A has also been shown to synergise with FSH in the stimulation of Sertoli cell proliferation (Buzzard et al. 2003) and exerts temporal-specific actions on the transformation of gonocytes to spermatogonia (Meehan et al. 2000). Follistatin antagonized the ability of activin A to aggregate Sertoli cell monolayers but did not inhibit the activin-induced stimulation of spermatogonia grown in co-culture with Sertoli cells (Mather et al. 1993).

Study of the action of follistatin on testicular function by targeted disruption of the follistatin gene in mice was not possible, since these mice had multiple defects in
other organs that resulted in death immediately after birth (Matzuk et al. 1995). However, transgenic mice overexpressing follistatin showed variable levels of disruption of spermatogenesis and Leydig cell hyperplasia, either acting through the neutralisation actions of the activins or the BMPs that can bind to follistatin (Guo et al. 1998). Evidence of a direct action of activin A on the testis emerged in transgenic mice overexpressing the activin βA subunit gene, which showed disruption of spermatogenesis (Tanimoto et al. 1999). Evidence of actions of BMP-4, 8-B and 8-A have emerged from studies of targeted disruption of these genes in mice indicating their importance in primordial germ cell generation, and the initiation and maintenance of spermatogenesis (Zhao et al. 1996, 1998, Lawson et al. 1999, Ying et al. 2000).

Given the actions of activin A and certain BMPs on the development of the testis, we set out to evaluate the development of testes from follistatin null mice by transplanting testes from these and wild-type mice to the external ear of castrated immunocompromised male mice, in order to determine if circulating follistatin 315, provided by the recipient could support testis development in the absence of locally produced follistatin by the transplanted donor mouse testicular tissue.

Materials and Methods

Experimental animals

C57/129 hybrid mice, heterozygous for the deleted follistatin allele were used to produce follistatin null offspring (Matzuk et al. 1995). The testes were collected from fetuses delivered by Cesarean section at day 18 of gestation. Graft recipients were 8–9-week-old recombinant activating gene (RAG) male mice that were immunodeficient and do not produce mature T or B lymphocytes. All mice were obtained from Monash Animal Services, Monash University, Australia and housed under a 12 h light:12 h darkness cycle at 22 °C. This study was approved by the Monash Medical Centre Animal Ethics Committee (MMCA 2000/43) and conforms to the conditions laid down by the NHMRC/CSIRO/AAC Australian Code of Practice for the Care and Use of Animals for Scientific Purposes (1997).

Collection of fetal testes for transplantation

Following decapitation of the fetus, the abdomen was opened and the testes, epididymis and part of the vas were removed and transferred into Dulbecco’s PBS (GibcoBRL, Life Technologies) at room temperature until transplantation. The tail of each donor fetus was removed for subsequent genotyping.

Transplantation and castration procedures

All procedures were performed in an specific pathogen-free room and sterile techniques were used for all procedures. Mice were anaesthetized by an i.p. injection solution of 0.5 ml Rompun (Xylazil 20, 20 mg/ml), 0.5 ml ketamine (100 mg/ml) and 9 ml PBS. A single dose of 0.3–0.45 ml was usually sufficient for most mice. The 8–9 week-old male mice were castrated through small scrotal incisions. Subsequently, following removal of the epididymis and vas, the fetal testes were transplanted into the recipient’s external ears using a technique similar to that previously reported for the rat (Johnson et al. 1996). A 1.5 mm opening was made on the dorsal surface of the pinna at a point about two-thirds of the distance from the tip of the ear. Through it, a channel was formed under the skin toward the ear tip by blunt, gentle dissection. The tip of the channel was pierced by a fine needle and the fetal testicular graft was inserted through this opening. The wound in the ear was self-sealing and did not require sutures.

Each recipient received one pair of testes from a single donor fetus and when the genotyping of the donor was complete, the RAG mice receiving grafts from mice heterozygous for the deletion of the follistatin gene were killed 24 h later. Only mice receiving testes from wild-type mice and mice homozygous for the deletion of the follistatin gene were allowed to survive until 7–8 weeks after transplantation.

Genotyping

A tail biopsy of 1.5 mm from each fetus was digested in 100 µL lysis buffer (10 mM Tris–HCl (pH 8.3), 50 mM NaCl and 0.2% Tween 20) plus 1 µl proteinase K (19 mg/ml) at 55 °C for 65 min, then at 98 °C for 12 min, and kept on ice to perform PCR later. Two pairs of PCR primers, which were the pair of hHPRT.3F (5′-TGCTGACCTGCTGGAAATTACA-3′) and hHPRT.3R (5′-CTGCATTGTATTGCCCAGTGT-3′) and the pair of Foldel.F (5′-CGCTGCAGGTCTGTGATAA-3′) and Foldel.R (5′-CTTTACAAGGGATCGTGG-3′), were used for differentiating the homozygous and heterozygous state for the deleted follistatin allele, and wild type. The PCR conditions were set up at the initial denaturation of 94 °C for 5 min, followed by 35 cycles of 94 °C for 30 s, 58 °C for 30 s and 72 °C for 30 s, and with final extension at 72 °C for 5 min. hHPRT.3F and hHPRT.3R were used for targeting the replacement cassette of the deleted mouse follistatin gene. The PCR products from the primer pair of hHPRT.3F and hHPRT.3R were 208 bp in size. Therefore, when there was a band of 208 bp, the genotype of the pup should be a heterozygote or a homozygote of the follistatin knockout. Foldel.F and Foldel.R were used for targeting the mouse follistatin gene. The PCR products from the primer pair of Foldel.F and Foldel.R were 157 bp in size. Therefore, when there...
was a band of 157 bp, the genotype of the pup should be
a wild-type or a heterozygous follistatin mutant. Thus,
the combination of these two pairs of primers enabled
the definition of the genetic status of the pups. The
possible results from PCR genotyping were shown in a
picture of the gel in Fig. 1.

**Histological analysis of testicular grafts**

The testes were removed from the external ear and placed
in Bouin’s fixative for 3–5 h and after transfer to 70%
ethanol, the tissue was dehydrated and embedded in
paraffin wax. The testes were serially sectioned at
a thickness of 5 μm, stained by the periodic acid Schiff’s
reagent and counter stained with 1% Harris’ haematoxylin for 1 min. Following dehydration, the sections
were mounted in DPX (BDH, Poole, Dorset, UK) and
cover-slipped (Fig. 2).

Multiple sections from each transplanted testis were
evaluated qualitatively to determine whether testicular
development had progressed from the stage at day 18 of
gestation when they were transplanted. Subsequently,
the sections from each testis were assessed semi-
quantitatively by determining the percentage of semi-
niferous tubules that showed full development of
spermatogenesis. The statistical significance of these
results was evaluated by the independent samples t-test.

**Results**

**Growth of the testicular grafts**

All grafts survived. The grafted testes increased in size in
their position on the dorsal surface of the external ear
with the increase in testis volume ranging from 50- to
100-fold.

In total, 22 male fetuses were collected for this study,
of which 7 were homozygous for the follistatin gene
deletion, 11 were heterozygous and 4 were wild type.
The 11 RAG mice receiving the grafts from the
heterozygous donors were killed as soon as the results
of genotyping were known. Unfortunately, four of the
seven RAG mice receiving grafts from the homozygous
mice died for unknown reasons some days after the
transplantation surgery. The final analysis was thus
performed on six follistatin null and eight wild-type
testis grafts which were evaluated when the recipients
were killed between 7 and 8 weeks post-transplantation.

**Histological assessment**

The morphological features of the follistatin null and
wild-type fetal testes at day 18 of gestation showed no
significant differences (Fig. 3a and d). The testes consisted
of seminiferous cords composed of immature Sertoli cells
with peripherally placed nuclei and gonocytes that lie
predominantly within the centre of the cords. Groups of
fetal Leydig cells, characterised by their large size and
ovoid nuclei, were present within the inter-tubular tissue.

At 7–8 weeks post-transplantation, there were no
significant differences between the grafted wild-type and
follistatin null testes (Fig. 3b and e). The grafted testes
had increased in volume and the seminiferous tubules
had expanded with the seminiferous epithelium showing
all types of germ cells from spermatogonia through to
mature, stage 19 spermatids. In many seminiferous
tubules from testes and from both genetic backgrounds,
there was evidence of luminal distension and disorgan-
ization of spermatogenesis with premature sloughing of
groups of post-meiotic germ cells (Fig. 3c and f). In some
tubules, only Sertoli cells could be found in the
epithelium. The loss of germ cells made assessment of
the stages of spermatogenesis difficult in both wild-type
and follistatin null testes. However, the presence of step
16 spermatids together with the appropriate basally
placed germ cells at stage VII of the cycle (Fig. 3b,
follistatin null) and the appearance of a typical stage I
(Fig. 3e, wild type) suggests that the typical cell
associations of spermatogenesis are maintained at this
ectopic site.

![Figure 1](https://example.com/fig1.png)

*Figure 1* PCR for genotyping the pups. This picture provides an example of the genotyping results. The PCR reactions were run with two pairs of primers: hHPRT.3F and hHPRT.3R; Foldel.F and Foldel.R. The PCR products from the primer pair of hHPRT.3F and hHPRT.3R were 208 bp in length, targeting the replacement cassette of the sequences in the knocked-out follistatin gene. The PCR products from the primer pair of Foldel.F and Foldel.R were 157 bp in length, targeting the sequences of the mouse follistatin gene. Thus, lanes 2, 3 and 7 represent wild-type pups, lanes 1, 5, 6, 8 and 9 represent the follistatin-deleted heterozygotes. Lanes 4 and 10 represent the follistatin knockout pups.

![Figure 2](https://example.com/fig2.png)

*Figure 2* Fetal testes that have enlarged in the external ears of the adult castrated RAG male mouse are shown (indicated by arrows).

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The percentages of seminiferous tubules containing germ cells were not significantly different between the two groups (follistatin null 85.9±3.7% vs wild type 83.7±5.1%). The number of seminiferous tubules with complete spermatogenesis also showed no differences between the two genotypic groups (follistatin null 18.6±1.3% vs wild type 19.8±1.7%).

Discussion
This study demonstrates that fetal testes from follistatin null mice can progress to complete spermatogenesis when transplanted into the external ear of castrated immunodeficient recipient mice. There were no obvious differences between the testes of the follistatin null mice and those of wild-type mice used as controls. Since the germ cells, Sertoli cells and Leydig cells of the follistatin null mice are unable to produce either follistatin 288 or 315, the results of this experiment indicate that the local production of these proteins is not critical to the completion of spermatogenesis. However, the host RAG mice are able to produce both forms of follistatin and, since follistatin 288 is predominantly tissue-bound to cell surfaces or basement membranes owing to its strong affinity for heparan sulphate proteoglycans, if follistatin is shown to be required to modulate the actions of the activins and BMPs during testicular development and spermatogenesis, then it would appear that this role can be played by the circulating form of follistatin, namely follistatin 315.

Clearly, for the grafted testes to grow 50–100-fold in volume, the blood vessels of the host RAG mice have invaded the testis and provided the necessary blood supply. These vessels are able to deliver circulating follistatin to the inter-tubular areas and, since the vascular endothelium can synthesize follistatin, this tissue could act as a local source of both forms of follistatin (Michel et al. 1996). However, as the seminiferous tubule is an avascular compartment, the follistatin can only be made available to the spermatogonia and Sertoli cells, both of which abut the basement membrane of the seminiferous tubule. Any requirement by the more luminally placed germ cells for follistatin can only be provided by transport by the Sertoli cells due to the presence of the inter-Sertoli cell tight junctions that comprise the basis of the blood–testis barrier. Gonocytes, spermatogonia and Sertoli cells, which have been shown to require follistatin to modulate the actions of activin A during testicular development, can access follistatin from the host vasculature. If follistatin plays a crucial role in the physiology of primary spermatocytes and spermatids, cells in which it has been localised (Meinhardt et al. 1998), then, in the follistatin null testis, this must be provided by transport through the Sertoli cell.

This study also demonstrates the feasibility of using the external ear as a site for successfully transplanting the testis in mice. While this technique has been successfully used in the rat (Johnson et al. 1996), this study represents the first report of this technique in mice. The successful completion of spermatogenesis at this site is probably due to the lower temperature of the pinna and its rich vascular supply.

This study also demonstrates that it is feasible to successfully generate sperm from genetically modified mice that die at birth and raise the possibility that these sperm could be used to generate homozygous mice by the use of assisted reproductive techniques. Alternatively, the successful transplantation of testes to the flank...
of nude mice as demonstrated by Honaramooz et al. (2002) may provide a technically less demanding approach.

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