

## THE UNIVERSITY of EDINBURGH

## Edinburgh Research Explorer

### Cell-specific ablation in the testis

Citation for published version: Smith, LB, O'Shaughnessy, PJ & Rebourcet, D 2015, 'Cell-specific ablation in the testis: what have we learned?', *Andrology*. https://doi.org/10.1111/andr.12107

### **Digital Object Identifier (DOI):**

10.1111/andr.12107

Link: Link to publication record in Edinburgh Research Explorer

**Document Version:** Publisher's PDF, also known as Version of record

**Published In:** Andrology

### **General rights**

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.



## ANDROLOGY

# ANDROLOGY

### **REVIEW ARTICLE**

### Correspondence:

Lee Smith, MRC Centre for Reproductive Health, University of Edinburgh, The Queen's Medical Research Institute, 47 Little France Crescent, Edinburgh EH16 4TJ, UK. E-mail: lee.smith@ed.ac.uk

#### Keywords:

animal models, germ cell transplantation, germ cells, gonadal development, Leydig cells, macrophages, paracrine factors, Sertoli cell, spermatogenesis, testis

Received: 19-Jun-2015 Revised: 19-Aug-2015 Accepted: 19-Aug-2015

doi: 10.1111/andr.12107

### **SUMMARY**

## Cell-specific ablation in the testis: what have we learned?

<sup>1</sup>L. B. Smith, <sup>2</sup>P. J. O'Shaughnessy and <sup>1</sup>D. Rebourcet

<sup>1</sup>MRC Centre for Reproductive Health, University of Edinburgh, The Queen's Medical Research Institute, Edinburgh, UK, and <sup>2</sup>College of Medical, Veterinary and Life Sciences, Institute of Biodiversity, Animal Health and Comparative Medicine, University of Glasgow, Garscube Campus, Glasgow, UK

Testicular development and function is the culmination of a complex process of autocrine, paracrine and endocrine interactions between multiple cell types. Dissecting this has classically involved the use of systemic treatments to perturb endocrine function, or more recently, transgenic models to knockout individual genes. However, targeting genes one at a time does not capture the more wide-ranging role of each cell type in its entirety. An often overlooked, but extremely powerful approach to elucidate cellular function is the use of cell ablation strategies, specifically removing one cellular population and examining the resultant impacts on development and function. Cell ablation studies reveal a more holistic overview of cell–cell interactions. This not only identifies important roles for the ablated cell type, which warrant further downstream study, but also, and importantly, reveals functions within the tissue that occur completely independently of the ablated cell type. To date, cell ablation studies in the testis have specifically removed germ cells, Leydig cells, macrophages and recently Sertoli cells. These studies have provided great leaps in understanding not possible via other approaches; as such, cell ablation represents an essential component in the researchers' tool-kit, and should be viewed as a complement to the more mainstream approaches to advancing our understanding of testis biology. In this review, we summarise the cell ablation models used in the testis, and discuss what each of these have taught us about testis development and function.

### INTRODUCTION

Testicular development and function is the culmination of a complex process of autocrine, paracrine and endocrine interactions between multiple cell types. Early studies that focussed upon dissecting these interactions generally relied on cell culture systems or systemic interventions in vivo using chemicals or surgical hypophysectomy to suppress endocrine and downstream testicular function. While these studies have given valuable insight into testis biology they suffer the serious drawbacks that cells do not behave normally in culture, whereas systemic interventions cannot assign impacts or roles to specific cell types or genes. A more detailed understanding of testis development and function has arisen from the use of transgenic mouse models, ablating the function of single genes to determine their individual roles. The development of conditional gene targeting has further refined this, permitting ablation of specific gene function in single cell types, and indeed the combination of these

reductionist approaches has identified many of the key genes essential for testis development and function. However, targeting genes one at a time does not capture the more wide-ranging role of each cell type in its entirety. An often overlooked, but extremely powerful approach to elucidate cellular function is the use of cell ablation strategies, specifically removing one cellular population and examining the impacts to reveal the importance of that cell type for development and function.

The major criticism levelled at cell ablation studies is the crudeness of the approach. It has been argued that removing an entire cell population, and thereby disrupting hundreds of signalling pathways simultaneously, provides little information regarding how the cell type in question influences other cellular populations. Whilst this is indeed true, and focussing on specific signalling pathways using this approach is flawed, evidence has demonstrated time and again that the power of cell ablation studies is to reveal a more holistic overview of cell-cell

Andrology, 1–15

1

distribution and reproduction in any medium, provided the original work is properly cited.

This is an open access article under the terms of the Creative Commons Attribution License, which permits use,

interactions. This not only identifies important roles for the ablated cell type, which warrant further downstream study, but also, and importantly, reveals functions within the tissue that occur completely independently of the ablated cell type. To date, cell ablation studies in the testis have specifically removed germ cells, Leydig cells, macrophages and recently Sertoli cells (Fig. 1). In this review, we will summarize the cell ablation models used in the testis, and discuss what each of these have taught us about testis development and function.

### APPROACHES TO CELL ABLATION

A range of options are available to those wishing to selectively ablate a single population of cells from an organ, but there are several key features shared by all approaches. The choice of method to be used should: (i) be specific, only affect the target cell type; (ii) be precisely controllable, for example so that timing of cell death, etc., is known, (iii) have high efficiency of ablation, completely or almost completely remove the target population, (iv) act quickly, such that impacts can be directly attributed to cell loss; and (v) have minimal off-target toxicity, such that other cell populations or organs are not affected. An overview of available options and their associated efficacy is shown in Fig. 2, and their impacts on testis architecture in Fig. 3.

Many models lacking a specific cell type have been generated as a serendipitous consequence of mutation or gene knockout. At the most basic level, knockout of Sry or Sox9 leads to sex reversal through failure of Sertoli cell specification, fundamentally demonstrating the requirement of Sertoli cells for masculinization [reviewed in (Kobayashi *et al.*, 2005; Wilhelm *et al.*, 2013)]. Within the developing testis knockout of other genes can lead to absence of specific cell types, for example knockout of PDGFR-alpha (Brennan *et al.*, 2003) impacts on foetal Leydig cell development, whereas knockout of desert hedgehog (DHH) (Clark *et al.*, 2000), or COUPTFII (Qin *et al.*, 2008) leads to absence of the adult Leydig cell population; W<sup>v</sup> mutations of cKit lead to near germ-free testes, through failure of primordial germ cell migration (De Franca *et al.*, 1994), whereas mutations such as the cell-specific loss of AR from Sertoli cells leads to loss of post-meiotic germ cells (Chang *et al.*, 2004; De Gendt *et al.*, 2004). In each of these cases, however, it is challenging to assign any downstream phenotypic impact specifically to loss of gene function or to the wider effects of cell ablation. As such, and with the exception of the  $W/W^v$  mouse, few of these gene knockout models have been used widely as a model of cell ablation outside the context of their own gene function. To widen the scope of cell ablation to different genotypes and species several alternative approaches have been developed that induce cell loss of a specific testicular cell population without the requirement for gene knockout.

### **GERM CELLS**

By far the most common approach to ablation of germ cells from the testis utilizes nascent chemotherapy agents such as busulfan (Myleran). The impact of busulfan on the spermatogenic cell population was first identified in the 1960s (Jackson et al., 1962), and its ability to apparently selectively remove the male germ cell population has been exploited widely in the past 50 years. This has been both in the pursuit of understanding about the role of germ cells in the testis, and also as a technical tool, important in preparation of recipients prior to germ cell transplantation (Brinster & Zimmermann, 1994; Brinster et al., 2003) [for review see (Brinster, 2002)], for fertility preservation and as a tool to investigate the spermatogonial stem cell niche. Busulfan is an alkylating agent that targets rapidly dividing mitotic spermatogonia resulting in apoptosis of this germ cell stage within 1 week, with a second wave of apoptosis in meiotic spermatocytes after 2 weeks (Choi et al., 2004) (Fig. 2). For these reasons, busulfan requires several weeks to completely ablate germ cells from the testis as the residual post-mitotic stages complete spermatogenesis and are released into the seminiferous tubule lumen (Medrano et al., 2014) (Fig. 3).

Exposure to radiation will also cause mitotic cells to undergo apoptosis and has been used as a technique to study the effects of germ cell ablation (Fig. 2). At high doses radiation exposure in

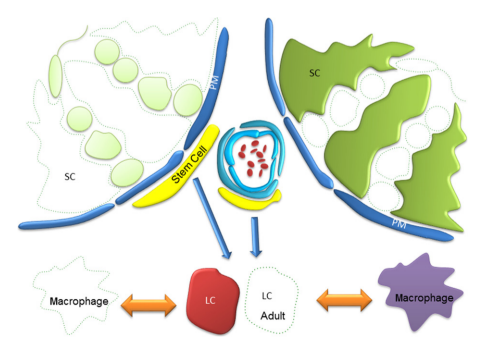
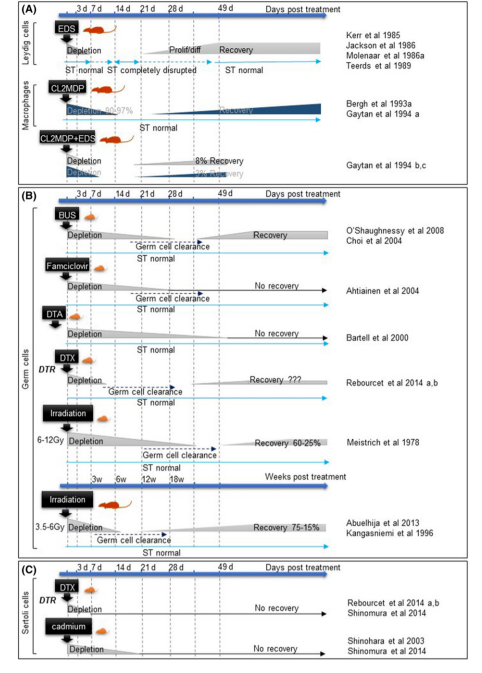


Figure 1 Schematic representation of the testis showing ablated cell populations. To date methodologies have been employed to selectively ablate germ cells, Leydig cells, Macrophages and recently Sertoli cells. By removing a single population and asking 'what changes' and 'what remains the same?', these studies have proven instrumental in defining our fundamental understanding of cell-cell communication in the testis, assigning specific roles to individual cell types whilst ruling out others. SC = Sertoli cell; PM = Peritubular Myoid cell; LC = Leydig cell; Stem cell = Leydig stem cell.

© 2015 The Authors. Andrology published by John Wiley & Sons Ltd on behalf of American Society of Andrology and European Academy of Andrology

**Figure 2** Schematic overview of depletion and recovery timings for each model of cell ablation. A schematic diagram comparing the rapidity of cell depletion and the duration of recovery days (d) after treatment, for each testicular cell type and method: (A) Leydig cells and macrophages when ablated independently or simultaneously; (B) Germ cells and (C) Sertoli cells. ST: seminiferous tubules, EDS: Ethylene dimethane sulphonate, CL2MDP: Liposome-entrapped dichloromethylene diphosphonate, BUS: busulfan, DTA: active diphtheria fragment A, DTR: diphtheria receptor, DTX: diphtheria toxin.

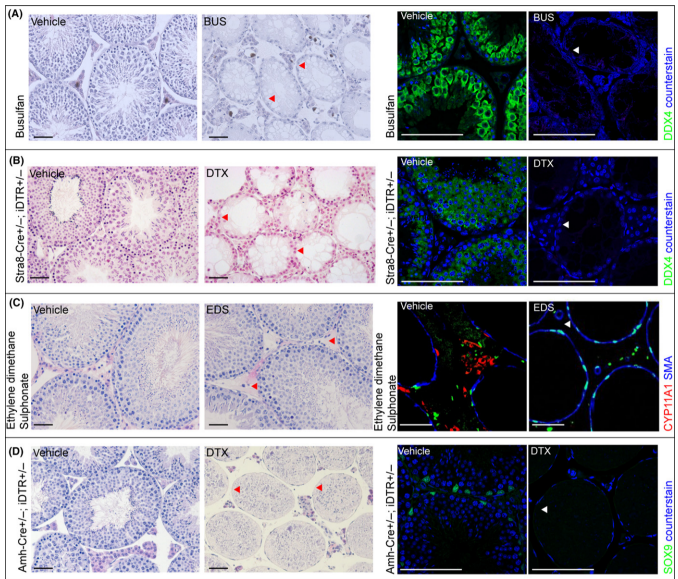


rat models can lead to infertility associated with a persistent failure to re-establish spermatogenesis (Kangasniemi *et al.*, 1996) and this has provided insight into how radiation-induced infertility can occur, with direct relevance to cancer patients undergoing radiotherapy. For example reciprocal germ cell transplant studies (irradiated spermatogonia into non-irradiated hosts or vice versa) reveal that, whereas germ cell loss is likely a direct result of impacts on germ cells, failure to re-establish spermatogenesis is a somatic cell problem rather than any direct impact on spermatogonial stem cells (Zhang *et al.*, 2007), suggesting radiation acts on both germ cells and somatic cells simultaneously.

The ability of very defined doses of radiation to selectively remove the germ cell population has also been explored as a method to generate recipient animals for germ cell transplants, especially in scenarios where genetic mutants lacking germ cells (e.g.  $W/W^{\vee}$ ) mice are unavailable or inappropriate (e.g. difficult to breed, immunocompatibility issues with donor cells, use of other species) (Zhang *et al.*, 2006). Even within mice, the removal of germ cells using busulfan is a strain-dependent balance between the required toxicity to kill sufficient germ cells and the sensitivity of the animal to the wider toxicological effects of the drug, particularly on haematopoiesis (Meistrich *et al.*, 1978; Abuelhija *et al.*, 2013). In this context, radiation provides a useful alternative to busulfan; the fact that it has not been taken up widely probably reflects the general move of researchers away from use of radiation wherever possible and the ready availability and convenience of a single busulfan injection which requires significantly less specific training, infrastructure or administrative oversight.

### ANDROLOGY

**Figure 3** Cell-specific ablation in the testis. Example of testis histology and immunolocalisation of cell-specific markers following Germ cell, Leydig cell or Sertoli cell ablation respectively (indicated by arrowheads). (A) Germ cell ablation in mouse using busulfan (DDX4 staining to mark germ cells); (B) Germ cell ablation in Stra8-Cre $\pm$ ;iDTR $\pm$  mice using DTX injection (DDX4 staining to mark germ cells); (C) Leydig cell ablation in rat using EDS (CYP11A1 staining to mark Leydig cells). (D) Sertoli cell ablation in Amh-Cre $\pm$ ;iDTR $\pm$  mice using DTX injection (SOX9 staining to mark Sertoli cells); (scale bar: 100  $\mu$ m).



An alternative model to either busulfan or radiation utilises a unique transgenic approach, expressing the Herpes Simplex virus Thymidine Kinase (TK) transgene from an inhibin-alpha promoter, which directs expression to Leydig and Sertoli cells, but not spermatogonia (Ahtiainen et al., 2004). The TK enzyme phosphorylates analogue nucleosides such as famciclovir, which are then incorporated into replicating DNA, acting as chain terminators and leading to eventual cell death. In the Inha/TK mouse, mitotically quiescent Sertoli and Leydig cells are resistant to the effects of TK, whereas the rapidly dividing spermatogonia cells are susceptible, probably through a bystander effect mediated by the gap junctions with Sertoli cells (Ahtiainen et al., 2004). As with busulfan, the ontogeny of cell death is gradual, taking 4 weeks to remove all germ cell populations through selective ablation of spermatogonia followed by natural development and sloughing of later germ cell stages until the testis is

devoid of germ cells (Fig. 2). Whilst time to germ cell clearance is long, the authors show that a 3-day treatment is sufficient to completely ablate all germ cell populations, and they observe no recovery 3 months post treatment. This could either be attributed to complete cell death of all spermatogonial stem cells, or impacts on Sertoli cells that prevent recovery of spermatogenesis. (Ahtiainen *et al.*, 2004).

Diphtheria toxin (DTX) has also been used in several models of germ cell ablation. In one model expression of the diphtheria toxin A-chain gene is directed to the male germline by fusion to 1 KB of the five prime flanking DNA of the rat male-germ cellspecific histone H1t gene (Bartell *et al.*, 2000). The H1t promoter fragment drives cell-specific expression of the diphtheria toxin A fragment in pre-spermatogonia. This results in widespread cell death, with just a few germ cells reaching the pachytene spermatocyte stage, and by 13 weeks of age the testis is almost

completely devoid of germ cells (Fig. 2). In contrast, females are completely fertile (Bartell *et al.*, 2000). Although the testes in these animals are extremely small, seminal vesicle weights are normal, suggesting that loss of germ cells does not impact androgen production.

In contrast to direct expression of the DTX A-chain inside germ cells, other transgenic approaches have been used which exploit the natural resistance of mice to DTX. In primates, DTX enters cells by binding to a cognate HBEGF receptor (also known as Diphtheria Toxin Receptor - DTR) and it is this internalisation followed by cleavage of the A-chain, which leads to cell death. In mice, DTX has low affinity for mHBEGF but transgenic expression of an inducible primate iDTR within mouse germ cells confers sensitivity to injected recombinant DTX. Breeding of iDTR to a germ cell Cre Recombinase line (Stra8-Cre) (Sadate-Ngatchou et al., 2008) results in expression of iDTR in male germ cells from postnatal day (pnd) 3 onwards (Rebourcet et al., 2014b). These animals show normal fertility but germ cell-specific cell death can be induced at any time of choosing through injection of DTX. Cell death is rapid, with apoptosis induced within 24 h and near complete clearance of germ cells from the seminiferous tubules within 7 days of a single injection (Fig. 2 and 3). To date this approach combines the fastest clearance of all germ cells with the lowest off-target toxicity although this approach does rely on the presence of multiple transgenes, which is impractical in most contexts. Furthermore, because of the variable nature of the Cre used (Bao et al., 2013), a small number (<1%) of spermatogonia fail to express the receptor and thus remain in the testis. However, the fact that these residual cells are able to repopulate the testis to produce complete spermatogenesis, demonstrates there is no long-term impact of germ cell ablation on testis function using this approach (Rebourcet et al., 2014a,b).

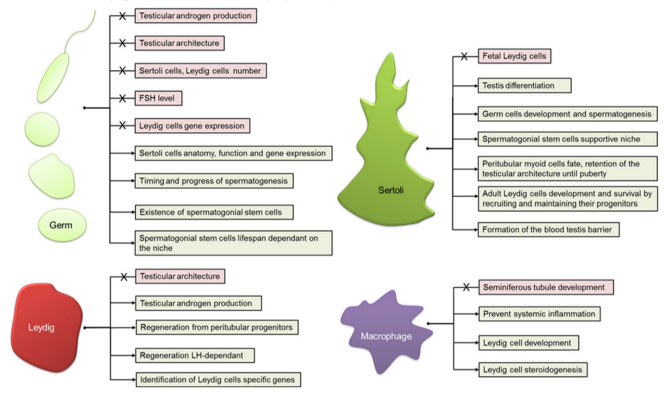
Germ cell ablation has significantly enhanced our understanding of testis development and function (Fig. 4). The most striking observation arising from germ cell loss is that the testis can form a normal architecture in the complete absence of germ cells. This suggests that germ cells play little role in the architecture of the developing testis. Indeed W/W<sup>v</sup> mice, which lack germ cells, have been used to demonstrate that germ cells play no role in the increase in Sertoli cell numbers observed in cases of hypothyroidism (Saxena et al., 2002), strongly suggesting that hypothyroidism acts through a direct effect on Sertoli cells [reviewed in (Holsberger & Cooke, 2005)]. This apparent lack of impact on testicular architecture is also observed if germ cells are ablated in adulthood (Fig. 3), with no effect seen on either Sertoli cell (De Franca et al., 1994; O'Shaughnessy et al., 2003) or Leydig cell numbers (O'Shaughnessy et al., 2003) and retention of general testicular architecture. Further to this, ablation of germ cells in adulthood does not impact on Leydig cell gene expression (O'Shaughnessy et al., 2008a) or androgen concentrations, with no change observed in either intratesticular testosterone (O'Shaughnessy et al., 2008a) or circulating testosterone concentrations (Gomes et al., 1973; Morris et al., 1987; De Franca et al., 1994), suggesting that germ cells play little role in control of testicular androgen production. In contrast to the lack of effect on Leydig cells, germ cell loss significantly impacts Sertoli cell anatomy (a significant reduction in smooth endoplasmic reticulum (De Franca et al., 1994)) and function (although not Sertoli cell number). By following the progressive nature of germ

cell loss after busulfan treatment, it is clear that the Sertoli cells are most influenced by post-meiotic germ cells, with significant changes in Sertoli cell gene transcript levels apparent only when these germ cells are lost from the seminiferous epithelium, ~15– 30 days post busulfan injection. Whilst there are also gene expression changes at earlier time points, these are not as widespread suggesting spermatogonia and spermatocytes exert less influence over Sertoli cell gene expression (O'Shaughnessy *et al.*, 2008b).

Perhaps one of the most intriguing observations is the relationship between germ cells and circulating follicle stimulating hormone (FSH) levels. The current paradigm dictates that FSH levels will increase in response to a reduction in germ cell number, via antagonism of Sertoli cell production of InhibinB, which acts in a negative feedback loop in the pituitary [reviewed in (Meachem et al., 2001)]. Whilst this concept is supported by the observation of a significant increase in circulating FSH concentrations following busulfan-induced germ cell loss in the rat (Gomes et al., 1973), and in the mutant W/W<sup>v</sup> mouse (De Franca et al., 1994), this is not observed in wild-type mice where in contrast, complete loss of germ cells either through busulfan injection (O'Shaughnessy et al., 2008a) or diphtheria toxin injection (Rebourcet, O'Shaughnessy, Smith, unpublished data) results in no increase in FSH levels whatsoever. It is difficult to reconcile these observations, although the fact that FSH levels are significantly increased in a mouse model of Sertoli cell ablation (Rebourcet et al., 2014a) suggests that the primary insult may be to the Sertoli cells. It could therefore be speculated that either the rat germ cells influence Sertoli cells to a greater extent than in the mouse, or alternatively, that rat Sertoli cells are more sensitive to perturbation by busulfan.

In addition to advancing our understanding regarding the control of testis development and cell-cell interactions, germ cell ablation, when combined with subsequent germ cell transplantation has also provided significant understanding of germ cell development. In a groundbreaking study published in 1994, Brinster and Zimmermann showed that spermatogenesis was possible following germ cell transplantation in immunocompatable mice (Brinster & Zimmermann, 1994). Key to the success of the study was the use of mice lacking germ cells, with both W/ W<sup>v</sup> mice and busulfan-treated mice successfully used. This revolutionary approach opened up the possibility of supporting spermatogenesis in response to adverse clinical conditions, for example fertility preservation in cancer patients. Along the way, however, use of germ cell ablation and transplantation has also enlightened our understanding of germ cell development and function, for example permitting the characterization of the pattern and kinetics of donor-derived spermatogonial stem cell colonisation. This includes the observation that Sertoli cells recognise and permit retrograde spermatogonial transit through the tight junctions of the blood-testis barrier (Nagano et al., 1999). Through this approach it has also been shown that crossspecies transplantation of rat spermatogonia into mice results in full 'rat' spermatogenesis (Clouthier et al., 1996), with the germ cells taking 52-53 days to fully mature, rather than the normal 35 days of mouse spermatogenesis (Franca et al., 1998). This means that it is the germ cells, rather than Sertoli cells, which control the timing and progress of spermatogenesis.

Identifying the stem cells amongst the spermatogonia lining the basement membrane of the seminiferous tubules using **Figure 4** What have cell-ablation models taught us? Overall testis differentiation, somatic cell development and testicular endocrine function is independent of Germ cell regulation. However, germ cell-ablation models have highlighted the intricate cross talk between germ cells and Sertoli cells. germ cell ablation combined with transplantation studies have also shown that germ cells control the timing and the progression of spermatogenesis, although the lifespan of the spermatogenesis. When ablated, Leydig cells are able to regenerate from peritubular progenitors under regulation from Luteinising hormone (LH) and other testicular secreted factors. Cell ablation studies have permitted the complete dissection of this regeneration process, which is believed to recapitulate normal adult Leydig cell development in most respects. Leydig cell-ablation studies have also aided the identification of Leydig cell-specific transcripts. The testis is an immune privileged organ. Macrophage ablation studies have shown that these cells are not essential to overall seminiferous tubules development. They prevent local systemic inflammation engendered by Leydig cell death. Macrophages and Leydig cells share an intricate physical and physiological relationship; macrophage ablation has revealed that this relationship is essential for Leydig cell development and regeneration. Sertoli cells are crucial components of the spermatogenis). Sertoli cell-ablation models have shown that, following their initial differentiation, Fetal Leydig cells are no longer dependent on Sertoli cells. In contrast the development and survival of the adult Leydig cell population is fundamentally dependent upon Sertoli cells.



morphological parameters has proven challenging. Germ cell ablation and transplantation has been used to estimate the number of spermatogonial stem cells through counting of individual colonies of expanding spermatogonia marked with B-galactosidase. Initial estimates using transplant into adult mice has produced an estimate of ~0.002% of total spermatogonia number (Nagano et al., 1999), however, transplant into younger recipients in which the blood-testis barrier (BTB) had yet to form increases colony number 10-fold suggesting the initial estimates were hindered by the ability for stem cells to cross the BTB in a retrograde direction and colonise available niches (Shinohara et al., 2001). The functional lifespan of spermatogonial stem cells has also been established using cell ablation and transplantation. These studies show that, whilst under normal circumstances stem cell number and proliferative capacity drops off with age, stem cells retain their proliferative capacity if transplanted into younger mice. Serial transplantation experiments demonstrate that this remains the case for more than 2 years in vivo in the mouse, significantly longer than the normal breeding window (Ogawa et al., 2003; Ryu et al., 2006), and highlighting the degeneration of the niche as the limiting factor in lifelong spermatogenesis.

Germ cell ablation and transplantation studies have also been used to establish when germline stem cells acquire spermatogonial stem cell activity. Ohta *et al.*, (2004) have shown that gonocytes collected from 13.5 days post coitum (dpc) embryos or later stages can colonise seminiferous tubules. Moreover, when transplanted into younger, 5–10 day old recipients, not only primordial germ cells from all foetal ages, but also epiblasts from 6.5 dpc embryos could colonise and go on to produce spermatozoa (Chuma *et al.*, 2005). Taking this to extreme levels, it is now possible (but challenging and with poor efficiency) to generate PGC-like cells from ES cells and transplant these into host testes to produce spermatids and live mice through assisted reproduction techniques (Hayashi *et al.*, 2011).

As discussed above, the development of knockout mouse models has revolutionised genetics and our ability to study complex biochemical and physiological mechanisms. Until the more recent development of conditional knockout technologies, however, one of the key issues when examining the role of genes in knockout animals was the inability to assign gene function to a specific cell type if the gene is expressed in several cells. Germ cell ablation and transplantation using mutant donor germ cells or mutant recipients has been successfully used to dissect the

Steel/cKit tyrosine receptor system, mutations in which lead to a block in spermatogenesis. In the testes of mice homozygous for mutations in either cKit (W/W<sup>v</sup>) (germ cell) or the kit ligand Steel factor (Sl/SLd) (Sertoli cell) the seminiferous tubules lack almost all germ cells (Ogawa et al., 2000). However, transplantation of wild-type germ cells into W/W<sup>v</sup> mice restores fertility in W/W<sup>v</sup> mutants, whereas transplantation of spermatogonial stem cells from Sl/Sld males into W/W<sup>v</sup> mice also restores fertility (Ogawa et al., 2000), both by overcoming the cell-specific impacts of their respective gene loss. As the expression of the ligand and the receptor differs (ligand-SC and receptor-GC), the germ cell loss defect in both mutants originate either from the precursor cells (W/W<sup>v</sup>) or from the microenvironment (Sl/SLd). Therefore, the outcome after transplantation of green fluorescent protein (GFP) tagged germ cells into the seminiferous tubules of Sl/SLd mice resulted in colonisation but no differentiation of the GFP germ cells, which remain as spermatogonia. In contrast, when they are re-transplanted into W/W<sup>v</sup> mice complete spermatogenesis was achieved (Ohta et al., 2000).

This initial proof of principle for assigning gene roles to either germ cell or somatic cell populations has been expanded to other genes, showing, for example that transplantation of spermatogonia from juvenile spermatogonial depletion jsd-mutant mice, which show an arrest of spermatogenesis after one round of meiosis, into wild-type host mice fails to restore fertility. In contrast, transplant of wild-type germ cells into a jsd host leads to full spermatogenesis, confirming that the *jsd* mutation is an intrinsic problem with the germ cells, and not the somatic environment (Boettger-Tong et al., 2000). In a similar approach, transplantation of germ cells from estrogen receptor-alpha (αERKO) knockout mice, which are infertile, into germ cell-depleted hosts leads to normal spermatogenesis, showing it is the somatic environment underpinning this particular infertility phenotype (Mahato et al., 2000). Similarly, germ cells from infertile testicular feminized (Tfm) mice (which have a null mutation in the androgen receptor (Ar)) are able to go on to produce spermatozoa once transplanted into a wild-type host mouse (Johnston et al., 2001). This approach has been widely exploited with investigations into the role of genes such as Etv5 (Morrow et al., 2007) Claudin11 and Rac1 (Takashima et al., 2011) and factors such as vitamin-A (McLean et al., 2002) in the regulation of spermatogonial stem cell niche populations [reviewed in (Kanatsu-Shinohara & Shinohara, 2013)].

To a large extent the development of conditional gene targeting and the wide availability of germ cell expressing Cre recombinase lines [reviewed in (Smith, 2011)] has superseded the requirement for such transplantation experiments to establish gene function. In fact the protocol has become so mainstream that today it is largely exploited for what it can do as a technology, for example xenogenic germ cell development for endangered species [which is itself not without problems (reviewed in (Paris et al., 2004)], or as an approach to generate transgenic domesticated animals (Zeng et al., 2013). Most recently germ cell ablation and transplantation has been associated with a drive towards the generation of artificial gametes (gametes generated by manipulation of their progenitors or of somatic cells) (Hendriks et al., 2015a), which raises significant moral and ethical questions that need to be debated by society in the coming years (Hendriks et al., 2015b).

### LEYDIG CELLS

The Leydig cells are the source of androgen production by the testis and these androgens are essential for male phenotypic differentiation, fertility and libido. It is clear that in eutherian mammals there are two populations of Leydig cells which arise sequentially during development (Roosen-Runge & Anderson, 1959; Lording & De Kretser, 1972). The foetal population of cells develops soon after testis differentiation and is essential for foetal synthesis of androgen and INSL3 which ensure masculinization of the foetus and testis descent (Nef & Parada, 1999). In all mammals so far studied a second, adult population of Leydig cells starts to develop in the pre-pubertal period [around days 7-10 in mice (Baker et al., 1999; Nef et al., 2000)]. This adult population of cells secretes androgens essential for male phenotypic and behavioural development and puberty and maintains fertility in the adult. The adult Leydig cells are completely dependent on LH support for both development and adult function; in the absence of LH few adult Leydig cells develop and circulating testosterone levels are barely detectable (O'Shaughnessy et al., 1998; Baker & O'Shaughnessy, 2001; Lei et al., 2001; Zhang et al., 2001). In contrast, the foetal Leydig cells in rodents appear to function largely independently of hormonal support (El Gehani et al., 1998; O'Shaughnessy et al., 1998; Baker et al., 1999; Lei et al., 2001; Zhang et al., 2006) although in other species LH (or hCG in humans) is required for foetal androgen production (O'Shaughnessy & Fowler, 2011).

Leydig cell ablation has proven incredibly informative in improving our understanding of testis development and function (Fig. 4). Studies of Leydig cell ablation started largely by chance through observations of the effects of ethane dimethane sulphonate (EDS) in the rat. In the 1960s Harold Jackson and colleagues were examining anti-spermatogenic agents as potential regulators of male fertility [see (Jackson, 1970)]. EDS is structurally very similar to busulfan and a single dose was shown, like busulfan, to cause disruption of the seminiferous epithelium over a similar time scale (Jackson, 1970). One significant difference, however, was that treatment with EDS led to a marked loss in weight of the prostate and seminal vesicles indicating loss of testosterone and a change in Leydig cell function (Jackson, 1970; Jackson & Jackson, 1984). Subsequently, in the mid-1980s it was shown that EDS acts by causing rapid ablation of the Leydig cells within 48 h of a single injection (Kerr et al., 1985; Molenaar et al., 1985; Morris et al., 1986) (Fig. 2). Other effects of EDS on the testis (e.g. loss of spermatogenesis) were shown to be due largely to the consequences of androgen withdrawal although some effects on Sertoli cell and peritubular myoid cell function have been reported (Verhoeven et al., 1989; Roberts & Griswold, 1990). EDS is a glutathione-dependent alkylating agent (Kelce & Zirkin, 1993; Morris, 1996), which induces apoptosis in Leydig cells through activation of the Fas system (Taylor et al., 1999; Kim et al., 2000) and possible upregulation of N-myc downstream-regulated gene 2 (NDRG2) (Li et al., 2012). It is not clear, however, how EDS acts to induce the apoptotic pathway or why EDS is specific to Leydig cells. There is also strong species specificity to the actions of EDS on Leydig cells with little effect of the toxicant seen in a number of species including the mouse, dog and monkey, whereas others such as the rat, guinea pig and frog show high sensitivity (Morris, 1996). This may provide a further clue to the specificity of the toxicant as it has been shown that in

rats the Leydig cells show specific expression of a number of metabolic enzymes which act to metabolise xenotoxicants (O'Shaughnessy *et al.*, 2014). It is possible that one of these enzymes may serve to activate EDS and species specificity may arise from variability in substrate specificity or enzyme isoforms expressed.

In the years since its actions were first identified EDS has proved an extremely valuable tool in our efforts to understand Leydig cell biology. Its usefulness is only limited by its failure to act in mice (except at very high doses which will probably have major systemic effects (Li et al., 2012)), which means it cannot be used in conjunction with the numerous gene knockout models available in that species. Much of the interest in EDS has centred on the key observation that the Leydig cell population in the adult will regenerate in 3-6 weeks following ablation (Kerr et al., 1985; Molenaar et al., 1986). This is important because it shows that Leydig cell stem cells persist in the adult testis and that the potential exists to regenerate Leydig cells in situations such as ageing where Leydig number and/or activity is reduced (Chen et al., 1996). In addition, Levdig cell re-generation after EDS appears to closely resemble normal development of the adult Leydig cell population thereby providing a model system for study of this process (Teerds & Rijntjes, 2007; Guo et al., 2013). The advantages of this model over normal development are that the interstitium is clear of Leydig cells and so there can be no confusion between developing and mature Leydig cells, other developmental events in the testis are not occurring simultaneously and hormone levels can be more easily manipulated in the more mature animals.

Study of Leydig cell re-generation after EDS has identified the origin of the differentiating Leydig cells, the role of gonadotrophins and has started to identify other factors which may be involved in the process. The identity of the adult Leydig cell precursor population has been contentious for a number of years with pericytes, peritubular cells, fibroblasts, and endothelial cells all suggested (Jackson et al., 1986; Kerr et al., 1987; Teerds et al., 1990; Davidoff et al., 2004; O'Shaughnessy et al., 2008b; Stanley et al., 2012). Recent data indicates, however, that the major source of adult Leydig cells after EDS treatment are peritubular cells (O'Shaughnessy et al., 2008b; Stanley et al., 2012) although a contribution from other sources cannot be ruled out. The role of gonadotrophins in Leydig cell regeneration was established in early EDS studies using hypophysectomised rats or testosterone implants. This work showed that Leydig cell regeneration after EDS is dependent on LH (Molenaar et al., 1986; Sriraman et al., 2003) although some early proliferation and differentiation appears to occur independent of LH (Teerds et al., 1989). Subsequent studies have shown that similar control mechanisms regulate normal adult Leydig cell development at puberty (Clark et al., 2000; Gnessi et al., 2000; Baker & O'Shaughnessy, 2001; Zhang et al., 2001, 2004; Baker et al., 2003). Other factors such as DHH, PDGF and anti-Müllerian hormone (AMH) have been implicated in the development of Leydig cells both during development and re-generation (Salva et al., 2004; Josso et al., 2006; O'Shaughnessy et al., 2008b) and further studies on EDS regeneration have also suggested that nerve growth factor (NGF) and Sertoli cells are involved in Leydig cell proliferation, differentiation and survival (Yan et al., 2000; Zhang et al., 2013).

While progress has clearly been made in our understanding of adult Leydig cell development and regeneration using EDS,

questions remain to be answered. For example data from mutant and knockout mice make it clear that LH is essential for Levdig cell differentiation but do changes in LH concentrations explain the onset and control of Leydig cell development? LH levels certainly rise after EDS treatment (Molenaar et al., 1986) and treatment of normal adult rats with LH or hCG for up to 5 weeks is reported to stimulate an increase in Leydig cell number (Christensen & Peacock, 1980; Teerds et al., 1988; Mendis-Handagama et al., 1998). Ablation of the germ cell population at the same time as EDS treatment, however, has been shown to cause more rapid regeneration of the Leydig cells without further affecting LH levels (Molenaar et al., 1986; Risbridger et al., 1987). It is also not clear that circulating LH levels rise prior to the start of normal adult Leydig cell development (Selmanoff et al., 1977; Michael et al., 1980; Jean-Faucher et al., 1983; Murphy et al., 1994). Finally, we have shown that Sertoli cell ablation prevents normal Leydig cell development (Rebourcet et al., 2014b) (see also section 6) indicating that the Sertoli cells must be involved in this process. As suggested above, understanding what regulates Levdig cell development might allow controlled regeneration of Leydig cells in circumstances where Leydig cell numbers and/or activity are reduced. This may be particularly relevant to ageing (Beattie et al., 2015) as Leydig cells which regenerate in aged rats after EDS treatment have similar activity to Leydig cells from young rats and they are able to maintain serum testosterone levels in the aged rats at the same level as those in young rats (Chen et al., 1996).

Leydig cell ablation by EDS causes a marked reduction in intra-testicular androgen levels and this technique has been used to investigate the role of androgens in the testis, particularly with reference to Sertoli cell function and spermatogenesis. Other techniques such as hypophysectomy or treatment with GnRH agonists or androgen receptor antagonists will have similar effects, however, and so data coming from EDS experiments are not unique in that respect. One of the more interesting observations using EDS was made in 1988 and showed that spermatogenesis can largely be maintained after EDS if the animal is treated with sufficient androgen (Sharpe *et al.*, 1988). This implies that androgen is the only Leydig cell product that is essential for spermatogenesis although a repeat of this study with more up-to-date and sensitive measures of spermatogenesis and Sertoli cell activity would appear to be warranted.

The ability to ablate a specific cell type in a tissue should allow the identification of transcripts or proteins, which are specific to that cell type, or that rely on that cell type. This technique has been used, for example to show that expression and activity of the steroidogenic enzymes, with the exception of HSD17B3, is limited to the Leydig cells (Murphy & O'Shaughnessy, 1992; O'Shaughnessy et al., 2008b). More recently, arrays have been used in two independent studies to identify a comprehensive set of Leydig cell-specific transcripts in the EDS rat model (O'Shaughnessy et al., 2014; Zhang et al., 2015). In both studies transcripts were identified that showed markedly reduced expression following Leydig cell ablation and in each case this included most of the known Leydig cell-specific transcripts. Perhaps surprisingly, however, there are quite marked differences in the lists of Leydig cell transcripts identified in the two studies. For example of the 67 known genes identified by Zhang and colleagues (Zhang et al., 2015) as Leydig cell exclusive only 23 are also identified by O'Shaughnessy and colleagues

(O'Shaughnessy et al., 2014). There are technical differences between the studies (e.g. the arrays used are different) but, perhaps more pertinent, the models are also slightly different. Zhang and colleagues (Zhang et al., 2015) used a 'standard' EDS model (normal adult rats treated once with EDS), whereas O'Shaughnessy and colleagues (O'Shaughnessy et al., 2014) used a germ cell-free rat model. The rational for using germ cell-free animals being that the reduction in androgen levels caused by EDS ablation of Leydig cells will cause germ cell apoptosis (Kerr et al., 1993) and, therefore, significant loss of germ cell-specific transcripts. This is turn may complicate identification of transcripts lost through direct ablation of the Levdig cells. Despite differences between studies, however, the approach to identifying cell-specific transcripts through cell ablation holds significant promise, and further studies should resolve differences in results so far published.

### MACROPHAGES

The resident testicular macrophages (characterised as CD68–, CD163+; 'ED2') form the largest population of immune cells in the testis, and form a close physical and physiological relationship with Leydig cells in the testicular interstitium [reviewed in (Winnall & Hedger, 2013)], with approximately one resident macrophage for every four Leydig cells (Bergh, 1985). Under non-inflammatory conditions, this population undergoes natural turnover through a slow infiltration of circulating (CD68+, CD163– 'ED1') monocytes/macrophages followed by a local conversion into the ED2 resident form (Sauter *et al.*, 2014).

Macrophage ablation has significantly enhanced our understanding of cell-cell communication in the testis (Fig. 4). In a series of elegant studies in the rat, the testicular function of the resident macrophage population has been investigated using injection of liposome-entrapped dichloromethylene diphosphonate into the testicular interstitium to ablate the macrophages (Bergh et al., 1993a). This treatment regimen in the adult reduces macrophage number in the injected testis by 90% 7 days post injection (Fig. 2). Results from these studies show that macrophages may play a previously unknown role in support of Leydig cell development and function as well as secreting factors that inhibit hCG-induced testicular inflammation (Bergh et al., 1993b). During pre-pubertal development, macrophage ablation results in an arrest in the size of the Leydig cell population until day 35, when macrophage repopulation permits ongoing Leydig cell development. Furthermore, the authors have demonstrated that, whereas exogenous hCG treatment for 6 days from day 21 results in a seven-fold increase in numbers of Leydig cells, no increase in Leydig cell number is observed in the contralateral, macrophage-ablated testis (Gaytan et al., 1994a), an observation confirmed in hypophysectomised and macrophage-depleted rats treated with hCG (Gaytan et al., 1995c). Importantly, in this study, treatment with FSH increased testicular weight and seminiferous tubule diameter independently of macrophage ablation, demonstrating that macrophages are dispensable for this aspect of testis function (Gaytan et al., 1995c). This first indication of an essential role for testicular macrophages in the development of the Leydig cell population has been examined further using the EDS model of induced Leydig cell ablation and regeneration, with or without concurrent macrophage ablation. Results from this study showed that macrophages are essential for regeneration of the Leydig cell population after EDS, with the size of the

population proportional to the number of macrophages, hinting at a physical, numerical and physiological relationship between the two populations (Gaytan *et al.*, 1994b,c). In a contrasting study, the role of macrophages in removal of apoptotic cells has also been investigated. When Leydig cell ablation was carried out 10 days after macrophage ablation there was a resultant influx of inflammatory lymphocytes and monocytes, not seen in the contralateral macrophage-containing testis. This suggests that resident macrophages may act to prevent a systemic inflammatory response to localised cell death in the testis (Gaytan *et al.*, 1995b).

Unilateral macrophage ablation in the adult testis results in increased LH, circulating testosterone and intra-testicular testosterone concentrations, with intra-testicular testosterone concentrations higher in the macrophage-retaining testis. This suggests that, although macrophages are essential for the development of the Levdig cell population, once established the macrophages enhance Leydig cell steroidogenesis, but are not essential (Gaytan et al., 1996). Indeed, treatment with hCG following bilateral ablation of the macrophage population leads to a 3.5-fold increase in serum testosterone, compared with an 8.6fold increase in intact controls (Gaytan et al., 1995a). A number of studies have also focussed on deciphering the mechanism involved in macrophage regulation of Leydig cell function. Using macrophage-conditioned media a lipophilic factor has been identified (Hutson et al., 1996) which appears to alter Leydig cell androgen synthesis through a StAR-independent pathway (Lukyanenko et al., 1998).

An alternative method of ablating the macrophage population in adulthood, – chronic 6-week treatment with anti-CSF1R antibody (which prevents development of replacement macrophages) – causes a small but significant reduction in testis weight (Sauter *et al.*, 2014). In this study, however, circulating LH and testosterone were unaffected, perhaps reflecting the development of a compensatory mechanism over the time course of treatment in contrast to the rapid induction of macrophage death caused by liposome injection. Alternatively, these data would support the suggestion that the mechanism by which macrophages are removed is responsible for the negative impacts on testicular function (Sauter *et al.*, 2014) and highlights the need for further study in this area.

In summary, these ablation studies suggest that macrophages play an essential role in the development of the adult Leydig cell population and are also important for support of steroidogenesis in adulthood (Hutson, 1998), but have little direct impact on the seminiferous tubule.

### SERTOLI CELLS

Sertoli cell differentiation following SRY/SOX9 cascade activation is the first step of testis development. The important role that these cells play in orchestrating testicular differentiation (promotion of foetal Leydig cell and peritubular myoid cell development) and subsequent masculinisation of the foetus is well established (Svingen & Koopman, 2013). During foetal and early postnatal life there is a period of intensive Sertoli cell proliferation which ceases at puberty as the cells undergo a fundamental change in activity (Sharpe *et al.*, 2003) (O'Shaughnessy, 2015). Inter-Sertoli cell junctional complexes form to generate the BTB, which creates a stable microenvironment for germ cell development/maturation. In adulthood, Sertoli cells act to maintain spermatogenesis with each Sertoli cell able to support a fixed number of germ cells such that the numbers of spermatozoa produced is dependent upon total Sertoli cell number (Berndtson & Thompson, 1990). Overall, Sertoli cells are essential for testis differentiation but whether they remain the central drivers of testis development beyond this point or, indeed, whether they play a major role in adult testis function, beyond spermatogenesis, has remained unknown for many years. Clearly the answer to these questions could be provided by cell ablation studies although this has proven difficult with respect to Sertoli cells until recent technological developments. These recent studies have uncovered many previously unknown roles for Sertoli cells in testis development and function (Fig. 4).

It has been shown that Sertoli cells will undergo apoptosis postnatally in mice lacking Sertoli cell expression of Dicer, which is required for micro RNA and small interfering RNA biogenesis (Papaioannou et al., 2009; Kim et al., 2010). Sertoli cell apoptosis starts at about postnatal day 3 in these animals although some Sertoli cells survive into adulthood. There is massive germ cell apoptosis in these animals and by 6 months of age the architecture of the testis has degenerated with only rare tubules remaining among an apparent abundance of interstitial tissue (Papaioannou et al., 2009). This model is interesting in so far as the role of Dicer in the Sertoli cells is concerned but it does not meet all the criteria for a good experimental model of Sertoli cell ablation -that is it cannot be precisely controlled and it does not act quickly. Nevertheless, some of the testicular changes seen in these animals (tubule degeneration, germ cell loss) mirrors/confirms changes seen with a targeted approach to Sertoli cells ablation described below.

An alternative approach to Sertoli cell ablation has been to use cadmium, a heavy metal which will deplete the seminiferous tubules of their contents and, potentially, ablate the Sertoli cells (Shinohara et al., 2003) (Fig. 2). This also does not meet all the criteria of the ideal cell ablation method as the impact of cadmium is not restricted to one testicular cell type (Marettova et al., 2015) but the Sertoli cells appear to be the most vulnerable to cadmium exposure and even low doses have specific effects on the Sertoli cells (vacuolisation, cell contraction, detachment, inhibition of protein synthesis, apoptosis) (Clough et al., 1990; Zhang et al., 2010). Given its effect on the Sertoli cells, cadmium has been used to explore the role of the Sertoli cells in development of the spermatogonial stem cell niche. Transplantation of Sertoli cells into cadmium- and busulfan-treated (Sertoli and germ cell depleted) seminiferous tubules (Shinohara et al., 2003) has permitted assessment of the potential of transplanted Sertoli cells to reconstitute a niche for spermatogonial stem cells and support active spermatogenesis. Using donor ROSA26-l29 males, it was possible to evaluate the colonization ability of immature and adult Sertoli cells and results showed that immature dividing Sertoli cells are more able to effectively colonize the denuded seminiferous tubules and support recovery of spermatogenesis in recipient testes (Shinohara et al., 2003).

The recent development of new transgenic mouse models expressing cytotoxic genes or specific toxin receptors has opened the door to more refined methods of Sertoli cell ablation (Palmiter *et al.*, 1987; Saito *et al.*, 2001; Buch *et al.*, 2005). Shinomura and colleagues (Shinomura *et al.*, 2014) and our group (Rebourcet *et al.*, 2014b) have independently applied similar approaches to drive the expression of diphtheria toxin A specifically in Sertoli cells from foetal life onwards, inducing Sertoli cell ablation from embryonic day 15. In addition, we have also combined an Amh-Cre line (Lecureuil *et al.*, 2002) and a iDTR line (Buch *et al.*, 2005) to permit ablation of Sertoli cells via injection of DTX (Rebourcet *et al.*, 2014a,b) (Fig. 2). The benefit of this inducible ablation model is that Sertoli cells can be ablated at different time points to assess temporal-specific roles of this population not only in supporting testis development but also in adult function (Rebourcet *et al.*, 2014a). These studies have emphasised the fundamental importance of Sertoli cells, revealing many previously undocumented roles that the cells play in the control of testis development and function (Rebourcet *et al.*, 2014a,b; Shinomura *et al.*, 2014).

As might be expected given the intricate relationship between Sertoli cells and germ cells, Sertoli cell ablation studies show that Sertoli cells are essential for germ cell survival, and that this is the case both in perinatal and in adult life. Injection of DTX leads to Sertoli cell death in the adult within 1 day and germ cell death follows rapidly, with no evidence of germ cells within the testis 10 days post DTX injection, including complete loss of mitotic spermatogonia (Rebourcet *et al.*, 2014a; Shinomura *et al.*, 2014). Only the reintroduction of immature Sertoli cells allows partial rescue of germ cell degeneration (Shinomura *et al.*, 2014). These data confirm the crucial role Sertoli cells play in the development and maintenance of the germ stem cell niche.

More surprisingly, Sertoli cell ablation in foetal or perinatal life demonstrates that the tubular architecture is not fully established until puberty. Ablation of Sertoli cells either from 15.5 dpc or in neonatal life induces de-differentiation of peritubular myoid cells, which no longer express smooth muscle markers. Recruitment of cells and re-establishment of a myoid signature can be achieved if single cell testis aggregates are recombined in xenografts (Rebourcet et al., 2014b), demonstrating that Sertoli cells actively enforce a myoid fate on peritubular cells throughout foetal and pre-pubertal development. In contrast, ablation of Sertoli cells during puberty (day 18) or in adulthood does not alter the tubular architecture suggesting that peritubular myoid cells fate is independent from Sertoli cells at this time, or that the joint deposition of the basement membrane by Sertoli and peritubular myoid cells is sufficient to stabilise the tubules and maintain myoid cell differentiation (Rebourcet et al., 2014b).

Our work has also revealed that Sertoli cells regulate the development of the adult Leydig cell population and are required for Leydig cell survival in adulthood. When Sertoli cells are ablated at postnatal day 2 or day 18 Leydig cell number in adulthood significantly decreases (70–90%), probably because of a reduction in the number of adult Leydig stem cells in the peritubular region. Perhaps more surprisingly, ablation of Sertoli cells in adulthood leads to a 75% decrease in LC number with the remaining cells concentrated in the sub-capsular region and around the rete testis/tubuli recti. The nature of the factor or factors linking the Sertoli cells to Leydig cell development and survival remain uncertain but are of particular interest as they have the potential to regulate the size and activity of the adult Leydig cell population.

Finally, ablation of Sertoli cells illustrates the key role that the peritubular myoid cells play as a component of the blood-testis barrier. When the Sertoli cells are ablated in adulthood there is no barrier to infiltration of biotin into the tubules but only about

<sup>© 2015</sup> The Authors. Andrology published by John Wiley & Sons Ltd on behalf of American Society of Andrology and European Academy of Andrology

10% of tubules show evidence of immune cell penetration of the tubules (Rebourcet *et al.*, 2014a). This shows that the peritubular myoid cells and underlying extracellular matrix form an effective barrier to macrophage invasion of the tubules. In addition, as peritubular myoid cells in the intact testis will normally act as a partial barrier to solute penetration of the tubule (Dym & Fawcett, 1970), failure to do this in the absence of Sertoli cells suggests that the function is Sertoli cell-dependent.

### CONCLUSION

The development and application of these various models of cell-specific ablation has significantly advanced our understanding of testis development and function. The dissection of the germ stem cell niche, the assignment of cell-specific gene function, interactions between germ cells and somatic cells and the exploitation for transplantation studies have all benefitted from cell ablation approaches. Cell ablation has also had a significant impact on our understanding of Leydig cell development through identification of a postnatal stem cell population and then modelling the regeneration of this population. Exploring the role of the enigmatic macrophages, revealing their significant influence over Leydig cells and the interplay between the two cell types would not be possible without using cell ablation techniques. Finally, and most recently, identifying the previously unknown roles Sertoli cells play in germ cell support, peritubular myoid cell development and Leydig cell development and lifelong support have opened new avenues of research which can now be explored using the more conventional tool-kit of the research scientist. With these studies describing the effects of Sertoli cell ablation, the remaining major cell type to be pursued in the testis is the peritubular myoid cell. It may prove difficult to target these cells specifically because of their similarity to smooth muscle cells but the effort is likely to be worthwhile as a number of recent studies have highlighted the importance of the peritubular myoid cells to testis function (Welsh et al., 2009, 2012; Nurmio et al., 2012; Qian et al., 2013; Chen et al., 2014). Other studies of interest would be the development of an inducible Leydig cell ablation model in the mouse to combine with the available gene knockout models and another testis macrophage ablation model to complement the earlier liposome studies.

While it is likely that specific gene ablation studies and other techniques will answer many of the remaining questions in testis biology, it can be argued that the types of cell ablation studies described here are essential precursors which lay the foundation for further advances in the field. Cell ablation as a technique lacks subtlety, but when used judiciously, it supports great leaps in understanding not possible via other approaches; as such it represents an essential component in the researchers' tool-kit, and should be viewed as a complement to the more mainstream approaches to advancing our understanding of testis biology.

### ACKNOWLEDGEMENTS

LBS is supported by the UK Medical Research Council Programme Grant Award (G1100354/1); and a Found Animals Michelson Grant Award (D1011-S05); LBS, POS and DR are supported by a BBSRC Project Grant Award (BB/J015105/1). We thank Dr Karen Kilcoyne for providing tissue sections from EDStreated rat testes.

### REFERENCES

- Abuelhija M, Weng CC, Shetty G & Meistrich ML. (2013) Rat models of post-irradiation recovery of spermatogenesis: interstrain differences. *Andrology* 1, 206–215.
- Ahtiainen M, Toppari J, Poutanen M & Huhtaniemi I. (2004) Indirect Sertoli cell-mediated ablation of germ cells in mice expressing the inhibin-alpha promoter/herpes simplex virus thymidine kinase transgene. *Biol Reprod* 71, 1545–1550.
- Baker PJ & O'Shaughnessy PJ. (2001) Role of gonadotrophins in regulating numbers of Leydig and Sertoli cells during fetal and postnatal development in mice. *Reproduction* 122, 227–234.
- Baker PJ, Sha JA, McBride MW, Peng L, Payne AH & O'Shaughnessy PJ. (1999) Expression of 3b-hydroxysteriod dehydrogenase type I and VI isoforms in the mouse testis during development. *Eur J Biochem* 260, 911–916.
- Baker PJ, Johnston H, Abel MH, Charlton HM & O'Shaughnessy PJ. (2003) Differentiation of adult-type Leydig cells occurs in gonadotrophindeficient mice. *Reprod Biol Endocrinol* 1, 4.
- Bao J, Ma HY, Schuster A, Lin YM & Yan W. (2013) Incomplete cremediated excision leads to phenotypic differences between Stra8-iCre; Mov10 l1(lox/lox) and Stra8-iCre; Mov10 l1(lox/Delta) mice. *Genesis* 51, 481–490.
- Bartell JG, Fantz DA, Davis T, Dewey MJ, Kistler MK & Kistler WS. (2000) Elimination of male germ cells in transgenic mice by the diphtheria toxin A chain gene directed by the histone H1t promoter. *Biol Reprod* 63, 409–416.
- Beattie MC, Adekola L, Papadopoulos V, Chen H & Zirkin BR (2015). Leydig cell aging and hypogonadism. *Exp Gerontol* 68, 87–91.
- Bergh A. (1985) Effect of cryptorchidism on the morphology of testicular macrophages: evidence for a Leydig cell-macrophage interaction in the rat testis. *Int J Androl* 8, 86–96.
- Bergh A, Damber JE & van Rooijen N. (1993a). Liposome-mediated macrophage depletion: an experimental approach to study the role of testicular macrophages in the rat. *J Endocrinol* 136, 407–413.
- Bergh A, Damber JE & van Rooijen N. (1993b) The human chorionic gonadotrophin-induced inflammation-like response is enhanced in macrophage-depleted rat testes. *J Endocrinol* 136, 415–420.
- Berndtson WE & Thompson TL. (1990) Changing relationship between testis size, Sertoli cell number and spermatogenesis in Sprague-Dawley rats. *J Androl* 11, 429–435.
- Boettger-Tong HL, Johnston DS, Russell LD, Griswold MD & Bishop CE. (2000) Juvenile spermatogonial depletion (jsd) mutant seminiferous tubules are capable of supporting transplanted spermatogenesis. *Biol Reprod* 63, 1185–1191.
- Brennan J, Tilmann C & Capel B. (2003) Pdgfr-alpha mediates testis cord organization and fetal Leydig cell development in the XY gonad. *Genes Dev* 17, 800–810.
- Brinster RL. (2002) Germline stem cell transplantation and transgenesis. *Science* 296, 2174–2176.
- Brinster RL & Zimmermann JW. (1994) Spermatogenesis following male germ-cell transplantation. *Proc Natl Acad Sci USA* 91, 11298–11302.
- Brinster CJ, Ryu BY, Avarbock MR, Karagenc L, Brinster RL & Orwig KE. (2003) Restoration of fertility by germ cell transplantation requires effective recipient preparation. *Biol Reprod* 69, 412–420.
- Buch T, Heppner FL, Tertilt C, Heinen TJ, Kremer M, Wunderlich FT, Jung S & Waisman A. (2005) A Cre-inducible diphtheria toxin receptor mediates cell lineage ablation after toxin administration. *Nat Methods* 2, 419–426.
- Chang C, Chen YT, Yeh SD, Xu Q, Wang RS, Guillou F, Lardy H & Yeh S. (2004) Infertility with defective spermatogenesis and hypotestosteronemia in male mice lacking the androgen receptor in Sertoli cells. *Proc Natl Acad Sci USA* 101, 6876–6881.
- Chen H, Huhtaniemi I & Zirkin BR. (1996) Depletion and repopulation of Leydig cells in the testes of aging brown Norway rats. *Endocrinology* 137, 3447–3452.

- Chen LY, Brown PR, Willis WB & Eddy EM. (2014) Peritubular myoid cells participate in male mouse spermatogonial stem cell maintenance. *Endocrinology* 155, 4964–4974.
- Choi YJ, Ok DW, Kwon DN, Chung JI, Kim HC, Yeo SM, Kim T, Seo HG & Kim JH. (2004) Murine male germ cell apoptosis induced by busulfan treatment correlates with loss of c-kit-expression in a Fas/FasL- and p53-independent manner. *FEBS Lett* 575, 41–51.

Christensen AK & Peacock KC. (1980) Increase in Leydig cell number in testes of adult rats treated chronically with an excess of human chorionic gonadotropin. *Biol Reprod* 22, 383–391.

Chuma S, Kanatsu-Shinohara M, Inoue K, Ogonuki N, Miki H, Toyokuni S, Hosokawa M, Nakatsuji N, Ogura A & Shinohara T. (2005) Spermatogenesis from epiblast and primordial germ cells following transplantation into postnatal mouse testis. *Development* 132, 117–122.

Clark AM, Garland KK & Russell LD. (2000) Desert hedgehog (Dhh) gene is required in the mouse testis for formation of adult-type Leydig cells and normal development of peritubular cells and seminiferous tubules. *Biol Reprod* 63, 1825–1838.

Clough SR, Welsh MJ, Payne AH, Brown CD & Brabec MJ. (1990) Primary rat Sertoli and interstitial cells exhibit a differential response to cadmium. *Cell Biol Toxicol* 6, 63–79.

Clouthier DE, Avarbock MR, Maika SD, Hammer RE & Brinster RL. (1996) Rat spermatogenesis in mouse testis. *Nature* 381, 418–421.

Davidoff MS, Middendorff R, Enikolopov G, Riethmacher D, Holstein AF & Muller D. (2004) Progenitor cells of the testosterone-producing Leydig cells revealed. *J Cell Biol* 167, 935–944.

- De Franca LR, Bartke A, Borg KE, Cecim M, Fadden CT, Yagi A & Russell LD. (1994) Sertoli cells in testes containing or lacking germ cells: a comparative study of paracrine effects using the W (c-kit) gene mutant mouse model. *Anat Rec* 240, 225–232.
- De Gendt K, Swinnen JV, Saunders PT, Schoonjans L, Dewerchin M, Devos A, Tan K, Atanassova N, Claessens F, Lecureuil C, Heyns W, Carmeliet P, Guillou F, Sharpe RM & Verhoeven G. (2004) A Sertoli cell-selective knockout of the androgen receptor causes spermatogenic arrest in meiosis. *Proc Natl Acad Sci USA* 101, 1327– 1332.
- Dym M & Fawcett DW. (1970) The blood-testis barrier in the rat and the physiological compartmentation of the seminiferous epithelium. *Biol Reprod* 3, 308–326.
- El Gehani F, Zhang FP, Pakarinen P, Rannikko A & Huhtaniemi I. (1998) Gonadotropin-independent regulation of steroidogenesis in the fetal rat testis. *Biol Reprod* 58, 116–123.

Franca LR, Ogawa T, Avarbock MR, Brinster RL & Russell LD. (1998) Germ cell genotype controls cell cycle during spermatogenesis in the rat. *Biol Reprod* 59, 1371–1377.

Gaytan F, Bellido C, Aguilar E & van Rooijen N. (1994a). Requirement for testicular macrophages in Leydig cell proliferation and differentiation during prepubertal development in rats. *J Reprod Fertil* 102, 393–399.

Gaytan F, Bellido C, Morales C, Reymundo C, Aguilar E & Van Rooijen N. (1994b) Effects of macrophage depletion at different times after treatment with ethylene dimethane sulfonate (EDS) on the regeneration of Leydig cells in the adult rat. *J Androl* 15, 558–564.

Gaytan F, Bellido C, Morales C, Reymundo C, Aguilar E & van Rooijen N. (1994c) Selective depletion of testicular macrophages and prevention of Leydig cell repopulation after treatment with ethylene dimethane sulfonate in rats. *J Reprod Fertil* 101, 175–182.

Gaytan F, Bellido C, Aguilar E & van Rooijen N. (1995a) Pituitarytesticular axis in rats lacking testicular macrophages. *Eur J Endocrinol* 132, 218–222.

Gaytan F, Bellido C, Morales C, Reymundo C, Aguilar E & van Rooijen N. (1995b) Response to Leydig cell apoptosis in the absence of testicular macrophages. *J Reprod Immunol* 29, 81–94.

Gaytan F, Bellido C, Morales C, van Rooijen N & Aguilar E. (1995c) Role of testicular macrophages in the response of Leydig cells to

gonadotrophins in young hypophysectomized rats. *J Endocrinol* 147, 463–471.

Gaytan F, Bellido C, Morales C, Garcia M, van Rooijen N & Aguilar E. (1996) In vivo manipulation (depletion versus activation) of testicular macrophages: central and local effects. J Endocrinol 150, 57–65.

Gnessi L, Basciani S, Mariani S, Arizzi M, Spera G, Wang C, Bondjers C, Karlsson L & Betsholtz C. (2000) Leydig cell loss and spermatogenic arrest in platelet-derived growth factor (PDGF)-A-deficient mice. J Cell Biol 149, 1019–1026.

Gomes WR, Hall RW, Jain SK & Boots LR. (1973) Serum gonadotropin and testosterone levels during loss and recovery of spermatogenesis in rats. *Endocrinology* 93, 800–809.

Guo J, Zhou H, Su Z, Chen B, Wang G, Wang CQ, Xu Y & Ge RS. (2013) Comparison of cell types in the rat Leydig cell lineage after ethane dimethanesulfonate treatment. *Reproduction* 145, 371–380.

Hayashi K, Ohta H, Kurimoto K, Aramaki S & Saitou M. (2011) Reconstitution of the mouse germ cell specification pathway in culture by pluripotent stem cells. *Cell* 146, 519–532.

Hendriks S, Dancet EA, van Pelt AM, Hamer G & Repping S. (2015a) Artificial gametes: a systematic review of biological progress towards clinical application. *Hum. Reprod. Update* 21, 285–296.

Hendriks S, Dondorp W, de Wert G, Hamer G, Repping S & Dancet EA. (2015b) Potential consequences of clinical application of artificial gametes: a systematic review of stakeholder views. *Hum. Reprod. Update* 21, 297–309.

Holsberger DR & Cooke PS. (2005) Understanding the role of thyroid hormone in Sertoli cell development: a mechanistic hypothesis. *Cell Tissue Res* 322, 133–140.

Hutson JC. (1998) Interactions between testicular macrophages and Leydig cells. *J Androl* 19, 394–398.

Hutson JC, Garner CW & Doris PA. (1996) Purification and characterization of a lipophilic factor from testicular macrophages that stimulates testosterone production by Leydig cells. *J Androl* 17, 502–508.

Jackson H. (1970) Antispermatogenic agents. Br Med Bull 26, 79-86.

Jackson CM & Jackson H. (1984) Comparative protective actions of gonadotrophins and testosterone against the antispermatogenic action of ethane dimethanesulphonate. *J Reprod Fertil* 71, 393–401.

Jackson H, Partington M & Fox BW. (1962) Effect of "Busulphan" ("Myleran") on the spermatogenic cell population of the rat testis. *Nature* 194, 1184–1185.

Jackson AE, O'Leary PC, Ayers MM & de Kretser DM. (1986) The effects of ethylene dimethane sulphonate (EDS) on rat Leydig cells: evidence to support a connective tissue origin of Leydig cells. *Biol Reprod* 35, 425– 437.

Jean-Faucher C, el Watik N, Berger M, De Turckheim M, Veyssiere G & Jean C. (1983) Ontogeny of the secretory pattern of LH and FSH in male mice during sexual maturation. *Int J Androl* 6, 575–584.

Johnston DS, Russell LD, Friel PJ & Griswold MD. (2001) Murine germ cells do not require functional androgen receptors to complete spermatogenesis following spermatogonial stem cell transplantation. *Endocrinology* 142, 2405–2408.

Josso N, Picard JY, Rey R & di CN (2006). Testicular anti-Mullerian hormone: history, genetics, regulation and clinical applications. *Pediatr Endocrinol Rev* 3, 347–358.

Kanatsu-Shinohara M & Shinohara T. (2013) Spermatogonial stem cell self-renewal and development. Annu Rev Cell Dev Biol 29, 163–187.

Kangasniemi M, Huhtaniemi I & Meistrich ML. (1996) Failure of spermatogenesis to recover despite the presence of a spermatogonia in the irradiated LBNF1 rat. *Biol Reprod* 54, 1200–1208.

Kelce WR & Zirkin BR. (1993) Mechanism by which ethane dimethanesulfonate kills adult rat Leydig cells: involvement of intracellular glutathione. *Toxicol Appl Pharmacol* 120, 80–88.

Kerr JB, Donachie K & Rommerts FFG. (1985) Selective destruction and regeneration of rat Leydig cells in vivo. A new method for the study of

seminiferous tubular-interstitial tissue interaction. *Cell Tissue Res* 242, 145–156.

- Kerr JB, Bartlett JMS, Donachie K & Sharpe RM. (1987) Origin of regenerating leydig-cells in the testis of the adult- rat - an ultrastructural, morphometric and hormonal assay study. *Cell Tissue Res* 249, 367–377.
- Kerr JB, Millar M, Maddocks S & Sharpe RM. (1993) Stage-dependent changes in spermatogenesis and Sertoli cells in relation to the onset of spermatogenic failure following withdrawal of testosterone. *Anat Rec* 235, 547–559.
- Kim JM, Luo L & Zirkin BR. (2000) Caspase-3 activation is required for Leydig cell apoptosis induced by ethane dimethanesulfonate. *Endocrinology* 141, 1846–1853.
- Kim GJ, Georg I, Scherthan H, Merkenschlager M, Guillou F, Scherer G & Barrionuevo F. (2010) Dicer is required for Sertoli cell function and survival. *Int J Dev Biol* 54, 867–875.
- Kobayashi A, Chang H, Chaboissier MC, Schedl A & Behringer RR. (2005) Sox9 in testis determination. *Ann N Y Acad Sci* 1061, 9–17.
- Lecureuil C, Fontaine I, Crepieux P & Guillou F. (2002) Sertoli and granulosa cell-specific Cre recombinase activity in transgenic mice. *Genesis* 33, 114–118.
- Lei ZM, Mishra S, Zou W, Xu B, Foltz M, Li X & Rao CV. (2001) Targeted disruption of luteinizing hormone/human chorionic gonadotrophin receptor gene. *Mol Endocrinol* 15, 184–200.
- Li T, Hu J, He GH, Li Y, Zhu CC, Hou WG, Zhang S, Li W, Zhang JS, Wang Z, Liu XP, Yao LB & Zhang YQ. (2012) Up-regulation of NDRG2 through nuclear factor-kappa B is required for Leydig cell apoptosis in both human and murine infertile testes. *Biochim Biophys Acta* 1822, 301–313.
- Lording DW & De Kretser DM. (1972) Comparative ultrastructural and histochemical studies of the interstitial cells of the rat testis during fetal and postnatal development. *J Reprod Fertil* 29, 261–269.
- Lukyanenko YO, Carpenter AM, Brigham DE, Stocco DM & Hutson JC. (1998) Regulation of Leydig cells through a steroidogenic acute regulatory protein-independent pathway by a lipophilic factor from macrophages. *J Endocrinol* 158, 267–275.
- Mahato D, Goulding EH, Korach KS & Eddy EM. (2000) Spermatogenic cells do not require estrogen receptor-alpha for development or function. *Endocrinology* 141, 1273–1276.
- Marettova E, Maretta M & Legath J. (2015) Toxic effects of cadmium on testis of birds and mammals: a review. *Anim Reprod Sci* 155, 1–10.
- McLean DJ, Russell LD & Griswold MD. (2002) Biological activity and enrichment of spermatogonial stem cells in vitamin A-deficient and hyperthermia-exposed testes from mice based on colonization following germ cell transplantation. *Biol Reprod* 66, 1374–1379.
- Meachem SJ, Nieschlag E & Simoni M. (2001) Inhibin B in male reproduction: pathophysiology and clinical relevance. *Eur J Endocrinol* 145, 561–571.
- Medrano JV, Martinez-Arroyo AM, Sukhwani M, Noguera I, Quinonero A, Martinez-Jabaloyas JM, Pellicer A, Remohi J, Orwig KE & Simon C. (2014) Germ cell transplantation into mouse testes procedure. *Fertil Steril* 102, e11–e12.
- Meistrich ML, Hunter NR, Suzuki N, Trostle PK & Withers HR. (1978) Gradual regeneration of mouse testicular stem cells after exposure to ionizing radiation. *Radiat Res* 74, 349–362.
- Mendis-Handagama SM, Watkins PA, Gelber SJ & Scallen TJ. (1998) The effect of chronic luteinizing hormone treatment on adult rat Leydig cells. *Tissue Cell* 30, 64–73.
- Michael SD, Kaplan SB & Macmillan BT. (1980) Peripheral plasma concentrations of LH, FSH, prolactin and GH from birth to puberty in male and female mice. *J Reprod Fertil* 59, 217–222.
- Molenaar R, de Rooij DG, Rommerts FF, Reuvers PJ & van der Molen HJ. (1985) Specific destruction of Leydig cells in mature rats after in vivo

Molenaar R, de Rooij DG, Rommerts FFG & van der Molen HJ. (1986) Repopulation of Leydig cells in mature rats after selective destruction of the existent Leydig cells with ethylene dimethane sulfonate is dependent on luteinizing hormone and not follicle-stimulating hormone. *Endocrinology* 118, 2546–2554.

Morris ID. (1996) Leydig cell toxicology. In: *The Leydig Cell* (eds AH Payne, MP Hardy & LD Russell), pp. 573–596. Cache River Press, Vienna.

Morris ID, Phillips DM & Bardin CW. (1986) Ethylene dimethanesulfonate destroys Leydig cells in the rat testis. *Endocrinology* 118, 709–719.

Morris ID, Bardin CW, Musto NA, Thau RB & Gunsalus GL. (1987) Evidence suggesting that germ cells influence the bidirectional secretion of androgen binding protein by the seminiferous epithelium demonstrated by selective impairment of spermatogenesis with busulphan. *Int J Androl* 10, 691–700.

- Morrow CM, Hostetler CE, Griswold MD, Hofmann MC, Murphy KM, Cooke PS & Hess RA. (2007) ETV5 is required for continuous spermatogenesis in adult mice and may mediate blood testes barrier function and testicular immune privilege. *Ann N Y Acad Sci* 1120, 144– 151.
- Murphy L & O'Shaughnessy PJ (1992). Development of steroidogenic enzyme-activity in the tfm mouse testis. *J Endocrinol* 135, P6.
- Murphy L, Jeffcoate IA & O'Shaughnessy PJ. (1994) Abnormal Leydig cell development at puberty in the androgen-resistant Tfm mouse. *Endocrinology* 135, 1372–1377.
- Nagano M, Avarbock MR & Brinster RL. (1999) Pattern and kinetics of mouse donor spermatogonial stem cell colonization in recipient testes. *Biol Reprod* 60, 1429–1436.
- Nef S & Parada LF (1999). Cryptorchidism in mice mutant for Insl3 *Nat Genet* 22, 295–299.
- Nef S, Shipman T & Parada LF. (2000) A molecular basis for estrogeninduced cryptorchidism. *Dev Biol* 224, 354–361.
- Nurmio M, Kallio J, Adam M, Mayerhofer A, Toppari J & Jahnukainen K. (2012) Peritubular myoid cells have a role in postnatal testicular growth. *Spermatogenesis* 2, 79–87.
- Ogawa T, Dobrinski I, Avarbock MR & Brinster RL. (2000) Transplantation of male germ line stem cells restores fertility in infertile mice. *Nat Med* 6, 29–34.
- Ogawa T, Ohmura M, Yumura Y, Sawada H & Kubota Y. (2003) Expansion of murine spermatogonial stem cells through serial transplantation. *Biol Reprod* 68, 316–322.
- Ohta H, Yomogida K, Dohmae K & Nishimune Y. (2000) Regulation of proliferation and differentiation in spermatogonial stem cells: the role of c-kit and its ligand SCF. *Development* 127, 2125–2131.
- Ohta H, Wakayama T & Nishimune Y. (2004) Commitment of fetal male germ cells to spermatogonial stem cells during mouse embryonic development. *Biol Reprod* 70, 1286–1291.
- O'Shaughnessy PJ. (2015) Testicular development. In: *Knobil and Neill's Physiology of Reproduction* (eds P Tony & Z Anthony), pp. 567–594. Academic Press, Amsterdam.
- O'Shaughnessy P & Fowler PA. (2011) Endocrinology of the mammalian fetal testis. *Reproduction* 141, 37–46.
- O'Shaughnessy PJ, Baker P, Sohnius U, Haavisto AM, Charlton HM & Huhtaniemi I. (1998) Fetal development of Leydig cell activity in the mouse is independent of pituitary gonadotroph function. *Endocrinology* 139, 1141–1146.
- O'Shaughnessy PJ, Fleming L, Baker PJ, Jackson G & Johnston H. (2003) Identification of developmentally-regulated genes in the somatic cells of the mouse testis using serial analysis of gene expression. *Biol Reprod* 69, 797–808.
- O'Shaughnessy PJ, Hu L & Baker PJ. (2008a) Effect of germ cell depletion on levels of specific mRNA transcripts in mouse Sertoli cells and Leydig cells. *Reproduction* 135, 839–850.

administration of ethane dimethyl sulfonate. *Biol Reprod* 33, 1213–1222.

- O'Shaughnessy PJ, Morris ID & Baker PJ. (2008b) Leydig cell regeneration and expression of cell signaling molecules in the germ cellfree testis. *Reproduction* 135, 851–858.
- O'Shaughnessy PJ, Monteiro A, Fowler PA & Morris ID. (2014) Identification of Leydig cell-specific mRNA transcripts in the adult rat testis. *Reproduction* 147, 671–682.
- Palmiter RD, Behringer RR, Quaife CJ, Maxwell F, Maxwell IH & Brinster RL. (1987) Cell lineage ablation in transgenic mice by cell-specific expression of a toxin gene. *Cell* 50, 435–443.

Papaioannou MD, Pitetti JL, Ro S, Park C, Aubry F, Schaad O, Vejnar CE, Kuhne F, Descombes P, Zdobnov EM, McManus MT, Guillou F, Harfe BD, Yan W, Jégou B, Nef S. (2009) Sertoli cell Dicer is essential for spermatogenesis in mice. *Dev Biol* 326, 250–259.

- Paris MC, Snow M, Cox SL & Shaw JM. (2004) Xenotransplantation: a tool for reproductive biology and animal conservation? *Theriogenology* 61, 277–291.
- Qian Y, Liu S, Guan Y, Pan H, Guan X, Qiu Z, Li L, Gao N, Zhao Y, Li X, *et al.* (2013) Lgr4-mediated Wnt/beta-catenin signaling in peritubular myoid cells is essential for spermatogenesis. *Development* 140, 1751–1761.
- Qin J, Tsai MJ & Tsai SY. (2008) Essential roles of COUP-TFII in Leydig cell differentiation and male fertility. *PLoSONE* 3, e3285.

Rebourcet D, O'Shaughnessy PJ, Monteiro A, Milne L, Cruickshanks L, Jeffrey N, Guillou F, Freeman TC, Mitchell RT & Smith LB. (2014a) Sertoli cells maintain Leydig cell number and peritubular myoid cell activity in the adult mouse testis. *PLoS ONE* 9, e105687.

Rebourcet D, O'Shaughnessy PJ, Pitetti JL, Monteiro A, O'Hara L, Milne L, Tsai YT, Cruickshanks L, Riethmacher D, Guillou F, Freeman TC, Mitchell RT, Smith LB. (2014b) Sertoli cells control peritubular myoid cell fate and support adult Leydig cell development in the prepubertal testis. *Development* 141, 2139–2149.

- Risbridger GP, Kerr JB & Dekrester DM. (1987) Influence of the cryptorchid testis on the regeneration of rat X leydig-cells after administration of ethane dimethane sulfonateX. J Endocrinol 112, 197.
- Roberts KP & Griswold MD. (1990) Ethylene dimethanesulfonate inhibits the function of Sertoli cells in culture at sublethal doses. *Endocrinology* 126, 1618–1622.

Roosen-Runge EC & Anderson D. (1959) The development of the interstitial cells of the albino rat. *Acta Anat* 37, 125–137.

Ryu BY, Orwig KE, Oatley JM, Avarbock MR & Brinster RL. (2006) Effects of aging and niche microenvironment on spermatogonial stem cell self-renewal. *Stem Cells* 24, 1505–1511.

Sadate-Ngatchou PI, Payne CJ, Dearth AT & Braun RE. (2008) Cre recombinase activity specific to postnatal, premeiotic male germ cells in transgenic mice. *Genesis* 46, 738–742.

Saito M, Iwawaki T, Taya C, Yonekawa H, Noda M, Inui Y, Mekada E, Kimata Y, Tsuru A & Kohno K. (2001) Diphtheria toxin receptormediated conditional and targeted cell ablation in transgenic mice. *Nat Biotechnol* 19, 746–750.

Salva A, Hardy MP, Wu XF, Sottas CM, MacLaughlin DT, Donahoe PK & Lee MM. (2004) Mullerian-inhibiting substance inhibits rat Leydig cell regeneration after ethylene dimethanesulphonate ablation. *Biol Reprod* 70, 600–607.

Sauter KA, Pridans C, Sehgal A, Tsai YT, Bradford BM, Raza S, Moffat L, Gow DJ, Beard PM, Mabbott NA, *et al.* (2014) Pleiotropic effects of extended blockade of CSF1R signaling in adult mice. *J Leukoc Biol* 96, 265–274.

Saxena DK, Yoshinaga K, Tanii I & Toshimori K. (2002) Are germ cell factors essential in the testicular enlargement after neonatal hypothyroidism recovery? A study using W/Wv mutant mice model. *Int J Androl* 25, 11–18.

Selmanoff MK, Goldman BD & Ginsburg BE. (1977) Developmental changes in serum luteinizing hormone, follicle stimulating hormone and androgen levels in males of two inbred mouse strains. *Endocrinology* 100, 122–127. Sharpe RM, McKinnell C, Kivlin C & Fisher JS. (2003) Proliferation and functional maturation of Sertoli cells, and their relevance to disorders of testis function in adulthood. *Reproduction* 125, 769–784.

Shinohara T, Orwig KE, Avarbock MR & Brinster RL. (2001) Remodeling of the postnatal mouse testis is accompanied by dramatic changes in stem cell number and niche accessibility. *Proc Natl Acad Sci USA* 98, 6186–6191.

Shinohara T, Orwig KE, Avarbock MR & Brinster RL. (2003) Restoration of spermatogenesis in infertile mice by Sertoli cell transplantation. *Biol Reprod* 68, 1064–1071.

Shinomura M, Kishi K, Tomita A, Kawasumi M, Kanezashi H, Kuroda Y, Tsunekawa N, Ozawa A, Aiyama Y, Yoneda A, Suzuki H, Saito M, Picard JY, Kohno K, Kurohmaru M, Kanai-Azuma M, Kanai Y. (2014) A novel Amh-Treck transgenic mouse line allows toxin-dependent loss of supporting cells in gonads. *Reproduction* 148, H1–H9.

Smith L. (2011) Good planning and serendipity: exploiting the Cre/Lox system in the testis. *Reproduction* 141, 151–161.

Sriraman V, Sairam MR & Rao AJ. (2003) Evaluation of relative roles of LH and FSH in regulation of differentiation of Leydig cells using an ethane 1,2-dimethylsulfonate-treated adult rat model. *J Endocrinol* 176, 151–161.

Stanley E, Lin CY, Jin S, Liu J, Sottas CM, Ge R, Zirkin BR & Chen H (2012). Identification, Proliferation, and Differentiation of Adult Leydig Stem Cells. *Endocrinology* 153, 5002–5010.

Svingen T & Koopman P. (2013) Building the mammalian testis: origins, differentiation, and assembly of the component cell populations. *Genes Dev* 27, 2409–2426.

Takashima S, Kanatsu-Shinohara M, Tanaka T, Takehashi M, Morimoto H & Shinohara T. (2011) Rac mediates mouse spermatogonial stem cell homing to germline niches by regulating transmigration through the blood-testis barrier. *Cell Stem Cell* 9, 463–475.

Taylor MF, Boer-Brouwer M, Woolveridge I, Teerds KJ & Morris ID. (1999) Leydig cell apoptosis after the administration of ethane dimethanesulfonate to the adult male rat is a Fas-mediated process. *Endocrinology* 140, 3797–3804.

Teerds KJ & Rijntjes E (2007). Dynamics of Leydig cell regeneration after EDS. A model for postnatal Leydig cell regeneration. In: *The Leydig Cell in Health and Disease* (eds P Anita & H Matthew), pp. 91–116. Humana, Totowa.

Teerds KJ, De Rooij DG, Rommerts FF & Wensing CJ. (1988) The regulation of the proliferation and differentiation of rat Leydig cell precursor cells after EDS administration or daily HCG treatment. *J Androl* 9, 343–351.

Teerds KJ, de Rooij DG, Rommerts FF, van den Hurk R & Wensing CJ. (1989) Stimulation of the proliferation and differentiation of Leydig cell precursors after the destruction of existing Leydig cells with ethane dimethyl sulphonate (EDS) can take place in the absence of LH. *J Androl* 10, 472–477.

Teerds KJ, de Rooij DG, Rommerts FF & Wensing CJ. (1990) Development of a new Leydig cell population after the destruction of existing Leydig cells by ethane dimethane sulphonate in rats: an autoradiographic study. *J Endocrinol* 126, 229–236.

Verhoeven G, Cailleau J & Morris ID. (1989) Inhibitory effects of alkane sulphonates on the function of immature rat Leydig, Sertoli and peritubular cells cultured in vitro. J Mol Endocrinol 2, 145–155.

Welsh M, Saunders PT, Atanassova N, Sharpe RM & Smith LB. (2009) Androgen action via testicular peritubular myoid cells is essential for male fertility. *FASEB J* 23, 4218–4230.

Welsh M, Moffat L, Belling K, de Franca LR, Segatelli TM, Saunders PT, Sharpe RM & Smith LB. (2012) Androgen receptor signalling in peritubular myoid cells is essential for normal differentiation and function of adult Leydig cells. *Int J Androl* 35, 25–40.

- Wilhelm D, Yang JX & Thomas P. (2013) Mammalian sex determination and gonad development. *Curr Top Dev Biol* 106, 89–121.
- Winnall WR & Hedger MP. (2013) Phenotypic and functional heterogeneity of the testicular macrophage population: a new regulatory model. *J Reprod Immunol* 97, 147–158.
- Yan W, Kero J, Huhtaniemi I & Toppari J. (2000) Stem cell factor functions as a survival factor for mature Leydig cells and a growth factor for precursor Leydig cells after ethylene dimethane sulfonate treatment: implication of a role of the stem cell factor/c- Kit system in Leydig cell development. *Dev Biol* 227, 169–182.
- Zeng W, Tang L, Bondareva A, Honaramooz A, Tanco V, Dores C, Megee S, Modelski M, Rodriguez-Sosa JR, Paczkowski M, *et al.* (2013) Viral transduction of male germline stem cells results in transgene transmission after germ cell transplantation in pigs. *Biol Reprod* 88, 27.
- Zhang FP, Poutanen M, Wilbertz J & Huhtaniemi I. (2001) Normal prenatal but arrested postnatal sexual development of luteinizing hormone receptor knockout (LuRKO) mice. *Mol Endocrinol* 15, 172– 183.
- Zhang FP, Pakarainen T, Zhu F, Poutanen M & Huhtaniemi I. (2004) Molecular characterization of postnatal development of testicular

- steroidogenesis in luteinizing hormone receptor knockout mice. *Endocrinology* 145, 1453–1463.
- Zhang Z, Shao S & Meistrich ML. (2006) Irradiated mouse testes efficiently support spermatogenesis derived from donor germ cells of mice and rats. J Androl 27, 365–375.
- Zhang Z, Shao S & Meistrich ML. (2007) The radiation-induced block in spermatogonial differentiation is due to damage to the somatic environment, not the germ cells. *J Cell Physiol* 211, 149–158.
- Zhang M, He Z, Wen L, Wu J, Yuan L, Lu Y, Guo C, Zhu L, Deng S & Yuan H. (2010) Cadmium suppresses the proliferation of piglet Sertoli cells and causes their DNA damage, cell apoptosis and aberrant ultrastructure. *Reprod Biol Endocrinol* 8, 97.
- Zhang L, Wang H, Yang Y, Liu H, Zhang Q, Xiang Q, Ge R, Su Z & Huang Y. (2013) NGF induces adult stem Leydig cells to proliferate and differentiate during Leydig cell regeneration. *Biochem Biophys Res Commun* 436, 300–305.
- Zhang YF, Yuan KM, Liang Y, Chu YH, Lian QQ, Ge YF, Zhen W, Sottas CM, Su ZJ & Ge RS. (2015) Alterations of gene profiles in Leydig-cellregenerating adult rat testis after ethane dimethane sulfonatetreatment. *Asian J Androl* 17, 253–260.