Leydig cell function in mice lacking connexin43

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Abstract

Connexin43 (Cx43) is the most abundantly expressed member of the connexin (gap junction protein) family and the only one so far identified in mouse Leydig cell gap junctions. Mice lacking Cx43 were used to investigate its role in testicular androgen production and regulation. Testes from term fetuses were grafted under the kidney capsules of castrated adult males. After 3 weeks, serum from host mice was analyzed for androgens. In order to test their response to stimulation, the grafted testes were incubated in vitro with varying concentrations of LH and their androgen end products analyzed. Incubation with radiolabeled progesterone was followed by high performance liquid chromatography to quantify the androgen-intermediate metabolites. Radiolabeled testosterone in the presence of NADPH was used to determine the activity of testosterone-metabolizing enzymes 17β-hydroxysteroid dehydrogenase (17βHSD), 5α-reductase (5αR), and 3α-hydroxysteroid dehydrogenase (3α HSD). Serum androgen levels did not differ between hosts carrying wild-type versus null mutant grafts although Cx43-deficient testes had more 17βHSD and 5αR activity than wild-type controls. Furthermore, the genotype of grafted testes did not influence LH-stimulated androgen production in vitro. These results indicate that the steroidogenic function of Leydig cells is not compromised by the absence of Cx43, perhaps because other gap junction proteins are present. Dye transfer experiments demonstrated that Cx43-deficient Leydig cells retain intercellular coupling, indicating that Cx43 is not the only protein contributing to their gap junctions. Thus, despite their prominence in Leydig cells, Cx43 gap junctions are not essential for androgen production.

Introduction

Cell–cell communication mediated by gap junctions plays important roles in development and physiological function, including that of endocrine organs (Levin 2001, Serre-Beinier et al. 2002). Gap junctions are specialized regions of contact between cells where intercellular membrane channels are concentrated. Each gap junction channel is composed of two hemichannels or connexons that dock up to make a cylindrical channel, linking the cytosol of one cell with the cytosol of a neighboring cell. Each connexon is formed by the oligomerization of six protein subunits known as connexins; to date, 19 of these have been annotated in the mouse, each encoded by a distinct gene (Söhl & Willecke 2003). Gap-junctional intercellular coupling allows cells to share low-molecular mass metabolites and second messengers that are <1000 Da, thus facilitating homeostatic and developmental processes (Harris 2001). The focus of the present study is gap junctions in the testes.

Connexin43 (Cx43) is the most widely distributed connexin in the body and is expressed very early in development (Ruangvoravat & Lo 1992, Yancey et al. 1992, De Sousa et al. 1993). It is the most abundant gap junction protein in the mouse testis, and so far the only one identified in Leydig cells (Perez-Armendariz et al. 1994, Varanda & de Carvalho 1994, Bravo-Moreno et al. 2001, Risley et al. 2002). Deletion of the Gja1 gene that encodes Cx43 causes a lethal cardiac malformation: mutant fetuses develop to term but die of asphyxiation soon after birth (Reaume et al. 1995). To study postnatal testis development in the absence of this connexin, previous work utilized a graft procedure that allows gonad development to continue under the kidney capsule of an adult mouse (Kuopio et al. 1989, Roscoe et al. 2001). Grafted testes from wild-type fetuses exhibited normal development for up to 3 weeks postnatal, but those lacking Cx43 suffered loss of spermatogonia, resulting in a ‘Sertoli cell-only’ condition in the seminiferous tubules (Roscoe et al. 2001). Electron microscopical analysis of the mutant testes...
revealed cells with normal Leydig cell morphology. Furthermore, preliminary experiments indicated that luteinizing hormone (LH)-stimulated androgen production still occurred in the mutant testes, although no attempt was made to assess the cells’ sensitivity to stimulation or the spectrum of androgens being produced.

Cx43 is also prominently expressed in the Leydig cells of human testes (Steger et al. 1999), making it a target of investigations aimed at understanding male infertility. Given the important roles played by Cx43 in the development and function of endocrine organs (Serre-Beinier et al. 2002, Klee et al. 2005), and the profound effect of its loss on spermatogenesis in the mouse (Roscoe et al. 2001), we considered it of interest to examine Cx43 null mutant Leydig cells in more detail to determine if their function had been altered.

Materials and Methods

Mice

Homozygous Cx43 knockout mice were obtained from matings of mice heterozygous for the mutant allele (Gja1<sup>+/−</sup>/Gja1<sup>−/−</sup>) on a CD1 background. Adult mice used as hosts for the grafted testes were Pkrdc<sup>−/−</sup>/Pkrdc<sup>−/−</sup> males (CB17/Icr Hsd<sup>1scid</sup>; Harlan Sprague–Dawley, Indianapolis, IN, USA) that weighed 20–22 g. The mice were bred in the Department of Animal Care and Veterinary Services at the University of Western Ontario and at the Robarts Research Institute, London, ON, Canada. They were maintained and handled in accordance with the principles set forth in the Guide for Care and Use of Laboratory Animals of the Institute for Laboratory Animal Research of the National Academy of Sciences.

Collection and grafting of fetal testes

On day 17.5 of gestation, the pregnant females were anesthetized with CO<sub>2</sub> and then killed by cervical dislocation. Fetuses were carefully dissected from the uteri, killed by decapitation, and fetal tail snips obtained.

These were used for genotyping by PCR. Each tail snip was digested at 58 °C for 8–24 h using 100 μl proteinase K (Invitrogen) in buffered solution. The digested mixture was diluted in the ratio of 1:10 with Nanopure water and heated at 95 °C for 20 min to inactivate the proteinase K. For each PCR, 1 μl of each digest was used as a template. Details of the primers, reagents, and amplification conditions were as described previously (Roscoe et al. 2001).

The testes from each fetus were removed and placed in 0.5 ml M199, pH 7.4 (Invitrogen) in a Falcon Petri dish and excess tissue removed. The testes were transferred to a six-well Millicell culture plate with isopore polycarbonate membranes, 3.0 μm pore size (Millipore Canada Ltd, Nepean, ON, Canada) over M199. They were cultured at 37 °C in 5% CO<sub>2</sub> and were used for grafting within 2 days.

Cx43 null mutant and wild-type testes were grafted as described previously (Roscoe et al. 2001). Briefly, the host animals were anesthetized with 0.04 ml per 10 g body weight of 2.5% ketamine hydrochloride (Ketaset, Ayerst Veterinary Laboratories, Guelph, ON, Canada) and 0.2% xylazine (Rompun, Bayer). The region to be incised was prepared, a lateral incision was made on the right dorsal area at the position of the kidney and the right kidney carefully exteriorized. While keeping the kidney moist at all times during the surgical procedure, a small puncture was made in the capsule and was used as an insertion point of the fetal testis. The two testes from the same fetus were grafted together but well separated to prevent them from merging and growing as one. The kidney was then sutured back into its abdominal position and the incision sutured appropriately.

Gonadectomy

For most experiments, the host male mice were left intact (i.e. not castrated) to avoid high circulating LH levels that would cause receptor desensitization. When assaying host serum for androgens, however, the host males were castrated to eliminate the contribution from the host testes. At 1 week post-grafting, the host males were anesthetized, shaved, and cleaned. The procedure was done through a small incision in the lower abdomen. The testes and the adherent fat tissue, easily seen through the aperture, were carefully exteriorized. After clamping and ligating the spermatic artery to minimize bleeding, each testis was detached and the remaining tissue returned to its abdominal location. At the same incision point, the procedure was repeated for the other testis. The incision was then sutured appropriately. After the surgical procedure, the mice were given buprenorphine analgesic (0.3 mg/ml, from Rickitt and Colman products, Hull, England, diluted in the ratio of 1:1000 with sterile water) at a dose of 0.018 ml/g body weight.

Harvesting blood and grafted testes

After 3 weeks of engraftment, the host mice were anesthetized and killed by cervical dislocation. About 800 μl blood from each mouse was obtained from the heart by cardiac puncture. This was left to clot and centrifuged at 2000 g for 5 min to give ~150 μl serum per sample, which was used for quantification of circulating androgens by RIA. Grafted testes were retrieved by peeling them from the kidney capsule. Any adhering tissue was detached and the testes immediately placed in M199 medium. The testes were either used immediately for in vitro experimentation or fixed for histological procedures.
Leydig cell counts

To compare Leydig cell numbers between mutant and wild-type testes, histological sections were immunostained with a polyclonal antibody raised in rabbits against human placental 3β-hydroxysteroid dehydrogenase (3βHSD), generously supplied by Dr Ian Mason, University of Edinburgh. This antibody was previously shown to be specific for Leydig cells in fetal mouse testes (Doody et al. 1990). A horseradish peroxidase (HRP)- or fluorescein-conjugated secondary antibody was used to detect bound primary antibody. A standard immunostaining procedure was used. Briefly, recovered grafts were fixed in Bouin’s fixative overnight and then transferred to 70% ethanol until needed. The testes were embedded in paraffin and sectioned at a thickness of 5 µm. The tissue sections were then deparaffinized using xylene and absolute ethanol. Slides in which an HRP-conjugated secondary antibody was to be used were then placed in 0.3% H2O2 in methanol for 30 min to remove any endogenous peroxidase activity. All slides were rehydrated by placing them in decreasing concentrations of ethanol before washing in PBS. Immunostaining was preceded by treatment of the hydrated slides for 45 min with blocking solution, 2% BSA (Life Technologies) in PBS, which was also used to dilute the primary and the secondary antibodies. The primary 3βHSD antibody (diluted in the ratio of 1:50) was placed on each slide at 4 °C overnight. Slides were then rinsed with PBS for 5 min before applying secondary antibody for 1 h. A goat anti-rabbit secondary antibody conjugated to Alexaflour 488 (1:1000 dilution) was used for fluorescent detection and the nuclei were counterstained with Hoechst 33258 (both from Molecular Probes, Eugene, OR, USA). The slides were mounted with AIRVOL mounting medium (Airproducts and Chemicals, Allentown, PA, USA), sealed with nail polish, and observed immediately. For HRP detection, the slides were processed using a Vectastain ABC kit (Vector Laboratories, Burlingame, CA, USA) according to the manufacturers’ instructions with an additional step of inhibiting endogenous peroxidase activity. HRP-conjugated goat anti-rabbit IgG (Vector Laboratories), diluted in the ratio of 1:250 and applied for 1 h at room temperature, was used as a secondary antibody. After washing with PBS and incubation with Vectastain for 30 min, 3,3′-diaminobenzidine was used to visualize the signal (FAST DAB tablets, Sigma-Aldrich, Oakville, ON, Canada). The slides were counterstained with hematoxylin and dehydrated through increasing concentrations of ethanol before mounting with Permount.

Leydig cells were quantified by counting the number of 3βHSD-immunostained cells in each testis section, taking care to avoid sections in which the seminiferous tubules had been cut obliquely. The number of tubules and Leydig cells was determined for the entire testicular cross-sections. The number of Leydig cells per tubule was compared with respect to genotype.

In vitro culture of testes

In addition to grafted wild-type and mutant testes obtained from late gestation fetuses, testes from adult mice were used to validate the assays. Grafted testes were retrieved at 3 weeks post-grafting. Each testis was decapsulated and teased out to increase perfusion of culture medium into the cells before placing it in an individual well of a Millicell culture plate. Incubation was done in M199 medium buffered with 0.1% HEPES at pH 7.4 (or pH 5 for measurement of type II 5α-hydroxysteroid dehydrogenase), 37 °C, and 5% CO2 and was conducted in medium alone (control) or medium plus ovine LH (NIH-LH26). The distribution of androgen metabolites produced in vitro was monitored by incubating the testes with [3H]progesterone and the enzyme activities were measured by incubating with [3H]testosterone (details below). The substrate concentrations for each enzyme were maximal to ensure that the substrate was not rate limiting. The reaction was started by addition of the testes and terminated by placing the incubation on ice. The medium was harvested immediately after centrifugation at 2000 g for 5 min and stored at −20 °C until required for assay. The testes were blotted, weighed, and stored at −70 °C for subsequent protein assay.

The sensitivity of the Leydig cells to LH was measured by subjecting the teased out, decapsulated grafted testes from non-castrated hosts to varying concentrations of LH (25–400 pg/ml) in M199 medium, pH 7.4. They were incubated for 4 h as described previously. One grafted testis from each animal was incubated in medium alone and served as the control, while the other graft was incubated in the presence of LH. RIA was performed on the harvested medium to determine androgen accumulation in response to LH.

Radioimmunoassays

Concentrations of circulating androgens in host blood and in culture medium were determined by RIA as described previously (Jansz & Pomerantz 1985, Roscoe et al. 2001). The anti-testosterone antibody we used cross-reacts 98% with dihydrotestosterone, 58% with 5α-androstane-3α,17β-diol, 24% with 5α-androstane-3β,17β-diol, and negligibly (<1.6%) with other androgens and estradiol (Jansz & Pomerantz 1985). Diethyl ether (2 × 2 ml) was used for extraction of steroids from the blood serum (60 µl). One hundred microliters (calculated to contain 6 pg) radiolabeled testosterone [1,2,6,7-3H(N)]-testosterone, specific activity 95.0 Ci/mmol (NEN Life Science Products, Boston, MA, USA) were added to the serum before extraction and used to
calculate steroid recovery from the samples. This was calculated at 78–82%. After drying, the extracted steroids were resuspended in 60 µl PBS with 0.1% gelatin (G-PBS) and used in duplicate RIA. The inter- and intra-assay coefficients of variation were 11 and 8% respectively.

### Androgen metabolite and steroidogenic enzyme assays

Conversion of the radioactive substrate [1,2,6,7,16,17\(^3\)H]-pregesterone, specific activity 93.0 Ci/mmol (NEN Life Science Products), to various androgen intermediate metabolites was determined in their profile. One hundred microliters (calculated to contain 0.5 µCi) radiolabeled progesterone were added to the incubation medium. Incubation was carried out for 4 h in the presence of a maximally stimulating LH dose (400 pg/ml) and a co-factor, 2 mM NADPH (Sigma-Aldrich Canada Ltd) were added to the incubation medium. The activity of 5\(\alpha\)-androstane-3\(\alpha\),17\(\beta\)-dihydrogenase or pH 7.4. A maximally stimulating LH dose (400 pg/ml) and a co-factor, 2 mM NADPH (Sigma-Aldrich Canada Ltd) were added to the medium. The activity of 17\(\beta\)HSD was determined by measuring conversion of [\(^3\)H]-testosterone (0.5 µCi) to [\(^3\)H]-androstenedione after 2 h of incubation at pH 7.4. Extraction and separation procedures were as described (Khalil & Walton 1985).

Standard HPLC was used to identify and measure the incubation end products (Khalil & Walton 1985). Steroids in the culture medium (0.5 ml) were concentrated on C\(_{18}\) reverse-phase cartridges (Phenomenex, Torrance, CA, USA) and the steroid fraction recovered with methanol (5 ml). A portion (1–2 ml) of the methanol solution was dried down in a steady stream of nitrogen, and the dried down extract redissolved in 250 µl mobile phase, 55% acetonitrile/water. Aliquots (200 µl) of each sample were analyzed on a C\(_{18}\) reverse-phase column (Bondclone, 300×3.9 mm) with 10 µm beads eluted with 55% acetonitrile/water for 30 min at a flow rate of 1 ml/min. Radiolabeled metabolites were detected using an on-line radioactivity detector and identified on the basis of their retention times compared with authentic standards, and quantified using [1,2,6,7\(^3\)H]-androstenedione as external standard (Khalil & Walton 1985). HPLC equipment consisted of a model 600A pump, model 710B automatic injectors, and a model 441 u.v. spectrophotometer, all from Waters Associates, Mississauga, ON, Canada. Radiolabeled steroid metabolites were detected and quantified by an in-line Beckman model 171 radioactivity detector (Beckman Instruments, Toronto, ON, Canada). Data from the UV and the radioactivity detectors were analyzed by computer using the MAXIMA 820 software package from Waters Associates.

### Leydig cell isolation

Leydig cell preparations were obtained as described previously with some modifications (Doody et al. 1990). Briefly, the grafted testes were retrieved from the host mice, decapsulated, and incubated with dissociation buffer at 36 °C for 15 min with agitation. The dissociation buffer consisted of M199 medium containing 0.36 mg/ml collagenase (type 1; Sigma) and 0.012 mg/ml DNase (type 1; Sigma). The enzymatic activity was terminated by placing the incubation tubes on ice and adding excess cold (4 °C) culture medium (three volumes). The seminiferous tubules were allowed to settle by gravity for 5 min and the medium, containing interstitial cells, aspirated and filtered through a 70 µm nylon mesh. The supernatant containing the interstitial cells was removed and centrifuged at 250 g. The cells were washed twice and plated in 35×10 mm tissue culture dishes (Falcon Plastics, Los Angeles, CA, USA) as 0.2 ml aliquots in culture medium supplemented with 10% fetal bovine serum. The cells were cultured overnight at 37 °C in a humidified atmosphere of 5% CO\(_2\) and 95% air to maintain the medium at pH 7.4. The next day, the medium was discarded and the cells were washed with culture medium to remove loosely attached cells.

To confirm the identity of the isolated cells as Leydig cells, the cells were immunostained for 3\(\beta\)HSD. Briefly, the plated cells were fixed for 20 min with 4% paraformaldehyde containing 0.1% glutaraldehyde. The cells were blocked with 2% BSA in PBS, treated with primary antibody (1:50), washed with PBS, and then treated with secondary antibody (1:1000 dilution of fluorescein-conjugated goat anti-rabbit IgG from ICN Biomedicals, Irvine, CA, USA). Hoechst 33342 was again used as a nuclear stain as described previously. The cells were mounted in Airvol, observed under phase contrast and fluorescence microscopy using a Zeiss (Thornwood, NY, USA) LSM 510 META confocal microscope mounted on an inverted Axiosvert 200 motorized microscope, and photographed. Digital images were prepared using Zeiss LSM and Adobe Photoshop 7.0 software. The images were collected with a charge-coupled device (CCD) camera (Hamamatsu Photonics, Hamamatsu City, Japan) using OpenLab software (distributed by Quorum Technologies, Inc., Guelph, ON, Canada). All but a small fraction of the cells (<10%) were positive for 3\(\beta\)HSD and thus the vast majority was considered to be Leydig cells. These cells were used for dye coupling assays.
**Dye coupling assay**

Dye transfer assays were carried out after the Leydig cells had been cultured for 24 h. Three milliliters of warm culture medium (37 °C) were added to each culture dish. A gap junction-permeant fluorescent dye, Lucifer yellow (LY; Molecular Probes) was backfilled via capillary action through a 1 mm thin-wall glass capillary (World Precision Instruments, Sarasota, FL, USA) pulled to 1 μm. The concentration of LY was 5% in ddH2O. This was injected into a single Leydig cell and any transfer of dye to adjacent cells was observed after 2 min. Fluorescent signals were imaged on a Zeiss inverted microscope equipped with a Sensicam CCD camera.

**Statistics**

Student’s t-test and one-way ANOVA were followed by parametric or non-parametric tests to determine the differences between the groups. The Kolmogorov–Smirnov goodness-of-fit test was used to determine the need for non-parametric tests. The Mann–Whitney test was applied to determine difference due to genotype when percentage change was assessed. Significant difference between the groups was inferred at $P<0.05$.

**Results**

**Leydig cell numbers**

Immunostaining cross-sections of 3 week testis grafts for 3βHSD revealed that the Leydig cells in Cx43-deficient testes have an altered distribution, they are less closely clustered than those in wild-type testes (Fig. 1A and B). Although this was not quantified, it was sufficiently obvious that a blinded observer could identify the mutant testes from immunostained tissue sections. However, counts of Leydig cells (C) and seminiferous tubules (D) did not reveal any significant difference between the genotypes and, consequently, the number of Leydig cells did not differ when normalized to the number of tubules in each cross-section (Mann–Whitney test). The same result was obtained when 18.5 dpc fetal testes were examined ($n=8$ for each genotype, data not shown), although the difference in Leydig cell clustering was not evident at that stage. Thus, although the mutant testes are generally smaller due to the paucity of developing germ cells (Roscoe et al. 2001), the total number of Leydig cells was not affected by the mutation, either in late fetal or in juvenile stages. In both genotypes, the number of Leydig cells per seminiferous tubule did not change ($P>0.05$, Student’s t-test) during the 3-week grafting period.

**Serum androgen levels**

In the previous study of Cx43-deficient testes (Roscoe et al. 2001), incubation of grafted testes in vitro with a supraphysiological level of LH or cAMP revealed no difference in androgen production between mutant and wild-type grafts. To confirm this in vivo, we examined androgen production by mutant and wild-type testes in the graft hosts. Serum androgen concentrations were determined after 3 weeks of postnatal development in mice castrated 1 week after the grafting surgery (Fig. 2). Preliminary experiments verified that the 2 week period between castration of the hosts and collection of the serum samples was sufficient for the androgens produced by the host testes to have been eliminated (data not shown). The results indicated that the androgen output of

![Image](https://www.reproduction-online.org/Reproduction%20(2006)%20132%20607–616%20Downloaded%20from%20Bioscientifica.com%20at%2009/03/2018%2007:29:16PM%20via%20free%20access)
the testis grafts was very low; the circulating androgen levels of the castrated, graft-bearing hosts were less than 30% of the level (2.15 ± 0.66 ng/ml) measured in intact, ungrafted males. Importantly, androgen levels in the graft-bearing hosts did not differ between graft genotypes (Student's t-test). Given that the two genotypes do not differ in Leydig cell numbers (Fig. 1), the results indicate that the ability of the Leydig cells to produce testosterone was not altered by the loss of Cx43.

**LH sensitivity**

Given that gap-junctional intercellular communication has been shown to be involved in cellular responses to extracellular stimuli in several instances (Serre-Beinier et al. 2002), we wanted to determine whether the LH sensitivity of the Leydig cells had been altered by the loss of Cx43. On the contrary, Fig. 3 shows that the response of grafted testes to LH concentrations ranging between 25 and 400 pg/ml was normal; there was a significant interaction between LH concentration and androgen output for both genotypes (Mann–Whitney test), but there was no significant difference between the genotypes for all the LH concentrations tested (Kruskal–Wallis test). Androgen output of wild-type adult testes was measured to provide a standard of comparison. The bars represent mean ± S.E.M.; the number of testes is indicated by the number in parentheses above each bar.

In order to determine if there was a change in secreted androgen products or testicular steroidogenic enzyme activities caused by the absence of Cx43, the end products of incubation were analyzed. HPLC was used to identify and quantify the end products released by mutant and wild-type grafted testes in vitro. Figure 4 shows that the predominant product from the grafted testes was androstenedione (HPLC retention time RT = 8.77), with no quantitative difference between the mutant and the wild-type testes. Testosterone (RT = 7.3) was also produced by all the testes, although in lower amounts. Other intermediate metabolites were either barely detectable and/or did not show consistency in the different samples, and are therefore not represented in the results. The metabolite distributions from the grafted testes were similar to the end products of 3 week-old juvenile males, although the ratio between androstenedione and testosterone was greater in the latter (Fig. 4).

Although the amounts differ, the metabolite distributions affirm that the grafted testes functioned as well as non-grafted testes of an equivalent age in terms of steroidogenesis, regardless of the genotype.

**Androgen end products**

The androgen end products released into culture medium from incubated rat or mouse testes are dependent on the stage of testicular development (Sheffield & O'Shaughnessy 1988, Ge & Hardy 1998).

**Steroidogenic enzyme activities**

Stage-dependent expression and activity of various enzymes involved in androgen biosynthesis and metabolism are implicated in causing variation in the major
end products from developing testes (Ge & Hardy 1998, O’Shaughnessy et al. 2002). There is transient expression and activity of isozymes of 17β-hydroxysteroid dehydrogenase (17βHSD), 5α-reductase (5αR), and 3α-hydroxysteroid dehydrogenase (3αHSD) involved in the metabolism of androstenedione to testosterone and the 5α-reduced testosterone (Shan et al. 1993, Ge & Hardy 1998, O’Shaughnessy et al. 2002). Figure 5 summarizes activity measurements for these enzymes in the grafted testes. The 3αHSD activity was detected at very low levels in wild-type testes but not in Cx43-deficient testes. In contrast, the Cx43-deficient testes exhibited significantly greater 17βHSD and 5αR activity. It was observed that the overall output of androgens from the grafted testes was very low. The results suggest that the grafts had not yet achieved biosynthetic characteristics of mature adult Leydig cells.

**Leydig cell dye coupling**

The absence of any noticeable effect of Cx43 deficiency on Leydig cell function could be interpreted in either of the two ways: that gap-junctional communication is not required for Leydig cell steroidogenesis, or that Leydig cells express one or more gap junction proteins in addition to Cx43 that can compensate for its absence. To explore the latter possibility, Leydig cells were isolated from mutant and wild-type grafted testes and their capacity to couple metabolically was assayed using a standard dye transfer test. To verify the identity of the isolated cells, they were immunostained for 3βHSD (Fig. 6). Virtually all of the cells in cultures of wild-type (Fig. 6A and B) or mutant (Fig. 6C and D) preparations stained positively for the enzyme. For cells of both genotypes, Lucifer yellow dye injection resulted in passage of the dye from the injected cell to its neighbors, indicating that Leydig cells lacking Cx43 retain gap-junctional communication (Fig. 6G and H, Table 1).

**Discussion**

Previous work involving the grafting of Cx43-deficient testes under the kidney capsules of adult mice revealed that spermatogenesis is severely compromised by the loss of this connexin (Roscoe et al. 2001). However, that study focused on gonad size, germ cell numbers, and germ cell development, showing that spermatogonia are reduced in numbers at birth and are not able to repopulate the seminiferous tubules during postnatal development. The present work examined steroidogenesis in the mutant testes and determined that the absence of Cx43, while reducing the clustering of Leydig cells, does not affect their numbers or impair their functional activity.

In addition to failing to find any effect of Cx43 absence on Leydig cell number, our data indicated that those numbers are constant between late fetal and 3-week juvenile stages. As indicated in previous research (Baker & O’Shaughnessy 2001), there is very little change in Leydig cell numbers between birth and 20 dpp. This is a period during which differentiation to fetal Leydig cells is minimal and differentiation into adult Leydig cells is just beginning. In addition, it has been shown that cellular death by apoptosis does not occur in Leydig cells during the neonatal, prepubertal, pubertal, and adult periods (Faria et al. 2003). On the other hand, the distribution of Leydig cells within the interstitium differed between genotypes and between developmental stages: the difference in clustering was obvious only in

![Figure 4 Analysis of androgens produced in vitro in response to LH (400 pg/ml). Fetal testes, either Cx43-deficient (Gja1−/−) or wild-type (Gja1+/+), were grafted into intact adult hosts for 3 weeks (left axis). No significant difference was observed between the genotypes (Kruskal–Wallace test followed by Dunn’s multiple comparison). Wild-type testes were removed directly from 3-week-old pups and their androgen output measured to provide a standard of comparison (right axis). The bars represent mean±S.E.M.; the number of testes is indicated by the number in parentheses above each bar.](image)

![Figure 5 Activities of androgen metabolizing enzymes in Cx43-deficient (Gja1−/−) or wild-type (Gja1+/+) grafted testes. The values are expressed as mean±S.E.M. (n=6). Activities of 17βHSD and 5αR were significantly higher in mutant testes (P=0.047 and 0.045 respectively; Kruskal–Wallace test followed by Dunn’s multiple comparison). 3αHSD was not detected in mutant grafts.](image)
the older testes. The more diffuse distribution of Leydig cells in Cx43-deficient testes could be indicative of an adhesion role for Cx43, which becomes prominent as testicular volume increases.

In the previous study (Roscoe et al. 2001), grafted testes showed better development when the host gonads were left intact but in the present study, the host gonads were removed 1 week after the grafting procedure so that the contribution of the grafted testes to serum androgens could be determined. Our results demonstrated that the circulating levels of testosterone did not differ between hosts carrying grafts of different genotypes. Thus, gap-junctional communication among Leydig cells via Cx43 channels is not essential for androgen production.

As another test of the functional competence of Cx43-deficient Leydig cells, grafted testes were recovered from their hosts and cultured in vitro with different concentrations of LH to determine their sensitivity to the gonadotropin. There was no significant difference in androgen production between the genotypes at any LH concentration. These results are in agreement with and extend previous results indicating that Cx43-deficient testes respond similarly to supraphysiological LH stimulation (Roscoe et al. 2001). If the in vitro results reflect the events occurring in vivo, then the mutant testes are equally responsive to LH as wild-type testes, even on a per-Leydig cell basis. Furthermore, we found that mutant juvenile testes respond to similar levels of LH as wild-type adult testes under the same assay conditions. Therefore, gap-junctional communication between Leydig cells mediated by Cx43 channels is not an important determinant of the cells’ capacity to respond to LH by producing androgens.

The end products of androgen production are dependent on the maturity of the Leydig cells and can be used to assess the stage of development of those cells (Ge & Hardy 1998, Mendis-Handagama & Ariyaratne 2001, O’Shaughnessy et al. 2002). Testosterone is the major androgen produced by mature adult Leydig cells (mALC) in mammalian species, whereas immature adult Leydig cells (imALC), prevalent at puberty, release mainly the 5α-reduced testosterone metabolites (Benton et al. 1995, O’Shaughnessy et al. 2002). The imALC differentiate from the newly formed adult Leydig cells (nALC) which show a predominant secretion of androstenedione. Our analysis of androgens produced by juvenile testes indicated a preferred secretion of androstenedione, which is in agreement with previous documentation of Leydig cells at this postnatal stage (O’Shaughnessy et al. 2000, Mendis-Handagama & Ariyaratne 2001). Grafted testes retrieved at an equivalent postnatal age also showed androstenedione as the primary end product, suggesting a population not yet advanced to the pubertal stage when there should be more 5α-reduced products of testosterone such as DHT and androstanediol. The particular androgens released from the testes depend on the balance between the synthesizing and the metabolizing enzymes. Progenitor Leydig cells show low 17βHSD and 5αR activity but active 3αHSD with the resultant release of androsterone as the major product (Shan et al. 1993, Ge & Hardy 1998).

**Table 1** Test for dye coupling between Leydig cells.

<table>
<thead>
<tr>
<th>Source of Leydig cells</th>
<th>Total cells injected</th>
<th>Dye transfer</th>
<th>No dye transfer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type adult testis</td>
<td>7</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>Grafted wild-type juvenile testes</td>
<td>14</td>
<td>14</td>
<td>0</td>
</tr>
<tr>
<td>Grafted Cx43-deficient juvenile testes</td>
<td>20</td>
<td>18</td>
<td>2</td>
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</tbody>
</table>
In the imALC, production of 5α reduced metabolites predominates due to increased expression and enzymatic activity of 5αR and 3αHSD with an accompanying rise in 17βHSD activity (Ge & Hardy 1998, O'Shaughnessy et al. 2002). Our results showed relatively higher 17βHSD activity than 5αR and 3αHSD, but overall conversion of androstenedione to testosterone and its 5α reduced products was very low. Hence, although there was detectable release of 3α-diol from wild-type and juvenile testes, and DHT from the Cx43-deficient testes, the amounts did not affect the overall testosterone and androstenedione levels. Likewise, the small but significant increase in 17βHSD and 5αR activity in the mutant Leydig cells had little effect on the distribution of the different androgens. Our results consistently support the presence of some fetal Leydig cells which release androstenedione as their principal products (O'Shaughnessy et al. 2000).

Despite the fact that Cx43 was the only connexin known to contribute to gap junctions coupling Leydig cells, our data clearly show that Cx43-deficient Leydig cells remain dye coupled. Thus, Leydig cells may express one or more additional connexins that are yet to be identified. Furthermore, the permeability properties of that/those connexins must be sufficiently similar to those of Cx43 to maintain steroidogenic function in its absence. Functional overlap between members of the connexin family has been demonstrated in several instances, as has the ability of one connexin to replace another during organ development (White 2003). Another possibility is that one or more members of the recently described pannexin family of gap junction proteins might be facilitating intercellular communication between Leydig cells (Panchin et al. 2000, Bruzzone et al. 2003). Northern blot screens for pannexin expression in rat testes detected transcripts encoding pannexin1 (Baranova et al. 2004, Ray et al. 2005). However, a more complete understanding of the roles of gap-junctional intercellular communication in Leydig cell function must await identification of the additional gap junction protein(s) that contribute to the channels and the physiologically important molecules that pass through them.

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