Differentiation of Testis Xenografts in the Prepubertal Marmoset Depends on the Sex and Status of the Mouse Host

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This study investigates the effects of the endocrine milieu of immunodeficient mouse host (intact vs. castrated male, intact male vs. intact female) on prepubertal marmoset (Callithrix jacchus) testicular xenografts. Previous marmoset xenografting studies used castrated nude mouse hosts which did not support efficient graft survival and maturation. Due to the distinct endocrine milieu in marmosets with a deletion of exon 10 in the LH receptor, we wanted to explore whether the most efficient xenograft development occurs in intact male mouse hosts compared to intact females or castrated males. We xenografted freshly isolated tissue from prepubertal marmosets (age range 4–6 months) into the back skin of three groups of nude mice (intact male, castrated male, and intact female). We collected serum for endocrine determinations and grafts after 20 weeks and determined hormonal/reproductive status, graft survival, somatic cell development and initiation of germ cell differentiation. Graft development, tubular integrity, and germ cell differentiation status in the grafts retrieved from different hosts was scored by morphometric analysis. The influence of the different endocrine status was compared between groups of hosts. Endocrine readouts and histological endpoints in xenografts substantiate that grafts were exposed to different microenvironments and responded with host specific developmental patterns. The intact male hosts supported the most significant progression of germ cell development. Our data provide evidence for the important role of the host milieu on survival and differentiation of marmoset xenografts. The xenografting model offers innovative avenues to exploit development and endocrine effects in the primate marmoset testis using limited numbers of non-human primates for the experimental settings.

Keywords: endocrine, sex, testis, xenograft, marmoset, spermatogenesis, castration, prepubertal

INTRODUCTION

Xenografting of testicular tissue can be considered co-transplantation of germ cells with its somatic microenvironment into a host. Meiotic differentiation in xenografted fragments is supported by complete restoration of vascular supply from the host which provides optimal nutrient and oxygen supply, facilitating spermatogenic and androgenic induction in testicular fragments (1, 2). The subcutaneous transplantation technique has been successfully used as an...
experimental tool to generate sperm from testicular fragments of various species (mouse, pig, goat, sheep, hamster, bull, rhesus monkey) (1, 3–9). Regarding the potential clinical applications pre-pubertal male cancer patients undergoing chemotherapy are exposed to high dose gonadotoxic treatment depleting spermatogonial stem cells in their testes, rendering many young cancer survivors eventually infertile for life. The study gains relevance from the fact that ectopic xenografting of human prepubertal testicular tissue is currently being developed as a clinically viable fertility preservation strategy. However, an optimized strategy for host environment has yet to be established (10–15). Experimental exploration using non-human primates revealed the generation of functional sperm in ectopically grafted macaque testicular xenografts (1, 5, 8, 9, 16–18). Past marmoset and human xenografting studies investigated the effect of donor tissue (maturation status), effect of gonadotropins as well as time post grafting and site of grafting on xenograft survival and maturation (1, 12, 16, 19–22).

Previous studies investigating host effect show conflicting evidence. In several studies intact hosts were observed to efficiently support complete spermatogenesis (23–27). However, efficient xenograft survival, improved sperm production, reduced degeneration in castrated hosts were reported by others (14, 26). Castrated recipients were continued to be used as hosts, as absence of testis in the recipient allowed host gonadotropins to rise, further stimulating graft Leydig cells to produce androgens. Host castration at the time of grafting disrupted the negative feedback axis increasing FSH levels in the host. Higher gonadotropin levels in castrated recipients stimulated Sertoli cell proliferation and development of grafts until the HPG axis was re-established between the grafted tissue and the host pituitary (14).

New world monkeys are considered valuable preclinical models in male reproduction research. Common Marmosets (Callithrix jacchus) demonstrate comparable testicular development and function as observed in humans (28). This New World monkey has comparable epithelial arrangement, testicular physiology and function. Their spermatogonial progenitor system is similar to man rendering the marmoset testis a valid model for pre-clinical research to study spermatogenesis and to develop clinically viable fertility preservation strategies (29, 30, 32–35). Pubertal activation (high serum levels of testosterone) in marmosets initiates after 6 months of age (31). Asynchronous germ cell proliferation is observed during pre- and postnatal period in marmosets. Similar to man marmoset testis contains gonocytes until few weeks after birth (29) and undergoes testicular quiescence (30).

Previous marmoset xenografting studies in castrated hosts revealed, in comparison to other donor species, poor graft survival and spermatogenic induction. This was explained by the presence of a distinct endocrine system in marmosets. However, the aspect of primate tissue development and host effect was only marginally explored (1, 5, 16, 36–38). Marmosets have a mutated Luteinizing hormone (LH) receptor which lacks exon 10 making the LH receptor unresponsive to LH but responsive to CG (39–41). This mutation renders xenografted marmoset fragments non-responsive to endogenous LH released from the mouse pituitary (16), rendering this model specifically valid to observe FSH and steroid-dependent effects (39–43).

We aimed to describe the effect of host sex and status on marmoset graft survival and maturation. Preceding marmoset grafting studies, in contrast to many other species, revealed poor differentiation of testis xenografts in castrated hosts (1, 16, 36). We therefore postulated that a female or intact male host microenvironment affects marmoset graft development in different ways. In this study we explored whether graft survival, development and the degree of spermatogenic induction is dependent on the status of the host (intact vs. castrate, male vs. female).

MATERIALS AND METHODS

Ethical Approval for Animal Use

Ethical approval for the use of marmosets (LANUV-NRW 8.87–50.10.46.09.018) and mice (LANUV-NRW 84-02.04.2012.A075) was obtained according to German federal law on the care and use of laboratory animals.

Marmoset Testicular Donors and Testicular Tissue Retrieval

Testes tissue from six prepubertal marmosets (Callithrix jacchus) bred at the institutional breeding facility, aged between 4 and 6 months were used for this study. All marmosets were housed in similar conditions in groups or families under a 12 h light/12 h photoperiod and were fed a diet containing fresh food and supplements, with food pellets (Altromin, Lage, Germany). They had unrestricted access to tap water. Before testes retrieval, the monkeys were anesthetized with ketaject (equivalent to 100 mg/ml ketamine per animal) and killed by decapitation. Body weights of all marmosets were recorded and blood was collected for serum testosterone analysis. All recorded parameters are shown in Table 1. Their total testicular weight ranged between 61 and 125 mg. All 12 testes retrieved from the 6 monkeys were

| TABLE 1 | Characterization of prepubertal marmoset monkey (Callithrix jacchus) donors. |
|---------|------------------|-----------------|-----------------|-----------------|------------------|
| Callithrix jacchus | Maturity state | Age (years/months/days) | Body weight (g/kg) | Total testicular weight (mg/g) |
| MC941 | Pre-puberty | 6 m 7 d | 270 g | 125.4 mg |
| MC945 | Pre-puberty | 6 m 4 d | 263 g | 77.8 mg |
| MC947 | Pre-puberty | 5 m 28 d | 253 g | 99.7 mg |
| MC969 | Pre-puberty | 4 m 25 d | 248 g | 72.5 mg |
| MC963 | Pre-puberty | 4 m 12 d | 192 g | 61.8 mg |
| MC964 | Pre-puberty | 4 m 12 d | 175 g | 68.9 mg |

Prepubertal marmoset reference values: Age: 4–6 m, body weight: 184±13 g, total testicular weight: 68.9–125.4 mg. Spg, Spermatogonia (43, 47)
Preparation of Testicular Fragments for Xenografting

Testicular tissue was dissected into 492 fragments (63–106 fragments per monkey) of uniform size (~1 mm³). The dissected tissue fragments were systematically divided into three groups. 12 fragments (one from each testis) were processed as pre-graft controls. These were fixed in Bouin’s solution for 24 h, transferred into 70% ethanol and embedded into paraaffin for sectioning prior to histological analysis. From all 6 marmoset donors, 20 testicular fragments were subjected to grafting. To avoid any bias and maintain homogeneity, these 120 testicular fragments were pooled before grafting. From this pool, fragments were randomly selected to be grafted into 20 mouse recipients receiving six grafts each (see Figure 1 and Table 2). Remaining fragments were stored on ice-cold Leibovitz-15 medium and cryopreserved using various methods for a parallel xenografting study investigating the cryopreservation of marmoset testicular tissue.

Surgical Castration of Immuno-Deficient Nude Mice and Xenografting Experiments

In total, 35 nude mice, aged between 5 and 6 weeks were used (NMRI-Foxn1nu / Foxn1nu; Janvier) as recipients for xenografting experiment. The mice were kept at the central animal facility of the medical faculty of the University Münster. Surgical castration of ten nude mice was performed immediately prior to the xenografting procedure. In total, 15 nude mice from intact female (n = 5), intact male (n = 5) and castrated male (n = 5) host groups were used as sham-grafted controls. The remaining 20 nude mice from the three host groups of intact female (n = 5), intact male (n = 10) and castrated male hosts (n = 5) were xenografted with fresh prepubertal marmoset testicular tissue. These mice were kept in the same facility with pellet food and water ad libitum. Mice were anesthetized using ketamine xylazine and six fragments per mouse were ectopically xenografted under the back skin of nude mice using Cancer Implant Needles (G13 Popper and Sons, Staunton, VA, USA) (4). All xenografted and sham-grafted mice were kept in groups of 4–5 mice per cage and were provided pellet food and water. Body weight of all mice was recorded weekly before cage change throughout the 20-week grafting period.

Serum Testosterone Analysis

Serum testosterone in serum samples of intact and castrated male mouse hosts was determined using a double-antibody radioimmunoassay protocol (44, 45). An iodinated tracer (testosterone-3-CM-histamine) by the chloramine-T/sodium meta-bisulphite method and an antiserum raised in rabbit against testosterone-3 was used. The detection limit of the assay was 0.68 nmol/l. A serum sample (200 µl) from each mouse was used for testosterone measurements.

BrdU Administration, Vaginal Smears Collection and Xenograft Retrieval After 20 Weeks

After 20-week grafting period, xenografted testicular fragments were retrieved from all host groups. Two hours prior to graft retrieval 100 mg/kg BrdU (5-bromo-2’-deoxyuridine) was injected into each mouse, according to body weight. Vaginal smears were collected from xenografted and shamgrafted female hosts before sacrifice to evaluate their cycling state. Mice were anesthetized using ketamine xylazine and were killed by cardiac puncture. Serum was prepared and stored at −20°C. Each mouse was sacrificed by cardiac puncture. Grafts were collected from the back skin of grafted mice. Ovaries, uterus and vagina were collected from xenografted and sham-grafted female mice. Testes and seminal vesicles were collected from xenografted and sham-grafted male mice. Retrieved marmoset grafts and mice organs were weighed and subsequently fixed in Bouin’s solution for further histological analysis.

Histological Analysis

PAS Staining

Retrieved xenografts and organs from xenografted and sham-grafted male and female mice were fixed in Bouin’s solution overnight and were transferred into 70% ethanol (v/v). Pre-graft testicular fragments and retrieved samples were embedded in paraffin and 5 µm thick sections were prepared for histological analysis. Every first and tenth section of each graft as well as sections from pre-graft control fragments were stained with Period acid-Schiff (PAS) / hematoxylin (46) and analyzed for tubule integrity and morphometric characteristics. The analysis of two independent cross-sections at >50 µm distance assured a screening of independent areas during morphometric scoring.

Immuno-Histochemical Analysis of Grafts

Immuno-histochemical analysis of graft sections from different host groups was performed to identify different germ cell and somatic cell populations. Sections were deparaffinized and rehydrated followed by antigen retrieval performed in citrate buffer (pH 6). Sections were cooled to room temperature, and after tris-buffered saline (TBS) wash they were incubated with 3% (v/v) H2O2 (hydrogen peroxide) for 15 min at room temperature (RT). Non-specific binding sites in the sections were blocked by incubation in the blocking buffer containing 25% chicken/goat serum with 0.5% (w/v) BSA in TBS for 30 min. Sections were incubated overnight at 4°C in different primary antibodies including Melanoma Antigen family A 4, MAGE A4 (received from Prof. G.C. Spagnoli from the University Hospital of Basel Switzerland; dilution 1:20), Boule-like protein, BOLL (166660; Santa cruz; dilution 1:200), ACROSIN (CSF10 Bisonda; dilution 1:200), Proliferating cell nuclear antigen, PCNA (ab92552; Abcam; dilution 1:200), and SRY-BOX9, SOX9 (AB5535; Abcam; dilution 1:50). Control sections were exposed to mouse and rabbit IgG antibodies (Sigma-Aldrich; Hamburg; Germany; dilution 1:50).
Sections were incubated with chicken anti-mouse biotin (Santa Cruz; SC2985; dilution 1:100), and chicken anti-rabbit biotin (Santa Cruz; SC2986; dilution 1:100). After incubation with biotinylated secondary antibodies, sections were incubated with streptavidin horseradish peroxidase (HRP) (Sigma-Aldrich; S5512; dilution 1:500) for 45 min. Staining with chromogen 3,3′-diaminobenzidine was performed to observe protein expression patterns. Sections were subsequently counterstained with hematoxylin, dehydrated, dehydrated and mounted. Immunofluorescence stainings were performed on graft sections retrieved from male hosts for ACROSIN and CD68 (Supplementary Table 2). Stained sections were visualized using an Olympus BX61 microscope connected to a Retiga 400R camera (Olympus, Melville, NY, USA). Images were captured for analysis using Cellsens imaging software (Olympus, Melville, NY, USA).

**FIGURE 1** Summary figure illustrates the study design (including number of donor testes, hosts, and testicular fragments xenografted in each host group), endpoints analyzed (to evaluate effect of grafting on hosts and effect of hosts on graft survival and differentiation) and highlights the key outcome from the analysis.
Estimation of Female Cyclicity by Histological Analysis

Fixed uteri and ovaries from sham-grafted and xenografted female mice were sectioned and stained with Period acid-Schiff/hematoxylin to evaluate the reproductive state of the female mice and to record the presence of corpora lutea, endometrium and ongoing folliculogenesis. Giemsa staining of vaginal smears of xenografted and sham-grafted nude mice was performed to estimate cycling status and estrous stages of female mice. Slides were fixed in methanol for 5 min. Giemsa stain was performed to estimate cycling status and estrous stages of female mice. Slides were fixed in methanol for 5 min. Giemsa stain was pipetted on top of the smears and slides were dried for 10 min. After distilled water (1x) and methanol (1x) wash, slides were dried for 10 min. To evaluate spermatogenic progression, tubule-wise blinded scoring of the most advanced germ cell type present in each tubule of all the grafts retrieved from the three host groups was performed. Tubules were categorized into three groups. Tubules containing no germ cells were scored as Sertoli cell only (SCO) tubules, tubules containing pre-meiotic or meiotic/postmeiotic cells as the most advanced germ cell type were scored as 2 independent groups.

Statistical Analysis

Statistical analysis of graft retrieval percentage (among different host groups) was performed using one-way ANOVA followed by Tukey’s multiple comparison tests. Paired T-tests were used to evaluate statistically significant differences in seminal vesicle weights, graft weights, and mean scores amongst different host groups. Values were considered significantly different if P < 0.05.

RESULTS

Recipient Data

Twenty mice were xenografted from which 16 survived during the 20-week grafting period. From all mice, marmoset xenografts were retrieved. Table 2 presents the number of sham-grafted controls and recipients per host group, means of body weight and organ weights (testes, seminal vesicles, ovaries, and uterus). A statistically significant difference was observed in seminal vesicle weight of intact and castrated male hosts.

Xenograft Analysis

One hundred and twenty testicular fragments were xenografted. 71 of 120 grafts were retrieved in total and subjected to histological evaluation revealing that 32 of 71 grafts contained seminiferous tubules. Table 3 depicts absolute and relative numbers of testicular grafts retrieved from different host groups. In addition the calculated retrieval percentages of grafts as well as the weights of grafts are listed. Lowest graft retrieval was recorded in castrated hosts. A significant difference was observed in the graft weight of intact male and female hosts (Table 3).

Validation of Cellular Morphology

Germ and somatic cell populations in the pre-graft controls of all six marmosets were histologically analyzed. Histological characteristics from all the pre-graft control sections demonstrate a prepubertal status (Supplementary Figure 2).
Sharma et al. Host Effect in Testis Xenografting

FIGURE 2 | The micrographs represent typical histological features to illustrate the parameters for scoring of morphological characteristics in testicular xenografts. PAS stained sections were evaluated for tubular survival (A–D), tubular appearance (E–H), epithelial pattern of Sertoli cells (I–L) and most advanced germ cells (M–P). Blinded scoring of each graft section was performed as depicted in the table. Cellular identity was validated by immuno-histochemical staining. Cells showing intense expression for SOX9, MAGEA4, BOLL, and ACROSIN were identified, indicating the presence of Sertoli cells (I–M), spermatogonia (N), spermatocytes (O), and spermatids (P), respectively. Scale bar 20 µm. Triangles depict representative positive cells for the respective markers.

TABLE 3 | Graft retrieval data from intact and castrated immunodeficient nude mouse host.

| Host parameters | Intact males | Castrated males | Intact females |
|-----------------|--------------|-----------------|---------------|
| Grafts retrieved | 36/42        | 13/24           | 22/30         |
| Graft retrieval (%) | 85.71 (A) | 54.16 (a,b) | 73.30 (B) |
| Graft weight (mg) | 3.2 ± 1 (a) | 1.7 ± 0.7       | 1.2 ± 0.6 (A) |

Different letters indicate statistically significant differences (a, b: p < 0.05; A, B: p < 0.02).

Identical histological features were observed in all marmosets. Immuno-histochemical and immunofluorescence staining was thereafter performed to validate the presence of different germ cell populations (pre-meiotic, meiotic, and post-meiotic germ cells) in retrieved xenograft sections as shown in Figures 2N–P and Supplementary Figure 3. Cell population in representative graft sections from all three hosts groups similar to the pre-graft control sections show intense expression of MAGEA4 validating the presence of spermatogonia, as depicted in Figure 2N. Cells in grafts from intact and castrated hosts express BOLL, validating the presence of spermatocytes as illustrated in the representative image in Figure 2O. Cells in grafts recovered from male hosts show strong expression for post-meiotic marker ACROSIN, validating the presence of spermatids as shown in Figure 2P and Supplementary Figure 3. Cells expressing SOX9 were observed in testicular xenografts recovered from all three host groups validating the presence of Sertoli cells in grafts as shown in Figures 2I–L.
Comparing Effect of Hosts on Morphology and Survival of Xenografts

Xenografts retrieved from different host groups were subjected to histological analysis, evaluated morphologically and scored (as shown in Figures 2–4) to determine qualitative changes in tubular survival, immune infiltration, tubular appearance, epithelial structure, and spermatogenic state. Overall mean scores (Mean ± SEM) for tubular survival, tubular appearance, epithelial arrangement of Sertoli cells, most advanced germ cell type and immune infiltration are represented in Figures 3, 4, respectively.

As shown in Supplementary Table 1, intact male hosts demonstrate improved (32%) tubular survival compared to female (21%) and castrated hosts (17%). Grafts recovered from intact female hosts were most primitive in tubular appearance (29%) compared to intact (14%) and castrated male host groups (0%). Epithelial arrangement in most of the grafts retrieved from castrated hosts appeared partially random (67%). In contrast to intact (82%) and castrated male hosts (83%), immune cell infiltration was increased in grafts retrieved from the female hosts (92%).

Comparing the Effect of Hosts on Spermatogenic Induction

Most advanced germ cell type analysis of each graft section shows that male hosts contain grafts with the most advanced germ cells up to early spermatids (Figures 3A,B,G, Supplementary Figure 3D,F and Supplementary Table 1). Intact female hosts contain significantly fewer grafts with differentiated germ cells compared to intact males.

Most advanced germ cell analysis of each seminiferous tubule of individual sections (two sections selected 50 µm apart) from each graft was performed to determine the most advanced cell type (SCO, spermatogonia, spermatocytes, spermatids) present. In total 1033 seminiferous tubules from 32 grafts were analyzed from all 3 host groups. 232 tubules were analyzed from 6 grafts...
of castrated male hosts, 347 tubules were analyzed from 14 grafts retrieved from intact male hosts and 454 tubules were analyzed from 12 grafts retrieved from the female hosts. The highest percentage of grafts recovered from female hosts (62%) contains spermatogonia as the most advanced germ cell type. Both intact and castrated male hosts show improved meiotic (46 and 30%, respectively) and post-meiotic (14 and 20%, respectively) spermatogenic progression compared to female hosts with low meiotic (21%) and no post-meiotic (0%) progression. Grafts retrieved from intact male hosts contain the highest percentage of tubules with post-meiotic germ cells (45.5%). Grafts recovered from female hosts contain the highest percentage of tubules with pre-meiotic germ cells (62.5%). Whereas grafts retrieved from castrated hosts contain highest percentage of tubules with SCOs (32.14%) as demonstrated in **Supplementary Table 1**.

**Androgen Level Evaluation in Hosts**

Testosterone measurements recorded from the serum samples collected from different host groups were heterogeneous and no significant difference in measurement was observed amongst individual host groups. Testosterone levels in sham-grafted control recipients from all three hosts were either low or below detection limits. Testosterone levels in 4 out of 5 castrated recipients were below detection limits, the fifth recipient had very low levels of testosterone. Among intact male recipients, high testosterone levels were recorded in 2 recipients (35.8

![Image of histological features](image.png)

**FIGURE 4** | The micrographs represent typical histological features to illustrate the features for scoring of morphological characteristics in testicular xenografts. PAS stained sections were evaluated for Extreme (A), Obvious (B), Partial (C), and No (D) immune cell infiltration. Graft sections stained with CD68 antibody show different degrees of host immune cell infiltration (E-H). Scores from the blinded scoring analysis of each graft recovered from three host groups are represented as Mean ± Standard error mean (SEM) in the bar graph above (I). Scale bar 100 µm.
and 33.6 nmol/l); while the remaining mice had very low or undetectable testosterone levels. High testosterone levels were recorded in 1 out of 5 intact female recipients (24.8 nmol/l); remaining mice had either very low or undetectable testosterone levels. However, statistically significant difference was observed in freshly retrieved seminal vesicle weights. Seminal vesicle weights represent a widely used bio-indicator of androgen levels in grafting studies as these provide useful and stable readouts of the androgen status compared to the highly fluctuating serum androgen levels in mice (1). Seminal vesicle weight of intact male recipients ranged from 372(±106) to 478(±92) mg, whereas those of castrated recipients was significantly lower, ranging from 17(±2.3) to 20.4(±18) mg (Table 2).

**Effect of Grafts on Reproductive State of Female Host**

Reproductive state of shamt-grafted and xenografted female mice was evaluated by the histological analysis of female reproductive organs (uterus and ovary). Estimation of female cyclicity was performed by Giemsa staining of vaginal smears collected from female mice before sacrifice. This data demonstrates that all the xenografted female mice were at different stages of their estrous cycle similar to the sham-grafted female controls (Supplementary Figure 1). These results are in line with morphological observations of uterine and ovarian sections of the grafted mice, which show the presence of corpora lutea, endometrium, and ongoing folliculogenesis (Supplementary Figure 1).

**DISCUSSION**

The xenografting model (ectopic placement of marmoset testicular tissue into nude mice) has been previously used (1, 16). It is also a central strategy in the present study and served as a valid and innovative translational research approach since it allowed the use of limited primate testicular material to generate valid and functional outcomes on spermatogenic induction and testicular development. Xenografting of tissue from a number of species has become a widely used tool to expose immature testicular tissue to different microenvironments (1, 5, 8, 14, 16, 27, 38, 47, 48). In this context, the role of gonadotropins, steroids and age of donor and host have been explored in a number of species (10, 14). In our study it facilitated the analysis of individual testicular grafts (N = 120) established from fragments of uniform size derived from 12 immature marmoset testes. Detailed scoring of primate testicular xenografts in combination with effect of hosts of different sex and status were analyzed in our study.

The species-specific role of FSH is still under debate (49). Its role has been described as limited in rodents; however, FSH plays a more active role in primates (50). Xenografting studies are applied as elegant and donor-sparing approaches to explore the testicular FSH response in detail. In man, prepubertal testicular tissue is rarely available to research studies for ethical reasons. However, xenografting studies using fetal human tissue have been performed. As recently reviewed (51) administering exogenous gonadotropin treatment to mouse hosts revealed the stimulation of epithelial maturation and spermatogenic induction in human testicular grafts. First trimester human fetal grafts from hCG treated hosts produced lower testosterone output while second trimester human fetal xenografts produced significantly increased levels of testosterone (52). Leydig cell maturation in grafts from untreated hosts was low whereas obvious Leydig cell differentiation was observed in grafts obtained from hCG treated hosts (52). Similar xenografting studies were performed using infant and juvenile macaque testes as donor material. Confirming the results obtained in human fetal xenografting studies, exogenous gonadotropin treatment resulted in enhanced testicular graft maturation from infantile (3 m, 6 m old) and juvenile (13 m old) rhesus testes (5, 8). No significant influence of human choriongonadotropin (hCG) alone or in combination with hCG and pregnant mare serum gonadotropins (PMSG) was reported on growth of rhesus testicular xenografts (5, 8). However, significantly higher seminal vesicle weights were recorded in exogenously treated xenografted hosts, indicating active Leydig cell function in the grafts (36). Hosts treated with higher dose (10 IU) of hCG demonstrate significantly increased body weights and seminal vesicle weights compared to non-treated and lower dose (1 IU) groups (36). These studies reveal that exogenous treatment of mouse hosts with gonadotropins affects steroid release and gonadal development from primate xenografts and can therefore be used as a preclinical experimental tool (51).

Unlike rodents, old-world primates, or man, the marmoset LH receptor, due to deletion of exon 10 in the LH receptor gene, does not respond to LH but to CG-like hormones (39–43). As mentioned in the introduction, xenografts from marmosets show poor progression of testicular development (1, 16). It was proposed that the mutated LH-receptor in marmosets is causative for the lack of post-meiotic differentiation in marmoset testicular xenografts. Since mice do not express CG, the mouse host does not stimulate the marmoset LH-responsive testicular Leydig cells and therefore the testosterone release is at baseline level. However an intact endocrine milieu is sufficient for stimulating full spermatogenesis in autologous grafts at the scrotal site in marmosets (37). Even co-grafting with hamster tissue did not overcome the blockade of development (16). In contrast to macaques and man, the marmoset tissue offers a peculiar endocrine situation providing options for delineating the LH and FSH actions since LH action is diminished in the mouse host. Before performing studies with substitution of exogenous hormones to the host we were interested to learn about the effects of different host environments on marmoset xenograft development. We showed that the endocrine status of the three different hosts had different effects on graft survival and spermatogenic progression. Taking into account the dysfunctional LH-testosterone axis in marmosets our selection of hosts created three specific hormonal microenvironments for the grafts. The intact recipients generated an endogenous androgen exposure with normal mouse FSH serum levels. The castrated hosts exposed the grafts to high mouse FSH serum levels under complete absence of endogenous testosterone. The endocrine
milieu of female hosts consists of variable levels of serum 
estrogens (as the females continued cycling) in combination 
with very low testosterone under normal mouse FSH serum 
levels. Observing unaffected cyclicity in intact female hosts leads 
us to assume that intact male hosts were not affected by the 
release of androgens from the xenografts. Comparable levels 
of seminal vesicle weights in sham-grafted castrated recipients 
substantiated that the androgen release from marmoset grafts 
was low as had been shown in previous xenografting studies 
using marmosets. All these findings confirm dysfunctionality 
of the LH-receptor in marmoset testicular tissue. We assume 
that our data will help to design appropriate experimental 
schemes to explore more specifically the role of FSH in 
primates.

Nude mice are historically widely used models to explore the 
developmental aspects of tumors. They are also used for studying 
the growth and differentiation of immature tissues. Over many 
years they became standardized models for (xeno-)grafting 
studies (14). Previous reports strongly suggest that host effects 
between individual mice are negligible (1, 14, 16, 53–55). 
Testicular xenografting studies investigating the effect of host 
castration (5), male or female host (48), co-grafting (16), single 
and group-housing (54), younger and older host recipients (55) 
report identical outcome and comparable effects on individual 
mice from experimental groups (14). Even studies investigating 
the effect of irradiation in testicular xenografts at a cellular 
level show similar effects in groups of hosts. Similar doses of 
irradiation revealed identical deleterious effects on germ and 
somatic cell populations in the grafts of each group (53). Likewise 
in the current study, body weights and organ weights were 
comparable between individual hosts in specific host groups and 
in the expected normal range. There was no evidence for a high 
variability due to individual hosts. Thus, neither in the current 
study, nor in previous xenografting studies using testis tissue, a 
“host effect” of individual nude mice on testis grafts could be 
detected.

Grafting in intact hosts (male and female), lead to recovery 
rates of grafts much better than reported in castrate recipients 
(current manuscript, 1, 16). We consider therefore the intact host 
environment to be effective for marmoset testis tissue survival 
and development. We assume that the presence of steroids 
irrespective of being androgens or estrogens plays a major role 
for survival. In the absence of steroids in castrates the survival 
of xenografts was rather poor. Surprisingly the intact hosts showed 
obvious differences in graft development with male hosts being 
superior to females in promoting spermatogenic induction. We 
can only speculate that either the absence of androgens or the 
fluctuations in FSH created this difference. In future studies 
exogenous hormone delivery using various host conditions may 
be an elegant strategy to explore the actions of hormones during 
spermatogenic induction.

Grafts retrieved from intact male hosts show most advanced 
tubular survival, seminiferous epithelial arrangement, tubular, 
and luminal enlargement, least immune cell infiltration and most 
prominent progression of spermatogenesis. Indeed all previous 
studies on xenografting of immature monkey tissue used 
castrated male hosts. Castration was considered advantageous 
since the loss of endogenous testes creates a microenvironment 
with high gonadotrophic drive considered to stimulate the 
maturity of grafted tissue. Another advantage of using 
castrated hosts is that any testosterone action reflects androgen 
action of the grafts. This might also be considered valid for our 
study as we cannot separate donor and host steroids in intact 
recipients. Since mice show extreme variation in serum steroid 
levels, the size of the recipient’s seminal vesicles was selected 
as the parameter for the release of androgens and therefore 
presented an easy readout of the activity along the endocrine axis 
(3, 4).

Grafts from all three hosts showed intense expression for 
MAGEA4. The staining expression patterns were similar to 
testes from age-matched marmoset controls and indicate the 
presence of spermatogonia in all host conditions. Unlike previous 
marmoset ectopic grafting studies reporting germ cell maturation 
up to the stage of early spermatocytes (1, 4, 16, 36), ectopic 
grafts from male hosts expressed BOLL (spermatocyte marker) 
and ACROSIN (spermatid marker) revealing the presence of 
spermatocytes and occasionally spermatids in grafts from both, 
intact and castrated male mice. Post-meiotic development stops 
at the stage of early spermatids. Interestingly female hosts did not 
support post-meiotic progression. The striking host-dependent 
effects were substantiated by analysis of the most advanced 
germ cell types showing significant differences in the degree of 
post-meiotic germ cell differentiation between intact male and 
female hosts. This observation is in agreement with a study on 
newborn piglet testicular tissue revealing that poor post-meiotic 
differentiation is observed in female hosts compared to males 
(25). In mice a syngeneic testicular grafting study showed efficient 
graft survival and good post-meiotic differentiation in female 
recipients (48). Female hosts seem to be good recipients for 
xenografts in terms of graft survival. However comparison with 
marmosets is difficult as in all other species the endogenous 
mouse LH induces production of androgens in the grafts 
leading to intra-testicular rise of male steroids. For instance, 
very high androgen levels have been recorded in pig xenografts 
(7). Species-specific differences may therefore lead to highly 
variable endocrine conditions in xenografted testis tissue. The 
marmoset may be an extreme model with no LH action present 
and therefore highly reduced testosterone stimulation. This 
observation is in agreement with data from previous marmoset 
xenografting studies achieving a low androgen status in castrated 
hosts irrespective of the age of donor tissue (1, 3, and 7 months 
of age) (16).

This is the first study comparing the effects of the host 
environment via xenografting of marmoset immature testes 
into intact and castrated male and female mouse immuno-
deficient hosts. We observed changes of xenograft survival, 
testicular maturation and spermatogenic induction. Intact male 
hosts support most efficiently the survival of as well as the 
spermatogenic induction and progression in prepubertal 
marmoset testicular xenografts. The effects were put into 
context with the peculiar regulation of androgens in marmosets. 
Current findings hold clinical relevance signifying that marmoset 
monkeys could potentially be employed as a preclinical model to 
understand the hormonal control of spermatogenesis.
AUTHOR CONTRIBUTIONS

Swati Sharma performed the animal study and data analysis, compiled the manuscript. RS-K assisted with animal studies, determined hormone levels, assisted with study design. STS acted as PI during study planning and performance, and assisted with paper writing.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fendo.2018.00467/full#supplementary-material

REFERENCES

1. Schlatt S, Kim SS, Gosden R. Spermatogenesis and steroidogenesis in mouse, hamster and monkey testicular tissue after cryopreservation and heterotopic grafting to castrated hosts. Reproduction (2002) 124:339–46. doi: 10.1530/rep.0.1240339
2. Schlatt S, Westernströer B, Gassei K, Ehmcke J. Donor-host involvement in immature rat testis xenografting into nude mouse hosts. Biol Reprod. (2010) 82:888–95. doi: 10.1095/biolreprod.109.082073
3. Schlatt S, Honaramooz A, Boiani M, Schöler HR, Dobrinski I, Schlatt S. Sperm obtained from the testes of neonatal mouse testis. Biol Reprod. (2003) 68:2331–5. doi: 10.1095/biolreprod.102.014894
4. Honaramooz A, Snedaker A, Boiani M, Schöler HR, Dobrinski I, Schlatt S. Sperm from neonatal mammalian testes is grafted into nude mice. Nature (2002) 418:787–81. doi: 10.1038/nature00918
5. Honaramooz A, Li MW, Penedo MC, Meyers S, Dobrinski I. Accelerated maturation of Primate Testis by Xenografting into Mice. Biol Reprod. (2004) 70:1500–3. doi: 10.1095/biolreprod.103.025536
6. Oatley JM, Avila DM, Reeves JJ, McLean DJ. Spermatogenesis and Germ cell transgene expression in xenografted bovine testicular tissue. Biol Reprod. (2004) 71:494–501. doi: 10.1095/biolreprod.104.027953
7. Arregui L, Rathi R, Zeng W, Honaramooz A, Gomendio M, Roldan ERS, et al. Xenografting of adult mammalian testis tissue. Anim Reprod Sci. (2008) 106:65–76. doi: 10.1016/j.anireprosci.2007.03.026
8. Rath R, Zeng W, Megee S, Conley A, Meyers S, Dobrinski I. Maturation of testicular tissue from infant monkeys after xenografting into mice. Endocrinology (2008) 149:5288–96. doi: 10.1210/en.2008-0311
9. Liu Z, Nie YH, Zhang CC, Cai YJ, Wang Y, Lu HF, et al. Generation of macaques with sperm derived from juvenile monkey testicular xenografts. Nature (2015) 26:139–42. doi: 10.1038/cr.2015.112
10. Rodriguez Sosa JR, Dobrinski I. Recent developments in testis tissue xenografting. Reproduction (2009) 138:187–94. doi: 10.1530/REP.09-00102
11. Schlatt S, Ehmcke J, Jahnukainen K. Testicular stem cells for fertility preservation: Preclinical studies on male germ cell transplantation and testicular grafting. Pediatr Blood Cancer (2009) 53:274–80. doi: 10.1002/pbc.22002
12. Sato Y, Nozawa S, Yoshiike M, Arai M, Sasaki C, Iwamoto T. Xenografting of testicular tissue from an infant human donor results in accelerated testicular maturation. Hum Reprod. (2010) 25:1113–22. doi: 10.1093/humrep/deq001
13. Wynn C, Curaba M, Vanable B, Langendocnt A, Donnez J. Options for fertility preservation in prepubertal boys. Hum Reprod Update (2010) 16:312–8. doi: 10.1093/humupd/dmp054
14. Arregui L, Dobrinski I. Xenografting of testicular tissue pieces: 12 years of an in vivo spermatogenesis system. Reproduction (2014) 148:R71–84. doi: 10.1530/REP-14-0249
15. Picton HM, Wynn C, Anderson RA, Goossens E, Jahnukainen K, Klieb S, et al. A European perspective on testicular tissue cryopreservation for fertility preservation in prepubertal and adolescent boys. Hum Reprod. (2015) 30:2463–75. doi: 10.1093/humrep/dev190
16. Wistuba J, Mundry M, Luetjens CM, Schlatt S. Co-grafting of hamster (Phodopus-Sungorus) and marmoset (Callithrix-jaccus) testicular tissues into nude mice does not overcome blockade of early spermatogenic differentiation in primate grafts. Biol Reprod. (2004) 71:2087–91. doi: 10.1095/biolreprod.104.033431
17. Jahnukainen K, Ehmcke J, Hergenrother SD, Schlatt S. Effect of cold storage and cryopreservation of immature non-human primate testicular tissue on spermatogonial stem cell potential in xenografts. Hum Reprod. (2007) 22:1060–7. doi: 10.1093/humrep/del471
18. Jahnukainen K, Ehmcke J, Nurnio M, Schlatt S. Autologous ectopic grafting of cryopreserved testicular tissue preserves the fertility of prepubeasent monkeys that receive sterilizing cytotoxic therapy. Cancer Res. (2012) 72:5174–8. doi: 10.1158/0008-5472.CAN-12-1317
19. Geens M, Block GD, Goossens E, Fredericks V, Steirteghem AV, Tournaye H. Spermatogonial survival after grafting human testicular tissue to immunodeficient mice. Hum Reprod. (2006) 21:390–396. doi: 10.1093/humrep/dei412
20. Schlatt S, Honaramooz A, Ehmcke J, Goebell PJ, Rubben H, Dhir R, et al. Limited survival of adult human testicular tissue as ectopic xenograft. Hum Reprod. (2006) 21:384–9. doi: 10.1093/humrep/dei352
21. Yu J, Cai ZM, Wan HJ, Zhang FT, Ye J, Fang JZ, et al. Development of neonatal mouse and fetal human testicular tissue as ectopic grafts in immunodeficient mice. Asian J Androl. (2006) 8:393–403. doi: 10.1111/j.1745-7262.2006.00189.x
22. Wynn C, Van Langendocnt A, Wese FX, Donnez J, Curaba M. Long-term spermatogonial survival in cryopreserved and xenografted immature human testicular tissue. Hum Reprod. (2008) 23:2402–14. doi: 10.1093/humrep/den272
23. Shinohara T, Inoue K, Ogonuki N, Kanatsu-Shinohara M, Miki H, Nakata K, et al. Birth of offspring following transplantation of cryopreserved immature testicular pieces and in-vitro microinsemination. Hum Reprod. (2002) 17:3039–45. doi: 10.1093/humrep/17.12.3039
24. Huan S, Sartini BL, Parks JE. Spermatogenesis in testis xenografts grafted from pre-pubertal Holstein bulls is re-established by stem cell or early spermatogonia. Anim Reprod Sci. (2006) 103:1–12. doi: 10.1016/j.anireprosci.2006.11.018
25. Abbasi S, Honaramooz A. Effects of recipient mouse strain, sex and gonadal status on the outcome of testis tissue xenografting. Reprod Fertility Dev. (2010) 22:1279–86. doi: 10.1071/RD10084
26. Reddy N, Mahla RS, Thath R, Suman SK, Jose J, Goel S. Gonadal status of male recipient mice influences germ cell development in immature buffalo testis tissue xenograft. Reproduction (2012) 143:59–69. doi: 10.1530/REP-11-0286
27. Schlatt S, Gassei K, Westernströer B, Ehmcke J. Modulating testicular mass in xenografting: a model to explore testis development and
endocrine function. Endocrinology (2010) 151:4018–23. doi: 10.1210/en.2010-0-0415
28. McKinnell C, Mitchell RT, Morris K, Anderson RA, Kelnar CJH, Wallace WH, et al. Perinatal germ cell development and differentiation in the male marmoset (Callithrix jacchus): similarities with the human and differences from the rat. Hum Reprod. (2013) 28:886–96. doi: 10.1093/humrep/des465
29. Mitchell RT, Cowan G, Morris KD, Anderson RA, Fraser HM, McKenzrie KJ, et al. Germ cell differentiation in the marmoset (Callithrix jacchus) fetal and neonatal lane closely parallels that in the human. Hum Reprod. (2008) 23:2755–65. doi: 10.1093/humrep/den295
30. Kelnar CJ, McKinnell C, Walker M, Morris KD, Wallace WH, Saunders PT, et al. Testicular changes during infantile ‘quiescence’ in the marmoset and their gonadotrophin dependence: a model for investigating susceptibility of the prepubertal human testis to cancer therapy? Hum Reprod. (2002) 17:1367–78. doi: 10.1093/humrep/17.5.1367
31. Irfan S, Wistuba J, Ehmcke J, Shahab M, Schlatt S. Pubertal and testicular development in the common marmoset (Callithrix jacchus) shows high individual variation. Prim Biol. (2015) 2:1–8. doi: 10.5194/pb-2-1-2015
32. Sharpe RM, Fraser HM, Broughton MF, McKinnell C, Morris KD, Kelnar CJ, et al. Role of the neonatal period of pituitary-testicular activity in germ cell proliferation and differentiation in the primate testis. Hum Reprod. (2003) 18:2110–7. doi: 10.1093/humrep/deg413
33. Li J-H, Donald JM, Golub MS. Testicular development, structure, function and regulation in common marmoset. Birth Defects Res. (2005) 74:450–69. doi: 10.1002/bdr.20057
34. McKinnell C, Mitchell RT, Walker M, Morris K, Kelnar CJ, Wallace WH, et al. Effect of fetal or neonatal exposure to monobutyl phthalate (MBP) on testicular development and function in the marmoset. Hum Reprod. (2009) 24:2244–54. doi: 10.1093/humrep/dep200
35. Mitchell RT, Saunders PT, Sharpe RM, Kelnar CJ, Wallace WH. Male fertility and strategies for fertility preservation following childhood cancer treatment. Endocr Dev. (2009) 15:101–34. doi: 10.1159/000207612
36. Wistuba J, Luetjens CM, Wesselman R, Nieschlag E, Simoni M, Schlatt S. Meiosis in autologous ectopic transplants of immature testicular tissue grafted to Callithrix jacchus. Biol Reprod. (2006) 74:706–13. doi: 10.1095/biolreprod.105.048793
37. Luetjens CM, Stukenborg JB, Nieschlag E, Simoni M, Wistuba J. Complete spermatogenesis in orthotopic but not in ectopic transplants of autologously grafted marmoset testicular tissue. Endocrinology (2008) 149:1736–47. doi: 10.1210/en.2007-1325
38. Ehmcke J, Gassei K, Westernströer B, Schlatt S. Immature rhesus monkey (Macaca mulatta) testis xenografts show increased growth, but not enhanced seminiferous differentiation, under human chorionic gonadotropin treatment of nude mouse recipients. Int J Androl. (2011) 34:e459–67. doi: 10.1111/j.1365-2605.2011.01179.x
39. Gromoll J, Wistuba J, Terwort N, Godmann M, Muller T, Simoni M. The carboxyterminal peptide of chorionic gonadotropin facilitates activation of the marmoset LH receptor. Exp Clin Endocrinol Diabetes (2009) 112:574–9. doi: 10.1055/s-2004-830409
40. Muller T, Simoni M, Pekel E, Luetjens CM, Chandolia R, Amato F, et al. Chorionic Gonadotrophin beta subunit mRNA but not luteinizing hormone beta subunit mRNA is expressed in the pituitary of the common marmoset (Callithrix jacchus). J Mol Endocrinol. (2004) 32:115–28. doi: 10.1677/jme.0.032115
41. Müller T, Gromoll J, Simula AP, Norman R, Sandhowe-Klaverkamp R, Simoni M. The carboxyterminal peptide of chorionic gonadotropin facilitates activation of the marmoset LH receptor. Exp Clin Endocrinol Diabetes (2009) 112:574–9. doi: 10.1055/s-2004-830409
42. Tannenbaum PL, Schultz-Darken NJ, Woller MJ, Abbott DH. Gonadotrophin-releasing hormone (GnRH) release in marmosets II: pulsatile release of GnRH and pituitary gonadotrophin in adult females. J Neuroendocrinol. (2007) 19:354–63. doi: 10.1111/j.1365-2826.2007.01535.x
43. Henke A, Luetjens CM, Simoni M, Gromoll J, Chorionic Gonadotropin beta-subunit gene expression in the Marmoset pituitary is controlled by steroidogenic factor 1, early growth response protein 1, and pituitary homeobox factor 1. Endocrinology (2007) 148:6062–72. doi: 10.1210/en.2007-0825
44. Chandolia RK, Weinbauer GF, Simoni M, Behre HM, Nieschlag E. Comparative effects of chronic administration of the non-steroidal androgens flutamide and casodex on the reproductive system of the adult male rat. Acta Endocrinologica (1991) 125:547–55. doi: 10.1530/acta.0.1250547
45. Chandolia RK, Luetjens CM, Wistuba J, Yeung C-H, Nieschlag E, Simoni M. Changes in endocrine profile and reproductive organs during puberty in the male marmoset monkey (Callithrix jacchus). Reproduction (2006) 132:355–63. doi: 10.1530/rep.1.01186
46. Brinkworth MH, Weinbauer GF, Schlatt S, Nieschlag E. Identification of male germ cells undergoing apoptosis in adult rats. J Reprod Fertility (1995) 105:23–33. doi: 10.1530/jrf.0.1050025
47. Spade DJ, Hall SJ, Safferini CM, Huse SM, McDonnell EV, Boekelheide K. Differential response to abiraterone acetate and di-n-butyl phthalate in an androgen-sensitive human fetal testis xenograft bioassay. Toxicol Sci. (2014) 138:148–60. doi: 10.1093/toxsci/kft266
48. Ma P, Ge Y, Wang S, Ma J, Xue S, Han D. Spermatogenesis following syngeneic testicular transplantation in Balb/c mice. Reproduction (2004) 128:163–70. doi: 10.1530/rep.1.00165
49. Schlatt S, Ehmcke J. Regulation of spermatogenesis: an evolutionary biologist’s perspective. Semin Cell Dev Biol. (2014) 29:2–16. doi: 10.1016/j.semcdb.2013.04.007
50. Plant TM, Marshall GR. The functional significance of FSH in spermatogenesis and the control of its secretion in male primates. Endocr Rev. (2001) 22:764–86. doi: 10.1210/edrv.22.6.0446
51. Hutka M, Smith LB, Mitchell RT. Xenotransplantation as a model for human testicular development. Differentiation (2017) 97:44–53. doi: 10.1016/j.diff.2017.09.001
52. Mitchell RT, Saunders PTK, Childs AJ, Cassidy-Kojima C, Anderson RA, Wallace WHR, et al. Xenografting of human fetal testis tissue: a new approach to study fetal testis development and germ cell differentiation. Hum Reprod. (2010) 25:2405–14. doi: 10.1093/humrep/dep183
53. Tröndle I, Westernströer B, Wistuba J, Terwort N, Schlatt S, Neuhaus N. Irradiation affects germ and somatic cells in prepubertal monkey testis xenografts. Mol Hum Reprod. (2017) 23:141–54. doi: 10.1093/molehr/gax003
54. Arregui L, Rath J, Megee SO, Honaramoos A, Gomendio M, Roldan E, et al. Xenografting of sheep testis tissue: a model for preservation of genetic material from endangered ungulates. Reproduction (2008) 136:85–93. doi: 10.1530/REP-07-0433
55. Ehmcke J, Gassei K, Schlatt S. Ectopic testicular xenografts from newborn hamsters (Phodopus sungorus) show better spermatogenic activity in aged compared with young recipients. J Exp Zool Part A (2008) 309A:278. doi: 10.1002/jez.459

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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