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Molecular Mechanisms in Spermatogenesis

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C. YAN CHENG, PhD, is a Senior Scientist at the Population Council's Center for Biomedical Research, inside the campus of the Rockefeller University in New York City. Dr. Cheng is a native of Hong Kong, graduated from the Chinese University of Hong Kong, and received his PhD in biochemistry and cell biology in the Laboratory of Professor Barry Boettcher at the University of Newcastle. He then received postdoctoral training in the laboratory of Drs. Wayne Bardin, Neal Musto and Glen Gunsalus at the Population Council in New York City. He has since stayed there, becoming a Senior Scientist in 1990. His studies in recent decades are focused mostly on the biology of the apical ectoplasmic specialization and the blood-testis barrier in the seminiferous epithelium. Working with a team of young scientists and collaborating with Professor Will M. Lee at the School of Biological Sciences, The University of Hong Kong, Dr. Cheng and his colleagues, Drs. Dolores Mruk, Helen Yan and Elissa Wong and Professor Will Lee have recently identified a novel autocrine regulatory loop in the seminiferous epithelium. This new autocrine-based loop utilizes laminin chains, polarity complex proteins and integrins to functionally link the apical ectoplasmic specialization, the blood-testis barrier and the hemidesmosome together to coordinate the events of spermiation and BTB restructuring to facilitate spermatid and preleptotene spermatocyte movement, respectively, that occur at stage VIII of the seminiferous epithelial cycle. These findings have now provided a framework for investigators in the field to design functional studies to understand the biology of spermatogenesis. Dr. Cheng is the recipient of several NIH grants and the CONRAD Program, and he serves as ad hoc reviewer for a number of leading journals in the field, such as Journal of Cell Biology, Nature Cell Biology, PNAS, Molecular Endocrinology, and Endocrinology.
In the past thirty years, significant advances have been made in the field of reproductive biology in “unlocking” the molecular and biochemical events that regulate spermatogenesis in the mammalian testis. It was possible because of the unprecedented breakthroughs in molecular biology, cell biology, immunology and biochemistry. I am fortunate to have personally witnessed such rapid changes in the field since I was a graduate student and a postdoctoral fellow in the late ’70s through the early ’80s. In this book, entitled *Molecular Mechanisms in Spermatogenesis*, I have included a collection of chapters written by colleagues on the latest developments in the field using genomic and proteomic approaches to study spermatogenesis, as well as different mechanisms and/or molecules including environmental toxicants and transcription factors that regulate and/or affect spermatogenesis.

The book begins with a chapter that provides the basic concept of cellular regulation of spermatogenesis. A few chapters are also dedicated to some of the latest findings on the Sertoli cell cytoskeleton and other molecules (e.g., proteases, adhesion proteins) that regulate spermatogenesis. These chapters contain thought-provoking discussions and concepts which shall be welcomed by investigators in the field. It is obvious that many of these “concepts” will be updated and some may be amended in the years to come. However, they will serve as a guide and the basis for investigation by scientists in the field. Due to the page limit, I could not cover all areas of interest in this monograph; instead, I tried to present this subject area with a balanced approach.

I hope this book will be helpful to young investigators who consider entering reproductive biology to get a balanced view of the latest developments in the field. For established investigators, these chapters will be helpful for their studies in the laboratory.

I am indebted to members of my laboratory who have provided insightful and critical discussion in the course of preparing this book. I am also grateful to all the staff at Landes Bioscience, in particular Cynthia Conomos, Celeste Carlton, Kristen Shumaker, and Megan Klein, who have helped me to work on this book from its inception through publication. Furthermore, I am grateful to my colleagues who have taken their time and worked with me these past two years on their chapters amidst the intensive day-to-day routines in their laboratories: teaching, administration, research and writing manuscripts and grant applications. Finally, I also want to thank my former mentors and friends Drs. Wayne Bardin, Barry Boettcher, Neal Musto,
Glen Gunsalus and Bruno Silvestrini for their critiques, help, encouragement and discussion during my graduate student and postdoctoral years in their laboratories in different parts of the world, who have introduced me to the fascinating areas of research in reproductive biology and animal/pharmaceutical models, set up a high standard of quality research, and unknowingly shaped my scientific personality and my approach to science.

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Spermatogenesis and Cycle of the Seminiferous Epithelium

Rex A. Hess* and Luiz Renato de Franca

Abstract

Spermatogenesis is a complex biological process of cellular transformation that produces male haploid germ cells from diploid spermatogonial stem cells. This process has been simplified morphologically by recognizing cellular associations or 'stages' and 'phases' of spermatogenesis, which progress through precisely timed and highly organized cycles. These cycles of spermatogenesis are essential for continuous sperm production, which is dependent upon numerous factors, both intrinsic (Sertoli and germ cells) and extrinsic (androgens, retinoic acids), as well as being species-specific.

Introduction

Spermatogenesis is the transformation of spermatogonial cells into spermatozoa over an extended period of time within seminiferous tubule boundaries of the testis. The seminiferous epithelium (Fig. 1) consists of germ cells that form numerous concentric layers penetrated by a single type of somatic cell first identified by Enrico Sertoli in 1865. The cytoplasm of Sertoli cells extends as thin arms around all the germ cells to nurture and maintain their cellular associations throughout the process of spermatogenesis. Germ cells multiply first by repeated mitotic divisions and then by meiosis, which involves the duplication of chromosomes, genetic recombination, and then reduction of chromosomes through two cell divisions to produce spherical haploid spermatids that differentiate into highly compacted spermatozoa for release into the tubule lumen. To study this complex and lengthy process, spermatogenesis has been organized by several different approaches, including the more popular method of 'Staging' or the recognition of germ cell association in time and the 'phases' of spermatogenesis (mitosis, meiosis and spermiogenesis). This review will examine the stages and their cycle in the production of sperm in several species, but the mouse will receive special emphasis, as it is currently the most commonly used species in research.

Cellular Components—Stages of Spermatogenesis

The seminiferous epithelium consists of only one somatic cell type, the Sertoli cell, but many different germinal cell types. The complexity of this epithelium was simplified when Leblond and Clermont were able to divide the epithelium into separate stages, according to the cellular associations observed in each tubular cross section. Stages of spermatogenesis are artificial definitions that are based upon rules established by the investigator. The original

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stages were defined according to changes observed in the Golgi region of spermatids, an area where the forming acrosomic system can be visualized by the periodic acid-Schiff’s reaction (PAS). In the mouse, XII stages are well defined by this method (Fig. 2). However, spermatogenesis is a continuum, which results in transitional areas being observed between two stages. In such cases, a preponderance of cell types can be used for stage identification. PAS staining and higher microscopic resolution is required to identify specific stages. However, for most research purposes, grouping stages into three categories is adequate and much easier for evaluation. For example, it is possible to group Stages I-V as ‘early’; Stages VI-VIII as ‘middle’; and Stages IX-XII as ‘late’ (Fig. 2).

Phases of Spermatogenesis

Mitosis

Spermatogonia are diploid germ cells (2n) that divide by mitosis and reside on the basement membrane (Figs. 1, 2). Currently, it is not possible to identify spermatogonial stem cells by routine microscopy, but different types of spermatogonia are recognized as type-A, intermediate, and type-B. In well-studied laboratory rodents, such as rats and mice, four classes of spermatogonia are present: undifferentiated type A spermatogonia (A single (A₁), A paired (A₂p), A aligned (A₃)); differentiated type A spermatogonia (A₁, A₂, A₃, A₄); intermediate spermatogonia (In); and type B spermatogonia (B). In these species, the different spermatogonial classes can be characterized by light and transmission electron microscopy according to the
Spermatogenesis and Cycle of the Seminiferous Epithelium

The presence and distribution of heterochromatin. It has also been suggested that undifferentiated spermatogonia, including Aai or stem cell, are located in niches of the seminiferous epithelium, which are regulated by the Sertoli cell.

Meiosis

B-spermatogonia divide by mitosis forming two preleptotene spermatocytes, cells representing the beginning of meiotic prophase (Fig. 2). These small cells rest on the basement membrane, but leptotene and zygotene spermatocytes become transit and move through the blood-testis-barrier (or Sertoli-Sertoli barrier). Preleptotene, leptotene and zygotene spermatocytes are located in specific stages and are identifiable by routine microscopy, although fixation artifact results in the leptotene and zygotene cells appearing to be attached to the basement membrane. Spermatocytes are found in all stages, because meiosis is a prolonged period of spermatogenesis that extends over approximately 14 days in the mouse. Thus, any attempt to isolate specific stages of spermatogenesis for molecular analysis, will include cells of this phase. Spermatocytes are the cells of meiosis and their regulation requires a special focus. Of special note, meiotic cell division occurs in and defines a single stage (XII). In the mouse, stage XII is found in approximately 10% of the seminiferous tubular cross sections and meiotic division is completed in approximately 1 day. This cellular division goes through three categories, all occurring in stage XII: (a) meiosis I, the division of 4n cells; (b) formation of secondary spermatocytes (2n), which are larger than step 1 spermatids, but rarely are found as the only spermatocyte in a tubular cross section; and (c) meiosis II, the division of 2n secondary spermatocytes to form haploid (1n) round spermatids. Studies in rats, buffalos, rams, and pigs revealed a striking increase in size for primary spermatocytes, from preleptotene to diplotene. This increase is followed by a dramatic decrease of cell size during spermiogenesis in such a way that, due to changes in chromatin and nuclear condensation, in rats, for instance, before spermiation spermatid nuclear volume reaches only 1/50th (~500 to 10^3 µm^3) of its initial volume.

Spermiogenesis

The transformation of spherical, haploid spermatids (1n) into elongate, highly condensed and mature spermatozoa that are released into the seminiferous tubule lumen is called spermiogenesis (Fig. 2). The differentiation of spermatids proceeds through at least 4 prolonged steps (or phases): Golgi, capping, acrosomal, and maturation. These steps are useful for the identification of specific stages in the cycle of the seminiferous epithelium.

Golgi

Golgi apparatus is very important during the early steps of spermiogenesis, as the formation of the acrosome is dependent upon this organelle's ability to produce vesicles and granules containing the enzymatic components of the acrosomic system that will cover the developing sperm nucleus. Differentiation of the first three steps of round spermatid formation involves a prominent Golgi apparatus that is identified by PAS staining. Step 1 spermatids have a small, perinuclear Golgi region without an acrosomal vesicle or granule. Subsequent steps 2-3 show proacrosomal vesicles and granules within the Golgi apparatus, with the formation of a single, large acrosomal granule within a larger vesicle that will indent the nucleus (Fig. 2).

Capping

Capping involves steps 4-5 round spermatids, where the acrosomic granule touches the nuclear envelope and the vesicle begins to flatten into a small cap over the nuclear surface. In steps 6-7, the acrosomic vesicle becomes very thin and the granule flattens. Step 8 is the last round spermatid, and the acrosome flattens over approximately 1/3 of the nuclear surface. In late stage VIII, step 8 nuclei begin to change shape.
Figure 2. Legend viewed on following page.
Figure 2, viewed on previous page. Mouse Stages in the cycle of the seminiferous epithelium (I-XII). Layers depicting the cellular associations are drawn with Sertoli cells separating each stage. Along the base are photos of early, middle and late spermatid nuclei, stained with the PAS reaction and hematoxylin. Spermatogonia (A, In, B); spermatocytes (Pl: preleptotene, L: leptotene, Z: zygotene, P: pachytene, D: diakinesis, Mi: meiotic division); round spermatids (1-8); elongate spermatids (9-16). Adapted with permission from Dr. Robert E. Braun.

**Stage I.** Two generations of spermatids are found in Stages I-VIII, round and elongate spermatids. In this stage, the round spermatid nucleus is smaller than in subsequent stages and contains a typical large central nucleolus. The Golgi is also small and lacks PAS+ granular material.

**Stage II.** Small PAS+ proacrosomal granules are seen in the center of the Golgi apparatus, which is attached to the nucleus of round spermatids.

**Stage III.** An acrosomic granule is well formed within the larger round Golgi vesicle, which forms an indentation of the round spermatid nucleus.

**Stage IV.** The acrosomic granule begins to flatten in this stage.

**Stage V.** The acrosomic system is clearly defined now and there is a straight line formed by the acrosomic granule lying on the PAS+ dark line that caps the round spermatid nucleus, surrounded by the vesicle. Along the basement membrane, B-type spermatogonia are prominent.

**Stage VI.** The acrosomic system begins to spread, but remains thick and the granules are distinct. In this stage, B-type spermatogonia undergo mitosis to form preleptotene spermatocytes. Elongate spermatids begin to migrate toward the lumen.

**Stage VII.** The acrosomic system spreads across the nucleus and becomes thinner, allowing the central acrosomic granule to bulge slightly above the acrosomic vesicle. Elongate spermatids are located at the luminal edge of the tubule, but the cytoplasm covers the sperm head and about 1/2 of the tail.

**Early VII.** There is more cytoplasm covering the mid region of elongate spermatids and no cytoplasmic lobe has formed. Numerous small preleptotene cell nuclei are found on the basement membrane.

**Middle VII.** The cytoplasmic lobe begins to form and elongate spermatid cytoplasm no longer covers the midpiece of tail. Large dark granules in cytoplasmic lobes are still absent.

**Late VII.** The cytoplasmic lobe is well formed and much of it is now between the sperm head and the basement membrane. Dark granules are beginning to appear distinct near the sperm head and sometimes below it. Preleptotene cell nuclei are enlarging as these cells transform into leptotene and chromatin begins to disperse into smaller, finer clumps.

**Stage VIII.** The acrosome is flattened and forms a cap that covers nearly half of the round spermatid nucleus. Many of the nuclei have migrated to the cytoplasmic plasmalemma and the acrosomic system may be oriented toward the basement membrane. Elongate spermatids are being released into the lumen through a process called spermiation, while excess spermatid cytoplasm forms large cytoplasmic lobes with large dark bodies beneath the head of step 16 spermatids.

**Stage IX.** Only one generation of spermatids is found in Stages IX-XII, the transition from round into elongate. Cross sections of step 9 spermatid nuclei are oblong, as they begin the elongation process, with the thin PAS+ acrosomic system off center and extending from the apex toward the caudal region of the nucleus. Cytoplasmic lobes fuse into very large residual bodies that are phagocytized by the Sertoli cell and disappear by Stages X-XI.

**Stage X.** The spermatid head forms a distinct protrusion with a sharp angle. Only the protrusion is covered by the PAS+ acrosome on the ventral side, while the dorsal side is covered to the caudal surface of the nucleus. Pachytene spermatocyte nuclei reach their maximum diameter prior to diplotene phase.

**Stage XI.** Step 11 spermatid nuclei become thinner, more elongated and begin to stain more intensely, indicating chromatin condensation. Diplotene spermatocyte nuclei become excessively large and begin to lose nuclear envelope as the cells enter diakinesis of meiosis I.

**Stage XII.** In this stage the most important identifying feature is the presence of meiotic and secondary spermatocytes. Step 12 spermatid nuclei are thinner and nuclear staining is intensely dark throughout except for the most caudal region. PAS+ acrosomic system forms a ventral and dorsal fin over the apical protrusion.
**Acrosomal**

Acrosomal steps 9-14 involve migration of the acrosomal system over the ventral surface of the elongating spermatid nucleus (Fig. 2). This migration of the acrosome is completed approximately by step 14 spermatid and is difficult to identify in typical histological sections, due to its presence in different planes of sections and angles or orientation. Thus, recognition of specific stages of spermatogenesis will typically rely on the acrosomal system observed in the round spermatids, rather than in the elongate cells. These spermatid steps also involve condensation of the chromatin, as the chromosomes are packed more tightly and stain more intensely with hematoxylin.

**Maturation**

Maturation steps 15-16 appear across Stages III-VIII and show fewer changes in nuclear shape and acrosomal migration. The nucleus continues to condense and the acrosome matures into a thin PAS+ structure that protrudes at the apex but covers nearly all the nucleus, except for that portion connected to the tail. Excess cytoplasm is removed in Stages VII-VIII, resulting in the formation of prominent cytoplasmic lobes and residual bodies, which contain unused mitochondria, ribosomes, lipids, vesicles and other cytoplasmic components.4,25,26

**The Cycle and Wave of Spermatogenesis**

Germ cells within each layer of the seminiferous epithelium change in synchrony with the other layers over time, producing the sequence of Stages described above (Fig. 2). The cells do not migrate laterally along the length of the seminiferous tubule; however, an unusual successive order of the Stages is observed, whereby sequential Stages occur with repetition along the length of the tubules, in a ‘wave’ of the seminiferous epithelium. That is, at least in the rodent, Stage I is followed by II, followed by III, etc. through Stage XIV, which is then repeated by Stage I. The Stages are found in ascending order from the rete testis to the center of the seminiferous tubule, where the Stages are reversed. The wave is produced by synchronous development of clonal units of germ cells through a mechanism of biochemical signaling that remains a subject of inquiry.

**Sperm Production**

The precise mechanisms by which spermatogonial stem cells (A) and other early proliferative spermatogonia (A_A) transform into differentiating spermatogonia (type A, In, and type B) and simultaneously renew their own population is now a major focus of reproductive biology. In addition to c-kit and vitamin A, which are important for differentiation of A into A1, other important factors are emerging as being involved in the regulation of spermatogonial stem cells. These proteins include the following: GFRα1, PLZF, OCT4, NGN3, NOTCH-1, SOX3, c-RET, RBM, EP-CAM, STRA8, and EE2.7,10,17,31,47,48

Spermatogonia give rise to spermatocytes after a fixed number of mitotic divisions that are characteristic of each species, as two to six differentiated spermatogonial generations have been observed in mammals (Table 1). Besides being useful for comparative studies among different species, the precise knowledge of the number of spermatogonial generations is essential for better understanding of regulatory mechanisms of spermatogenesis. Compared to many other well-known self-renewing cell systems in the body, spermatogenesis is thought to have the greatest number of cell divisions during its expansion. For instance, in mice, rats, and pigs, about ten generations of spermatogonia are necessary to form preleptotene spermatocytes from one spermatogonia stem cell (A_A_A_A_A_A_A_A_A_A_A_A_B); whereas in humans this figure is much lower and estimated to be only 4 mitotic divisions. As will be shown later, both the kinetics and rate of germ cell loss have an impact on the number of sperm produced.

Knowledge of the spermatogenic cycle length is fundamental for determining the spermatogenic efficiency and performing comparative studies among species. The total duration of
Spermatogenesis and Cycle of the Seminiferous Epithelium

Spermatogenesis based on 4.5 spermatogenic cycles ranges from approximately 30 to 78 days in mammals (8.6-8.9 for each cycle and 39-40 days for total duration in mice) (see reviews by refs. 4,23,50-51), and is under the control of the germ cell genotype, according to a study using xenogenic (rats to mice) spermatogonial transplantation. 52 Also, similar results were found utilizing porcine and ovine testis xenografts. 53 Although strain or breed differences can be found among members of the same species, the length of the spermatogenic cycle has been generally considered to be constant for a given species and is not phylogenetically determined. However, it is suggested in the literature that the temperature and some drugs may influence the duration of spermatogenesis, 54-56 probably altering the cell cycle. 57,58 In most mammals, each spermatogenic cycle lasts around 9 to 12 days, whereas the total duration of spermatogenesis lasts nearly 40 to 54 days. Particularly in humans, the entire spermatogenic process is very long and lasts more than 70 days. As a general pattern for mammals, and probably related to the synchronized development of different germ cell types per seminiferous tubule cross-sections (Stages), each phase of spermatogenesis (spermatogonial, spermatocyte, and spermatid) lasts approximately one third of the duration of the entire process.

Germ cell loss (apoptosis) occurs normally during spermatogenesis in all mammals investigated, 59 playing a critical role in determining total sperm output. However, the greatest influence on germ cell production is the capacity for mitosis, and the number of generations of spermatogonial divisions, which will dictate, at least in part, the number of cells that enter meiosis. Taking into account the number of generations of differentiated spermatogonia and

Table 1. Number of differentiated spermatogonial generations and germ cell ratios1

<table>
<thead>
<tr>
<th>Species</th>
<th>Spermatogonial Generations</th>
<th>Meiotic Index (％)2</th>
<th>Overall Rate of Spermatogenesis3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bull</td>
<td>6 (A1.3, In, B1.2)</td>
<td>3.6 (10)5</td>
<td>65 (75)</td>
</tr>
<tr>
<td>Buffalo</td>
<td>6 (A1.3, In, B1.2)</td>
<td>3.4 (15)</td>
<td>74 (71)</td>
</tr>
<tr>
<td>Ram</td>
<td>6 (A1.3, In, B1.2)</td>
<td>3.1 (23)</td>
<td>37 (85)</td>
</tr>
<tr>
<td>Goat</td>
<td>6 (A1.3, In, B1.2)</td>
<td>2.8 (30)</td>
<td>91 (65)</td>
</tr>
<tr>
<td>Boar</td>
<td>6 (A1.4, In, B)</td>
<td>3.2 (20)</td>
<td>68 (73)</td>
</tr>
<tr>
<td>Peccary</td>
<td>6 (A1.4, In, B)</td>
<td>3.2 (20)</td>
<td>74 (71)</td>
</tr>
<tr>
<td>Wild boar</td>
<td>6 (A1.4, In, B)</td>
<td>2.7 (33)</td>
<td>29 (89)</td>
</tr>
<tr>
<td>Dog</td>
<td>6 (A1.4, In, B)</td>
<td>3.4 (15)</td>
<td>51 (80)</td>
</tr>
<tr>
<td>Rat</td>
<td>6 (A1.4, In, B)</td>
<td>3.4 (15)</td>
<td>97 (62)</td>
</tr>
<tr>
<td>Mouse</td>
<td>6 (A1.4, In, B)</td>
<td>2.3-3.1 (23-43)</td>
<td>44-84 (67-83)</td>
</tr>
<tr>
<td>Gerbil</td>
<td>5 (A1.3, In, B)</td>
<td>2.8 (30)</td>
<td>34 (73)</td>
</tr>
<tr>
<td>Capybara</td>
<td>5 (A1.3, In, B)</td>
<td>2.1 (48)</td>
<td>21 (84)</td>
</tr>
<tr>
<td>Agouti pac a</td>
<td>5 (A1.3, In, B)</td>
<td>3.2 (20)</td>
<td>31 (76)</td>
</tr>
<tr>
<td>Dasyprocta sp</td>
<td>5 (A1.3, In, B)</td>
<td>3.0 (25)</td>
<td>28 (78)</td>
</tr>
<tr>
<td>Chinchilla</td>
<td>5 (A1.3, In, B)</td>
<td>3.0 (25)</td>
<td>49 (62)</td>
</tr>
<tr>
<td>Jaguar</td>
<td>5 (A1.3, In, B)</td>
<td>2.8 (30)</td>
<td>45 (65)</td>
</tr>
<tr>
<td>Cat</td>
<td>5 (A1.3, In, B)</td>
<td>2.8 (30)</td>
<td>19 (85)</td>
</tr>
<tr>
<td>Rabbit</td>
<td>5 (A1.2, In, B1.2)</td>
<td>3.3 (18)</td>
<td>39 (69)</td>
</tr>
<tr>
<td>Marmoset</td>
<td>4 (A1.2, B1.2)</td>
<td>3.4 (15)</td>
<td>25 (60)</td>
</tr>
<tr>
<td>Man</td>
<td>2 (A pale, B)</td>
<td>1.3 (68)</td>
<td>3.2 (80)</td>
</tr>
</tbody>
</table>

1 Data from our laboratory and compiled from the literature (see reviews in França and Russell, 1998; França et al. 2002; França et al, 2005). 2 Number of spermatids per each primary spermatocyte. 3 Number of spermatids formed per each differentiated type A1 spermatogonia. 4 Type A spermatogonia (A); intermediate spermatogonia (In); and type B spermatogonia (B). 5 Numbers in parentheses show the percentage of germ cell loss based on the theoretical yield.
the two meiotic divisions prior to the formation of haploid spermatids, only 2-3 spermatozoa out of 10 are produced from each differentiated type A₁ spermatogonia in most mammalian species (see overall rate of spermatogenesis in Table 1).23,60 Thus, significant germ cell loss occurs during the spermatogonial phase, called ‘density-dependent regulation’, primarily during mitotic divisions of type A₂ to A₄ spermatogonia, possibly mediated by the p53 tumor suppressor protein, as well as Bcl-2, Bax and Fas. One possibility, as yet untested, is that the degeneration is a homeostatic mechanism to limit germ cells to the number that can be supported by available Sertoli cells. Apoptosis is also frequent during meiosis (Table 1), especially in humans, and is probably related to chromosomal damage. Also, it should be mentioned that missing generations of spermatocytes and spermatids in the seminiferous epithelium, plus apoptosis, contribute to the low efficiency of human spermatogenesis.60,61

The Sertoli cell has several important roles in spermatogenesis, including the following: support and nutrition of the developing germ cells; compartmentalization of the seminiferous tubule by tight junctions, which provides a protected and specialized environment for the developing germ cells; controlled release of mature spermatids into the tubular lumen (spermiation); secretion of fluid, proteins and several growth factors; and phagocytosis of the degenerating germ cells and phagocytosis of the excess cytoplasm (residual body) that remains from released sperm.2 The Sertoli cell also mediates the actions of FSH and luteinizing hormone (LH)-stimulated testosterone production in the testis, apparently in a stage-dependent manner.62 Although it is strongly suggested that FSH plays a major role in the initiation, maintenance and restoration of spermatogenesis in primates, it appears that in most mammalian species testosterone has this important role in maintaining ‘quantitatively’ normal spermatogenesis, whereas FSH plays a qualitative role and is not strictly necessary for fertility.62 Recent investigations of the Sertoli cell specific knockout of androgen receptor (SCARKO) mouse found that spermatogenesis rarely advanced beyond diplotene spermatocytes.63 Thus, at least in this species, androgens are crucial for late meiosis and spermiogenesis.

The relative mass of seminiferous tissue determines how much space is devoted to sperm production. In general, species whose testes have a high proportion of seminiferous tubular tissue produce more sperm per unit mass (Fig. 3).23,50,60 Regardless of other factors, the number of Sertoli cells is now well established as being one of the most important determining factors that defines maximum sperm production.64-73 In all mammalian species investigated, no Sertoli cell proliferation has been observed after puberty. Thus, the perinatal and prepubertal period, when the size of the Sertoli cell population is established, ultimately dictates the magnitude of testis size and sperm production. This occurs because Sertoli cells have differing capacities to support germ cell development and each Sertoli cell is able to support only a relatively fixed number of germ cells in a species-specific manner (Fig. 3).23,50,60 Thus, animals with more Sertoli cells have more germ cells per testis, and the number of Sertoli cells per gram of tissue combined with the number of spermatids per Sertoli cell is positively correlated with sperm production per gram of testis.

There also appears to be species-specific regulation of the total Sertoli cell population.66,71,74-84 Volume density of Sertoli cells in the seminiferous epithelium changes considerably in mammals (from ~15% in mice to ~40% in humans) and is inversely related to the efficiency of sperm production. Thus, in contrast to humans, species with a lower proportion of Sertoli cells in the seminiferous epithelium, such as mice, rabbits, rats, hamsters, and pigs are among those with the highest Sertoli cell and spermatogenic efficiencies (Fig. 3).23

Daily sperm production per gram of testicular parenchyma is a measure of spermatogenic efficiency in sexually mature animals and is useful for species comparisons. In mammalian species, four to sixty million spermatozoa are produced daily per gram of testis tissue (Fig. 3), and in humans for instance approximately 1,500 spermatozoa are produced with each heartbeat. Usually, species that have shorter spermatogenic cycle lengths have higher spermatogenic efficiency (Table 1; Fig. 3). However, the higher efficiency of spermatogenesis observed in some mammalian species results from the combination of higher Sertoli cell support capacity
Figure 3. Comparative species testicular data for percentage of seminiferous tubules (%), Leydig cell or interstitial space (%), Sertoli cells (millions)/gram of testis parenchyma, spermatids per Sertoli cell, and daily sperm production per gram of testis (millions).
for germ cells and greater number of Sertoli cells per gram of testis. Data shown in Figure 3 for
the domestic boar and wild boar illustrate this assumption, because the lower Sertoli efficiency
observed for the wild boar is compensated for by the higher Sertoli cell number per gram of
testis, resulting in similar daily sperm production per gram of testis in both species. Higher
seminiferous tubule volume density (%) in the testis, lower Sertoli cell volume density (%) in
the seminiferous epithelium, greater number of spermatogonia generations, and lower germ
cell loss during spermatogenesis, also correlate significantly with spermatogenic efficiency.

**Regulation of the Cycle**

Stages in the cycle of the seminiferous epithelium are established early in the postnatal
period. For example, cellular associations suggesting specific stages have been found as early as
day 10, which is about the same time that androgen receptors (AR) begin to be expressed in
early Sertoli cells, suggesting that Sertoli cells regulate the formation of stages. Transplanta-
tion data also support this conclusion. Using the green fluorescence protein mouse (GFP), the
same stage of spermatogenesis was observed throughout a single colony, although different
colonies were in different stages, 2 months after transplantation. After 3 months, the colonies
were much larger and some had merged into a single colony. Most interestingly, these merged
colonies exhibited synchronization, as the entire colony became one stage. It was suggested that
the transplanted germ cells were probably sensitive to Sertoli cell factors that caused the fused
colonies to become one stage.

Another animal model to address the establishment of stages and cycles is the vitamin A
deficient rat, which results in an arrest of spermatogenesis, with type A1 spermatogonia differ-
entiation inhibited. Resupplementation with retinol re-establishes spermatogenesis, but
the seminiferous epithelium throughout the entire testis is synchronized within 2-3 stages.
Synchronization was found to be stable for more than 10 cycles of the epithelium and
repopulation of the epithelium appeared to be due primarily to the completion of mitotic
activity by type A1 spermatogonia, which were arrested in the G2 phase of their cycle. Thus,
in this model, it appears that the regulation involves both Sertoli and germ cell responses to
vitamin A. Sertoli cells appear to maintain the correct stages, although synchronized, over
time, while the germ cells respond to retinol to continue the correct cellular cycle by comple-
tion of G2. An investigation of the retinoic acid receptor knockout mouse (RARα-/-) further
revealed that vitamin A may be involved in the initial establishment of stages and their long-term
regulation, which also appears to be stage-specific, as the first wave of spermatogenesis was
arrested at step 8-9 spermatids and preleptotene and leptotene spermatocytes in stage VIII-IX
were delayed in the first three waves.

Sertoli cells do appear to regulate the cellular associations or ‘stages’ within the epithelium;
therefore, it has been logical to hypothesize that Sertoli cells may also regulate the ‘duration of
the cycle of the seminiferous epithelium’. Morphological intimacy between Sertoli and germ cells was
first observed in the 19th Century and today we know that up to 50 different germ cells may
contact a single Sertoli cell and that a single germ cell can be associated with several Sertoli cells.

Based upon this ‘Mother cell’ concept, early studies using transplant technology hypothesized
that if rat germ cells were transplanted into the mouse testis, the rat germ cells may acquire the
mouse testis duration of the cell cycle, 8.6 days versus 12.9 days for the rat. However, a
subsequent experiment demonstrated “the complete domination of rat germ cell genotype in
differentiation timing.” Thus, it appears that the germ cell determines duration of the sperma-
togenic cycle, while the resident Sertoli cell is responsible for maintenance of cellular associations or
stages through the production and secretion of important factors and providing proper physical
and functional environment for spermatogenesis development.

It has been known for many years that the first wave of spermatogenesis proceeds faster than
does the adult seminiferous epithelial cycle. Stage frequency in cross sections is the same on
days 13, 23, 30 and in adult mouse testes and the same was found in the rat; however,
mean duration of the cycle from 10 to 30 days was approximately 1 day shorter than in the
adult rodents.\textsuperscript{85,102,103} When germ cells from the GFP mouse were transplanted into the adult testis, the rate of growth was 2x faster during the first 2 months compared to the third month post transplant.\textsuperscript{8} It is often pointed out that this reduction in the rate of the spermatogenetic cycle during development is correlated with testicular descent; therefore, it is possible that a higher intra-abdominal temperature may result in acceleration of the cell cycle and mitotic events, as observed in fish (tilapias) maintained at elevated temperatures.\textsuperscript{58} Although this explanation may have some credibility, other data suggest that the first wave may be different simply because the spermatogonia are filling clonal niches and establishing an epithelial wave. For example, the first wave has a unique regulation that is dependent on a subpopulation of neurogenin 3 (Ngn3) negative spermatogonia that differentiate into the first wave germ cells, while Ngn3\textsuperscript{+} cells are reserved for stem cells and subsequent waves of spermatogenesis.\textsuperscript{28}

Finally, there are numerous studies showing stage and Sertoli cell specific expressions of proteins and it appears that different stages have different dependences upon androgens and FSH, with these factors having a greater influence just before spermiation in stage VII-VIII.\textsuperscript{62,104-109} However, understanding the individual contribution of factors to the maintenance of the cycle and stages of spermatogenesis will require careful analysis and interpretation, because disruption of individual factors will often lead to an initial stage-specific and/or cell-specific effect, but the long term consequence is secondary degeneration of the entire process of spermatogenesis.\textsuperscript{110,111} This very important aspect of mammalian spermatogenesis is still poorly understood and should be a focus of intensive research in the coming years, mainly because appropriate animal models are now available for dissecting molecular regulation of the cycle of the seminiferous epithelium.

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Testicular Development and Spermatogenesis:
Harvesting the Postgenomics Bounty

Antoine D. Rolland, Bernard Jégou and Charles Pineau*

Introduction

Spermatogenesis is a sophisticated process facilitating transmission of the genetic patrimony and, thus, perpetuation of the species. Mammalian spermatogenesis is classically divided into three 3 phases. In the first—the proliferative or mitotic phase—primitive germ cells or spermatogonia undergo a series of mitotic divisions. In the second—the meiotic phase—the spermatocytes undergo two consecutive divisions to produce the haploid spermatids. In the third—spermiogenesis—spermatids differentiate into spermatozoa. The entire process is regulated by paracrine, autocrine and endocrine pathways, an array of structural elements and chemical factors modulating somatic and germ cell activity (for reviews, see refs. 1-4). The communication network linking the various cellular activities during spermatogenesis is highly complex and sophisticated.5,6

Determination of the function and regulation of genes and their products is one of the key objectives of human biology. The advances in molecular biology and genomics of the last 20 years have greatly improved our global knowledge of spermatogenesis, by identifying numerous genes essential for the development of functional male gametes (for reviews, see refs. 7,8). Significant progress has recently been made in the large-scale analysis of testicular function, deepening our insight into normal and pathological spermatogenesis. Several laboratories have built on rapid progress in genome sequencing and microarray development, by carrying out genome-wide expression studies, leading to the identification of hundreds of genes differentially expressed within the testis (for review see ref. 9). The development of tools for high-throughput protein identification has allowed a few laboratories to undertake differential protein profiling expression studies and/or the systematic analysis of testicular proteomes from various species, based either on the entire organ10 or on isolated cells.11-13

This chapter reviews the current state of large-scale gene expression analyses of spermatogenesis, from gonad development during sex determination to hormonal regulation. It also deals with the advantages and limitations of transcriptomics and proteomics for studies of the expression program of testicular germ cells. Finally, the concept of systems biology—which involves integrative ‘omics’ (i.e., combining genomics, transcriptomics and proteomics) together with bioinformatics and modeling—is discussed.

Gene Expression Profiling Technologies: The Underlying Differences

Several different technologies are now available for studying gene expression by assessing mRNA and protein levels. However, there are fundamental differences between these technologies.
Transcriptome and Transcriptomics

The complete set of ribonucleic acid (RNA) transcripts produced by the genome in any given organism is called the transcriptome. Transcriptomics—the global analysis of gene expression, also called genome-wide expression profiling—is now widely used to investigate the genes and pathways involved in various biological processes. This approach is based on the principle that genes with similar patterns of expression may have related functions and may be regulated by the same genetic control mechanism. Common technologies for the genome-wide or high-throughput analysis of gene expression include spotted and oligonucleotide microarrays and tag-based approaches, such as serial analysis of gene expression (SAGE), cap-analysis gene expression (CAGE) and gene identification signature (GIS) methods. Microarray methods are dependent on the choice of sequences to be screened at the outset, whereas sequencing-based approaches require no such prior sequence selection. The various transcriptomic technologies used in these two approaches allow scientists to study tens of thousands of genes simultaneously, rather than considering one gene at a time.

Proteome and Proteomics

The word “proteome”—a contraction of “protein” and “genome”—was first coined by Marc Wilkins at the 4th 2D Gel Electrophoresis Meeting in Sienna in 1994.14 This term encapsulates the complex and dynamic nature of protein production, at reference points spanning from individual cells to organisms. Genomes are essentially identical in the different cells of an organism, whereas proteomes and transcriptomes vary between cells, over time and as a function of environmental stimuli and stress.

Proteomics research deals with the temporal dynamics of protein production in a given biological compartment at a given time. Until recently, “proteins” were considered solely as the direct products of genes for the purposes of this definition. The definition of proteomics has recently been altered so that this field now covers not only direct gene products, but also proteins undergoing structural alterations due to cell metabolism and turnover (posttranslational modifications).15 The experimental basis of proteomics, which has become one of the most important areas of research in the postgenomics era, is not new. Nonetheless, proteomics has undoubtedly benefited from unprecedented advances in genome sequencing, bioinformatics and the development of robust, sensitive, reliable and reproducible analytical techniques.

Transcriptomics Versus Proteomics, Finding the Right Balance

The recent completion of the first high-quality drafts of the mouse16 and human17,18 genomes has provided scientists with access to a wealth of relevant sequence information essential for the systematic and comprehensive characterization of gene product function. These genome sequencing projects surprisingly revealed that mammalian genomes contain far fewer protein-coding genes than previously thought. The mouse and human genomes have each been found to contain about 22,000 genes (Ensembl19 release 43), corresponding to 22,000 functional proteins according to the original one gene-one protein dogma of molecular biology. However, alternative splicing can routinely generate 100,000 proteins from 22,000 genes.20,21 If we include posttranslational modifications (e.g., phosphorylation, glycosylation and proteolysis),22,23 then the 22,000 genes may give rise to a million proteins,24,25 each with different functions. This complexity of multilayered gene expression mechanisms is partly responsible for the frequently reported discrepancies between mRNA and protein abundance.26-28 Thus, although transcriptomics probably still has a greater throughput capacity than proteomics, it is clear that protein diversity cannot be fully characterized by gene expression analyses alone. Great attention must be paid to selecting the most appropriate methods for large-scale experiments, according to whether the biological question addressed relates more to the transcriptional and/or splicing mechanisms underlying a particular process or to role in the process considered of the proteins generated and their subtly different isoforms.
Gonad Development During Sex Determination

The primary event in mammalian sexual development is the differentiation of the bipotential gonad into either a testis or an ovary. The transient expression of a single gene, Sry (sex-determining region, chromosome Y), in the supporting cell lineage of mice, from embryonic day (ed) 11.5 onwards, is necessary and sufficient to direct the differentiation of an XY gonad into a testis. This suggests that ovary development may be the “default” pathway of gonad differentiation (Fig. 1). Small et al.\(^{29}\) investigated the molecular mechanisms underlying gonad differentiation, using high-density oligonucleotide microarrays to compare the gene expression profiles of male and female mouse gonads from 11.5 to 18.5 days post-coitum (dpc). They reported the differential expression of thousands of genes during the development of ovaries or testes and between gonads at various time points. The genes identified as differentially expressed included all those previously identified as involved in this process, such as Sry itself, Sox9, Dax1, Sf1 and Wt1. Major differences between the sexes were found for genes encoding proteins involved in meiosis, steroidogenesis and apoptosis, consistent with the initiation of meiosis I in the ovary, the differentiation of Leydig and Sertoli cells and the burst of cell apoptosis observed in the testis. Nef et al.\(^{30}\) carried out a similar study on transgenic mice synthesizing green fluorescent protein (GFP) in the somatic compartment of the genital ridge of both male and female embryos. They specifically isolated somatic cells, including both supporting cell (future Sertoli and granulosa cells) and steroidogenic cell (future Leydig and theca cells) precursors, from 10.5 to 13.5 dpc. This made it possible to monitor very early events directly associated with the Sry-signaling pathway, without germ-cell “contamination”. This is important because of the potential effects of the expression program of germ cells. At 10.5 dpc, only nine genes on the sexual chromosomes displayed differential expression between XX and XY gonads, indicating that sex determination had not yet occurred. Thereafter, the number of differentially expressed genes increased, eventually reaching more than two thousand by 13.5 dpc. Three genes encoding Cdk inhibitors were found to be more strongly expressed in the ovary than in the testis. This led the authors to suggest that the higher levels of proliferation of testicular somatic cells immediately after Sry expression—a crucial event in male sex differentiation—might be a direct or indirect consequence of the inhibitory effect of Sry on Cdk inhibitor production, rather than a reflection of the activation of gene products involved in proliferation. Another identical study investigated gene expression in differentiating mouse gonads at 10.5 and 11.5 dpc.\(^ {31}\) The results obtained were consistent with those of Nef et al.,\(^ {30}\) but with a larger number of dimorphic genes identified at 11.5 dpc, probably due to differences in normalization and data processing. Nef’s study was remarkable for the identification of several differentially expressed genes for which human orthologs mapped to loci associated with sexual disorders. These three studies demonstrate the existence of strong expression programs in both the developing testis and the ovary, calling into question the notion that ovary differentiation is the “default” pathway of gonad differentiation. One 2D gel-based proteomic study recently investigated gonad development during sex determination.\(^ {32}\) The authors compared whole gonads from mice 13.5 dpc and demonstrated differential expression between male and female gonads for 36 protein spots (6% of the total), three of which they went on to identify. The smaller proportion of proteins than of genes displaying differential expression\(^ {30}\) (10%) may be due to the larger amounts of protein required for visualization on a 2D gel. This would favor the detection of ubiquitous proteins, which are likely to be produced in larger amounts. Although only three proteins displaying differential expression between male and female gonads were identified, none of the corresponding transcripts was demonstrated as differentially expressed in any of the transcriptomic experiments described above. One protein displayed a specific phosphorylation pattern in male extracts. This pattern clearly could not be picked up through a transcriptome-based approach, and indicated the presence of potentially more abundant or active kinases in male gonads.
Testicular Development and Spermatogenesis

Gene Expression Profiling in Spermatogonial Cells

The molecular identity of spermatogonial cells and the signaling events triggering their renewal or entry into spermatogenesis remained largely unknown until recently. Several studies aimed to establish reference proteome maps for these cells, with the aim of gaining insight into...
their biology (Fig. 2). These studies included proteomic analyses of cultured primordial germ cells from chicken and freshly isolated rat spermatogonia. Another study provided information about discrete low-copy number proteins, through the prefractionation of protein cell extracts on 2D gels with a narrow pH range. This work is currently being pursued, with reverse-phase HPLC used to separate the extracts into subproteome pools before protein identification (Couvet et al, unpublished). Global approaches of this type, designed to decipher a static proteome, should be rewarding in the long term. Indeed, the availability of genome sequence data has generated an urgent need for systematic protein identification for elucidation of the encoded protein networks governing cellular function. Large-scale protein-protein interaction maps have generally been based on results obtained with the yeast two-hybrid system, which detects only binary interactions (for review, see ref. 35). However, the advent of highly sensitive protein identification methods based on mass spectrometry has made it feasible to identify protein complexes directly, at the proteome-wide scale (for review, see ref. 18).

The identification of several spermatogonial markers and the recent development of culture systems in rodents have made it possible to carry out gene expression profiling on spermatogonial cells in various developmental states (Fig. 2). Hamra et al used enriched preparations of rat spermatogonial stem cells (SSC) (type I collagen-non-binding/laminin-binding germ cells) cultured on different feeder cell lines to identify genes associated with the maintenance (on MSC-1 cells) or loss (on STO cells) of stem cell activity. As many as 248 genes were found to be downregulated during the loss of stem cell activity, their level of transcription remaining stable whilst this activity was maintained. These genes are therefore probably involved in

![Gene expression profiling of SSC undergoing self-renewal or differentiation in rodents](image)

Figure 2. Pathways for the self-renewal and differentiation of spermatogonial stem cells (SSC). The daily production of millions of spermatozoa in mammals is ensured by the presence within the male germ line of stem cells able to maintain their own stock and to differentiate and continuously initiate new waves of spermatogenesis. The balance between proliferation, differentiation and the maintenance of spermatogonial stem cells is governed by intrinsic factors, such as PLZF, and by signals from the testicular stem cell niche, including Sertolian signals, such as GDNF or ERM. GDNF: glial cell line-derived neurotrophic factor; ERM: Ets-related molecule; PLZF: promyelocytic leukemia zinc finger.
self-renewal rather than differentiation. The authors focused on a subset of 115 genes for which mouse homologs also displayed downregulation during germ cell differentiation in vivo, including Bcl6b (see below, ref. 37). Mean expression levels for these genes, referred to as the "stem cell index" (SCI), were strongly correlated with SSC activity, and with the expression of individual genes, such as Erg3, expressed only in undifferentiated stem cells. Other studies have investigated specific pathways related to single factors involved in SSC self-renewal and spermatogonial differentiation, as demonstrated by targeted disruption in the mouse. Zfp145-null mice lack a transcriptional repressor specifically expressed in spermatogonia in the testis (promyelocytic leukemia zing finger, PLZF) and are unable to maintain their spermatogonia. An analysis of gene expression in isolated spermatogonia (α6-integrin-positive cells) from one-week-old mutant mice identified more than 230 genes as differentially expressed with respect to the wild type. Analyses of gene expression have also been used to investigate the signaling pathway triggered by the sertolian factor Ets-related molecule (ERM) in mutant mice with impaired spermatogenesis during adulthood. The authors studied total testes from four-week-old mice—no obvious phenotype being visible at this age—and identified specific alterations in the pattern of expression of many spermatogonial genes, thereby demonstrating an effect of ERM on the SSC/spermatogonial expression program. They also showed that a large number of genes were differentially expressed in isolated Sertoli cells. The genes concerned were found to encode secreted factors previously reported to regulate the hematopoietic stem cell niche, and which also seemed to regulate SSC behavior. The SSC self-renewal and differentiation pathways mediated by one such factor, the glial cell line-derived neurotrophic factor (GDNF), have also been analyzed by gene expression profiling experiments. Hofmann et al sorted GRFα-1 (the GDNF coreceptor)-positive germ cells from six-day-old mice after isolation by sedimentation under gravity and differential plating. They monitored the gene expression profiles of these cells cultured in the presence or absence of GDNF for 10 hours. They identified more than a thousand genes differentially expressed in the presence and absence of GDNF and focused on one upregulated gene, that encoding fibroblast growth factor-receptor 2 (FGFR2). They found that bFGF amplified the proliferative effects of GDNF on SSC in culture and concluded that GNDF rendered germ-line stem cells more responsive to bFGF. Oatley et al cultured Thy1-positive cells from adult mice on STO feeders (which do not maintain SSC activity), with or without GDNF/GFRα-1. They identified 199 genes that were downregulated 18 hours after the elimination of GNDF/GFRα-1 from the medium ("self-renewal"-associated genes) and 79 genes that were upregulated in these conditions ("differentiation"-associated genes). They found 193 and 63 genes to be upregulated and downregulated, respectively, following the reintroduction of GNDF/GFRα-1 (2, 4 and 8 hours later). They then focused on the transcriptional repressor gene Bclb6, one of the six genes both downregulated following the elimination of GNDF/GFRα-1 and upregulated following its reintroduction, regardless of the reintroduction time (these genes also included Erg3, see above ref. 36). The pattern of changes in Bclb6 expression demonstrated that the protein encoded by this gene played a significant role in SSC maintenance. In vitro, cultured SSC treated with Bclb6-siRNA were smaller, fewer in number and had a lower colonization capacity than untreated cells, whereas, in vivo, Bclb6-null mice were found to have a higher proportion of tubules displaying impaired spermatogenesis than wild-type mice. These studies have identified many factors as involved in SSC signaling pathways, but they also highlight the importance of these factors for other stem cell lineages. These findings suggest that the expression of a specific combination of factors, with its own specific regulatory system, rather than the expression of germ line-specific factors may account for the unique ability of SSC to promote spermatogenesis (see below).

Postnatal Testis Development and Spermatogenesis

The development of germ cells through spermatogenesis is probably the aspect of male reproduction most frequently investigated in large-scale experiments. Spermatogenesis—an
amazing cellular differentiation process leading to the daily production of millions of spermatidzoa—involves the coordinated expression of specific genes and the generation of specific gene products at each step of the process, together with continuous communication between developing germ cells and testicular somatic cells.\(^1\)

Basic strategies have been used to address this issue through the global characterization of genes and proteins expressed in the testis or germ cell-enriched samples of animals, from invertebrates through to humans (Fig. 3). These strategies include the systematic identification of chromatin-associated proteins in *C. elegans* germ cells,\(^41\) insoluble chromatin-associated proteins in mouse elongated spermatids,\(^42\) testicular proteins in pig,\(^10\) and mouse\(^43\) and analysis of germline gene expression in *C. elegans,\(^44\) mouse\(^45,46\) and human\(^47\) testis. Some of these studies used filtering strategies to identify germ line-specific, male germ line-specific or testis-specific genes and proteins, but none was able to provide meaningful results deepening our understanding of spermatogenesis, given the amount of data generated and the various cell types from which they originated.

Several groups have also undertaken more sophisticated analyses, based on different strategies, to identify genes or proteins preferentially or specifically expressed at each stage of spermatogenesis. One of these strategies involved comparisons of different categories of purified germ cells, combined with a SAGE experiment in mice,\(^48\) GeneChip microarray experiments in mice\(^49-51\) and rats,\(^52,51\) or differential proteomic analysis in the rat\(^13\) (Fig. 3). Another involved comparisons of total testis samples from animals of various ages during the first wave of spermatogenesis, in mice, and involved differential display analysis,\(^53\) spotted PCR microarray experiments\(^54,55\) and GeneChip microarray experiments\(^49,56\) (Fig. 3). Both strategies have advantages and disadvantages. The use of isolated cells makes it possible to detect the differential expression of transcripts produced in only small amounts that could not be detected in total testis samples, but the time-consuming isolation procedures required may also alter the expression pattern. Conversely, the use of total testes overcomes the problem of artifacts related to sample preparation and can allow more precise profiling by increasing the number of time points analyzed. However, the observed changes in expression cannot be unambiguously attributed to any particular cell type. Indeed, during the postnatal testis development, the changes in expression observed in such experiments result not only from the different types of germ cell, but also from the various somatic cells, which may display changes in expression pattern during this period.\(^57,58\) This matter was partially resolved by Shima et al, who used both approaches in parallel.\(^49\) They first identified transcripts particularly abundant in each cell type (isolated type A and B spermatogonia, pachytene spermatocytes, round spermatids, Leydig cells, Sertoli cells and peritubular cells) and then monitored levels of these transcripts during the ontological development of the testis. They found that the pattern of gene expression during postnatal development was consistent with the cell type of origin for most transcripts. For example, somatic and premeiotic genes displayed a rapid decrease in expression due to their “dilution” by the genes expressed in maturing germ cells, whereas meiotic and postmeiotic genes displayed a dramatic increase in expression from puberty onwards. This suggests that neither changes in expression during isolation nor the diversity of cell types in total testis samples pose a real problem for the monitoring of spermatogenesis. Correlations have also been reported between expression profiling experiments, not only between purified germ cells and total testis samples, but between species.\(^9\) Indeed, the high reproductibility of commercial GeneChips makes cross-species comparisons and the inclusion of external data possible (see below ref. 51). Furthermore, PCR microarrays can be used to identify new genes or transcripts not yet picked up. This advantage is clearly illustrated by two studies using spotted PCR microarrays consisting of testis-subtracted and germ cell-enriched libraries to investigate the testicular transcriptome of mice carrying Y-chromosome deletions.\(^59,60\) These studies identified several new X- and Y-linked spermatid transcripts as up- or downregulated. These transcripts and the corresponding
Testicular Development and Spermatogenesis

The use of these different approaches has led to the identification of thousands of genes differentially expressed during germ cell development. This massive body of data is of great interest for an overall understanding of spermatogenesis, but its mining for the identification of key factors remains a huge challenge, as high-throughput in vivo gene inactivation is not yet feasible in rodents. Many other strategies are possible, including identification of the genes operating downstream from relevant transcription factors and focusing on testis-specific genes or genes conserved throughout evolution. We recently compared mouse, rat and human spermatogenesis transcriptomes. Several thousands of genes were found to be differentially expressed in the three species, and about one thousand orthologs were identified. It was not
possible to prepare some samples for humans, so we instead based our comparison of expression patterns on the high degree of correlation between rodent profiles. This filtering strategy identified 888 orthologs with patterns of expression conserved, not only in rodents, but also in humans, suggesting that strong selection constraints operate, allowing these orthologs to achieve their functions at a given step. We also compared our data with those available for 17 somatic tissues from mice and found that most of the testis-specific genes were actually meiotic and postmeiotic genes. These findings indicated that the functional identity of both Sertoli cells and spermatogonia was probably more closely related to the expression of a specific combination of genes rather than to the expression of very specific genes.

In addition to these analyses of testis and germ cell expression patterns at the transcriptome or proteome levels, several studies have also investigated correlations between such data or have tried to take the regulation of translation into account in their experiments. Multidimensional protein identification technology (MudPIT) identified more than 1600 proteins in a human tissue-profiling experiment, and it was possible to compare the expression profiles of 683 of these proteins unambiguously with those obtained in microarray experiments. Surprisingly, the gene expression profiles of all organs tissues clustered together, as did the protein expression profiles of these organs indicated that transcriptome or proteome patterns from different organs were more similar than were transcriptome and proteome patterns for the same organ. Differences in sensitivity between methods may bias data comparisons, accounting for these results. However, the authors also found that the testis displayed the weakest correlation between transcriptome and proteome data of any of the eight organs studied, with a correlation coefficient of 0.138, whereas the liver displayed the strongest correlation, with a correlation coefficient of 0.432. This weak correlation may be the consequence of particular aspects of gene/transcript regulation during spermatogenesis, such as mRNA storage in free RNP particles with repressed translation or delays between transcription and translation. This issue was specifically addressed in a microarray experiment monitoring the movement of mRNAs between RNPs and polysomes during meiotic and postmeiotic mouse testis development. More than 700 translationally regulated transcripts (with redistribution of at least 20% of mRNAs between the free RNPs and the polysomal fractions) were identified. Most of the transcripts identified displayed an upregulation of translation during late spermiogenesis, a common regulatory mechanism compensating for the cessation of transcription from mid-spermiogenesis onwards. A small cluster of meiotic mRNAs translated only in postmeiotic cells was also identified. The translational regulation of the genes identified in this study may not necessarily lead to a significant difference in RNA and protein expression profiles. For example, an mRNA may be produced in large amounts but inefficiently translated at one stage, whereas it may be produced in small amounts but efficiently translated at another. It is entirely possible for there to be larger amounts of protein present in the first of these cases than in the second, despite the much lower translation efficiency. Several genes identified by Iguchi as displaying translational regulation were found to have similar mRNA and protein expression patterns in our transcriptome and proteome data for male germ cells in the rat (Table 1). Conversely, certain genes with divergent RNA and protein patterns were not found to be translationally regulated. These apparent discrepancies may be accounted for by differences in the species studied. However, some of our rat data profiles were confirmed by in situ analyses in mice. The observed discrepancies may also result from additional mechanisms affecting the ratio of transcript to protein, such as protein stability and turnover.

Focus on Spermatozoa

Transcriptomics usually generates larger datasets than proteomics, but spermatozoa are a special case, for which proteome-based studies have probably generated more relevant data. Most of the studies described in this review were carried out on rodents or non mammalian models. Human samples have been widely studied only in investigations concerning sperm constituents and fecundation. Such studies are therefore the only studies in which human
### Table 1. A first set of genes whose mRNA and protein expression levels were compared in transcriptomic and proteomic differential analyses of rat spermatogenesis

#### Correlated RNA and Protein Expression Patterns

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"Correlated" genes displayed similar mRNA and protein expression levels while "uncorrelated" genes displayed discrepancies in mRNA and protein expression levels in between 2 germ cell types. Adapted from references 13 and 51.
Testicular Development and Spermatogenesis

physiopathology has been addressed directly. Spermatozoa are transcriptionally inactive and contain only very small amounts of RNA. Nevertheless, several groups have investigated the human sperm transcriptome, through both microarray-based experiments, and SAGE library construction (Fig. 4). These studies highlighted an unexpected diversity of mRNA species and led to two major conclusions: (1) The RNA content of spermatozoa is representative of spermatogenesis and can therefore be used to assess sperm quality and/or to explore certain cases of male sterility and (2) the spermatozoon may deliver more than just its DNA to the oocyte. It may also concern coding and noncoding transcripts playing an important role in early zygote development. However, it remains difficult to discriminate between the thousands of sperm transcripts that will be translated into “true spermatozoon proteins” and residual transcripts used at earlier steps of spermatogenesis.

The possibility of recovering large numbers of spermatozoa in highly pure preparations has led to many groups trying to decipher the proteome of mature male gametes. Various strategies have been used (Fig. 4). Martínez-Heredia et al aimed to map the human sperm proteome, using 2DE combined with MALDI-TOF MS, leading to the resolution of over 1000 protein spots and the unambiguous identification of 98 proteins. More detail of the human sperm proteome emerged from the differential extraction of proteins followed by nano-LC-MS/MS analysis. This approach led to the identification of 1760 proteins (of the 2300 predicted by the authors to be present in human sperm), constituting the largest catalog of proteins potentially involved in or important for fertilization and a myriad of potential contraceptive targets.

In an experiment similar to that carried out by Martínez-Heredia, but with greater success, 382 of 600 2DE protein-spots, corresponding to 342 unique proteins, were identified in the *Drosophila melanogaster* sperm proteome. These proteins include several conserved throughout evolution, from invertebrates to mammals, and crucial for sperm function.

In addition to these attempts to characterize the complete sperm proteome, several studies have involved the use of sample prefractionation, making it possible to investigate the protein content of diverse subcellular compartments of spermatozoa, leading to localization of the proteins identified. This, in turn, makes it possible to formulate hypotheses concerning the processes in which these proteins are involved. Protein localization and the formulation of hypotheses relating to protein function are two prerequisites for proteome analysis, as defined by Anderson. Such analyses have focused on the flagellum and/or fibrous sheath, acrosomal content and sperm head membrane (Fig. 4). Sixty proteins from the accessory structures of mouse sperm flagellum—the outer dense fibers, fibrous sheath and mitochondrial sheath, recovered by sucrose density gradient centrifugation of SDS-resistant tail structures—were identified by 2DE combined with MALDI-TOF/TOF-MS/MS. Four of eight proteins identified were also found in the human sperm fibrous sheath. Both the acrosomal content (soluble proteins released after the acrosomal reaction) and membrane constituents (surface-biotinylated proteins of intact sperm and proteins from acrosomal vesicles released after acrosomal reaction) were investigated in mouse by 1DE combined with HPLC-MS/MS identification, so as to focus on proteins likely to be involved in fertilization. Several hundred proteins were identified, 114 of which were predicted to be transmembrane or signal peptide-containing proteins. One third of male mice with the corresponding gene deletions were found to be sterile or subfertile, confirming the pertinence of this strategy.

Proteomics has also been used for the investigation of posttranslational modifications during sperm maturation and capacitation and for the identification of proteins responsible for some of these modifications (Fig. 4). Thus, 2D-DIGE has been applied to rat spermatozoa from the cauda and caput epididymis, to highlight changes in protein profile during the transit of the sperm through the epididymis. Significant differences were observed for 60 protein spots, and eight proteins were identified by MALDI-TOF MS, including one protein undergoing serine phosphorylation as demonstrated by 2D western blotting. Non capacitated and in vitro capacitated mouse spermatozoa were also compared, to investigate membrane protein redistribution during capacitation. In total, 27 proteins were shown to be dissociated from lipid rafts and,
Figure 4. The main components and maturation of spermatozoa. Once released from the seminiferous epithelium, spermatozoa are transferred to the epididymis. During their transit in the epididymis, spermatozoa become motile. However, they are not yet competent for fertilization. Within the female reproductive tract, they finally acquire the ability to fertilize eggs, during a time-dependent process called capacitation, making it possible for them to undergo the acrosomal reaction and to bind to the oocyte membrane.
therefore, potentially involved in the signaling pathways associated with the initiation of capacitation. These signaling pathways included tyrosine phosphorylation, a major process that has also been specifically addressed in proteomic studies. Combining anti-phospho-tyrosine 2D western blots, IMAC, post-IMAC dephosphorylation and LC-MS/MS analysis, Ficarro et al reported the identification of more than 60 phosphorylation sites in 15 proteins from human capacitated sperm and 16 additional proteins undergoing tyrosine phosphorylation during capacitation. Proteins potentially responsible for these tyrosine phosphorlation events were investigated in capacitated bull sperm. A cytosolic fraction enriched in tyrosine kinase activity was generated by poly-Glu-Tyr affinity chromatography, with MALDI-TOF MS, QTOF MS and LC-MS/MS analyses subsequently used to identify 126 proteins.

Finally, several groups have used antisperm antibodies (ASA) from seminal plasma to identify proteins potentially involved in immunological diseases causing infertility. Six and four proteins, from membrane extracts of normal human and mouse sperm, respectively, were identified on 2D western blots with human ASA. A similar approach was also used to identify seven testicular proteins recognized by sera from rats subjected to experimental autoimmune orchitis.

**Hormonal Regulation of Spermatogenesis**

Spermatogenesis is regulated by two gonadotropins released from the anterior pituitary in response to GnRH stimulation: luteinizing hormone (LH) and follicle-stimulating hormone (FSH). LH induces the synthesis of testosterone by Leydig cells, influencing the subsequent development of peritubular and Sertoli cells, and then germ cells, whereas FSH directly induces Sertoli cell division and differentiation, thereby increasing the spermatogenetic capacity of the testis (Fig. 5).

**FSH**

In vitro cultures of rat Sertoli cells and transcriptomic experiments have led to the identification of several hundreds of genes regulated by FSH treatment. Major effects, essentially involving upregulation, were observed within the first eight hours, with fewer genes shown to be altered 24 hours after treatment. Expression profiles for several genes previously shown to be regulated by FSH were confirmed in the microarray experiment. However, as cultures at time 0 were used as a control (rather than untreated cells at each time point), it remains possible that the changes in expression profile observed for several genes reflects natural changes in gene expression by Sertoli cells during culture. Sadate-Ngatchou et al evaluated the testicular effects of FSH in vivo, using hypogonadal (hpg) mice, a model in which circulating LH and FSH are undetectable. As in the in vitro experiment, most of the effects were observed shortly after the FSH treatment (within 4 hours) and corresponded to an overall increase in gene expression. Hundreds of transcripts displayed differential expression profiles, including some encoding proteins with relevant molecular functions, such as DNA and RNA metabolism, particularly after four hours. By contrast, transcripts encoding proteins involved in cell adhesion, cell growth, cell communication and signal transduction displayed differential expression profiles throughout the analysis. Testicular regulation by FSH was also investigated by immune suppression, with 60 genes identified as differentially expressed in 18 dpp rats after four days of anti-FSH antiserum injections. These genes included some already known to be regulated by FSH and/or identified in the other transcriptomic studies, and additional genes involved in the cell cycle. There is a marked correlation between in vitro and in vivo effects, as FSH receptor expression is strongly restricted to Sertoli cells. However, it is impossible, in studies carried out in vivo, to distinguish between genes directly altered in Sertoli cells and genes indirectly altered in other cell types (particularly germ cells) in response to Sertoli cell stimulation. This is particularly true of immunosuppression experiments, in which expression profiling is carried out several days after the reduction/suppression of circulating FSH is observed.
Androgens

The hpg mouse model has been used to investigate the regulation of spermatogenesis by androgens. As hpg mice lack LH, they have little or no detectable testosterone (T). Two experiments were carried out to evaluate the effects of T on immature testes of hpg mice not previously exposed to androgens (24-hour time course experiment after a single injection of T) and more mature testes progressing through spermatogenesis (a single experiment 4 hours after an injection of T in hpg mice previously treated with T for a period of five days, followed by a 14-day washout period). Unlike the effects of FSH, the early effects of T (within the first 12 hours) on testicular expression profiles were weak, consisting of an overall decrease in expression, whereas later effects (after 24 hours) were more marked and resulted in an increase in gene expression. Interestingly, most of the genes affected by T also showed a decrease in expression, but very few genes were retrieved in both experiments, suggesting that the primary effects of androgens in the testis at the onset of spermatogenesis are different from those in more mature testes in which spermatogenesis is already occurring. Early transcriptome alterations (4-16 hours) following testosterone injections were followed in neonatal mouse (8dpp) testes in another study by the same group. Several hundreds of genes differentially expressed in treated and untreated animals were identified,
but only a few correlations with the results of the previous experiment were observed, probably due to the lack of FSH in hpg mice and/or of the non physiological amounts of T administered to neonatal mice.

In a different approach, ABP-transgenic mice, a model of chronic androgen depletion, have been used for the identification of genes regulated by testosterone.\(^90\) This study highlighted several hundreds of genes as differentially expressed in 30 dpp WT and ABP-transgenic mice, most of these genes being upregulated in the transgenic mice. Statistical analyses identified several pathways/functions displaying significant alterations, including genes encoding proteins involved in cell adhesion, proteolysis and peptidolysis, the GPCR signaling pathway, cytokine and growth factor activity. Finally, the authors investigated possible interactions between the genes highlighted in a functional network analysis, and identified several proteins (including interleukins 2, 4, 6 and 10, MYC, AGT and MAPK8) likely to be key factors in the effects mediated by androgens, based on the large number of interactions in which they were engaged.

As the androgen receptor (Ar) is expressed in peritubular cells, Leydig cells and Sertoli cells, the changes in gene expression reported in the three studies above cannot be assigned to any particular cell type. Several groups have therefore addressed the issue of spermatogenesis regulation by androgens in mutant mice displaying Sertoli cell-selective knockout of the androgen receptor Ar (Arinvflox(ex1-neo)/Y;Tg (Amh-Cre) mice,\(^91,92\) SCARKO mice\(^93\)). In one study, the testicular transcriptomes of two-month-old wild-type (WT) and mutant mice were compared and about 60 genes with altered expression patterns (up- or downregulated by a factor of at least two) were highlighted.\(^91\) A downregulated gene, claudin 3 (Cln3), was then studied, as this gene had a putative androgen response element in its promoter and was therefore thought likely to be a direct target of androgens. The Cln3 protein is specifically expressed and located in newly formed tight junctions (TJs) and mutant mice seemed to display greater permeability to small molecules despite the persistence of other TJ components. It was therefore suggested that androgens may regulate the permeability of the blood-testis barrier.

In another study, Denolet et al compared the testicular transcriptome of 10 days post partum (dpp) SCARKO mice and their control littermates.\(^93\) They identified several hundreds of genes differentially expressed in these two groups of mice at this age, about 40 of which displayed a change in expression by a factor of at least two, most of which were underexpressed in SCARKO mice. A significant proportion of the downregulated genes were related to MAPK activity and serine-type endopeptidase activity, whereas a significant proportion of upregulated genes were associated with cell cycle regulation, cell growth, cell adhesion and signal transduction. The expression profiles of genes identified as differentially expressed at 10 dpp during prepubertal development (from 8 to 20 dpp) in WT and SCARKO mice were also monitored. All genes displaying transient upregulation in the WT during this period were found to be downregulated, as early as 8 dpp, in SCARKO mice. This suggests that androgens play an important role in initiating meiosis.

Although these last three studies all addressed the specific question of the role of androgens in Sertoli cells,\(^91-93\) they all highlighted several genes potentially only indirectly regulated by androgens, as the animals used for these experiments were several days to several weeks older than the age at which Ar is first detected in the Sertoli cells (3 to 5 dpp). Screening for potential androgen response elements (AREs) within the genomic sequences of genes found to be differentially expressed in wild-type, \(Arinvflox(ex1-neo)/Y\) and \(Arinvflox(ex1-neo)/Y;Tg (Amh-Cre)\) mice was carried out to address this problem.\(^92\) This study identified more than 3000 putative AREs within 59 genes, 108 of which were also conserved in 32 human orthologs. Other experiments with neonatal and hpg mice focused on very early events more likely to be directly mediated by androgens, but the cell types (i.e., Leydig cells, peritubular cells or Sertoli cells) responsible for these events could not be distinguished.\(^88,89\) Very little overlap was observed between these studies for these issues. Many studies have shown that the effects of androgens during spermatogenesis are probably mediated by gene repression, consecutively modifying junction dynamics and tubular remodeling.
Evolution and Reproduction

The genome sequences available for several species have proved a highly useful resource for phylogenetic studies and taxonomic classification, and have made it possible to investigate genome-wide changes in gene expression. Khaitovich et al used probes targeting identical sequences in the human and chimpanzee genomes (94% of probes) to investigate gene expression patterns in five organs within and between these two species. They found that a neutral model with negative selection and divergence time as the major factors was a useful null hypothesis for studies of both genome and transcriptome evolution. The testes—which displayed the most rapid divergence of both transcriptome and expressed gene sequences between the two species, suggesting positive selection in terms of both expression and sequences—and the brain—which displayed the lowest level of divergence—were exceptional in terms of their gene expression patterns. It was also shown that brain genes displayed the lowest diversity of expression within species (i.e., divergence between individuals) of the organs tested, closely followed by testicular genes. It was also found that expression in the testis was associated with the largest number of significant reductions in diversity in organs other than the testis, suggesting that strong selective constraints on genes, rather than a weak influence of the environment, accounted for the low level of diversity of gene expression in the testis. A surprising result was obtained in a similar experiment carried out with different mouse strains. In mice, the testis was again found to be the organ with the highest divergence in gene expression between species and with the lowest diversity within both species and subspecies. However, in this study, the testis also displayed the lowest level of divergence between subspecies. The authors therefore suggested that the early phase of speciation may not be driven by regulatory changes in genes that are potential targets of sexual selection, and that the divergence in these genes is established only during a later phase of the speciation process. In addition to such studies investigating quantitative changes in gene expression, the issue of qualitative changes in gene expression has been addressed by comparing splicing events in mouse and human tissues. By combining the comparison of genomic sequences, the mapping of transcripts to genomic sequences and microarray expression data, the authors identified conserved and divergent alternative splicing events in more than ten thousand mouse and human orthologs. They found a strong correlation between the levels of conserved splice forms in mouse and human genes, suggesting that the functionally important splice forms were established largely before the divergence of humans and rodents and have since been maintained. These results confirm previous results in humans, showing levels of alternative splicing to be highest in the testis and brain, but the testis was found to display the largest number of divergent splices, whereas the brain had the highest number of conserved splices. These results suggest that strong evolutionary forces acting on the brain and testis have led to opposite evolutionary profiles but have tended to have similar effects on genomic sequence, gene expression and alternative splicing levels.

The available annotated genome sequences make it possible to map genes expressed in several processes to chromosomal locations. This approach has again generated particularly relevant observations in the field of male reproduction. In their investigations of the correlation between sequence and divergence in the pattern of gene expression in apes, Khaitovich et al observed that only testis-divergent genes were significantly clustered together on a particular chromosome, the X chromosome. They also found that genes expressed in the testis and located on the X chromosome displayed significantly higher levels of sequence divergence than other genes. No such significant enrichment of the X chromosome in testis-divergent genes could be found between mouse species or subspecies, probably due to the small sample size considered. The use of statistical tests is essential in chromosome assignment studies, as gene density varies within and between chromosomes. For example, genes identified during a SAGE analysis of human sperm were considered to be non randomly distributed on chromosomes, with the X chromosome being the third poorest in terms of testicular genes based on the density of expressed genes as a function of the number of expressed genes per million base pairs, but this analysis did not take into account the overall density of genes on each chromosome. By contrast, several studies have
analyzed the chromosomal location of testicular genes, using various statistical tests: a proteomic analysis of sperm chromatin in *D. melanogaster,* an analysis of large-scale microarray-based gene expression data and of the distribution of ESTs in various mouse libraries and a SAGE analysis of mouse total testis, all of which also concluded that X-linked genes were underrepresented in the testis or male germ line. However, these results must be interpreted with caution, given the conflicting results published concerning the expression of X-related genes at the different stages of spermatogenesis considered independently. We observed a complete exclusion of X-related gene products from the meiotic clusters of rodents and humans, whereas these genes were preferentially expressed in somatic Sertoli cells, spermatogonia and spermatids, at least in rodents. These results confirm and extend previous gene expression profiling findings for rat testicular cells, developing mouse testis and Spo11 mutant mouse testis. Another experiment, using the same microarrays as Chalmel et al, addressed the specific issue of the expression of genes on the sex chromosomes in mouse germ cells. However, as the authors compared the mean expression of X-related genes between samples rather than comparing the number of genes expressed with that expected by chance, they concluded that the X chromosome underwent persistent inactivation after meiosis. These studies seems to indicate that two main counteracting evolutionary forces affect the distribution of male-biased genes, one force removing these genes from the X chromosome and the other force adding them. Finally, several studies have investigated the location of testicular genes on chromosomes. They have reported, as in many other organs, the existence of a large number of clusters of genes or enriched loci. These clusters may be of physiological relevance and are suggestive of potential enhancers or common regulatory elements. However, careful consideration of the results is again required, taking into account whether statistical tests were used, and further promoter analyses and validations are required.

The Complexity of Data Analysis: Towards Holistic Biology of Spermatogenesis

Recent advances in genomics and the results obtained with new technologies for large-scale gene expression analyses have greatly improved our knowledge of male reproduction, by making it possible to investigate the molecular mechanisms underlying this process throughout the genome. The studies reviewed here have led to the identification of factors likely to be of importance for particular steps in testicular development or required for male fertility. However, most of these studies have failed to exploit the large amounts of data generated by such high-throughput approaches to provide useful and meaningful results to enhance our overall understanding of the cellular events involved. Several strategies have been used, focusing on a restricted group of relevant genes or proteins likely to be selected for further investigations. Such strategies include tissue-profiling experiments and cross-species comparisons for the detection of testis-specific genes or genes conserved through evolution, the specific and conserved expression profiles of which may be correlated with essential functions (Fig. 6). These filtering processes have been demonstrated to be efficient for identifying the key factors from very long lists of candidates, but they cannot bridge the gap between the identification of thousands of coexpressed genes or proteins and an understanding of the connections between them, providing the ultimate explanation of the correct progression of a complete biological process.

Elucidation of the complete set of transcriptional regulation mechanisms leading to the coordinated expression of a full set of functional products and identification of the protein network interactions occurring during normal testis and germ cell development are crucial to an understanding of pathological disorders of the human testis and their origins. In the last few years, considerable effort has been made to integrate data from large-scale experiments and to develop tools enabling researchers not only to describe a group of genes or proteins with similar expression profiles, but also to develop new hypotheses from their analyses. One way of analyzing such experiments is to use the gene descriptions (annotations) of the Gene Ontology (GO) Consortium for functional data mining. The GO Consortium provides
the scientific community with predictions concerning gene function, the process in which the gene product is involved and subcellular components with which the gene product is associated, using a controlled and structured vocabulary (ontologies). This makes it possible to evaluate whether a particular set of genes or proteins has annotations enriched in GO terms, and thus to demonstrate objectively that specific functions are significantly associated with a given process. This approach is still quite descriptive, but it may facilitate the identification of unexpectedly important pathways and the prediction of functions for uncharacterized genes. Furthermore, it can provide additional insight into observed transcriptional profiles and/or may facilitate the identification of genes or proteins belonging to the same complex from sets of coexpressed genes or proteins. Such multifaceted analysis was recently proposed as a way of deepening analyses of data from a gene expression profiling study of the male germ line. These authors used in silico promoter analysis to evaluate the occurrence of known transcription factor binding sites (TFBSs) within the regulatory sequences of genes coexpressed during mouse spermatogenesis. Combining TFBS predictions, DNA conservation and high-quality expression data, they found that the cAMP response element of the spermiogenesis-related factor *Crem* was indeed significantly more abundant in the genomic regions of loci specifically expressed in the mouse testis-specific post meiotic cluster than would be expected by chance (Fig. 6B). The systematic extension of such analyses should make it easier to identify and to discover relevant regulatory elements involved in the establishment of the germline expression program. The authors also investigated the use of data for a large protein network now available for validating the biological significance of clusters of coexpressed genes in terms of protein complexes. They focused on a small group of genes selected by filtering on the basis of testis-specific expression and conserved expression in mammals and for which at least one interacting factor had been identified. This analysis yielded 87 interacting factors for the initial set of 15 genes, corresponding to genes expressed in the testis and other genes not detected on microarrays (Fig. 6C). The large number of interactions detected for genes shown to be important for male reproduction also clearly demonstrates the relevance of this approach for the confirmation and extension of expression data. Finally, transcript and protein expression profiles were compared during male germ cell development, leading to the detection of several genes displaying apparently delayed transcription and translation during spermiogenesis. An analysis similar to that for genomic sequences could be carried out to identify known or new motifs within untranslated regions in such transcripts, making it possible to predict the involvement of specific RNA-binding proteins in these translation regulation mechanisms during spermatogenesis. We are currently initiating such a project in our laboratory. As an example, the discrepancy between mRNA and protein expression levels during germ cell differentiation was confirmed for minichromosome maintenance protein 7 (MCM7) and explained in part by the identification of two additional transcripts in meiotic and post-meiotic germ cells which roles remain to be elucidated (Fig. 7). In this context, a recent study by Liu et al reporting the identification of a number of genes encoding mRNAs specifically subject to alternative 3’-processing during meiosis and postmeiotic development, is of particular interest.

The study by Chalmel et al thus paves the way toward a systems-based analysis of male sexual reproduction, by highlighting the possibility of bridging the gap between DNA sequence, transcriptional activity, and translation regulation, right up to the reconstitution of functional protein complexes, at each step of spermatogenesis. Modeling of the entire process of spermatogenesis will, however, remain a major challenge, as it must include regulation by various hormones and continuous, complex communication between all the cell types present in the testis. The magnitude of this challenge has recently increased, with the arrival of data from the ENCODE project concerning exhaustive analyses of the transcription features of about 1% of the human genome. This study has revealed the pattern of gene expression to be much more complex than initially expected, with dispersed regulation and pervasive transcription, together with an abundance of non coding RNAs, dramatically modifying current notions concerning the nature of genes and their expression.
Figure 6. Data mining: Going beyond expression profiles. (Figure adapted from ref. 103). A) Filtering strategy. Cross-species comparison and tissue profiling identified 80 testis-specific orthologs with conserved expression patterns in mouse, rat and human spermatogenesis. Genes differentially expressed during spermatogenesis were identified for each species. Differentially expressed genes corresponding to orthologs represented on microarrays for all 3 species were identified, using the HomoloGene database. Finally, expression data for 17 mouse somatic tissues downloaded from the GEO web server were used to select testis-specific genes. SE, SPG, SC, ST, TU and TT correspond to Sertoli cell, spermatogonia, pachytene spermatocyte, early spermatid, seminiferous tubule and total testis samples, respectively. SO, MI, MEI and PM correspond to the somatic, mitotic, meiotic and postmeiotic expression clusters, respectively. B) Promoter analysis. Automated multi-step promoter analysis showed mouse testis-specific postmeiotic gene promoter regions to be specifically enriched in the CRE motif. We show here the results for Capza3, a gene also identified as present within the interaction node of conserved and testis-specific genes (panel C). C) Interaction network analysis. Data from IntAct, MINT and BioGRID were used to monitor interactions between conserved and testis-specific gene products. Some proteins important for male reproduction were found within these networks and were found to associate with a large number of interacting factors. As not all the interacting factors were detected on microarray analyses, interaction network analysis might help to extend expression data.
Figure 7. Transcriptomic and proteomic differential expression profiles for minichromosome maintenance protein 7 (MCM7) during rat spermatogenesis. Adapted from reference 111. A) mRNA levels in the 3 germ cell types, Sertoli cells and total testis from 2 independent experiments—Affymetrix data; (B) Relative levels of proteins in the 3 cell types—2D-DIGE data; (C) MCM7 northern blot of rat testis mRNA and for different isolated testicular cell types and total testis; (D) Western blot analysis of MCM7 protein in isolated rat testicular cells. TT: total testis; SC: Sertoli cell; SPG: spermatogonia; SPC: pachytene spermatocyte; SPT: early spermatid; CL: Late spermatid cytoplasmic lobes/residual bodies; PC: peritubular cell; LC: Leydig cell. (E) Immunolocalization of MCM7 protein in the adult rat testis. A transverse testis section showing strong labeling of the nuclei of intermediate and type B spermatogonia and preleptotene spermatocytes (iSPG, BSPG, and pISP, respectively) and a weaker signal in the nuclei of pachytene spermatocytes (pSPC) nuclei. Note that as spermatocyte differentiation proceeds (stages III to VII), the intensity of MCM7 labeling decreases, and that early spermatids are not labeled. No signal was observed in Sertoli cells, early and elongated spermatids (S, rSPT, and eSPT, respectively). Roman numerals indicate stages of the seminiferous epithelial cycle in rat species 112. Experiments were carried out with preimmune serum as a negative control.
Conclusions and Future Directions

The large series of “Omics” datasets for the testis collected to date constitute a valuable asset for scientists and doctors working in the field of male reproduction. They may provide us with insight into the molecular events controlling spermatogenesis and a more detailed understanding of human reproductive disorders, making it possible to develop new therapeutic strategies. The contribution made by many groups, through projects carried out on a small scale, as reported here, is vital. The rational compilation of this enormous set of data in a repository system, such as the GermOnline database would itself be a very useful step before the hypothesis-driven mining of the collected data with dedicated bioinformatics tools.

However, as Richard Ivell warned more than a decade ago, “All that glisters is not gold”, and it should be borne in mind that common testis gene transcripts are not always what they seem. Furthermore, simply determining gene expression or protein levels may not be meaningful. The way in which genes or proteins interact with other genes/proteins in response to internal or external signals remains a key issue. Microarray profiling experiments bridge the gap between DNA sequence annotation data and information on protein structure, function and network interaction. They do so by providing data on mRNA levels and exon composition in different cell types, at different developmental stages. In the near future, it should be possible, using novel all-exon and tiling arrays covering all known exons and the complete genomes of rodents and H. sapiens, to determine properties of the testicular somatic and germ cell transcriptomes, including transcript length, concentration and alternative splicing patterns. These experiments should also make it possible to identify novel non-coding RNAs not detected by current approaches.

Proteomics has already provided important new information concerning basic somatic and germ cell function and organization within the testis and has expanded our understanding of spermatogenesis. Technological innovations, improving the sensitivity of instruments and protein quantification, and concerning posttranslational modifications, are rapidly occurring. This will no doubt attract new biologists and clinicians to this field. However, starry-eyed newcomers to proteomics technologies should proceed with caution in this field, which remains highly technical. They would be prudent to rely on expert platforms to provide them with top-quality biological material, allowing them to concentrate on the biological questions they wish to address.

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CHAPTER 3

Estrogens and Spermatogenesis

Chandrima Shaha*

Introduction

For many years androgens were thought to be the primary hormones required for proper functioning of the male reproductive system, and testes were shown to be the major producers of the hormone. In the 1970s it was recognized that in addition to androgens testes were also a source of estrogens, and the intratesticular concentration of estrogens was higher than levels present in the serum of females of reproductive age.1,2 Since then there has been an interest in the study of the role of estrogens in the male. However, in recent years a great interest has been regenerated to decipher the role of estrogens in the control of male fertility because of various developments. For example, discovery of the estrogen receptor β (ERβ) in the male3 was an important development, and the evidence emerging from mice with targeted disruption of estrogen receptors showing defects in male fertility4 was another provocative evidence of the importance of estrogens in male reproduction. In addition, the description of reduction in sperm counts and increase in the incidence of testicular tumors in men to which environmental estrogens were causally linked5 were also important observations on pathology-inducing effects of estrogens. It is now established that estrogens are involved in numerous physiological processes in the male, for example, bone turnover, behavior and the cardiovascular system, but controversy exists as to whether male fertility over the past five decades has truly shown a decline6 due to the relatively low levels of estrogens in the environment that humans are exposed to.7 As a result of this renewed interest, the role of estrogens in male reproductive physiology is rapidly being redefined. It is therefore pertinent that a comprehensive evaluation of the data on the role of estrogens in the male is made. However, because of the ambiguity in the definition of precise function of the estrogens in the male, extensive research into the effects of the hormone on the male reproductive system is warranted.8

The most direct studies on effects of estrogens have been through administration of estrogens to animals and investigating their reproductive potential. For these studies, variations in doses have been shown to demonstrate diverse effects. Administration of estrogens and xenoestrogens during fetal and neonatal development has been reported to be associated with a series of male reproductive disturbances, such as impairment of sperm production, cryptorchidism, epididymal defects, weakened fertility, and an increased incidence of testicular cancer.9-13 When the gene knockout technology was developed, it was shown that mice lacking a functional estrogen receptor α (ERα) receptor gene were infertile and this was the first definitive demonstration of the importance of estrogen in male fertility.14-16 The cause of infertility in these mice were primarily due to a defect in efferent ductule development and function.17 In addition, mice lacking a functional aromatase gene (aromatase knockout, ArKO), aromatase being the enzyme necessary for the conversion of testosterone to estrogens were, reported to be

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infertile. Thus, lessons provided by knock-out (KO) mice for ERα and/or estrogen receptor β (ERβ) as well as aromatase, provided compelling evidence for a role of estrogens in spermatogenesis and male fertility. In this chapter, a review of current knowledge on estrogen production in the testis, estrogen receptors and the effects of estrogens in the testis is discussed.

**Estrogens and Antiestrogens**

Estrogenic activity is a property shared by a number of polycyclic compounds containing, in most cases, a phenolic ring and another oxygenated cycle located at the opposite end of the molecule (Fig. 1A). There is a large internal hydrocarbon moiety that contributes to an optimal orientation of these two polar functions for selective H-bonding with specific amino acid residues of the ligand-binding pocket of estrogen receptor (ER). Other than steroid hormones, flavones, isoflavones and coumestanes also are potent estrogens. All these chemicals are linear, planar molecules. The planar feature of the estrogens gives a closed conformation to the receptor upon binding. This structural property distinguishes these strong ER agonists from a variety of angular ligands, gem-diphenylethylenes, triphenylethylenes, diarylimidazolines and diarylpiperazines, which maintain the ligand-binding pocket in an open conformation. Linear, planar molecules are currently referred to as type I estrogens, while angular molecules are categorized as type II estrogens (Fig. 1A,B). Grafting of a reactive group, i.e., amido (ICI 164,384) or sulfoxide (ICI 182,780; RU 58,668) via a long alkyl side chain onto estradiol at positions of 7α and 11α totally abrogates its estrogenicity and converts it into a “pure” antiestrogen (Fig. 1C). Selective estrogen receptor modulators or SERMs are synthetic compounds mainly derived from type II estrogens. The large majority of SERMs contains a dialkylaminoethyl side chain (Fig. 1D) which is responsible for their antagonistic activity. Like the long alkyl side chain characterizing pure antiestrogens described above, dialkylaminoethyl side chains of SERMs protrude from the binding pocket, allowing for interactions with critical amino acid residues in the receptor.

**Overview of Spermatogenesis**

Spermatogenesis is a complex process of male germ cell proliferation and maturation from diploid spermatogonia through meiosis to mature haploid spermatozoa. Spermatogonia arise from the primordial germ cells, which migrate into the genital ridge during fetal life. Under the influence of the Y-chromosome-bearing stromal cells of the developing gonad, they differentiate into gonocytes, the male germ cell precursors and undergo mitotic arrest. After birth, they are reactivated and differentiate into spermatogonia that go through a cycle of division and differentiation with cellular apoptosis keeping the number of cells in check. Spermatogenesis in the adult takes place within the seminiferous tubules of the testis which consist of the seminiferous epithelium, comprising of germ cells and supportive somatic Sertoli cells that stand on the basement membrane which is surrounded by the peritubular myoid cells. The interstitial tissue between the seminiferous tubules contain androgen-producing Leydig cells and interstitial macrophages. Developing germ cells in different stages of differentiation form intimate associations with the Sertoli cells. The germ cells are arranged in defined cellular associations that constitute the cycle of the seminiferous epithelium, and each particular association of germ cells is referred to as a stage. The cycle of the seminiferous epithelium is the time interval between the appearance of the same stage at a certain point of the tubule. Spermatogonia, the stem germ cells in the testis include type A spermatogonia, and type B spermatogonia. Spermatogonia undergo continuous mitoses to produce a large number of germ cells available for entry into meiosis. After the last mitosis of type B spermatogonia, preleptotene primary spermatocytes are formed and these cells initiate meiotic division. After completion of the meiotic division, the differentiation of round spermatids into the mature elongated spermatid take place through the process of spermiogenesis. This process involves formation and development of the acrosome and flagellum, condensation of the chromatin, change in shape of the nucleus, and removal of the cytoplasm before release of the spermatid during spermiation.
Figure 1. Structures of estrogens and antiestrogens. A) Structures of type I estrogens which are linear, planar molecules that include estradiol, coumestrol and diethylstilbestrol. B) Structures of type II estrogens which are angular molecules that includes bisphenol A, Cyclophenyl and 2,3-diarylpiperazine. C) Structures of pure antiestrogens, RU58668 and ICI 182,780. Grafting of a reactive group, i.e., sulfoxide (ICI 182,780; RU 58,668) via a long alkyl side chain (“spacer”) onto estradiol at 7α or 11β totally abrogates its estrogenicity and converts it into a “pure” antiestrogen. D) Structure of Selective Estrogen Receptor Modulators, namely raloxifene and 4-hydroxytamoxifen. These are synthetic compounds mainly derived from type II estrogens. The large majority of Selective Estrogen Receptor Modulators contains a dialkylaminoethyl side chain which is responsible for their antagonistic activity.
The Gonadotropins and the Hypothalamic-Pituitary Gonadal Axis

Gonadotropins are the major endocrine regulators of spermatogenesis. Luteinizing hormone (LH) acts on the Leydig cell to stimulate the secretion of androgens, namely testosterone, which in turn acts on androgen receptors in the seminiferous epithelium to control spermatogenesis. Follicle stimulating hormone (FSH) receptors on the Sertoli cells react to FSH and regulate spermatogenesis by stimulating the production of numerous Sertoli cell factors. The role of androgens is very well demonstrated in the hypogonadal (hpg) mouse where the androgens have been shown to fuel all phases of germ cell development. Transgenic mice with targeted disruptions of the FSH receptor gene or the FSH β-subunit gene are fertile and display all stages of germ cell development, suggesting that FSH is not an absolute requirement for fertility. However, in these mice testes are smaller and less sperm are produced with many defective spermatozoa. Therefore, full fertility relies on the delicate balance of the hypothalamic-pituitary-testis axis. It is important to understand the functioning of the hypothalamic-pituitary gonadal axis in the context of estrogens because estrogens provide the feedback stimulus along with androgens (Fig. 2). The reproductive hormonal axis consists of three main components: (A) the hypothalamus, (B) the pituitary gland, (C) the testis. This axis functions in a strongly regulated manner to produce optimal concentrations of circulating steroids required for normal male sexual development, sexual function and fertility.

Figure 2. The hypothalamic-pituitary axis. A diagram illustrating the orientation of the hypothalamic-pituitary axis and the feedback loops with inhibin and the steroid hormones secreted by the testis. The paracrine routes in the testis are also indicated. The cell types expressing ERα and ERβ are indicated as well.
Hypothalamus

The integrating center of the reproductive hormonal axis is the hypothalamus (Fig. 2) that is the site of production of the peptide hormone gonadotropin-releasing hormone (GnRH) which is delivered to the adenohypophysis of the pituitary gland by a short portal venous system where it stimulates the synthesis and release of gonadotropic hormones LH and FSH.7 The release of GnRH is seasonal (peaks in the spring), circadian (highest testosterone levels are in the a.m.) and pulsatile (peaks occur every 90-120 minutes). The GnRH production from hypothalamus is under feedback control from gonadal hormones.

Pituitary

LH and FSH are glycopeptides consisting of two peptide chains (α and β) synthesized by the pituitary cells and are secreted into the general circulation thereby transporting to the testis.43 LH and FSH share a common α peptide chain (α chain) with thyroid-stimulating hormone and human chorionic gonadotropin and differ from each other by the presence of a specific β chain, the latter providing specificity of biologic action.44 The pituitary also secretes prolactin which affects testicular function indirectly by inhibiting GnRH release from the hypothalamus and therefore LH and FSH secretion from the pituitary. Prolactin also directly inhibits pituitary gonadotrophic cells and the Leydig cells of the testes.

Testis

Testicular control is achieved by LH, FSH and androgens and many other peptides and growth factors (e.g., inhibin, activin, insulin-like growth factor 1, transforming growth factors) that are secreted locally in the seminiferous tubular microenvironment.45

Feedback Control of Gonadotropins

Negative-feedback of GnRH release is exerted by testosterone and estrogens through steroid receptors present in the hypothalamic neurons and in the pituitary.46 Although the concentration of estradiol in the blood of men is relatively low compared with testosterone, it is a much more potent inhibitor of LH and FSH secretion (approximately 1000-fold). Testosterone acts primarily to feedback at the level of the hypothalamus whereas estrogens provide feedback to the pituitary to modulate the gonadotropin secretion response to each GnRH surge. Several disease states can occur with excess estrogens. Pituitary gonadotropin secretion is suppressed by peripheral estrogens.47 Inhibin and activin are factors produced in the testis which have several functions. In the adult testis, paracrine signals from germ cells are important for Sertoli cell inhibin production. While inhibin seems to have a dual, endocrine (regulation of FSH secretion in pituitary) and para/autocrine role in testicular function, the activin actions seem to be of the local type in the testis and numerous extragonadal sites.48,49

Biosynthesis of Estrogens and Sites of Estrogen Biosynthesis in the Testis

Estrogen biosynthesis is catalyzed by a microsomal member of the cytochrome P450 superfamily, namely aromatase cytochrome P450 (P450arom, the product of the CYP19 gene) (Fig. 3). Aromatase is a terminal enzyme which transforms irreversibly androgens into estrogens and it is present in the endoplasmic reticulum of numerous tissues. The P450 aromatase is a microsomal enzymatic complex composed of two proteins: a ubiquitous NADPH-cytochrome P450 reductase and a cytochrome P450 aromatase, which contains the heme and the steroid binding pocket. Six isoforms of cytochrome-dependent mono-oxygenases (CYP) are involved in the biosynthesis of various steroid hormones, starting from cholesterol—CYP11A, CYP17, CYP19, CYP11B1, CYP21B, and CYP11B2.50 The rate-limiting step in all steroid hormone biosynthesis is the cleavage of the side chain of cholesterol by CYP11A to form the C21 steroids, pregnenolone and progesterone (Fig. 3). Hydroxylation and subsequent cleavage of the two-carbon side chain of the C21 steroids by the CYP17 (17-α hydroxylase activity/C17-20
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lyase activity) yields the C19 steroids, androstenedione and dehydroepiandrosterone. While testosterone is formed from the precursor, androstenedione, estrogens are ultimately formed by aromatization of androstenedione and testosterone, catalyzed by the CYP19 (aromatase). In addition to the cytochrome P450, a series of hydroxysteroid dehydrogenases (3'-HSD and 17β-HSD) participate in the biosynthesis of the steroid hormones. The 3β-HSD converts pregnenolone to progesterone, whereas the 17β-HSD converts androstenedione to testosterone. 17β-HSD type 1 and CYP19 catalyze the end steps in 17β-estradiol biosynthesis through androstenedione (Fig. 3). Estrogens are extensively metabolized by a number of oxidative and conjugative reactions that can lead to their deactivation and subsequent elimination. Alternatively, oxidation and conjugation reactions of estrogens may generate metabolites that have distinct biological activities, including altered hormonal properties; genotoxicity, through the formation of reactive species that modify cellular DNA and protein or chemotherapeutic properties as anti-estrogens. They are also potentially anti-angiogenic.

The identification of sites for estrogen biosynthesis is done by identifying aromatase activity and this activity is even visible in embryonic testes. For example, aromatase activity is present in the fetal rat testis around day 19 of gestation. At this stage of development, basal aromatase activity is found in both the immature Leydig cells and Sertoli cells. In fact in the neonates, the Sertoli cells are more active in producing estrogens than Leydig cells suggesting that these cells are an important source of estrogen in the postnatal testis. It is during this time that FSH receptors first appear on the Sertoli cells but germ cells do not contain detectable aromatase.

![Figure 3. Synthesis of estradiol. Figure shows the primary routes of estrogen biosynthesis in the testis. The first and rate-limiting step in the synthesis of all steroid hormones is conversion of cholesterol to pregnenolone that involves the cleavage of a 6-carbon group from cholesterol. CYP 11A, the cholesterol side chain cleavage cytochrome P450 is used for the purpose. Pregnenolone is converted to progesterone through the use of 3 β-hydroxysteroid dehydrogenase. Hydroxylation and subsequent cleavage of the two-carbon side chain of the C21 steroids by the CYP17 (17α-hydroxylase activity/C17-20 lyase activity) yields the C19 steroids, androstenedione and dehydroepiandrosterone. Testosterone is formed from the precursor, androstenedione and estrogens are ultimately formed by aromatization of androstenedione and testosterone, catalyzed by the CYP19 (aromatase).](image-url)
activity at the early neonatal stage. By adulthood, rodent Leydig cells express a high level of aromatase which is stimulated by LH and steroids. However, it has been recognized that apart from the hormone producing Leydig cells and Sertoli cells, germ cells are also a source of estrogen in the adult. Mature rat germ cells express a functional aromatase with production of estrogens equivalent to that of Leydig cells but there is a variation in aromatase expression in the different germ cell types. For example, the amount of P450arom transcripts is higher in pachytene spermatocytes as compared to round spermatids. In humans, in addition to Leydig cells, aromatase is present in ejaculated spermatozoa and in immature germ cells. Therefore, in addition to the somatic cells of the testis, the spermatogenic cells also appear to be the sources of estrogens. Thus, the evidence for the presence of multiple sites of estrogen biosynthesis in the testis reflects the importance of estrogen in testicular functioning.

Distribution of Estrogen Receptors in the Testis

Estrogen receptors are ligand activated molecules that serve as transcription factors and mediate pleiotropic effect of estrogens on various tissues. For a summary of various aspects of estrogen receptor functioning, several other reviews (see refs. 60-62) are recommended. These receptors orchestrate both transcriptional and nongenomic actions in response to estrogens, xenoestrogens and signals coming from growth factor signaling pathways (Fig. 4). When estrogen binds to its receptors they dimerize and interact with DNA sequences to regulate gene transcription. Estrogens can also stimulate or repress transcription by binding to DNA-associated transcription factors. Recently, estrogen has been reported to induce multiple cytosolic signaling processes, such as activation of Src, Ras, Raf, protein kinase C (PKC), protein kinase A (PKA), potassium channels, intracellular calcium levels and nitric oxide (for reviews, see refs. 66-69). Since activation of these signaling molecules depends on cell types studied and the conditions used, the precise nongenomic signaling pathways of estrogen and their functional significance are still controversial.

Two types of distinct receptors are expressed, namely, ERα and ERβ. ERs are members of the steroid hormone superfamily of nuclear receptors, which share a common structural architecture and consist of three independent functional domains: the N-terminal or A/B domains, the C or DNA-binding domain, and the D/E/F or ligand-binding domain. Binding of a ligand to the ER causes a series of downstream events, including receptor dimerization, receptor-DNA interactions mediated by EREs present in the promoter region of target genes, recruitment of and interaction with transcription factors, and the formation of a preinitiation complex. Ligand-receptor interactions ultimately cause changes in target gene expression. The N-terminal domain of nuclear receptors encodes an activation function called AF-1, which mediates protein-protein interactions to induce transcriptional activity. The C-terminal or ligand-binding domain contains the AF-2 interacting surface that mediates ligand binding and receptor dimerization to stimulate transcriptional activity. Thus, AF-1 and AF-2 are both involved in mediating the transcriptional activation function of ERs. The most conserved region of the α and β forms of ER is the DNA-binding domain; the positions of the cysteine residues that coordinate the two zinc fingers of the C domain are conserved in both (ERα and ERβ). There is general agreement that ERs function as dimers, and coexpression of ERα and ERβ in the same cell causes the formation of homodimers (ERα/ERα and ERβ/ERβ) or heterodimers (ERα/ERβ), which affect ligand-specificity. The relative amounts of ERα and ERβ in a given tissue are key determinants of cellular responses to estrogen and other ER agonists and antagonists. The interactions between ERs and EREs are mediated by other factors, including the ability of ERβ to modulate ERα transcriptional activity and recruitment of several protein coactivators and repressors by both ER subtypes. The effects could also occur through nontranscriptional mechanisms mediated by protein-protein interactions occurring between ERs and growth factors e.g., IGF-1 and EGF. As mentioned earlier, there is evidence for the presence of a small pool of ERs localized to the plasma membrane. For example, BSA-conjugated estradiol, which is unable to gain entry into the cytosol and acts at the plasma membrane, decreases testicular
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Estrogens and androgen production in vitro. Membrane ER is thought to signal mainly by coupling to GTP-activating proteins and through pathways involving second messengers (e.g., calcium) and kinase cascades. The integration of several pathways implies that estrogen action in any particular tissue and organ is the result of activities mediated by both the genomic and nongenomic pathways.

An overview of the distribution of estrogen receptors in the testis provides information on the cells that are potentially capable of responding to estrogen. Both types of estrogen receptors ERα and ERβ are present in the testis. While ERα dominates in some specific tissues and is mainly involved in reproductive events, ERβ is the more generally expressed estrogen receptor. In the testis, ERβ is the more abundant receptor and is typically found in nearly every cell type of the interstitium and the seminiferous tubule, except for the elongated spermatids. ERβ is expressed in the Sertoli cells, Leydig cells, and germ cells, as well as in the epididymis, prostate, and seminal vesicles. ERα is present predominantly in the Leydig cells, efferent ducts, and epididymis of rats and monkeys. ERα is primarily present in the efferent ducts that transport spermatozoa suspended in the fluid secreted by the Sertoli cells from the testis to the epididymis. Although there are reported differences in ERα localization in the epididymis of various species, its presence in efferent ductule epithelium has remained constant across species. ERα protein is abundant in epithelial cells of the efferent ductule with intense immunohistochemical staining of the nonciliated cell nucleus and the ciliated cells showing considerable variability in staining. ERα protein is also present in the fetal testis. Immunohistochemical studies show the presence of ERα in the mouse undifferentiated gonad between days 10-12 after birth, suggesting that estrogen may have a role very early in the differentiation process.
Leydig cells within the rodent fetal testis contain ERα until birth. ERs are expressed in the Leydig cells at a stage in development when the androgen receptor is not yet expressed highlighting a role for estrogen at this stage. There is evidence for both ERβ mRNA and protein as early as day 16 of gestation in the gonocytes, Sertoli cells and Leydig cells. However, it is the gonocytes that express ERβ in higher abundance than the other testicular cells. At birth, the testis expresses both ERα and ERβ that is localized to the seminiferous epithelium, Sertoli cells and developing germ cells. As we have seen in the earlier section that a variety of cells in the testis synthesize estrogen, studies represented in this section show that multiple cell types express estrogen receptors as well which indicates that estrogens influence a number of cell types including the germ cells themselves.

**Estrogen Receptor and Aromatase Knockout Mice and Their Phenotypes**

Mice with targeted disruption of estrogen receptors provide us interesting view of the role of estrogens in the males and there is a great difference between disruption of ERα, ERβ and a combination of both. For example, ERα disruption (ERKO mice) cause increased plasma testosterone levels and are infertile. Upon histological examination they show seminiferous tubular swelling and loss of spermatogenesis. The sperm made in the testes of these animals are nonfunctional. The testicular LH and FSH receptors are up-regulated, but gonadotropin levels are normal. ERβ persists in the ERKO mice, showing that there is no interrelationship between expressions of the two ER types. Infertility in these mice is primarily due to distension of the rete testis and efferent ducts resulting in progressive loss of testicular weight with prominent exfoliation of germ cells. Therefore, estrogens have a role in fluid reabsorption in the efferent ducts and the initial segment of the epididymis. As regards sexual behavior, the ERKO mice have normal amount of mountings with lesser levels of aggression and lower number of intromissions and ejaculations.

ERβ knockout (BERKO) mice were first generated by inserting a neomycin resistance gene into exon 3 of the coding gene, using homologous recombination in embryonic stem cells. BERKO mice exhibit phenotypes distinctly different than those of αERKO mice. They develop normally and as young adults, they are indistinguishable grossly and histologically from their littermates. Unlike males lacking ERα, male BERKO mice are fully fertile and reproduce normally. The mice exhibit no major alterations in the function of the male reproductive system. Therefore, it appears that ERβ is not essential for normal testicular functioning.

The reproductive phenotype of αβ-ERKO mice is very close to that of α-ERKO mice and it is characterized by infertility and enlarged seminiferous tubules. ArKO mouse lacks aromatase products (namely, estradiol and estrone). At around 18 week of age these mice show a specific postmeiotic defect coinciding with an elevation in apoptosis and a reduction in fertility. The ArKO mice also present with a significant reduction in copulatory behavior in adulthood, reiterating the importance of estrogen in male reproduction.

The mechanisms involved in the development of infertility is different in α-ERKO male mice if compared with ArKO, because reduced fluid reabsorption occurs in α-ERKO male mice and in ArKO mice early arrest of spermatogenesis suggests a failure in germ cell differentiation probably due to the lack of estrogens in the testicular environment. It appears that estrogen activity in the male reproductive tract differs with regard to both the types of ERs involved in the pathway of estrogenic action, and the site of action through the male reproductive tract because β-ERKO male mice are fully fertile as compared to the other two. Accordingly, in ArKO male mice, the failure of germ cell differentiation that is probably related to the lack of estrogen action on seminiferous epithelium while ERα disruption and related arrest of fluid reabsorption take place in the efferent ductules of α-ERKO mice. Spematogenesis is conserved in very young ArKO mice because a small quantity of estrogens, from external sources, probably is sufficient to promote germ cell maturation for a brief period, and therefore, the degree of infertility is less severe in ArKO mice than in α-ERKO. Later, the continuous lack of
estrogens causes sperm abnormalities with advancing age in ArKO mice, since estradiol is probably necessary to maintain spermatogenesis and promote normal sperm maturation, both in the seminiferous epithelium and through the reproductive tract. Therefore, ERα appears to be the most important component for estrogens action in male reproduction because α-ERKO mice are infertile while the β-ERKO mice are fertile.

Effects of Estrogen Administration on Testicular Function

Estrogen administration in animals has provided a substantial amount of literature on how estrogens affect the different cellular components of the testis and what are the functional consequences. The confusion about the involvement of estrogen in spermatogenesis is due to the fact that estrogen action is important at numerous levels in male reproductive physiology including effects on the hypothalamic-pituitary-testis axis, Leydig cells, Sertoli cells, germ cells, and epididymal function. Thus the extensive range of effects that estrogens have in the male reproductive tract, cause difficulties in the interpretation of experimental findings. The following section discusses the effects of estrogen administration and deprivation on processes that are required for normal spermatogenesis and fertility.

Estrogen Exposure and Spermatogenesis

A major component of the negative feedback action of androgens on gonadotropin secretion is mediated by estrogen. Studies in humans show that administration of estradiol enhance gonadotropin suppression induced by a testosterone-based contraceptive, indicating estrogen’s role as a negative feedback regulator of gonadotropin secretion that disrupts spermatogenesis. Neonatal exposure to either estrogens or estrogen-like compounds promotes changes in gonadotropin secretion in rodents. Neonatal estrogen exposure can have important long-term effects on the hypothalamic-pituitary-testis axis and thus spermatogenesis. A single high dose of estradiol benzoate to 1-day-old male rats causes a reduction in both GnRH secretion and pituitary responsiveness to GnRH. In male hamsters, neonatal exposure to diethylstilbestrol (DES), a synthetic estrogen during neonatal period impairs the action of androgens on target organs. In rats neonatal administration of DES delays the establishment of the blood-testis barrier affecting spermatogenesis. It has also been shown that neonatal exposure of rats to low levels of estrogens can advance the first wave of spermatogenesis at puberty and also affect the development of excurrent ducts of the rat testis. Other than germ cells, neonatal estrogen treatment results in dose-dependent alterations in Sertoli cell numbers, efficiency of spermatogenesis, and germ cell apoptosis in adulthood. Therefore, the above studies show that neonatal exposure to estrogen affects spermatogenesis but depending on the time and dose of exposure this effect can be reversible.

Given that the appropriate concentrations of LH and FSH as well as a tightly regulated onset of secretion during the neonatal and pubertal periods, is fundamental to whether normal spermatogenesis proceeds, neonatal estrogen exposure can have important long-term effects on the hypothalamic-pituitary-testis axis and thus spermatogenesis. Interestingly, whether or not estrogen administration to juvenile mice will interfere with the hypothalamo-pituitary-gonadal axis appears to be strain-dependent, which could lead to confusion when interpreting the literature on the interaction between estrogen, the regulation of pituitary hormone production, and fertility.

Adult monkeys treated with an aromatase inhibitor show a decrease in the conversion of round to elongated spermatids and a decrease in sperm output from the testis, also suggesting that estrogen is important for spermatid differentiation in the primates. A different scenario appears to be relevant in boars. It has been shown that boars treated with aromatase inhibitors show a delayed lumen formation, lower testicular weight, fewer detergent resistant spermatids, and fewer Sertoli cells, but by 7 to 8 months, these boars recover and show larger testes with more Sertoli cells. Total Leydig cell volume increases in proportion to testis size. Therefore, reducing endogenous estrogen is consistent with a delay in testicular maturation/puberty that
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allows for a longer window for the proliferation of Sertoli cells and maturation of Leydig cells, resulting in larger testes and higher spermatid production.103

Effects of Estrogens on Sertoli Cells

Through the development of the testis, estrogen has varied effects on the Sertoli cells. The proliferation of Sertoli cells occur from day 16 of fetal life in the rat and at birth, there are 1 million cells and around 15 days there are about 40 million cells. After postnatal day 15, proliferation ceases,104 differentiation commences and the number of the Sertoli cells in the testis remains stable throughout adulthood.105 During this period, Sertoli cells produce considerable amounts of estrogen, leading to the suggestion that estrogen is involved in the division of the Sertoli cells. Aromatase activity is highest in Sertoli cells from prepubertal rats, declines as Sertoli cells mature, and is hormonally regulated, principally by FSH. The data show that estrogen has a stimulatory effect on Sertoli cell division yet a negative effect on Sertoli cell differentiation and development, because, estrogen production is high in proliferating Sertoli cells but lower in differentiated cells106 and FSH-induced aromatase activity starts to decline towards the end of division. The ability of Leydig cells to produce testosterone increases from about day 14 after birth,107 which inhibits Sertoli cell aromatase activity,108 suggesting that androgens from the maturing Leydig cells may participate in the down-regulation of aromatase during the switch from Sertoli cell division to differentiation. Germ cells which are starting to develop during the switch between Sertoli cell division and differentiation, induce a decrease Sertoli cell aromatase activity.109

Effects of Estrogen on Germ Cells

The presence of estrogen receptors during various stages of germ cell development indicates that there could be a direct effect of estrogens on germ cells. For details on expression of estrogen receptors in germ cell a review by ref. 110 is referred to. The presence of aromatase in germ cells demonstrate the capability of these cells to synthesize estrogens that could be affecting the same cells in an autocrine mode. In a study aimed at investigating the effect of estradiol and diethylstilbestrol on the testis from 14.5-day-old rat fetuses in culture, alterations in the germ cells in terms of changes in viability occurred after estradiol and DES exposure.111 In the human testis, estradiol appears to be a potent germ cell survival factor when studied in in vitro cultures.112 Studies using rat gonocytes in culture show that estrogen stimulates gonocyte proliferation.113 The high aromatase activity in the Sertoli cells during proliferation of gonocytes is an indication of estrogen action where differentiating spermatogonia during the early neonatal period have been shown to contain ERβ.114

Estrogens and Spermatozoa

In the last stage of mammalian spermiogenesis, the bulk of spermatid cytoplasm is extruded in tubular lumen while a small cytoplasmic mass is retained around the sperm mid-piece as cytoplasmic droplet. This droplet moves to the end of the tail and finally sheds from mature spermatozoa.115 Human ejaculate can contain spermatozoa with excess residual cytoplasm which has been retained around the sperm mid-piece due to an incomplete maturation process. Both estrogen receptors are found in the excess residual cytoplasm of immature sperm, while sperm tails show only ERα.116 It has been shown that specific allelic combinations of the ERα, which confer a stronger estrogen effect, may negatively influence human spermatogenesis.117 Concomitant expression of ERβ and ERα in human ejaculated spermatozoa has been reported showing a differential distribution of the two ER subtypes, the former being prevalently located in the midpiece, but the latter being in the tail. In the same study, ERα was shown to interact with the p55 regulatory subunit of PI3K, whereas ERβ interacts with Akt1.118 Additional studies have also demonstrated presence of ERβ on sperm.119 Interestingly, while possessing estrogen receptors, human ejaculated spermatozoa also express cytochrome P450 aromatase indicating the cell could also serve as a source of estrogen.120 ERβ contains two silent polymorphisms, RsaI (G1082A) and
AluI (G1730A), the frequency of the heterozygous RsaI AG-genotype is three times higher in infertile men showing that ERβ may have modulating effects on human spermatogenesis.121

Studies on Xenoestrogens

Synthetic estrogens, also called xenoestrogens, are a diverse group of compounds in the environment that mimic the action of the natural hormone 17β-estradiol in estrogen-dependent tissues. There is considerable controversy as to whether environmental chemicals that mimic estrogens can adversely affect the endocrine and reproductive systems. Evidence has accumulated over several decades now that estrogen is essential for spermatogenesis and that intratesticular concentrations of estrogen are very high.82 It has also been realized that estrogen like chemicals present in the environment adversely affects male reproductive function. The compounds thought to be responsible for such changes include industrial chemicals (e.g., polychlorobiphenyls, alklyphenols, pesticides (e.g., DDT derivatives, methoxychlor, kepone), pharmaceutical agents (e.g., DES, tamoxifen, raloxifene), phthalates (e.g., di-2-ethylhexylphthalate, di-n-butyl phthalate), and phytoestrogens (e.g., genistein, daidzen).122,123 The extent of exposure to these chemicals on members of a population differs as occupations in agriculture, petrochemicals and the construction industry entails higher exposure. Since estrogen receptors are present in the pituitary and spermatogenic cells,82 estrogen like chemicals can act as agonists or antagonists for the hormone and interfere with spermatogenesis. Therefore, agents able to mimic estrogens can potentially alter the action of the hormone on spermatogenic cells leading to functional impairment of the male gamete. Reports of impaired spermatogenesis as a consequence of exposure to agents with estrogenic activity termed as endocrine disruptors are well documented in rats.124 In addition to such examples, it is well established that estrogen administration to experimental animals during adulthood can impair sperm production and maturation.125 The mechanism by which these xenoestrogens exert their action on the cell could occur by either agonistic or antagonistic actions on ERα and ERβ126 resulting in changes in steroid hormone receptor gene expression, altered steroid hormone metabolism, change in cross-talk between ERs and other signaling systems.127

Exposure of neonatal testis, populated by fetal-type Leydig cells to endocrine-active compounds may have far-reaching consequences. Fetal rat testis shows inhibition of development of spermatogonia, Leydig cells and Sertoli cells on estrogen exposure.111,128 Bisphenol A (BPA), [2,2-(4,4-dihydroxydiphenol)propane] is an estrogenic compound that is widely used in the manufacture of polycarbonate plastics, which serve as containers for foods and beverages and as a constituent of dental sealants. The core structure of BPA resembles that of the natural estrogen, consisting of two phenolic rings joined by a bridging carbon. Exposure of the human population to BPA is significant given its widespread use in consumer products. BPA and its metabolites have been measured in the blood of normal men.129 Administration of low doses of the industrial and estrogenic chemical BPA reduces spermatogenesis in mice130 and suppresses androgen biosynthesis by mature rat Leydig cells. In mice, neonatal exposure to a relatively large dose of BPA causes damage to the motility and morphology of sperm.131 BPA induces apoptosis in Sertoli cell cultures isolated from 18-day-old rats.132 4-tert-octylphenol is an environmental pollutant with estrogenic activity and is directly toxic to cultured rat spermatogenic cells and Sertoli cells.133 Therefore, BPA, a xenoestrogen appears to have the potential to affect multiple cell types in the testis. Chronic exposures of mice to 0.5 or 50 μg/ml bisphenol A decreases ERβ and increases ERα gene expression in germ cells134 but a single injection of estradiol benzoate at high doses (500 μg) causes the opposite effect in prepubertal rats, i.e., decreased ERα mRNA levels and increased ERβ expression.135 Parabens are p-hydroxybenzoic acid ester compounds widely used as preservatives in foods, cosmetics, toiletries and pharmaceuticals. These compounds exert a weak estrogenic activity as determined by in vitro estrogen receptor assay and in vivo uterotrophic assay. In mice, a decrease in daily sperm production occurs when treated with the above compound.136
Tumors of germ cell origin comprise about 95% of all testis cancers. Interaction between genetic and environmental factors, including inappropriate exposures to endocrine-active chemicals can lead to hypospadias, testicular cancer, abnormal spermatogenesis and undescended testis. Exposure to estrogens early in life can lead to defects in tissue differentiation in the fetal period. Although the effects of exposures to environmental chemicals in adulthood are typically transient, chemical exposures that alter gene activity during development disrupt differentiated function in hormone-responsive tissues of the adult. Testicular cancer incidence is highest in younger men (20-40 years old), and the etiology of this disease could be related to estrogen exposure during the prenatal period. For example, in a study of young adult twins in England, the overall risk of testicular cancer was higher for dizygotic than for monozygotic twins. These observations support a role for estrogens in the etiology of this disease, which has been increasing in most countries. The possibility that xenoestrogens may cause adverse effects in the reproductive tract was first highlighted by reports on adolescent sons born to pregnant women who had taken the highly potent synthetic estrogen DES. These individuals developed a variety of testicular and epididymal abnormalities in adulthood. Phytoestrogens are plant-derived compounds with estrogenic activity. They are common in both human and animal diets, particularly through soy-based foods. Adult male rats, fed a high phytoestrogen diet for 3 days, demonstrate significantly reduced fecundity which is reversible. In these instances, the expression of ER increases in the initial segment of the epididymis, but decreases in the cauda epididymis following 3 days on the high phytoestrogen diet. Therefore, phytoestrogens can also affect estrogen receptors on spermatozoa and eventually fertility.

**Estrogen Deprivation and Spermatogenesis**

The functional consequences of estrogen deprivation in adulthood have also been investigated by the administration of the anti-estrogens. For example, Faslodex (ICI 182,870) compound binds to both ER and ER and reduces the stability of ER in vitro. Rats and mice treated with Faslodex (2 to 150 days) shows efferent duct dilation, a progressive decrease in testis weight, decreased sperm concentration and decreased fertility indicating massive effects of antiestrogens in germ cells. These changes are similar to those observed in the ER KO mice where they occur after a reduction in expression of ER with no change in expression of ER. 

**Estrogens and Germ Cell Apoptosis**

The physiological significance of the spontaneous germ cell apoptosis that occurs during spermatogenesis is possibly related to maintenance of the proper number of germ cells per Sertoli cells because these cells are terminally differentiated cells with no capacity for renewal, and are able to support only a certain number of germ cells. Therefore, germ cell death during development is to limit the number of germ cells to match the supportive capacity of the Sertoli cells. Apoptosis may also serve to eliminate germ cells with mutations in the DNA. In meiotic spermatocytes, there appears to be a quality-control system or checkpoint for monitoring chromosome synopsis. This control system is thought to recognize unrepaired double-strand DNA breaks in unsynapsed chromosomes during meiotic metaphase and to induce apoptosis of the affected cell. During prepubertal development, a wave of extensive germ cell apoptosis occurs in the rodent testis. This early germ cell apoptosis, which mainly affects spermatogonia and spermatocytes, appears to be essential for functional spermatogenesis in adulthood. In the adult testis, spermatogenesis is accompanied by spontaneous germ cell degeneration resulting in the loss of up to 75% of the potential number of mature spermatozoa. In the human testis, spontaneous germ cell apoptosis involves all three classes of germ cell, i.e., spermatogonia, spermatocytes, and spermatids. Various apoptotic signaling pathways are operative in the testis. The Fas system has been suggested to play a role both in maintaining the immune-privileged nature of the testis and in regulating testicular germ cell apoptosis. FasL has been found in mouse, rat, and human Sertoli cells and is generally assumed to be constitutively expressed by the Sertoli cells. Some reports have shown FasL
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Presence in germ cells also.\textsuperscript{125,174} Fas expression, in turn has been demonstrated in the germ cells of the rat and human testes\textsuperscript{125,162} and in some reports Fas presence has also been shown in the Sertoli cells.\textsuperscript{163}

Estrogen exerts different effects in specific cell types, for example, in estrogen receptor positive MCF-7 human breast cancer cells, estrogen strongly stimulates cell proliferation and does not induce cell death.\textsuperscript{164} In contrast, for several cell types, stably transfected with ER, estrogen is highly toxic and induces cell death.\textsuperscript{165,166} Recent studies demonstrate that estrogens also exert nongenomic effects\textsuperscript{167} possibly related to its ability to activate signaling pathways such as nitric-oxide synthase in endothelial cells.\textsuperscript{168} Estrogens have been linked to apoptosis in a variety of cell lines\textsuperscript{168,169,170} but studies on spermatogenic cell apoptosis in response to estrogens are relatively few and consequently very little information is available on estrogen induced changes. Apoptosis is of great relevance for successful production of spermatogenic cells as excess cells need to be removed for the proper maintenance of testicular homeostasis.\textsuperscript{171} As in other tissues, selection of the apoptotic pathway depends on the physiological and pathological state of the spermatogenic cells and members of the Bcl-2 family of proteins have been implicated in apoptosis induced by certain agents along with activation of the Fas-FasL system under various conditions.\textsuperscript{125,172,173}

While earlier studies implicated FasL from Sertoli cells for engaging Fas receptor on spermatogenic cells during apoptosis, later studies provided evidence in support of both Fas and Fasl being expressed in spermatogenic cells as well.\textsuperscript{125,174} Estradiol given subcutaneously to adult male rats for 10 days show a decrease in apoptosis of germ cells.\textsuperscript{175} Similarly, low concentrations of 17ß-estradiol (10\textsuperscript{-9} and 10\textsuperscript{-10} mol/L) effectively inhibits male germ cell apoptosis induced by deprivation of survival factors to human seminiferous tubules in culture.\textsuperscript{176} A study of the effect of estradiol on testicular function in hpg mice with slow-release estradiol implants, which achieved circulating estradiol concentrations of approximately 40 pg/ml show full qualitatively normal spermatogenesis after 70 days of treatment illustrating that spermatogenesis could be restored in the hpg mice with estradiol.\textsuperscript{177} Studies from our laboratory show a distinct increase in Fas-FasL expression in rat spermatogenic cells upon exposure to DES leading to germ cell death. This increase was confined to the spermatid population, which correlated with increased apoptosis seen in the haploid cells. Testosterone supplementation was able to prevent DES-induced Fas-FasL up-regulation and apoptosis in the spermatogenic cells. DES-induced germ cell apoptosis did not occur in Fas-deficient lpr/lpr mice.\textsuperscript{125} Subsequently, using an in vitro model we provided evidence for the amplification of the death-inducing signals through mitochondria resulting in cytochrome c release. The activation of the apoptotic pathway occurred through estrogen receptors because estrogen antagonists prevented the activation of apoptosis. Since a part of this study was done with germ cell cultures independent of Sertoli cells, the importance of the independent capability of cells of the spermatogenic lineage to respond to estrogens was established.\textsuperscript{178} For the role of Fas/FasL in the testis, a review by ref. 179 is recommended. Estradiol treatment increases germ-cell apoptosis mainly at stages IV-X of the spermatogenic cycle, rather than at stage VII when apoptotic germ-cell death is triggered by gonadotrophin withdrawal in adult rats.\textsuperscript{180} Testis regression in Syrian hamsters is associated with an important increase of apoptosis 3 days post DES administration primarily affecting spermatocytes and, to a much lesser extent, spermatogonia.\textsuperscript{181} A combined regimen of gossypol plus methyltestosterone and ethinylestradiol as a contraceptive induces germ cell apoptosis in rats.\textsuperscript{182} In another model of seasonally breeding bank vole testis, males treated with a high dose of estradiol or its antagonist ICI 182,780, disruption of testicular structure and tubular atrophy showed increased apoptosis of germ cells.\textsuperscript{183} Zearalenone, a non-steroidal estrogenic mycotoxin induces testicular germ cell apoptosis in a time-dependent and stage-specific pattern, peak frequency of apoptosis gradually progressing at stages I-VI of seminiferous tubules with time after dosing, suggesting that the damaged germ cells, especially spermatogonia and spermatocytes, gradually underwent the processes leading to apoptosis.\textsuperscript{184}
Future Directions

There are many important questions regarding the role of estrogens in male fertility that are still to be completely answered. For example, is there a physiological role for estrogen in male reproduction through the pituitary-gonadal axis or is there a local effect? Can exposure to low doses of estrogen or estrogenic substances interfere with spermatogenesis and male fertility? What kind of estrogen receptors is most important for testicular function and in which cell types? It is also necessary to identify putative causal agents by the systematic screening of environmental chemicals and chemicals present in human foods to assess their ability to disrupt the endocrine system. In addition, it will be necessary to develop methods to measure cumulative exposure to estrogen mimics.

Sperm counts in Western countries might have fallen by approximately 50% since the 1930s to 1940s and was an issue about environmental effects on the male. The initial study by Carlsen and colleagues was subjected to much criticism and reanalysis. Independent reanalysis reached exactly the same conclusions as had the original study, and a relatively recent updated analysis that included semen analysis data up to 1996 (101 studies in all) again confirmed the trends and conclusions of the original study. However, these studies are all based on meta-analyses of retrospective data, and therefore, fresh systematic studies are needed to readdress the questions of sperm count and environmental exposure to estrogenic chemicals.

It is known that disorders of sexual differentiation, a hormonally mediated process that occur in the fetus in utero are associated with a high risk of developing testis cancer in young adulthood. In utero DES exposure has been linked to testis cancer. Estrogen exposure in the first trimester of pregnancy is particularly important as it is associated with a small but significant increase in risk of developing testis cancer. However, this view has been contradicted. In general, it seems fair to conclude that exposure to exogenous estrogens in early pregnancy results in only a modest increase in risk of developing testis cancer. It has also been speculated that increased risk of testis cancer might stem from increased exposure to endogenous (maternal) estrogens. The premalignant germ cells, the carcinoma in situ cells from which testis cancer arises, have their origins in fetal life. Subnormal androgen exposure and increased estrogen exposure can be the potential stimulus for these cells to be predisposed towards cancerous growth later in life. Few reviews recommended for readings on endocrine disruption and male reproduction see refs. 201-206.

Therefore, it is substantiated by literature that there is a strong possibility of estrogens being responsible for several reproductive disorders. To conclusively prove the role of estrogens where different doses play different roles under different cellular conditions further research is necessary with specific goals. With our increased understanding of the role of estrogens in male fertility, we should be able to design small molecule modulators of estrogen action to interfere with deleterious effects of estrogens.

List of Abbreviations

ArKO, aromatase knockout; ERα, estrogen receptor α; ERβ, estrogen receptor β; ER, estrogen receptor; SERMs, Selective estrogen receptor modulators; hypogonadal; GnRH, gonadotropin-releasing hormone; FSH, Follicle stimulating hormone; LH, Luteinizing hormone; P450arom, aromatase cytochrome P450; HSD, hydroxysteroid dehydrogenases; PKC, protein kinase C; PKA, protein kinase A; ERKO, ERα knockout; BERKO, ERβ knockout; DES, diethylstilbestrol; BPA, Bisphenol A.

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Selenium, a Key Element in Spermatogenesis and Male Fertility
Carla Boitani* and Rossella Puglisi

Abstract
Selenium is essential for normal spermatogenesis of mammals and its critical role is mainly mediated by two selenoproteins, namely phospholipid hydroperoxide glutathione peroxidase (PHGPx/GPx4) and Selenoprotein P. PHGPx/GPx4 is the major selenoprotein expressed by germ cells in the testis, having multiple functions and representing the pivotal link between selenium, sperm quality and male fertility. Selenoprotein P is a plasma protein that is required for selenium supply to the testis. In the last years, nutritional studies and experimental animal models lacking/overexpressing a specific PHGPx isoform and selenoprotein P have highly expanded our understanding on how the male reproductive system depends on selenium. The focus of this review is to report and discuss the most relevant and recent findings in this field. Clinical data have pointed to a correlation between abnormal PHGPx content in sperm and disturbance of human male fertility. However, additional evidence is still required to draw any definitive conclusions about therapeutical strategies for improving fertility by selenium administration.

Introduction
Selenium (Se) is an essential trace element that plays an important role in a number of physiological processes in animals and humans. This element is incorporated into proteins as selenocysteine (Sec, the 21st amino acid), thanks to a peculiar translation reprogramming that allows mRNA UGA codon to be specifically recognized by the selenocysteinyl-tRNA, instead of canonically functioning as stop signal. One or more Sec residues are consistently found in the primary structure of all selenoproteins and they are essential catalytic site components in a variety of selenoenzymes.

The testis represents a specific and privileged target of Se. This element, in fact, appears to be essential for maintaining a normal spermatogenesis and for male fertility. In case of Se deficiency, regulatory mechanisms strive to maintain an adequate level of this element in the male gonad and, when selenium is administered again, the Se is supplied to the testis with priority over other tissues.1 Given the strict dependence of normal sperm production on Se, a particular interest has been addressed to selenoprotein P (SEPP1), a plasma and extracellular selenoprotein predominantly produced by liver, that carries Se to male germ cells.2 In the mammalian testis, almost the entire Se content is associated with the enzyme phospholipid hydroperoxide glutathione peroxidase (PHGPx/GPx4), a member of the large subfamily of the glutathione peroxidases (GPx) selenoproteins, most of which are antioxidant enzymes that...
reduce hydroperoxides at the expense of glutathione (GSH). However, PHGPx has a number of unique features compared to other family members, making the role of this enzyme in the testis worth of particular interest.

In this review, we provide a brief overview of the importance of selenium and selenoproteins in mammalian spermatogenesis and male fertility, and report the most relevant findings in this field.

The Selenoproteins of the Male Gonad

A comprehensive and comparative analysis of all known members of the selenoprotein family has recently been performed in the mouse testis by real-time PCR.\(^3\) As mentioned above, PHGPx/GPx4 mRNA is by far the most abundant among those coding for selenoproteins and large evidence has recently highlighted the different roles this gene plays in the male gonad. Other selenoprotein transcripts approximate levels that are 10-fold lower than that of PHGPx. Among these, the protein product of the Thioredoxin/Glutathione Reductase (TGR) gene, a member of the thioredoxin reductase (TR) family, is expressed in post-puberal testis and is particularly abundant in elongating spermatids at the site of mitochondrial sheath formation, while it is absent in mature sperm. It was recently proposed that TGR cooperates with PHGPx by acting as a novel disulfide bond formation system at the level of structural protein components of the sperm.\(^4\) As for Selenoprotein V (Sel V), Northern blot and in situ hybridisation analyses showed a low, yet testis-specific, expression of this molecule restricted to the seminiferous tubules,\(^5\) the physiological function of which is still unknown. Low abundance transcripts for Selenoprotein W, Selenoprotein K, Selenoprotein 15 and Selenoprotein S also appear in the testis; however, the specific roles of these proteins have not been characterized yet. SEPP1, a typically plasma selenoprotein and also synthesized in the testis, only in the Leydig cells,\(^6\) was recently shown to be required for sperm development by the sterility phenotype of the male Sepp1 knock out mice.\(^7\) In addition, a number of selenoprotein-synthesis key factors are also expressed in the testis, including the Selenocysteine Lyase, an enzyme (not containing Se) that specifically catalyzes Sec decomposition to alanine and elemental Se, and the Selenophosphate Synthetase (SPS2), allowing Se utilization in selenoprotein biosynthesis.\(^8\) Table 1 lists the selenoproteins expressed in the mouse testis.

<table>
<thead>
<tr>
<th>Selenoprotein</th>
<th>mRNA</th>
<th>Protein</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>PHGPx/GPx4</td>
<td>++++</td>
<td>++++</td>
<td>-Antioxidant</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>-Structural protein of sperm midpiece</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>-Structural protein of sperm mitochondrial sheath</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>-Involved in sperm chromatin condensation</td>
</tr>
<tr>
<td>TGR</td>
<td>+</td>
<td>+</td>
<td>-Disulfide bond formation</td>
</tr>
<tr>
<td>SEPP1</td>
<td>+</td>
<td>+</td>
<td>-Selenium delivery and antioxidant</td>
</tr>
<tr>
<td>Sel V</td>
<td>+</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>Sel W</td>
<td>+</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>Sel K</td>
<td>+</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>Sep 15</td>
<td>+</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>Sel S</td>
<td>+</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>SPS 2</td>
<td>+</td>
<td>n.d.</td>
<td>Selenium donor in selenoprotein biosynthesis</td>
</tr>
</tbody>
</table>

n.d., Not determined. Relative abundance: ++++, very high; ++, modest; +, low.
PHGPx/GPx4

Expression of PHGPx in the Testis

The selenoprotein phospholipid hydroperoxide glutathione peroxidase (PHGPx), first purified from pig liver by Ursini and coworkers in 1982, was included among peroxidases, with particular reference to GSH-peroxidases superfamily, because of its ability to reduce phosphatidylcholine hydroperoxides.9 Nowadays we know that this enzyme has unique properties compared with other members of the family, including the ability to reduce the intracellular membrane phospholipid hydroperoxides and to use the thiol groups of proteins as substrates, besides those of GSH,10 particularly when the GSH intracellular level is low.

The Phgpx gene encodes for three isoforms having specific subcellular localization in mitochondria, cytosol and nucleus, respectively, and differing on their N-terminal amino acid sequence. The N-termini of the mitochondrial (mPHGPx) and cytosolic (cPHGPx) variants derive from the same exon 1a by different translation sites, whereas the N-terminus of the nuclear isoform (nPHGPx) is generated by the alternative exon 1b, which in turn is driven by another promoter located in the first intron of the gene.11

It is interesting to note that the testis exhibits the highest specific activity of PHGPx so far measured in mammalian tissues, being almost two orders of magnitude higher than that of brain and liver.11,12 Based on this observation, the interest in the presence and function of different variants of this enzyme in the testis has rapidly increased. During mouse embryogenesis, both mitochondrial and cytosolic PHGPx mRNAs are expressed at higher levels with respect to that of the nuclear isoform. However, the expression of both nuclear and mitochondrial PHGPx is down-regulated between E15.5 and E17.5.13 During postnatal life, PHGPx expression in the testis is initiated at puberty and is hormone-dependent. In both hypophysectomized rats and testosterone-deprived rats, in fact, testicular PHGPx activity and mRNA appeared to significantly decrease and were partially restored by treatment with hCG and testosterone, respectively.14,15 Our group has studied the cellular distribution of different PHGPx isoforms within the male gonad in both rat and mouse. We investigated the patterns of mPHGPx and nPHGPx expression during male germ cell differentiation at both mRNA level and protein level, using highly enriched fractions of pachytene spermatocyte germ cells, as well as fractions of steps 1-8 round spermatids. We demonstrated that nPHGPx is switched on in the post-meiotic phase and that, by contrast, mPHGPx is expressed from mid-late pachytene spermatocytes onwards.16 Moreover, we also observed the presence of abundant catalytic activity of PHGPx and/or nPHGPx in purified populations of pachytes spermatocytes and round spermatids, the haploid cells showing the highest activity.16 The presence of PHGPx was also demonstrated in elongated spermatids as well as epididymal spermatozoa by immunohistochemistry with antibodies recognizing the three isoforms.10,17

In spite of PHGPx abundance in the testis, still very little is known about mechanism(s) underlying the massive up-regulation of the gene in this tissue. The existence of two distinct promoters resulting in alternative transcription of mPHGPx/cPHGPx and nPHGPx was conclusively demonstrated by reporter gene analysis.11 Functional cis-regulatory elements were identified within the proximal promoter region of nPHGPx.18 By using highly purified rat spermatid cells, Tramer et al19 were able to demonstrate that nPHGPx expression is mediated by the transcription factor CREM-t, binding a genetic element localized in the first intron of Phgpx gene. In addition, reporter gene assays carried on in a somatic cell line showed that CREM-t expression activates the promoter region of Phgpx intron1a.19

Evidence for More Than One Function for PHGPx in the Testis

The high expression of PHGPx in the testis underlines the relevance of this gene to spermatogenesis. All three isoforms of this enzyme efficiently catalyze the reduction of phospholipid hydroperoxides by oxidation of glutathione, clearly indicating that they are involved in germ cell protection against oxidative damage. In particular sperm cells, which are
provided of a very special membrane enrichment in phospholipids with polyunsaturated fatty acids (PUFA), depend on several scavenging systems, including PHGPx, for protection from oxidative stress. Consistent with this idea, spermatozoa contain a large amount of PHGPx. On the other hand, both in vitro and in vivo evidence has been accumulating in the last years, pointing to other roles for PHGPx beyond the classical antioxidant one. Both mPHGPx and cPHGPx represent almost 50% of the capsule material embedding the spermatozoa mitochondrial helix, where these molecules are apparently enzymatically inactive and form protein aggregates by disulfide bridge cross-linking. The role of PHGPx as structural protein may explain the impairment of sperm motility associated with morphological alterations of sperm midpiece that was observed in Se-deficient animals. In line with this conclusion, spermatozoa of infertile men with asthenozoospermia showed a significant, albeit controversial, decrease of rescued PHGPx specific activity and expression, thus emphasizing the clinical relevance of PHGPx enzyme(s).

As for the nuclear variant, it was proposed to be involved in the stabilization of condensed chromatin during sperm maturation. Indeed, knock-out mice specifically lacking nPHGPx, although viable and fertile, displayed defective chromatin condensation in caput epididymis sperm. This defect was apparently overcome during subsequent epididymal sperm maturation, suggesting that nPHGPx is dispensable for the maintenance of male fertility in the mouse. In agreement with this, the sperm of Se-deficient mice displayed abnormal heads. This complex event is paralleled by loss of glutathione and the thiol-oxidase activity of PHGPx is preferentially addressed to protamines, the nuclear proteins required for sperm head condensation. In contrast, the deletion of all exons of the Phgpx gene results in embryonic lethality between 7.5 and 8.5 dpc of homozygous mice, pinpointing the vital relevance of this gene. Functional silencing of the different PHGPx isoforms during in vitro embryogenesis by short interfering RNA technology, provided a possible explanation for the early intrauterine death of Phgpx-deficient mice. In fact, targeted knockdown of mPHGPx strongly impaired hindbrain development and induced cerebral apoptosis, whereas a silenced expression of the nuclear isoform led to a delayed heart development. These data suggest that mPHGPx and/or cPHGPx are more important for murine embryogenesis than nPHGPx.

Gain-of-function approaches were used to study the protective role of PHGPx against oxidative stress in vivo. Indeed, transgenic mice overexpressing PHGPx in all tissues were more resistant to diquat-induced liver damage, due to the decrease in lipid peroxidation compared to wild-type animals. In line with this finding, mice having a single copy of Phgpx gene displayed an increased sensitivity to oxidative stress produced by gamma irradiation, paraquat and hydrogen peroxide. However, more informative models to investigate specific PHGPx functions in spermatogenesis in vivo consist in conditioned knock-out and/or transgenic animal models that lack/overexpress a specific isoform in a cell/stage-specific manner. In this direction, we investigated the physiological effects of mPHGPx overexpression during early male germ cell differentiation by generating transgenic mice bearing the rat mPhgpx coding sequence driven by the mouse Synaptonemal Complex Protein 1 promoter. This strategy allows the transgene to be specifically activated in the testis from the zygotene to diplotene stages of the first meiotic division. Because endogenous mPHGPx transcripts first appear at the middle-late pachytene stage, the expression of the mPHGXP transgene was not only increased in amount, but also developmentally anticipated. Transgenic mice were fully viable and developed normally, indicating that mPHGPx overexpression did not affect embryogenesis and postnatal life and ruling out the possibility of nonspecific effects due to transgene insertion per se. Interestingly, primary spermatocytes specifically underwent an increase in apoptosis frequency, according to testis- and stage-specific expression of the transgene, which eventually resulted in a severe loss of haploid germ cells and tubular epithelium disorganization. In line with these features, adult transgenic male mice also displayed a reduction in fertility. This study thus suggested that mPHGPx expression is tightly regulated in pachytene spermatocytes and that
any spatial-temporal increase in mPHGPx expression results in a damage to spermatogenesis and eventual loss of haploid cells.34

Selenoprotein P

Selenoprotein P (SEPP1) is a peculiar selenoproteome member, differing from other selenoproteins because of its unique content of several Sec residues (up to 17, depending on the species). SEPP1 is an extracellular protein that is typically found in the plasma and that is mainly secreted by liver, even though it is also produced by other tissues, including testis.6 The hypothesis of SEPP1 function as Se transportation/distribution protein was conclusively demonstrated by studies of Sepp1 gene inactivation in mice.35,36 In fact, Sepp1 knockout mice exhibited a decreased tissue distribution of Se within the organism, being the brain the most affected organ and followed by kidney and testis. In agreement with these findings, these mice had a number of neurological and sterility problems similar to those of mice bearing a specific deletion of the Se-rich C-terminal domain of SEPP1, pinpointing testis and brain as the most Se-sensitive tissues. Consistent with this feature, the expression of selenoproteins, including PHGPx and Sel W, appeared to be significantly decreased in brain and testis, when Sepp1 gene was deleted.3 The Sepp1 knockout mouse model has provided further insight into the mechanisms of Se importance to spermatogenesis. Indeed, mature spermatozoa of these mice displayed a number of flagellar structure defects, including truncated mitochondrial sheath, extrusion of axonemal microtubules and outer dense fibers from principal piece, as well as a hairpin-like bend formation at the midpiece-principal piece junction.7,38 Sperm defects of Sepp1-null mice were indistinguishable from those of wild-type males that had been fed with a low selenium diet (see also next section). However, sperm defects of Se-deficient wild-type mice were reverted by a normal Se diet, whereas in the case of Sepp1 knockout mice, dietary Se supplementation had no effect on sperm phenotype and did not restore fertility,7 reinforcing the notion that the testis strictly depends on Se supply by circulating SEPP1. Accordingly, it was very recently shown that a liver-specific expression of a transgene coding for human SEPP1 in Sepp1 knockout mice can rescue the abnormal testicular phenotype and fertility of these animals, provided they are maintained under normal Se diet.39

It is currently accepted that preferential Se delivery to testis by SEPP1 is mediated by apolipoprotein E receptor 2 (ApoER2), a receptor expressed in the testis, exclusively by Sertoli cells.40 Interestingly, Sertoli cells display a SEPP1 localization at the level of vesicle-like structures at the basal region suggesting SEPP1 internalization via receptor-mediated endocytosis. Moreover, coimmunoprecipitation analysis demonstrated an interaction of testicular ApoER2 with SEPP1. These morphological and biochemical findings are also supported by the genetic evidence that ApoER2-null mice display reduced levels of testis, but not liver, Se, defective spermatozoa and male infertility.40,41 Thus, both SEPP1 and ApoER2 are essential for maintaining normal Se levels in the testis and are functionally linked with each other in the pathway(s) providing Se to male germ cells. However, it is still not known how the Se content of SEPP1 is delivered to spermatogenic cells. In addition, SEPP1 may also be produced in the testis itself. In fact, selenoprotein P mRNA was found to be expressed in rat male gonads, only in Leydig cells.6 However, the physiological function of SEPP1 in this cell type still remains to be elucidated. Another function of SEPP1 is to serve as oxidant defense agent. In fact, it was reported that injections of Se into Se-deficient rats resulted in the appearance of SEPP1 in plasma, correlating closely with an increase in protection against the diquat-induced liver necrosis and lipid peroxidation.42,43

Nutritional Considerations

The nutritional importance of Se for male reproduction has been known for decades and thoroughly assessed by feeding rats with a Se-deficient diet, eliciting the appearance of a number of reproductive disorders, including reduced fertility, sperm impaired motility and abnormal tail morphology.22,44 When a low Se diet was administered for several generations, severe
testicular atrophy that could be reversed by a Se-adequate diet was observed.\textsuperscript{45} As mentioned above, morphological studies of spermatids and epididymal spermatozoa have pinpointed the sperm flagellum as the major target of Se deficiency,\textsuperscript{38} as well as altered shape of spermatozoa head,\textsuperscript{27} associated with incomplete chromatin condensation.\textsuperscript{28}

The effect of Se-deficient, Se-adequate and Se-excess diets on spermatogenesis has extensively been investigated in a series of studies in the mouse, conclusively showing that either a Se deficiency or excess in the diet resulted in an increase in oxidative stress that negatively affected male germ cell number and differentiation and fertility,\textsuperscript{46-49} such effect being modulated by the redox-sensitive and cell proliferation controlling transcription factors, NFkappaB and AP1.\textsuperscript{47,48,50} It thus appears that abnormal Se levels in the diet, namely either deficiency or excess, are both detrimental to animal health and fertility. Moreover, the idea that high levels of antioxidants impair the progression of male germ cell differentiation is also strengthened by the finding of increased germ cell apoptosis in male transgenic mice that specifically overexpressed mitochondrial PHGPx in male meiotic cells.\textsuperscript{34} Consistent with this finding, the dietary administration of a mixture of the anti-oxidant agents vitamins C and E to mice increased the occurrence of sperm head abnormalities and caused a reduction in sperm production.\textsuperscript{51} Unfortunately, however, the alteration of male reproductive potential by increased/decreased Se does not give a clear-cut insight on this issue, in light of the very large number of selenoprotein-coding genes and the multiple body districts where these genes are expressed.\textsuperscript{52}

Clinical Implications

In contrast to experimental animals, the impact of dietary Se on male fertility in man is not proven yet. A study performed in Scotland reported a sperm motility increase in patients that had received an oral Se supplementation, although a positive response to treatment vs. placebo was only observed in 56% of them.\textsuperscript{53} However, owing the very limited tolerance to both a defective and an exceeding dietary Se level in mammals, including humans, any therapeutical strategy of improving fertility by Se administration in man requires that the Se status is accurately assessed before the treatment is initiated.

In light of the structural role that mPHGPx plays on sperm flagellum in the rat, the possibility that a reduction in sperm mPHGPx impairs fertilization in men was thoroughly addressed by several authors. Interestingly, sperm from infertile patients with oligoasthenozoospermia displayed a dramatic decrease in the level of mPHGPx expression, which was also associated with reduced sperm motility and defects in mitochondrial morphology and function.\textsuperscript{23} In addition, Foresta et al.\textsuperscript{24} studying sperm PHGPx activity after a reducing “rescuing” procedure in patients with different etiologies of infertility, found that these patients consistently had a lower level of enzymatic activity than healthy controls. They thus proposed that PHGPx is included among clinical tests for infertility diagnosis. In contrast, no significant difference in PHGPx activity was observed between normo- and hypo-motile human sperm when the enzyme activity was assayed under native conditions.\textsuperscript{25} PHGPx activity may also depend on the genetic background. This issue was addressed by analyzing the whole \textit{Phgpx} gene for polymorphisms in groups of infertile and fertile men and discovering the presence of 11 variant sites in 5 out of 42 infertile patients.\textsuperscript{54} The majority of these variant sites, however, was likely irrelevant to fertility, because they were located in introns, and only 1 of exon variants actually led to an Ala93-Thr exchange that reduced PHGPx activity in the porcine \textit{gpx-4} homologue. A more recent study\textsuperscript{55} did not find any correlation between a low level of sperm PHGPx and the presence of \textit{Phgpx} gene variants in oligoasthenozoospermic patients. Therefore, even though the possibility that \textit{Phgpx} polymorphism seldom represents an actual cause of infertility cannot be ruled out, the actual relationship between quantitative/qualitative PHGPx alteration and infertility in man still remains to be further investigated.
References

Extracellular Matrix and Its Role in Spermatogenesis

Michelle K.Y. Siu* and C. Yan Cheng

Abstract
In adult mammalian testes, such as rats, Sertoli and germ cells at different stages of their development in the seminiferous epithelium are in close contact with the basement membrane, a modified form of extracellular matrix (ECM). In essence, Sertoli and germ cells in particular spermatogonia are “resting” on the basement membrane at different stages of the seminiferous epithelial cycle, relying on its structural and hormonal supports. Thus, it is not entirely unexpected that ECM plays a significant role in regulating spermatogenesis, particularly spermatogonia and Sertoli cells, and the blood-testis barrier (BTB) constituted by Sertoli cells since these cells are in physical contact with the basement membrane. Additionally, the basement membrane is also in close contact with the underlying collagen network and the myoid cell layers, which together with the lymphatic network, constitute the tunica propria. The seminiferous epithelium and the tunica propria, in turn, constitute the seminiferous tubule, which is the functional unit that produces spermatozoa via its interaction with Leydig cells in the interstitium. In short, the basement membrane and the underlying collagen network that create the acellular zone of the tunica propria may even facilitate cross-talk between the seminiferous epithelium, the myoid cells and cells in the interstitium. Recent studies in the field have illustrated the crucial role of ECM in supporting Sertoli and germ cell function in the seminiferous epithelium, including the BTB dynamics. In this chapter, we summarize some of the latest findings in the field regarding the functional role of ECM in spermatogenesis using the adult rat testis as a model. We also highlight specific areas of research that deserve attention for investigators in the field.

Introduction
Spermatogenesis is a precisely regulated process by which one spermatogonium (diploid, 2n) divides and differentiates into 256 spermatids (haploid, 1n) via 14 stages of the seminiferous epithelial cycle with six mitotic and two meiotic divisions in adult rat testes. In order to complete these intriguingly regulated events, there are extensive junction restructuring in the seminiferous epithelium at both the blood-testis barrier (BTB; note: BTB is a testis-specific structure composed of side-by-side arranged tight junctions [TJ], the basal ectoplasmic specialization [ES], the basal tubulobulbar complexes [TBC], both are testis-specific adherens junction [AJ] types, and the desmosome-like junctions [DJ]) between adjacent Sertoli cells; and anchoring junctions, such as apical ES, apical TBS, DJs and gap junctions (GJ), between Sertoli and germ cells (see Fig. 1). This thus permits developing germ cells, such as preleptotene...
and leptotene spermatocytes, traverse the BTB at stage VIII of the seminiferous epithelial cycle for further development into round, elongating, and elongated spermatids, yet these cells must remain attached to the nourishing and supporting Sertoli cells. In light of these extensive junction restructuring events during spermatogenesis to accommodate the timely migration of germ cells across the epithelium, it is not entirely unexpected that the morphological layouts of TJ and anchoring junctions in the testis are relatively unique versus other epithelia. Furthermore, unlike other blood-tissue barriers, such as the blood-brain and the blood-retina barriers, which are constituted by endothelial TJs of the microvessels in the corresponding organs namely brain and eyes, respectively, the BTB is contributed almost exclusively by adjacent Sertoli cells near the basement membrane of the seminiferous tubules, and the TJ-barrier in the microvessels in the interstitium contribute little, if any, to the BTB function. Interestingly, the peritubular myoid cell layer in rodent testes was shown to prevent the penetration of electron dense markers, such as lanthanum, colloidal carbon or thorium, into the seminiferous epithelium in almost ~85% of the tubules examined, even though myoid cells in primate testes were much less effective to restrict the penetration of these markers across the BTB. Collectively, these findings illustrate the myoid cell layer in the tunica propria contributes to the BTB integrity, at least in rodent testes.

The BTB, which physically divides the seminiferous epithelium into the basal and the adluminal (apical) compartments, segregating virtually the entire events of post-meiotic germ cell development and maturation from the systemic circulation, is located closely to the basement membrane (a modified form of extracellular matrix, ECM) (Fig. 1). This morphological layout is in sharp contrast to other epithelia where TJ is located at the apical portion of the cell epithelium, to be followed by the adherens belt (composed of AJ) and desmosomes. Such physical intimacy between the BTB and the basement membrane thus illustrates the possible role of ECM on junction dynamics at the BTB in the testis. Indeed, it was reported that infertile patients with aspermatogenesis were shown to have abnormal basement membrane structures. Recent studies have also demonstrated the crucial role of ECM components, such as collagens and laminins, in junction dynamics since these proteins were shown to work in concert with proteases, protease inhibitors, cytokines (e.g., TNFα), and focal adhesion (FA) components found at the ES to regulate the steady-state levels of integral membrane proteins at the cell-cell interface. In this chapter, we intend to highlight the recent advances of how ECM proteins and their partners regulate junction dynamics in the testis.

**Unique Features of Extracellular Matrix (ECM) in the Testis**

ECM, largely composed of glycoproteins and polysaccharides, fills the extracellular space at the cell-cell contact sites. In rodent testes, a specialized form of ECM, constituted largely by type IV collagen and laminins, along with heparan sulfate proteoglycan and entactin, forms the basement membrane (~0.15 μm thick), which encloses each seminiferous tubule and is in contact with the base of Sertoli cells and spermatogonia (Fig. 1). One interesting feature of the basement membrane is that it is adjacent to the blood-testis barrier (BTB), where tight junctions (TJ) coexist with adherens junctions (AJ) (Fig. 1), such as basal ectoplasmic specialization (ES) and basal tubulobulbar complex (TBC), and desmosome-like junctions (DS), unlike other blood-tissue barriers (e.g., blood-brain barrier and blood-retina barrier) where TJs are furthest away from the ECM, and are localized to the apical portion of the epithelium/endothelium, to be followed by AJ, desmosomes and gap junctions.

**Functions of the Blood-Testis Barrier (BTB)**

The BTB divides the seminiferous epithelium into the basal and adluminal compartments and thus creates a unique microenvironment for spermatogenesis (Fig. 1). It maintains an immunological barrier by sequestering post-meiotic germ cell development from the systemic circulation, regulates the passage of molecules into the adluminal compartment or vice versa and confers cell polarity. As such, developing germ cells depend exclusively on Sertoli cells...
Figure 1. A schematic drawing illustrating the latest model on the regulation of junction dynamics in adult rat testes, including the blood-testis barrier (BTB) and the ectoplasmic specialization (ES).

For instance, junction restructuring events that occur at the blood-testis barrier (BTB) and the apical ectoplasmic specialization (apical ES) in the seminiferous epithelium during spermatogenesis apparently are regulated via intriguing interactions between cytokines (e.g., TNFα), proteases (MMP-2, MMP-9, MT1-MMP), protease inhibitors (TIMP-1, TIMP-2), collagens, laminins, adaptors, kinases and phosphatases. Legend continues on following page.
Testicular Development and Spermatogenesis

Figure 1, continued from previous page. As described in the text, TNFα regulates the homeostasis of the proteases and protease inhibitors in the basement membrane, which in turn affects the collagen ultrastructural network, perhaps forming biologically active fragments that regulate BTB and/or ES dynamics. However, it remains to be shown if NC1 domain of collagen α3(IV) is indeed responsible for the transient “opening” of the BTB to accommodate preleptotene spermatocyte migration across the barrier that occurs at stage VIII of the seminiferous epithelial cycle, which should be vigorously examined in future studies. Recent studies have shown that similar mechanism(s) is also operating at the apical ES to regulate the transient opening and/or closing of the apical ES to facilitate spermatid movement during spermatogenesis and perhaps also the cellular events that occur at spermiation at late stage VIII of the epithelial cycle, which also involve the participation of proteases and protease inhibitors. This figure was prepared based on recent findings in the field as described in the text.

for structural, anchoring and nutrient supports. Although BTB confers one of the tightest blood-tissue barriers in mammalian body, it is highly dynamic in nature since it must ‘open’ (or ‘disassemble’) at stage VIII of the epithelial cycle in adult rat testes on the apical portion of the migrating preleptotene and leptotene spermatocytes and then ‘close’ (or ‘reassemble’) at the basal portion of the cell to facilitate cell migration while maintaining the barrier integrity. Without this, spermatogenesis cannot complete. At present, the mechanism(s) governing this timely BTB restructuring is not entirely clear. However, recent studies have shown that ECM components, such as collagen IV, are working in concert with proteases, protease inhibitors and cytokines (e.g., TNFα) to regulate TJ dynamics in the testis.8,9,14

Collagen IV

Its Expression and Localization in the Testis

Type IV collagen and laminins are the building blocks of the basement membrane in the testis.7,23,25 Type IV collagen network is formed by the association of monomer, which is a triple helical structure composed of three α chains.25,26 Each monomer is characterized by an N-terminus noncollagenous 7S domain (~15 amino acid residues), a middle collagenous domain (~1400 residues of Gly-Xaa-Yaa repeats) and a carboxyl terminal noncollagenous (NC1) domain (~230 residues). There are six genetically distinct α chains including ubiquitous α1(IV) and α2(IV) chains and more restricted α3(IV)-α6(IV) chains.26,27 α1(IV)-α5(IV) chains are present in rodent testes.28-30 Moreover, α3(IV) and α4(IV) chains are the major (~80%) collagen chains found in the basement membrane of bovine testes,31 implicating the unique structural and/or functional role of α3(IV) and α4(IV) chains in the seminiferous tubule basement membrane in the testis. Collagen α1(IV) and α2(IV) chains are products of Sertoli and myoid cells,28,32,33 whereas α3(IV) is a product of Sertoli and germ cells in the rat.34

Roles in TJ Dynamics

There is mounting evidence that collagen functions perhaps not just as a scaffolding protein.23 For instance, recent studies have shown that Sertoli cell TJ-barrier assembly in vitro was associated with a transient but significant increase in collagen α3(IV) indicating de novo synthesis of collagens is associated with TJ assembly, suggesting the involvement of collagen α3(IV) in TJ dynamics.14 Furthermore, the presence of an anti-collagen antibody in Sertoli cell cultures during TJ-barrier assembly reversibly disrupted the TJ-barrier, further supporting that the interference of an ECM function affects TJ dynamics. Although the underlying mechanism is presently unclear, subsequent studies have shown that these effects were mediated, at least in part, by cytokines, such as TNF-α, which regulate ECM homeostasis via proteolysis.14
TNFα

**TNFα and its Receptor and Testicular Function**

ECM harbors a pool of cytokines, such as TNFα, which can be released when ECM proteins, such as collagens, are degraded. TNFα (a ~50 kDa trimeric protein, consisting of three identical subunits of 17 kDa each) is produced mainly by activated monocytes and macrophages in the systemic circulation, and is crucial to inflammation, cell proliferation and apoptosis. In the testis, TNFα is a product of germ cells (e.g., round and elongating spermatids), macrophages, and Sertoli cells. Its receptors, p55 and p75, are two structurally related, but functionally distinct receptors found in epithelial cells, including Sertoli cells; however, the p55 TNFα receptor (TNFR p55) in Sertoli cells is the main receptor for TNF signaling. In the testis, TNFα plays a crucial role in regulating germ cell apoptosis, Leydig cell steroidogenesis as well as junction dynamics (Fig. 1). For instance, it is now known that in adult rat testes, the number of Sertoli cells, at ~40 million cells, remain relatively stable throughout the adulthood since by day 15 post-partum, Sertoli cells cease to divide. As such, these limited number of Sertoli cells cannot support an unlimited number of developing germ cells. Indeed, it has been shown that each Sertoli cells support ~40-50 developing germ cells in adult rodent testes, and that as much as 75% of the developing germ cells undergo apoptosis and/or spontaneous degeneration, failing to become mature spermatozoa, which is the mechanism being used in the seminiferous epithelium to regulate the precise number of developing germ cells. Interestingly, TNFα was shown to reduce germ cell spontaneous degeneration in rat and human seminiferous tubules cultured in vitro, illustrating its germ cell survival promoting effect.

**Roles in TJ Dynamics**

The role of TNFα on BTB dynamics has been elucidated by both recent in vitro and in vivo studies and a summary of the results on these studies are depicted in Figure 1. In a recent in vitro study, the presence of recombinant TNFα was shown to perturb the Sertoli cell TJ-permeability barrier dose-dependently and specifically since the disrupted TJ-barrier can be resealed upon its removal. This in vitro effect of TNFα on the Sertoli cell TJ-barrier was also confirmed by an in vivo study. In this study, transient and reversible BTB disruption was shown when adult rats were treated with 2 μg recombinant TNFα per testis, which is comparable to its endogenous intratesticular level (~0.5 μg per testis when estimated by a solid-phase immunoblot assay), via an intratesticular injection and assessed by electron microscopy, fluorescent microscopy and a functional assay that monitors the diffusion of a fluorescent dye (fluorescein thioisocyanate, FITC, Mr 389) from the systemic circulation to the seminiferous epithelium behind the BTB. These in vitro and in vivo studies, along with the observations that the expression of TNFα is stage-specific, being highest at stages VII-VIII, coinciding with the events of preleptotene and leptotene spermatocyte migration across the BTB, further support the hypothesis that TNFα secreted by Sertoli and germ cells into the microenvironment at the BTB at stage VIII contributes to the transient BTB “opening” to assist preleptotene spermatocyte migration. This effect of TNFα in “opening” the BTB perhaps is working in concert with its germ cell survival promoting ability so that the migrating preleptotene spermatocytes that likely to take place in “clones” would not undergo spontaneous degeneration. For instance, it is known that germ cell maturation and development occur in “clones” via inter-cellular bridges as they traverse the seminiferous epithelium. Perhaps it is important in future studies to design functional experiments to assess if the level of TNFα at the BTB microenvironment is sufficient to induce BTB restructuring while promoting germ cell survival.

**Regulation of TJ Dynamics: Effects on TJ-Proteins and ECM Proteins**

Furthermore, TNFα apparently exerts its effects on the Sertoli cell TJ-permeability barrier function by regulating the expression of TJ-proteins (e.g., occludin) as illustrated by both in vitro
and in vivo studies,\textsuperscript{12,14} thereby determining the steady-state protein levels of the integral membrane proteins at the BTB. Besides regulating TJ-proteins, TNFα was shown to induce Sertoli cell collagen α3(IV), matrix metalloprotease (MMP)-9 and tissue-inhibitor of metalloproteases (TIMP)-1, and to promote the activation of pro-MMP-9 to proteolytically active MMP-9.\textsuperscript{14} MMPs and TIMPs are proteases and protease inhibitors respectively, that work synergistically to regulate ECM remodeling.\textsuperscript{49} As such, these results suggest that the activated MMP-9 induced by TNF-α may be used to breakdown the existing collagen network by cleaving collagen IV, separating the middle collagenous domain from the N-terminal 7S and the COOH-terminal NC1 domains in the ECM. Such cleavage process possibly affects the scaffolding function of ECM,\textsuperscript{49} thus inducing a loss of other basement membrane proteins (e.g., laminins) and cytokines (e.g., TNFα and TGF-β), which, in turn, contributes to TJ disruption and BTB restructuring because Sertoli cells can no longer attach to an intact ECM. Furthermore, the released biologically active fragments, the NC1 domains, can bind to the middle collagenous domain, inhibit the assembly of intact collagen IV network.\textsuperscript{50} Also, these biologically active fragments can have a negative feedback effect that inhibits collagenase production and thus affecting collagen degradation.\textsuperscript{51} As such, the induced collagen α3(IV) and TIMP-1 by TNFα may be a negative feedback mediated by the biologically active fragments, so as to replenish the collagen network in the disrupted TJ-barrier and limit the activity of MMP-9. Obviously, this hypothesis must be vigorously examined in future studies. In short, the following question must be address. First, can the collagen α3(IV) NC1 synthetic peptides regulate Sertoli cell MMP-9 and TIMP-1 production and/or their activation using Sertoli cells cultured in vitro? If they can, can they also regulate Sertoli cell TJ-permeability barrier when administered in vitro, or perhaps in vivo? Second, can these in vitro studies be reproduced in vivo that an administration of the NC1 domain peptides intratesticularly that leads to a disruption of the BTB function by disrupting the levels of MMPs and TIMPs in the basement membrane?

After reviewing the involvement of collagen in TJ dynamics, the following section introduces the crucial role of laminin, another ECM component, in ES dynamics.

**Ectoplasmic Specialization (ES)**

ES is a testis-specific, actin-based adherens junction residing in the basal (defined as basal ES) and apical (defined as apical ES) compartment of the seminiferous epithelium.\textsuperscript{3,52,53} Both basal and apical ES consist of a layer of hexagonally packed actin bundles sandwiched between the plasma membrane of the Sertoli cell and the cisternae of endoplasmic reticulum. Basal ES is localized at the Sertoli-Sertoli cell interface at the BTB, present side-by-side with TJ, desmosome-like junctions and gap junctions. Apical ES is found between the heads of developing elongating/elongate spermatids (step 8 and beyond in rat and mouse testes) and Sertoli cells which persists until replacing by apical tubulobulbar complex (apical TBC) restricted to the concave side of the elongated spermatid heads just a few hours before spermiation that occurs at late stage VIII of the seminiferous epithelial cycle in adult rat testes.\textsuperscript{20,52}

**Basal ES**

Cadherins\textsuperscript{54} and nectin-2\textsuperscript{55} are two AJ transmembrane proteins that are currently found at the basal ES. Recent study has shown that there is an engagement/disengagement mechanism between basal ES and TJ proteins via their corresponding peripheral adaptors, catenins and ZO-1,\textsuperscript{36} perhaps being used to reinforce the BTB conferring its barrier function, making the BTB as one of the “tightest” blood-tissue barriers in the mammalian body. Such mechanism was suggested to facilitate preleptotene/leptotene spermatocyte migrate across the BTB at stage VIII of the epithelial cycle that the TJ and basal ES proteins become “disengaged” during BTB restructuring to facilitate germ cell migration across the barrier. However, much study is needed to elucidate the intriguing cross-talk mechanism(s) between basal and apical ES since the “opening” (or restructuring) of the BTB near the basement membrane and the disruption of the apical TBC at spermiation at the luminal edge of the
epithelium take place almost simultaneously and since both ultrastructures are present at the opposite ends of the Sertoli cell epithelium, it is not entirely unexpected these events are intimately regulated in the Sertoli cell. Indeed, recent studies have shown that FAK (focal adhesion kinase) is restricted to the basal ES at the BTB where its activated and phosphorylated form, pFAK, is restricted to the apical ES, suggesting that protein kinases that are found at the apical and basal ES are likely play a crucial role to coordinate cross-talk between different cellular events that occur at the opposing ends of the Sertoli cell epithelium.

Apical ES: A Hybrid Cell-Matrix-Cell Junction Type

Besides cadherin-catenin, nectin-afadin protein complexes that are found at both basal and apical ES, apical ES also consists of integrin-laminin complex, which is usually restricted to the focal contact in cell-matrix interface in other epithelia. Such hybrid cell-matrix-cell junction type is suggested to be involved in the rapid junction remodeling facilitating the orientation and movement of spermatids at spermiation. While most of the previous studies on ES were largely focus on its morphology, recent studies have shifted the focus to identify the putative components in the ES in order to explore the mechanisms that regulate ES restructuring during the epithelial cycle. These findings will be summarized and discussed herein; in particular, recent data regarding the involvement of laminin-integrin complex and its downstream effectors in facilitating germ cell migration are discussed.

Integrin: The First FA Component Found at the Apical ES

Integrin, a heterodimeric transmembrane receptor composed of α and β subunits, is the first integral membrane protein positively identified at the ES. To date, there are 18 α subunits and 8 β subunits present in mammals. Among them, α1, α3, α4, α5, α6, α9, β1, β2 and β3 integrins have been identified in testes. α1, α3, α9, β1 subunits are detected in the basement membrane of the seminiferous epithelium, whereas α1, α4, α5, α6 and β1 subunits are found at the apical ES. When β1 integrin subunit was first detected at the ES in 1992, this study first demonstrated the presence of a ECM-associated protein at the nonbasement membrane site namely the ES, since studies from other epithelia have shown that integrins are largely restricted to focal contacts and hemidesmosomes at the sites of cell-matrix anchoring junctions. For instance, β1 integrin is a known receptor for a wide variety of ECM including collagens, fibronectin and laminin in other epithelia. However, there is no ECM protein present in nonbasement membrane at the time when integrin was first reported at ES. Until recently, more than 10 years after integrin being detected at ES, laminin α3β3γ3 and other FAC component proteins were detected and structurally linked to integrin at the apical ES. Such findings, along with the presence of proteases activity at the apical ES site, illustrating that apical ES is utilizing the most efficient migration device usually restricted to cell-matrix anchoring junctions to facilitate germ cell movement across the epithelium during spermatogenesis.

Laminin 333 and α6β1 Integrin form a Bona Fide Complex at the Apical ES

Laminins are heterotrimers composed of one each of the α, β, and γ chains. To date, 5 α-subunits, 4 β-subunits, and 3 γ-subunits have been found in mammalian tissues, which can give rise to at least 16 different functional laminins. By binding to their transmembrane receptors, integrins, at the cell/matrix anchoring junctions, also known as focal contacts constituted by focal adhesion complexes (FAC), laminins and integrins provide not only adhesion between epithelial cells and basal lamina, they also effect signaling through the downstream effectors, the FAC, leading to cell migration during normal and in pathological conditions, such as tumor invasion. Laminin γ3 was the first laminin subunit found at the apical ES by immunofluorescent microscopy and was shown to form a bona fide complex with β1 integrin by communoprecipitation studies. Subsequently, this γ3 chain was found to form a functional laminin protein complex with the α3 and β3
chains, known as laminin 333, which are restricted to the elongating/elongated spermatids and also interact with β1 integrin at the apical ES.57 Perhaps the most important of all, the pivotal role of laminin 333 at the apical ES was demonstrated by the perturbed adhesion between Sertoli and germ cells (mostly spermatids), leading to germ cell loss from the epithelium following treatment of adult rat testes with the laminin blocking antibodies, including anti-laminin α3 or γ3 IgG.57

Proteolysis of Laminin by MMP-2 and MT1-MMP Regulate Apical ES Dynamics

At apical ES, not unlike the cell-matrix interface, proteolysis may also present to regulate its restructuring. For instance, the remodeling of laminin can occur via the effects of proteases since MMP-2, MT1-MMP (a membrane anchored metalloprotease that can activate MMP-2) and TIMP-2 were found to colocalize with laminin γ3 and β1 integrin at the apical ES in adult rat testes.13,74 Furthermore, MMP-2 and MT1-MMP were shown to be activated when germ cells, especially spermatids, were detached from the epithelium in vivo after treating rats with Adjudin, formerly called AF-2364 [1-(2,4-dichlorobenzyl)-1H-indazole-3-carbohydrazide], which is a potential male contraceptive derived from indazole-3-carboxylic acid and has the capability to selectively disrupt adherens junction between Sertoli cells and germ cells.75-77 Perhaps the most important of all, the use of a specific MMP-2 and MMP-9 inhibitor, (2R)-2-[(4-biphenylsulfonyl)amino]-3-phenylpropionic acid, could effectively delay the loss of spermatids from the epithelium induced by Adjudin, indicating the potential role of proteolysis in apical ES disassembly.13 Such proteolysis of laminin by MMPs leading to the production of laminin fragments at apical ES may be essential for spermatid movement and spermiation since laminin-5 fragments has been shown to affect migration in both breast epithelial cells and prostate cancer cells (see Fig. 1).78-81 This possibility should be vigorously tested in future studies.

FA Complexes at the Apical ES

The laminin integrin complex at the apical ES confers its cell-matrix FA property. Such property is further confirmed by the discovery of numerous FA components (see Table 1), including β3-integrin, vinculin, c-Src, Csk, ILK, phosphatidylinositol 4,5-bisphosphate (PI(4,5)P2), phospholipase C (PLC)-γ, Fyn and Keap1 in the ES site.2,53,64,82-86 Recent findings in our laboratories further explore another vital FA component, phosphorylated focal adhesion kinase (pFAK), and its downstream effectors, the p85 subunit of phosphatidylinositol 3-kinase (PI3K), protein kinase B (PKB), p21 activated kinases (PAKs) and Crk- associated substrate (CAS), at the apical ES.15,16 FAK is a nonreceptor protein tyrosine kinase (PTK) that may be a crucial linker for β1 integrin, recruiting ES components to apical ES.15 When FAK interacts with β1 integrin, FAK undergoes autophosphorylation at Tyr-397, creating high-affinity-binding site for multiple molecules, including (i) SH2-domain-containing molecules, such as Src family protein kinases, (ii) effector proteins, such as PI3K and PLC-γ, and (iii) adaptor proteins, such as growth-factor-receptor-bound protein (Grb)7 and Nck-2.87-89 Furthermore, the newly recruited Src-family kinases at the apical ES can further enhance FAK catalytic activity by inducing phosphorylation of FAK at Tyr-576 and Tyr-577 in the kinase domain activation loop. Two other FAK-associated proteins, CAS and paxillin, can also be phosphorylated by Src-family kinases, leading to Rho family GTPase-mediated cell motility.87-95

Recent studies using both in vitro and in vivo models, including Adjudin and androgen suppression models, to study ES dynamics have illustrated the involvement of several signaling pathways which are initiated by β1 integrin/pFAK during apical ES restructuring. These pathways include (i) the integrin/pFAK/c-Src/pERK,15 (ii) the integrin/pFAK/PI3K/pPKB/PAK/pERK16 and (iii) the integrin/pFAK/c-Src/Cas/CrkDock180 (Siu and Cheng, unpublished observations) (see Fig. 1). All these three pathways have the ability to modulate cell adhesion, migration, tissue remodeling and development, and tumor cell metastasis as shown in studies of other epithelia.87-95 These signaling pathways were shown to be triggered within a few hours after treating adult male rats with a single or multiple doses of Adjudin (40-50 mg/kg b.w.)
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<tr>
<th>Proteins</th>
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<th>Binding Partners</th>
<th>Functions/ Properties</th>
<th>Phenotypes in Knock-Out Mice</th>
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<td>Laminin γ3</td>
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<td>β1 Integrin, pFAK&lt;sup&gt;397&lt;/sup&gt;, c-Src, MMP-2, MT1-MMP</td>
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<td>Embryonic lethality on E5.5</td>
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<td>pFAK&lt;sup&gt;397&lt;/sup&gt;</td>
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<td>MMP-2, MT1-MMP</td>
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Table 1. Continued

*This table was prepared based on the following articles and/or reviews (1-3, 7-9, 13, 15, 16, 57, 58, 68, 69, 87). Due to the page limit, many original articles were not cited, however, these references can be found in the cited reviews and/or articles listed herein.

Abbreviations used: n.k., not known; CAR, coxsackie- and adenovirus receptor; Crk, an oncogene identified in a chicken sarcoma called chicken tumor virus number 10, encoding an activator of PTK; Csk, carboxyl-terminal Src kinase, a PTK that phosphorylates a Tyr residue in src family kinases; Dock180, CED-5 (cell death abnormal-5)/180 kDa protein downstream of chicken tumor virus number 10 (Crk); ERK2, externally regulated kinase-2, a mitogen activated protein (MAP) kinase; FAK, focal adhesion kinase; Fer kinase; the Fujinami sarcoma/feline sarcoma (fps/fes) proto-oncogene encoding a 94 kDa nonreceptor PTK called Fps/Fes kinase; ILK, integrin-linked kinase; MMP-2, matrix metalloprotease-2; MT1-MMP, membrane-type 1-matrix metalloprotease; MTMR2, myotubularin related protein-2; PI(4,5)P₂, also called PtdIns(4,5)P₂, phosphatidylinositol 4,5-bisphosphate; PTK, protein tyrosine kinase; PI3K, phosphoinositide (or phosphatidylinositol) 3-kinase; P130 Cas, Crk-associated protein encoded by the Crkas gene; PKB, protein kinase B, also known as Akt, a Ser/Thr protein kinase, a product of the normal gene homolog of v-akt, the transforming oncogene of AKT8 virus; PLC-γ1, phospholipase C-γ1; TIMP-2, tissue inhibitor of metalloproteases-2; PAK, p21-activated kinase, a Ser/Thr protein kinase; PDK1, 3-phosphoinositide-dependent protein kinase 1; PTEN, phosphatase and tensin homolog deleted on chromosome 10, a protein tyrosine phosphatase that shares homology with tensin, and a tumor-suppressor gene located on chromosome 10q23; p120ctn, p120 catenin; c-Src, a nonreceptor PTK of the transforming gene of Rous sarcoma virus; WASP, Wiskott-Aldrich Syndrome protein.
either via i.p. or by gavage, which also matched quite nicely with the subsequent germ cell depletion events, especially spermatids at the apical ES, at 6-8 h after treatment.\textsuperscript{15,16} Perhaps the most important of all, pretreatment of rats with anti-β1 integrin antibody,\textsuperscript{16} PP1 (a c-Src inhibitor),\textsuperscript{59} wortmannin (a PI3K inhibitor),\textsuperscript{16} or U0126 (an ERK inhibitor)\textsuperscript{96} via intratesticular injection were shown to delay the Adjudin-mediated spermatid loss from the epithelium, further confirming the involvement of these signaling pathways in the regulation of apical ES restructuring. Furthermore, the integrin/pFAK/c-Src/pERK pathway has recently been validated and expanded by another in vivo model, the androgen suppression model, in which rats were treated with androgen and estrogen implants to suppress the intratesticular androgen level, leading to the alteration of the Sertoli-germ cell apical ES function and the subsequent germ cell sloughing.\textsuperscript{60} All of these findings thus illustrate that the cell-cell anchoring junction in the testis is indeed a hybrid cell-cell and cell-matrix junction type.

Furthermore, recent studies have also demonstrated the presence of TJ component proteins at the apical ES, which include the coxsackie and adenovirus receptor (CAR)\textsuperscript{97-99} and JAM-C (junctional adhesion molecule-C).\textsuperscript{100} These results thus illustrate that the apical ES is also having the structural and perhaps the functional properties of the TJ. While the precise physiology underlying these observations is not entirely clear, it is increasingly clear that the apical ES is adopting some of the best features found in AJ, focal contacts and TJ to regulate the rapid junction restructuring event pertinent to spermatogenesis.

**Concluding Remarks and Future Perspectives**

As briefly reviewed herein, there are mounting evidence illustrating the pivotal role of the basement membrane, a modified form of ECM, on the junction restructuring events that occur at the Sertoli-Sertoli and/or Sertoli-germ cell interface at the BTB and ES, many of which are mediated via the effects of cytokines (e.g., TNFα and TGF-β3) on the steady-state levels of the integral membrane proteins at these sites. Interestingly, some of these effects are likely mediated via the homeostasis of the proteases and their endogenous inhibitors, which in turn, affects the structural and physico-chemical properties of the basement membrane and/or protein levels at the cell-cell interface. It is obvious that much new information will be added in the years to come and some of the postulates put forth here and depicted in Figure 1 schematically will be updated and/or rewritten. Perhaps it is also important that future studies should include a detailed analysis on the peritubular myoid cells and their role on the BTB function, spermatogonial stem cell renewal, and perhaps ES restructuring such as the use of Sertoli-myoid cell and Sertoli-germ-myoid cell cocultures. The recent deployment of molecular, biochemical and cellular techniques to study junction dynamics in the testis has yielded some unprecedented opportunities for investigators to identify new leads to develop male contraceptives. They also offer exciting opportunities to understand the impact of environmental toxicants on male reproductive physiology.

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Chapter 6

Inflammatory Networks in the Control of Spermatogenesis
Chronic Inflammation in an Immunologically Privileged Tissue?

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Abstract

Spermatogenesis is a complex, organized process involving intimate interactions between the developing germ cells and supporting Sertoli cells. The process is also highly regulated. Studies suggest that regulation in the seminiferous epithelium involves molecules normally associated with either immune or inflammatory processes; in particular, interleukin 1α (IL1α), IL6, tumor necrosis factor (TNFa), activin A and nitric oxide (NO). While there is considerable evidence that these inflammatory mediators have effects on spermatogonial and spermatocyte development as well as critical supportive functions of the Sertoli cells, which are undoubtedly of considerable importance during testicular inflammation, there remains some skepticism regarding the significance of these molecules with respect to normal testicular function. Nonetheless, it is evident that expression of these regulators varies across the cycle of the seminiferous epithelium in a consistent manner, with major changes in production coinciding with key events within the cycle. This review summarizes the evidence supporting the hypothesis that inflammatory cytokines play a role in normal testicular spermatogenesis, as well as in the etiology of inflammation induced sub-fertility. The balance of data leads to the striking conclusion that the cycle of the seminiferous epithelium resembles a chronic inflammatory event. This appears to be a somewhat paradoxical assertion, since the testis is an immunologically privileged tissue based on its well-established ability to support grafts with minimal rejection responses. However, it may be argued that local immunoregulatory mechanisms, which confer protection from immunity on both transplanted tissues and the developing spermatogenic cells, are equally necessary to prevent local inflammation responses associated with the spermatogenic process from activating the adaptive immune response.

Background

Spermatogenesis and the resulting cycle of the seminiferous epithelium are complex and highly organized processes that involve intimate and dynamic interactions between the developing germ cells and their supporting Sertoli cells. The repeating cell associations of the seminiferous epithelium are a consequence of spermatogonia entering the process of spermatogenesis at regularly spaced intervals, which are considerably shorter than the time required for the
entire spermatogonic process, and proceeding through the process at a tightly controlled and predictable rate. In the human, for example, it takes approximately 64 days for a spermatogonium to mature into a structurally complete sperm and to be released from the seminiferous epithelium. During this period, 4 rounds of differentiation of the spermatogonial stem cell population, or waves of spermatogenic initiation, occur. Collectively, this leads to the establishment of a complex stratified epithelium comprising 4 separate generations of spermatogonic cells each at different levels of maturation. Spermatogenesis takes about 35 days in the mouse and 48-50 days in the rat, but in all mammalian species several rounds of spermatogonial differentiation during this time period produce multiple germ cell generations within each seminiferous tubule cross-section. The regular timing of these events means that the generations form distinct and recurring cellular associations, referred to as stages of the cycle of the seminiferous epithelium. In the human, 6 such stages have been described, while in the mouse the number is 12 and in the rat, 14. How this high degree of coordination is maintained remains largely unknown. In particular, what triggers the stem cells to divide and produce the next generation of developing cells at regular intervals? Conversely, what prevents the spermatogonial stem cells and their offspring from differentiating randomly and continuously? Overall, what is the mechanism that coordinates the process across and along the seminiferous epithelium to ensure that the orderly progression of cellular associations, or stages, is maintained?

Mounting evidence suggests that inflammatory regulators play a key role in the initiation of the spermatogenic wave and in many other aspects of germ cell development. These regulators include the well-characterized cytokines interleukin-1 (IL1) and IL6, but also non-proteinaceous mediators of inflammation and immunity, such as nitric oxide (NO). Furthermore, the evidence suggests that such mediators are produced in the testis under normal conditions by somatic cells, including the Sertoli, peritubular and Leydig cells, and the germ cells, rather than by immune cell types.

On the other hand, it is well established that the testis is an immunologically privileged tissue. This is demonstrated by the prolonged survival of grafts into the testicular interstitial tissue, and the ability of cotransplanted testicular cells to confer protection from immunological rejection in non-testicular sites. The mechanisms responsible for immune privilege of the testis remain incompletely understood, but most evidence suggests that the somatic cells of the testis, and the Sertoli cell in particular, play a key role in the regulation of this property. Moreover, it is evident that the most numerous immune cell type within the testis, the resident macrophages of the testicular interstitium, produces extremely low levels of pro-inflammatory cytokines and mediators when challenged with the potent inflammatory mediator lipopolysaccharide (LPS). The fact that these cells also display anti-inflammatory properties and produce cytokines, such as IL10 and transforming growth factor β (TGFβ), when stimulated, suggest that they possess an immunosuppressive phenotype. Such 'alternatively activated' macrophages are generally associated with sites of reduced immune responses. While lymphocytes and inflammatory monocyte-like macrophages also circulate through the testicular interstitium, the unique immune status of the resident macrophages almost certainly serves to reduce the onset and severity of inflammatory and subsequent immunological responses within the testis. Although still a matter for conjecture, these controlled immune/rejection and inflammatory responses are assumed to be in place to benefit the developing spermatogenic cells, which might otherwise be recognized by the host immune system as foreign and come under immunological attack, due to their highly immunogenic properties.

It would appear, therefore, that the testis has something of a split immunological personality. On the one hand, there is clear evidence that inflammatory mediators are produced constitutively within the seminiferous epithelium, yet the testis exhibits considerable resistance to the activation of adaptive immune responses. In other words, it appears that testicular immunological privilege coexists with a seminiferous epithelium that otherwise exhibits characteristics of a chronically inflamed tissue! It will be argued in this review that this arrangement is essential for successful spermatogenesis and that disturbances in the balance between these two
immunological ‘compartments’ of the testis result in either immune-mediated damage or spermatogenic failure, leading to germ cell death. Since dysregulation of the networks involved in these processes during systemic or reproductive tract inflammation almost certainly contributes to infertility, a better understanding of this aspect of testicular function is essential.

Production, Regulation and Actions of Inflammatory Mediators in the Seminiferous Epithelium

In general, inflammation occurs when cells of the mononuclear phagocyte lineage (monocytes and macrophages) become activated. This may be triggered by specific pathogenic molecules (e.g., endotoxins such as LPS), phagocytosis of opsonized (antibody- or complement-coated) particles or immune complexes, and/or various intracellular components released by tissue damage. Activation induces the production of cytokines, acute-phase proteins, proteases and complement components, reactive oxygen and nitrogen species, and lipid mediators, such as prostaglandins and platelet activating factor. Non-myeloid cells are less effective than monocytes/macrophages, but also may initiate a response if they share some of the essential receptors and signaling pathways. In this regard, the Sertoli cell is particularly interesting because of several features it shares with cells of the monocyte/macrophage lineage, not least its ability to respond to LPS, cyclical phagocytic activity during spermatogenesis, and ability to produce a range of inflammatory mediators.

The most intensively studied of the inflammatory mediators produced in the seminiferous epithelium are the cytokines IL1α, IL6, tumor necrosis factor (TNFα) and activin A, and the highly reactive nitrogen molecule, nitric oxide (NO). It is on these molecules that this review will focus. While there is evidence that other inflammatory molecules also are involved (e.g., IL2, IL18, interferons and eicosinoids), they have been excluded because the evidence for their role in regulating spermatogenesis in the adult remains incomplete or speculative. This is not meant to imply that future studies will not bring such roles to light. Moreover, it should be recognized that some of the mediators discussed here, as well as other immunoregulatory cytokines not discussed (e.g., the TGFβs), have effects on fetal and postnatal testicular development, an aspect of testicular biology that also has been excluded from the scope of this review.

Interleukin 1

Interleukin 1 is the most comprehensively studied of all of the cytokines normally expressed in the testis. IL1 is produced in two forms, α and β, which share approximately 25% sequence homology and are encoded by separate genes. Both forms act through the same receptor complex (IL1R), and exert essentially the same effects across a broad range of immunological and inflammatory processes. Signaling occurs principally (although not exclusively) through activation of the MyD88/TRAF and MAP kinase/Jnk pathways, regulating many pro-inflammatory genes through stimulation of the transcription factors, NFκB and AP-1 (Fig. 1). The IL1s are synthesized as 31 to 33 kDa precursor proteins, which are enzymatically cleaved to produce active 17 kDa forms. In the case of IL1α, both the long and short forms are biologically active, but for IL1β the precursor protein is inactive. The precursor of IL1β is cleaved by IL1 converting enzyme (ICE or caspase 1) during the process of secretion into the extracellular space, whereas IL1α is cleaved either by the calcium dependent membrane associated cysteine protease (calpain) or by extracellular proteases. As both IL1α and β lack a signal sequence, however, their mechanisms of secretion remain poorly defined. Nonetheless, IL1β is copiously secreted by activated monocyte/macrophages and is the main secreted form during inflammation. IL1α is also found in secretions, but is more commonly found in association with the cell membrane and is thought to act as an autocrine or paracrine growth factor involved in direct cell-to-cell communication. In general, the majority of IL1α usually remains within the cell, from whence it may be released following cell damage.
There is a third member of the IL1 cytokine group that is homologous with IL1α and IL1β and binds to IL1R, but lacks the ability to transduce a signal. This molecule acts as an antagonist of IL1 action and is called, naturally enough, IL1 receptor antagonist (IL1ra).48 Within the normal testis, the balance of experimental data suggests that IL1α is expressed by the Sertoli cells, where it first appears around day 15-20 in the developing rat testis, and is differentially expressed throughout the cycle of the seminiferous epithelium in the adult (Fig. 2).5,28,31,49 Specifically, IL1 mRNA and protein production occurs at a more or less constant level throughout the cycle, with the exception of stage VII in the rat, when production is low or non-detectable. Both in vitro, and in vivo data suggest that the Sertoli cell production of IL1α is driven by the presence of spermatogenic cells, but it is most effectively simulated by the phagocytosis of residual bodies produced during spermiation at stage VIII of the cycle.31 In contrast to studies clearly showing that inflammation stimulates IL1α production by the Sertoli cells in vitro,28-30 the production of IL1α in the intact rat testis is not stimulated during inflammation in vivo,15,50 indicating that its production is constitutively and possibly maximally up-regulated under normal conditions.

Within the seminiferous epithelium, the IL1R has been localized to both Sertoli and spermatogenic cells.51 In vivo and in vitro data suggest that IL1 stimulates DNA synthesis in intermediate and type B spermatogonia as well as preleptotene spermatocytes,52-55 and acts as
Figure 2. Inflammatory mediator expression during the cycle of the seminiferous epithelium in the rat. Cyclical production by the Sertoli cells (IL1α, IL6 and activin A) and germ cells (TNFα and iNOS) and expression of the FSH receptor (FSHR) are superimposed on critical events during the cycle, such as spermiation, bundling of the spermatids by the Sertoli cells and tight junction (TJ) reorganization to allow meiotic germ cells to pass through the blood-testis barrier. While DNA synthesis occurs throughout the cycle, there are two major peaks of DNA synthesis, at stages IV-VI and at stages VIII-X. The transillumination appearance of the tubules is also shown—the length of each stage of the cycle is presented in proportion to the time taken for each stage to occur. The figure is based on integrated mRNA and protein data from several studies,9,28,32,49,53,84,123,163,191 and is intended to show relative changes only—some details are approximations based on the available data. Figure modified from original.32
an autocrine regulator of several Sertoli cell functions involved in the support of spermatogenesis, including the production of lactate and transferrin.\textsuperscript{56,57} It should be noted that Sertoli cells also secrete a 24 kDa testis-specific form of IL1α, which is the product of an alternative mRNA transcript.\textsuperscript{58} The 24 kDa variant lacks the calpain cleavage site and consequently is not post-translationally cleaved. The physiological significance of this variant is not entirely clear, although it does possess biological activity.\textsuperscript{58,59} It is worth noting here that expression of IL1α also has been reported in isolated late pachytene spermatocytes and round spermatids germ cells,\textsuperscript{60,61} although these data are in conflict with other studies.

Although immunohistochemistry has localized IL1β to both the seminiferous epithelium and interstitial tissue of normal mice,\textsuperscript{60} quantitative studies in the rat suggest that IL1β expression is comparatively very low in the normal testis.\textsuperscript{6,15,49} During inflammation induced by LPS in intact rats, IL1β expression is up-regulated in the testis, but to a much lesser extent than normally occurs in other tissues, such as the liver.\textsuperscript{15} In vitro and in vivo studies suggest that production arises from the Leydig cells and a subset of monocyte-like macrophages in the testis, rather than from cells of the seminiferous epithelium.\textsuperscript{15,50,62,63} Testicular interstitial fluid IL1β concentrations, as measured by ELISA, also increased following injection of LPS in adult rats, but surprisingly, there was no increase in the overall IL1 bioactivity of testicular interstitial fluid. This was assumed to be due to the constitutively high production of IL1α by the Sertoli cells.\textsuperscript{8} A similar observation had been reported previously,\textsuperscript{50} wherein the authors suggested that the lack of change in IL1 bioactivity may have been due to a compensatory increase in IL1ra produced by the Sertoli cell.\textsuperscript{64} Testicular interstitial fluid fractionation studies, however, established that very little of the IL1β entering the interstitial fluid following LPS stimulation had been cleaved to its active form, suggesting a possible deficiency in the activity of ICE/caspase 1.\textsuperscript{15} It appears that pro-inflammatory cytokine activity may be regulated at both the transcriptional and post-translational level in the testis.

Outside the seminiferous tubules, both IL1α and IL1β have direct inhibitory effects on gonadotropin-stimulated androgen production by the Leydig cells.\textsuperscript{59,65,66} In the mouse, evidence suggests that the major site of inhibition occurs at the level of 17α-hydroxylase/C17-20 lyase (P450c17) expression, with affects of higher doses on the cholesterol side chain cleavage enzyme (P450scc) and 3β-hydroxysteroid dehydrogenase (3β-HSD).\textsuperscript{66,67} In the rat, inhibition appears to involve P450scc, specifically.\textsuperscript{68}

**Interleukin 6**

Interleukin 6 is a member of an important family of mediators involved in the regulation of the acute-phase response to injury and infection, which exert their action via binding to specific receptors that associate with a common membrane signal transducer gp130, leading to the activation of the Jak/Stat and MAP kinase cascades.\textsuperscript{69,70} Although evidence suggests that the Leydig cells actually may be the major source of this cytokine in the testis,\textsuperscript{7,71} IL6 is produced by isolated rat Sertoli cells in response to stimulation by follicle-stimulating hormone (FSH) and testosterone, or by phagocytosis and other inflammatory stimuli, including IL1α, IL1β, TNFα and LPS (Fig. 1).\textsuperscript{7,8,28,29,71,72} Within the seminiferous epithelium, IL6 is produced in a coordinated manner (Fig. 2), and FSH differentially stimulates IL6 secretion during the cycle of the seminiferous epithelium.\textsuperscript{28,73} Both gp130 and IL6R mRNA are expressed in rat Sertoli cells, and are stimulated by IL1 and IL6, but only the IL6R subunit is stimulated by FSH.\textsuperscript{74} The data from many studies indicate that that IL1 and IL6 are integrated in a complex network of endocrine and local regulatory mechanisms within the seminiferous epithelium. Stimulation of Sertoli cell IL1α mRNA production, in turn, stimulates IL6 secretion through activation of leukotriene production via the lipoxygenase pathway.\textsuperscript{28,29} This results in an endogenous cyclical pattern of secretion that corresponds with the changes in the stages of the spermatogenic cycle, similar to that of IL1α (Fig. 2). IL6 has been found to act as an inhibitor of meiotic DNA synthesis in preleptotene spermatocytes\textsuperscript{75} and increases basal and FSH-induced transferrin and cyclic GMP secretion by Sertoli cell.\textsuperscript{75,76} On the other hand, IL6 reduces κ
opioid receptor mRNA levels in the Sertoli cell. In models of experimental autoimmune orchitis (EAO), IL6 has been implicated as playing an ameliorative or protective role within the seminiferous epithelium.

**Tumor Necrosis Factor α**

Tumor necrosis factor α is a 17 kDa glycosylated polypeptide, principally produced by activated monocytes and macrophages, which binds as a trimer to either of the two TNF receptors (TNFR1 and TNFR2), and plays a central role in the initiation of the inflammatory response by stimulating the production of IL1 and IL6 (Fig. 1). As its name suggests, TNFα can also exert a cell-death signal via TNFR1, through interaction with the TNF-receptor-associated death domain protein (TRADD) or the Fas-associated death domain protein (FADD), and activation of the caspase-dependent apoptotic pathway (Fig. 1). Whether TNFα has a stimulatory, pro-inflammatory effect, or a destructive effect depends on the receptor subtype engaged and the expression of specific adaptor proteins within the target cell.

In situ hybridization studies in mice have confirmed the presence of TNFα mRNA in round spermatids and pachytene spermatocytes, as well as in testicular interstitial macrophages. Moreover, bioactive TNFα was produced by the round spermatids in vitro and mRNA for the corresponding receptor was located on both Sertoli and Leydig cells. FSH stimulates TNFα receptor subunit protein expression in the Sertoli cell. Expression of TNFα by germ cells within the seminiferous epithelium, like that of IL1α and IL6 in the Sertoli cell, is cyclical (Fig. 2). There is no evidence that TNFα is produced by the Sertoli cell, but treatment of isolated testicular macrophages with LPS induces its secretion.

Within the seminiferous epithelium, TNFα appears to play a complex role the regulation of Sertoli cell function and spermatogenesis. TNFα reduces spontaneous germ cell degeneration in cultured human and rat seminiferous tubules, suggesting a germ cell survival effect mediated via the Sertoli cell. On the other hand, in vitro studies indicate that TNFα disrupts Sertoli cell tight-junction assembly by inhibiting production of the junction protein, occludin and inducing the expression of matrix metalloprotease-9 and its inhibitor tissue inhibitor of metalloproteases-1. Likewise, TNFα has been reported to increase plasminogen activator inhibitor-1 (PAI1) expression in rat testicular peritubular cells, indicating that it plays a key role in controlling testicular protease activity. Similar to IL1, TNFα stimulates basal lactate production by cultured Sertoli cells, but TNFα generally antagonizes the actions of FSH on Sertoli cell function, including the stimulation of aromatase activity and lactate production. Conversely, Delfino and colleagues have shown that TNFα stimulates androgen receptor expression in Sertoli cells via up-regulation of NFκB, which binds to several enhancer motifs in the androgen receptor promoter. These many studies suggest that TNFα produced by the germ cells exerts a complicated paracrine effect on the adjacent Sertoli cell to alter its function at various stages during the cycle of the seminiferous epithelium.

In testicular pathology, TNFα has been implicated as a major causative agent in the development of EAO. In rats with EAO, there is a significant increase in the number of TNFα-positive testicular macrophages and the number of TNFR1-positive germ cells. The majority of TNFα-positive germ cells were apoptotic, suggesting that TNFα could act to trigger germ cell apoptosis in this model, acting together with other local cell death regulatory systems such as the Fas-Fas ligand system. TNFα also stimulates IL6 and leukocyte adhesion molecule expression in Sertoli cells. 

Outside the seminiferous epithelium, TNFα is an effective inhibitor of Leydig cell steroidogenesis acting through the NFκB signaling pathway. Inhibition of LH/hCG binding by TNFα has been reported, but the majority of studies in the mouse suggest that inhibition occurs primarily at the level of steroidogenic gene expression, particularly P450scc, P450c17 and 3β-HSD. In studies on porcine Leydig cells, the inhibitory affect of TNFα also was reported to involve a decrease in steroidogenic acute regulatory protein (StAR) mRNA and protein levels. In non-stimulated or hCG-treated intact or hypophysectomized rats,
intratesticular delivery of TNFα induced a rapid and sustained reduction in StAR protein expression and testosterone biosynthesis.\textsuperscript{105}

\textbf{Activin A}

The activins are homodimers or heterodimers of several homologous subunits (designated $\beta_A\beta_E$), and are members of the much larger TGFβ family of dimeric cytokines. Homodimers of $\beta_A$ form activin A, which is widely expressed and has been extensively studied.\textsuperscript{106} Relatively little is known about the distribution and actions of the other activin forms, which appear to be both less abundant and less widely distributed. Heterodimers of either the $\beta_A$ or $\beta_B$ subunits with an homologous $\alpha$ subunit are feedback inhibitors of FSH secretion from the pituitary, which are produced by the testis and are called inhibin A and inhibin B, respectively.\textsuperscript{107} Conversely, activin A and B act as stimulators of pituitary FSH secretion. Increasing data, however, illustrate the fact that activin A in particular is also a paracrine growth factor and inflammatory regulator.\textsuperscript{108,109} Unlike the other mediators discussed in this review, which have predominantly or exclusively pro-inflammatory actions, the functions of activin A tend towards immunoregulation or immunosuppression, although activin A does appear to play a key role in early inflammation.\textsuperscript{106,109-111}

The regulation of activin A production is still poorly characterized, but its synthesis and secretion is stimulated by IL1 in several cell types, including the Sertoli cells and peritubular cells of the testis.\textsuperscript{112} This regulation probably involves signaling via the p38 MAP kinase/Jnk pathway through the transcription factor AP1, since the promoter of the $\beta_A$ subunit includes AP1 sites, but no NFkB sites,\textsuperscript{113-115} although this assumption is yet to be formally proven (Fig. 1). In the Sertoli cell, $\beta_A$ production is negatively regulated by FSH through the protein kinase A pathway, and is stimulated by LPS.\textsuperscript{30,112} Binding of activin A to target cells causes dimerization of membrane type II and type I signal-transducing receptors and activation of the Smad family of transcription factors.\textsuperscript{116-118}

Activin A bioactivity can be regulated at the transcriptional and translational level, as well as at a post-translational level through the binding of $\beta_A$ subunit to apparently functionally inactive $\beta$ subunits (such as the $\beta_C$ subunit)\textsuperscript{119} or to the $\alpha$ subunit of inhibin.\textsuperscript{106} Activity is also controlled post-secretion by the activin-binding protein, follistatin.\textsuperscript{120,121} Follistatin binds activins, as well as a small number of related TGFβ family members, with very high affinity, competing with their ability to interact with their receptors and essentially neutralizing biological activity.

Under control conditions in the adult testis, immunohistochemical and in situ hybridization studies indicate that the $\beta_A$ subunit of activin is largely associated with the Sertoli cell, but it appears to have a more widespread distribution within germ cells, extending into spermatogonia, spermatocytes and spermatids.\textsuperscript{32,122,123} It is also found in macrophages and mast cells within the interstitial tissue of adult rats.\textsuperscript{32} Peritubular cells from immature rats in culture produce activin A,\textsuperscript{112,124} but the protein was not detectable by immunohistochemistry in peritubular profiles of adult rat testis,\textsuperscript{32} suggesting that activin A production by these cells may decline with age. This possibility is given extra weight by the observation that, in prepubertal rat testes, peritubular cells rather than Sertoli cells appear to be the main source of activin A.\textsuperscript{122} Given the ability for activin subunits to form dimers with differential activity, it is critical to measure the whole protein, rather than just the assess the presence of a particular subunit in isolation. This has been facilitated in the case of activin A by the availability of a specific sandwich-type ELISA that detects the dimeric protein.\textsuperscript{125,126} A combination of whole protein and mRNA studies on activin A/$\beta_A$ subunit expression in several species have established that activin A protein is present at all stages of the seminiferous cycle, but undergoes a distinct cyclical pattern of production (Fig. 2).\textsuperscript{32,123}

Experiments with adult rat seminiferous tubule cultures indicate that IL1\textalpha may be driving activin A production at all stages of the cycle where IL1\textalpha is produced, but that the ability of Sertoli cells to respond to IL1\textalpha changes throughout the cycle.\textsuperscript{32} This may involve changes in
IL1R levels, production of endogenous IL1ra by the Sertoli cells, and/or a cycle-specific switch in the inhibitory regulation of activin A by FSH. Regardless of the fine detail of this control, activin A secretion by the seminiferous epithelium displays a very large surge at stages VIII-XII in the rat, immediately following spermiation and the resumption of IL1α production that follows stage VII (Fig. 2). By the end of stage XII, however, activin A production has returned to basal levels. It is likely that the bioavailability of activin A is also regulated at this time by the production of follistatin, which reaches maximal levels of expression within the cycle at stages X-XIV.

The surge of activin A at stages VIII-XII coincides with and then extends beyond a burst of DNA synthesis associated with meiotic division in preleptotene spermatocytes and a round of type A spermatogonial division. The ability of activin A to stimulate spermatogonial and preleptotene spermatocyte DNA synthesis has been established by several in vitro experiments, using tubule fragment cultures or Sertoli-spermatogonial cell cocultures. However, other studies have shown that activin A inhibits and follistatin stimulates FSH-induced spermatogonial proliferation in testis fragment cultures from younger rats. Moreover, activin A delays meiosis in cultures of rat primary spermatocytes, and in many other cell types activin A has been shown to inhibit cell growth and induce apoptosis. To complicate the matter further, activin A can block production and down-stream effects of IL1 and IL6 during inflammatory responses, and may have similar effects on endogenous IL1α and IL6 within the seminiferous epithelium. At present, it is difficult to predict exactly what role the surge of activin A might play in the control of germ cell mitosis and meiosis during stages VIII-XII, and studies to address this issue are warranted.

As with activin A, inhibin B is produced in a stage regulated manner in the rat seminiferous epithelium (Fig. 2). Further, the production of inhibin B is regulated by the presence of developing germ cells and by FSH and IL1α in a reciprocal pattern with activin A. For example, the incubation of cultured stage VIII tubules with IL1ra resulted in a reduction in activin A secretion and a corresponding increase in inhibin B production. In rat Sertoli cell cultures, IL1α and IL1β stimulated βA mRNA production and activin A protein secretion, while concurrently reducing inhibin B protein secretion and transcription of both βB and inhibin α mRNA. Conversely, FSH inhibits activin A secretion by the Sertoli cell, but stimulates inhibin B secretion. While still only partially revealed, these data highlight the complex and dynamic cytokine and regulatory hormone network that operates within the seminiferous epithelium.

**Nitric Oxide**

The nitric oxide synthases (NOS) are a group of three related enzymes, neuronal NOS (nNOS or NOS1), inducible NOS (iNOS or NOS2) and endothelial NOS (eNOS or NOS3), which catalyze the conversion of L-arginine to L-citrulline and NO. At low levels (<1 μM), NO acts as a regulatory molecule, but at high levels NO causes damage to DNA, proteins and lipids through free radical generation. The NOS enzymes are homodimeric proteins composed of identical monomers of ~130-160 kDa and are encoded by three separate genes. Both nNOS and eNOS are constitutively expressed enzymes whose activity is regulated through a calcium-calmodulin mediated mechanism. In contrast, iNOS is a constitutively activated form of NOS that is regulated at the transcriptional level by a range of inflammatory mediators, including LPS, IL1 and TNFα (Fig. 1). Consequently, iNOS has both physiological and pathophysiological actions.

All three NOS forms have been identified in testicular tissue from several species. They appear to be involved in the regulation of normal male fertility at multiple levels and are pathogenic under some circumstances. At the cellular level, NOS has been found in Sertoli (nNOS, eNOS, iNOS), Leydig (nNOS, eNOS, iNOS) and peritubular cells (iNOS), spermatogenic cells (eNOS, iNOS) and testicular macrophages (iNOS). It is important to note that macrophage expression of iNOS is confined to the minority of monocyte-like macrophages of the rat testis, and it is not expressed by the majority resident macrophages. In the normal rat
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Seminiferous epithelium, iNOS is particularly expressed by elongating spermatids at stage IX of the cycle, and in pachytene spermatocytes at stages IX-XII, with relative lower levels of expression in the Sertoli cells and peritubular cells across all stages.9 The nNOS gene also produces a testis-specific isoform, TnNOS, which has been localized specifically to Leydig cells and is implicated in the control of steroidogenesis through its ability to regulate steroidogenic enzyme levels.144-147 Regardless of its source, NO has been shown to inhibit Leydig cell steroidogenesis directly, and treatment with NOS inhibitors counteracts the decrease in testosterone associated with sepsis or stress.148-151 The mechanism of action probably involves oxidative damage through generation of reactive nitrogen species, such as the peroxynitrite anion.141,142 In addition to the regulation of androgen production and germ cell number, NO production has been implicated in the control of the formation and disassembly of the Sertoli cell junctions that constitute the blood-testis barrier, as well as the junctional complexes involved in Sertoli-germ cell adhesion.152

Surprisingly, male mice with deletions of each NOS form are fertile, although only the fertility of iNOS null mice has been studied in great depth. Potentially, considerable redundancy exists in all of these processes whereby the substitution of another NOS enzymes, as well as reactive species other than NO, may occur in the genetic absence of each individual NOS form. Significantly, iNOS null mice have increased testis weights (131% of control) as a result of decreased pachytene and round spermatid apoptosis, which leads to a 65% increase in daily sperm output.155 The susceptibility of pachytene spermatocytes to iNOS is entirely consistent with the expression of iNOS in normal wildtype rat testes, and is strongly indicative of a key role for iNOS in determining the germ cell carrying capacity of Sertoli cells.9,153

Not surprisingly, given the response of the NOS forms to stress in other tissues, iNOS expression and NO production in the testis is greatly up-regulated during inflammatory events induced by the injection of LPS,9 testicular torsion154 or testicular heating.153 NO over-production leads to stage-specific germ cell damage and loss by both apoptosis and necrosis, as well as changes in testicular blood flow and interstitial fluid content.9,155

Cytokines in the Regulation of Normal Spermatogenesis—What Should We Believe?

On the face of it, the large body of data outlined in the preceding section would seem to indicate that pro-inflammatory mediators must play an important role in the functions of the seminiferous epithelium. However, because these molecules are associated with inflammation and can be artefactually induced in many cell types by inflammation, stress or even the very act of cell isolation, doubts have been expressed regarding their actual contribution to normal testicular function. This attitude is not entirely unjustified. The sensitivity of many cells to stimulation by LPS and other endotoxic agents and the failure of many earlier studies in particular to eliminate the complication of endogenous endotoxin contamination has led some researchers to question the validity of the observations that these mediators are expressed by testicular cells under normal conditions. In other words, it is suspected that the production of inflammatory mediators by testicular cells in the absence of an apparent inflammatory stimulus may be due to stressing of the cells during their isolation, endotoxin contamination of in vitro preparations, or even an underlying pathology in the animals used as a source of tissue. For example, conflicting results concerning IL1α localization in spermatogenic cells may be attributable to such uncontrolled experimental variables.6,61 The problem is compounded by the use of RT-PCR to detect these mediators because of the sensitivity of such methods. Indeed, using RT-PCR alone, one can “demonstrate” the presence or absence of an mRNA species by simply increasing or decreasing the number of cycles, respectively. Fortunately, the development of quantitative RT-PCR methods in recent years has gone a long way towards reducing this problem. Nonetheless, a reliance on mRNA data alone ignores well documented variations in cytokine translational efficiency and the requirement for post-translational processing to produce a bioactive protein (Fig. 3). Finally, a number of immunohistochemical
studies using antibodies of poorly defined specificity, often with inadequate controls or using poorly-fixed testicular tissue, may have contributed to confusion in the literature as well. Skepticism has been compounded by the fact that, with the exception of activin A, which is fetal lethal, breeding studies show that male mice with deletions of IL1, IL6, TNFα and iNOS, or of the relevant receptors, are all fertile. Accordingly, in spite of considerable evidence that IL1α is produced by the Sertoli cell and has regulatory effects on spermatogonial proliferation and development, mice lacking the IL1R, and hence unresponsive to IL1, display relatively normal fertility. Yet it cannot be ignored that numerous studies have shown by quantitative mRNA and protein methods using both in vivo and in vitro approaches that the mediators discussed in this review are produced by testicular cells and that they have effects on spermatogenesis. Moreover, most of the findings have been consistent and reproducible in different research laboratories over many years, and it seems unlikely that endotoxin contamination or sick animals can account for all the observations. In fact, some studies have gone to considerable lengths to completely eliminate these possibilities. While there is still much room for debate concerning the details, there are sufficient data to declare that IL1α, TNFα, IL6, activin A and iNOS/NO have a case to answer with regards to their role in normal testicular function.

As outlined above, the expression of IL1α, TNFα, IL6, activin A and iNOS occurs in a regulated manner across the cycle of the seminiferous epithelium (Fig. 2). These proteins show distinct patterns of cyclical production and their patterns of production coincide with key events in Sertoli cells and spermatogenic cells. A critical series of events occurs at, and immediately after, stage VIII in the rat: the release of sperm from the epithelium (spermiation), a peak

Figure 3. Measurement of IL1β. This cytokine displays a particularly complex regulation, involving a biologically inactive precursor which is activated by the enzyme IL1 cleaving enzyme, ICE or caspase 1, at the time of its secretion from the cell. The mechanisms of production, processing and secretion are poorly understood, but appear to be relatively inefficient, and there can be large discrepancies between the levels of mRNA expression and the amount of bioactive protein secreted by the cell. Moreover, significant secretion of the inactive precursor without processing may occur, as has been observed in the testis and other systems. As a result, what is measured can make very big differences to the conclusions reached. Even quantitative methods which detect mRNA may tell us relatively little about the actual levels of protein produced. Methods which detect the protein may also detect the precursor, leading to an over-estimation of the bioactive protein. Bioassays provide the most informative measure of the physiological levels of IL1, although even this measurement may be compromised by the presence of anti-inflammatory cytokines that inhibit its activity in various assays.
of DNA synthesis by preleptotene primary spermatocytes cells and type A spermatogonia prior to meiotic and mitotic division, respectively, and the reorganization of tight junctions of the blood-testis barrier to allow the meiotic cells to enter the adluminal compartment. These events also coincide with a recovery of the responsiveness of the Sertoli cell to FSH. At the same time, there is a resumption in production of IL1 by the Sertoli cells and of TNFα and iNOS by the germ cells, a rise in Sertoli cell IL6 production, and a transient peak of activin A production. These responses are highly reminiscent of an inflammatory event. Could the epithelium be experiencing a cyclical inflammatory burst, associated with sperm release and phagocytosis of the residual bodies?

Significantly, nuclear localization of the key pro-inflammatory transcription factor, NFκB in the Sertoli cell and the germ cells also shows a cyclical pattern within the seminiferous epithelium. There is a good relationship between nuclear NFκB levels in spermatocytes and both TNFα and iNOS expression by these cells. Curiously, however, there does not appear to be a close concordance between the content of NFκB in the Sertoli cell nucleus, which is elevated at most stages of the cycle but appears to decline during stages VIII and IX, and the production of IL1α or IL6 by this cell. This is, perhaps, less surprising in the case of activin A and inhibin B, since NFκB almost certainly does not regulate their expression. However, it should be borne in mind that the overlying regulatory influence of the developing germ cells, as well as FSH and testosterone in the control of several of these mediators may distort the cyclical expression patterns observed. For example, production of IL6 by the Sertoli cell is stimulated by FSH and, although it is regulated by IL1α and nuclear NFκB, its production appears to be most closely aligned with changes in the expression of the FSH receptor (Fig. 2).

More detailed studies on the regulation of these and other transcriptional regulators and signaling pathways involved in inflammatory cytokine production throughout the cycle of the seminiferous epithelium will be required to resolve these issues.

Additional questions remain regarding the precise biological effects exerted by some of these mediators within the seminiferous epithelium. It appears that different in vitro models can lead to very different conclusions. This is best exemplified by the fact that TNFα reduces germ cell apoptosis in seminiferous tubule cultures and, hence, could be considered a germ cell survival factor. However, TNFα is also implicated in the breakdown of Sertoli cell tight junctions, stimulation of apoptosis by Fas-Fas ligand activity in spermatogenic cells, and the onset of orchitis. It would appear that the role of TNFα in regulation of spermatogenesis may depend upon its context, i.e., when and for how long it is produced, how much of it is produced and which receptors are available to respond to it. In a similar manner, activin A has been identified as both a stimulator and inhibitor of spermatogonial development by different experiments.

Nonetheless, even though there are many questions still to be answered, it is possible to propose a model of cytokine network signaling in the Sertoli cell based on the available data (Fig. 4). It can be postulated that release of sperm and resorption of residual bodies at stage VIII-IX of the cycle triggers an inflammatory response in the Sertoli cell as evidenced by an up-regulation of IL1α. IL1α in turn stimulates a surge of spermatogonial proliferation and a new generation of germ cells entering the spermatogenic cycle. IL1α also induces the production of IL6, which acts to regulate the number of spermatocytes progressing through meiosis. The role of activin A in this model is not entirely clear, but this cytokine may cooperate with IL1α to stimulate spermatogonial proliferation, or with IL6 to control meiotic progression, or it may even act to block the activity of IL1α and IL6. As a result of this network, the entry of spermatogonia into spermatogenesis occurs in short bursts that are timed to coincide with the release of sperm into the lumen, while at the same time the entry into meiosis is being modulated. A parallel network involving the spermatocytes and spermatids is also triggered at this time, possibly involving IL1α from the Sertoli cells, stimulating the production of TNFα and iNOS/NO, which induce the disassembly of the intercellular tight junctions to allow the meiotic cells to transit the blood-testis barrier. Variants of these regulatory networks may also...
operate at other times and in different ways throughout the remainder of the cycle, for example, in controlling the proliferation of intermediate and B type spermatogonia during stage V.4,55

Given the considerable evidence for a direct role for cytokine networks in the control of spermatogenesis under normal conditions, therefore, how can we explain the fact that the relevant 'knockout' mice are fertile? Three main issues need to be considered here:

Redundancy of Action

Most cytokines and enzymes involved in inflammatory pathways share overlapping functions and signaling pathways, as in the case of IL1 and TNFα (Fig. 1), or belong to families with similar properties and range of actions (e.g., the NOS forms). Absence of a critical gene, particularly during development, may even stimulate appropriation of other genes with similar function. It is common knowledge that deletions of even fundamental regulators like IL1, TNFα, or IL6 results in relatively subtle immunity phenotypes.165 Rather than focusing on the individual players, the inflammatory response itself should be considered as being responsible for spermatogenic regulation. Consequently, the retention of fertility in mice lacking the IL1R, and presumably insensitive to either IL1α or IL1β,157 may be due to the fact that IL1 is just one of several inflammatory mediators with overlapping functions induced by phagocytosis in the Sertoli cells. Moreover, the possibility for alternative or non-classical actions of IL1α in the seminiferous epithelium, which bypass the classical ILR must also be considered.166,167

Superficial Assessments of Fertility

Few of the knockouts relevant to the present discussion have been carefully examined for subtle testicular phenotypes, which may occur: the iNOS null mouse is a notable exception.153 Null mice may be considered to be fertile, because they are capable of breeding in an animal house environment, but further examination may expose testicular or germ cell anomalies.168 The genetic background of the animal also should be considered, since phenotypes observed in one strain of knockout mice can disappear when backcrossed onto another background.
“Hidden” Phenotypes

Fertility or other physiological phenotypes may not manifest under normal animal house conditions, or in the absence of some stressor or other intervention, such as ageing or infection. A good example of this is the IL6 null mouse,\textsuperscript{159} which has apparently normal testicular function, but actually possesses an aromatase deficiency that can lead to anomalous steroidogenic responses (Fig. 5).

In summary, we have outlined a complex model of spermatogenic regulation involving inflammatory networks that leads to the hypothesis that the cycle of the seminiferous epithelium is actually a cyclical inflammatory event. As such, spermatogenesis may be viewed as a localized inflammatory process wherein the Sertoli cell functions in the role of the monocyte/macrophage. With this concept in mind, it is particularly important that the testis maintains a tight immunoregulatory environment to suppress local antigen-specific immune responses. Although outside the scope of this review, it is clear that the testicular resident macrophages, in concert with the Sertoli and Leydig cells, conspire to inhibit immune responses, which might otherwise be triggered by the ongoing inflammatory process within the seminiferous tubules (reviewed in detail by Hedger and Hales, ref. 169).

Inflammation and Testis Function—Role of Inflammatory Mediators and Implications for Fertility

While the role of inflammatory mediators in normal spermatogenesis may be disputed, there is no doubt that they play a critical role in the suppression of spermatogenesis by inflammation. Normal testicular function can be directly inhibited by local or systemic illness, infection and chronic inflammatory disease in men,\textsuperscript{170-172} and similar decreases in gonadal function occur in experimental animal models of chronic inflammation and systemic immune activation.\textsuperscript{149,173-178} In spite of a common assumption that such reproductive failure is related to the negative effects of a raised body temperature on spermatogenesis, there is very little evidence to support this contention in febrile

![Diagram](image)

Figure 5. The “hidden” phenotype of the IL6 null mouse model. Infection of mice with Taenia crassiceps leads to a testicular feminization syndrome, as a result of up-regulation of aromatase activity. While testicular function and fertility in IL6 null mice is virtually indistinguishable from that of the wild-type mouse, these mice show no feminization following infection directly implicating IL6 in the control of aromatase activity in the testis of both normal and infected animals. Thus, a role for IL6 in testicular function and pathology is only evident if the appropriate experiment is carried out. This model provides an excellent example of a significant phenotype that appears only under conditions of modified function.
patients, or in experimental animals. In fact, it is much more likely that specific inflammatory events are directly involved. As a logical extension of the key role of inflammatory mediators in normal male spermatogenesis and steroidogenesis, it is not surprising that inflammatory events, either within the testis itself or in the rest of the body, compromise testis function. Indeed, based on the normal expression pattern of such mediators, one would expect certain aspects of spermatogenesis to be particularly sensitive to shifts in their expression.

Another general perception is that reduced fertility following an illness is due to decreased androgen production. Serum androgen levels are generally decreased with illness, and certainly contribute to a lack of libido and well-being. The experimental models that have helped to elucidate this aspect of immune-testis interactions have involved administration of the inflammatory mediator LPS or studies on the effects of agents and treatments that alter the vasculature of the testis. These models involve an increase in both systemic levels and testicular expression of inflammatory mediators in addition to effects on androgens and spermatogenesis. There is no doubt that IL1, TNFα and NO, in particular, have largely negative effects on Leydig cell steroidogenesis, and that this inhibition involves both central and direct effects on the Leydig cell itself. Consequently, inflammation leads to suppression of the hypothalamic-pituitary-Leydig cell axis, and a corresponding decline in androgen levels, and all the subsequent clinical implications. However, it is not certain that this is the main cause of damage to spermatogenesis during inflammation. For one thing, there are distinct species differences in the dynamics and severity of the inhibition of the endocrine axis in the LPS-induced inflammation models. While the mouse displays a rapid onset and prolonged suppression of steroidogenesis after treatment with LPS, well above the threshold necessary to sustain spermatogenesis in this species. Moreover, serum testosterone levels actually appear to rise in the boar following LPS treatment due to an increase in pulsatile LH secretion.

In contrast to the many studies on endocrine parameters, there have been few detailed studies of the direct effects of inflammation on spermatogenesis itself. Our own studies, using an acute LPS induced systemic inflammation model in the adult rat have shown a selective up-regulation of pro-inflammatory molecule production in the testis, including IL1\(\beta\), TNFα, IL6 and iNOS/NO, but not IL1\(\alpha\) or activin A, significant recruitment of circulating monocytes from the systemic circulation, but no change in resident macrophage numbers, and a complex inhibition of pituitary secretion of LH, glucocorticoids and direct effects on the Leydig cell. Within 24 hours of LPS administration, a maturational delay became evident in the leptotene/zygotene spermatocytes (stages IX-XIII). This was followed within six days by an increase in the sloughing of these cells and the associated (more luminal) round spermatids (at stages I-VIII) and an associated increase in apoptosis of all spermatocytes at stages IX-XIII. Significantly, the severity of the illness also had effects on the observations. In the earlier study, where illness among the animals appeared to be more severe, there was a more pronounced spermatocyte and spermatid loss, spermatogonial apoptosis at stages I-V and changes in vascular permeability, including micro-hemorrhage. In fact, vascular changes could account for the increased spermatogonial apoptosis at stages I-V in this group, since these cells are particularly sensitive to an interruption in testicular blood flow. On the other hand, at lower doses of LPS, even those that caused maximal inhibition of intratesticular testosterone, little or no germ cell loss was observed. Remarkably, even following a high dose of LPS, spermatogenesis had returned to qualitatively normal levels by 28 days post-injection. This recovery suggests that a compensatory reduction in normal germ cell attrition and possibly an increase in proliferation may occur following an acute inflammatory event, restoring the spermatogenic capacity of the testis to its pre-inflammation levels. It should be noted, however, that this model involves an acute inflammatory episode only. Very little quantitative data exists concerning the effects of on-going systemic or testicular inflammation similar to that occurring in a significant proportion of the human male population suffering chronic inflammatory conditions such as arthritis.
Altogether, the data point towards a direct effect of inflammation on the seminiferous epithelium, and that the effects of acute inflammation are most pronounced and earliest on the seminiferous epithelium at stages IX-XIII, i.e., immediately following spermiation. Importantly, the effects of LPS-induced inflammation on spermatogenesis in the rat were not consistent with withdrawal of either testosterone, which initially affects the release of mature spermatids into the tubule lumen and the integrity of the junctions between the Sertoli cells and mid-phase round spermatids at stages VII-VIII, or of FSH, which particularly affects spermatogonia at stages XIV-III.\(^{189,190}\) While there is no doubt that loss of hormonal support may contribute to spermatogenic damage in some models, the observations suggest that alternative explanations must be sought to explain spermatogenic failure in the acute inflammatory model. A potential cause of damage in this model is the action of LPS and pro-inflammatory mediators on the seminiferous epithelium itself. This could involve increased levels of circulating mediators as well as local production by leukocytes and somatic cells in the testis. In addition to systemic increase in these mediators, LPS causes the up-regulation of testicular IL1\(\beta\), TNF\(\alpha\), IL6 and iNOS expression.\(^{9,15,50,86,184}\) Although the absolute levels of expression of IL1\(\alpha\) and activin A do not appear to change, their distribution and timing may be affected. As discussed in the previous section, these inflammatory mediators are produced in a regulated fashion during the normal cycle of the seminiferous epithelium and have direct and complex effects on both Sertoli cell and spermatogenic cell function. Changes in IL1, activin A and IL6 would disrupt the signaling processes involved in controlling meiotic progression and spermatogonial proliferation. Similarly, local increases in TNF\(\alpha\) and NO would disrupt Sertoli tight junction integrity and germ cell attachment. All of these factors, as well as direct effects of LPS on the Sertoli cell itself may affect critical supportive functions of the Sertoli cells. Consequently, it is particularly significant that the region of the seminiferous epithelium that is first affected by LPS induced inflammation is the stages immediately following spermiation (IX-XIII), where these cytokine networks appear to play their most complex roles (Fig. 2). Overall, these data suggest that inflammation directly affects spermatogenesis through interfering with these critical inflammatory networks of the seminiferous epithelium.

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Inflammatory Networks in the Control of Spermatogenesis

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Transcription Regulation in Spermatogenesis

Wing-Yee Lui* and C. Yan Cheng

Introduction

Spermatogenesis is a highly coordinated process in which diploid spermatogonia (2n) differentiate into mature haploid (1n) spermatozoa in the seminiferous epithelium. In this process, spermatogonia undergo several mitotic divisions and either enter a stem cell renewal pathway, or commit themselves for further development. Diploid spermatocytes subsequently undergo two meiotic divisions and result in the production of haploid round spermatids. They then enter the process of spermiogenesis in which profound morphological and biochemical restructuring, such as the formation of acrosome and flagellum occur, and give rise to mature spermatozoa.

The cyclic and synchronous nature of spermatogenesis leads to specific pattern of cellular associations at a given segment in the tubules in which germ cells at particular stages of differentiation will associate with one another. Such cellular associations have been classified into the stages of the seminiferous epithelium. There are twelve (stages I-XII) and fourteen stages (stages I-XIV) of the seminiferous epithelium in mouse and rat, respectively1,2 according to their cellular associations. Such differentiation pattern apparently requires precise regulation of specific genes at a given stage. In order to have a better understanding how transcription factors exert their regulatory function to modulate cellular and stage-specific gene expression during spermatogenesis, we summarize herein some of the recent findings in the study of transcription regulation during spermatogenesis into five categories: (i) general transcription factors, (ii) nuclear receptor superfamily of transcription factors, (iii) other transcription factors involved in testicular functions, (iv) testis-specific gene transcription, and (v) transcriptional regulation of cell junction dynamics. The chapter is not intended to be exhaustive, rather, it serves as a guide for future studies based on latest findings in the field.

Transcription Regulation in Spermatogenesis

General Transcription Factors

Regulation of stringent stage-specific gene expression in testicular cells and the massive wave of transcriptional activity in germ cells following meiosis are governed by a highly specialized transcriptional mechanism.3 Such temporal and restricted pattern of gene transcription is achieved by the presence of germ cell-specific transcription factors (Table 1). In addition, various general transcription factors, in term of their expression levels and their testis-specific isoforms, are differentially regulated in germ cells and in testes. It is believed that the differential expression of general transcription factors also play a crucial role to ensure proper and efficient transcription in germ cells throughout spermatogenesis.4 For instance, TFIIB (a transcription factor that serves as a positioning factor for polymerase), TATA-binding protein (TBP)

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and RNA polymerase II were found to be accumulated in early haploid germ cells. Their levels in haploid germ cells are much higher than in somatic cells. Adult rodent testes contain 80-200 molecules of TBP mRNA per haploid genome-equivalent, whereas adult spleen and liver contains 0.7 and 2.3 molecules of TBP mRNA per haploid genome-equivalent, respectively. Such organization of transcription factors enable early spermatids accumulate enough mRNA for their development until the final stages of spermiogenesis.

Table 1. List of the genes encoding the transcription factors whose deletion in the mice generate defects in males

<table>
<thead>
<tr>
<th>Gene Disrupted</th>
<th>Male Phenotype</th>
<th>Female Phenotype</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>AR</td>
<td>Complete arrest at pachytene spermatocyte stage Fertile Yeh, 2002^23</td>
<td>Feminine-like appearance</td>
<td></td>
</tr>
<tr>
<td>RARα</td>
<td>Complete arrest and severe degeneration of the seminiferous epithelium Fertile Lufkin, 1993^34</td>
<td>Small testis with a decrease in serum testosterone concentration</td>
<td></td>
</tr>
<tr>
<td>RXRβ</td>
<td>Partial arrest at primary spermatocyte stage Fertile Kastner, 1996^35</td>
<td>Partial arrest at primary spermatocyte stage Structural abnormalities in spermatozoa</td>
<td></td>
</tr>
<tr>
<td>GCNF</td>
<td>Embryonic lethality Embryonic lethality Chung, 2001^156</td>
<td>Embryonic lethality</td>
<td></td>
</tr>
<tr>
<td>TR2</td>
<td>Functional testis having normal sperm number and motility Fertile Shyr, 2002^73</td>
<td>Functional testis having normal sperm number and motility</td>
<td></td>
</tr>
<tr>
<td>TR4</td>
<td>Delay in the first wave of spermatogenesis Prolonged stages XI to XII of spermatogenesis Reduced fertility Mu, 2004^72</td>
<td>Delay in the first wave of spermatogenesis Prolonged stages XI to XII of spermatogenesis Reduced fertility</td>
<td></td>
</tr>
<tr>
<td>CREM</td>
<td>Complete arrest at pachytene spermatocyte stage Fertile Nantel, 1996; Blendy, 1996^157</td>
<td>Complete arrest at pachytene spermatocyte stage Fertile Nantel, 1996; Blendy, 1996^157</td>
<td></td>
</tr>
<tr>
<td>CREB (α and β isoforms)</td>
<td>Fertile</td>
<td>Fertile</td>
<td>Hummler, 1994^158</td>
</tr>
<tr>
<td>CREB (α, β and δ)</td>
<td>Die shortly after birth Die shortly after birth Rudolph, 1998^159</td>
<td>Die shortly after birth</td>
<td></td>
</tr>
<tr>
<td>Rhox5</td>
<td>Subfertile Increased frequency of apoptotic meiotic spermatocytes Fertile Pitman, 1998;^95 MacLean, 2005^26</td>
<td>Subfertile</td>
<td></td>
</tr>
<tr>
<td>Sperm-1</td>
<td>Subfertile Fertile Pearse, 1997^97</td>
<td>Exhibit progressive loss of spermatogonia and increase in apoptosis with age Costoya, 2004^102</td>
<td></td>
</tr>
<tr>
<td>Plzf</td>
<td>Embryonic lethality Embryonic lethality Narita, 1997^161</td>
<td>Embryonic lethality Embryonic lethality</td>
<td></td>
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<tr>
<td>WT1</td>
<td>Conditional knockout mice show impaired spermatogenesis and predicted to be fertile Gao, 2006^106</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>GATA-1</td>
<td>Embryonic lethality Embryonic lethality Pevny, 1991^160</td>
<td>Embryonic lethality Embryonic lethality</td>
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<tr>
<td>GATA-4</td>
<td>Embryonic lethality Embryonic lethality Narita, 1997^161</td>
<td>Embryonic lethality Embryonic lethality</td>
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<tr>
<td>GATA-6</td>
<td>Embryonic lethality Embryonic lethality Koutsourakis, 1999^162</td>
<td>Embryonic lethality Embryonic lethality</td>
<td></td>
</tr>
<tr>
<td>MSY2</td>
<td>Infertile Infertile Yang, 2005^139</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>CAF1</td>
<td>Infertile Infertile Nakamura, 2004^143; Berthet, 2004^144</td>
<td>Infertile</td>
<td></td>
</tr>
</tbody>
</table>

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In addition to the unique expression pattern of various general transcription factors in germ cells, the presence of their testis-specific isoforms may play a specialized function in spermatogenesis. ALF or TFIIA-α is a testis-specific isoform of TFIIA which may have specificity for a subset of transcriptional activators.6,7

**Nuclear Receptor Superfamily**

**Lipophilic Hormone Nuclear Receptors**

**Androgen Receptor**

Androgens are crucial steroid hormones in male reproduction and their actions ranging from regulating sexual differentiation, sexual maturation, spermatogenesis to production of gonadotropins.8-12 Androgens exert their effects through the androgen receptor (AR). AR is a ligand-inducible transcription factor (110 kDa) that regulates the expression of target genes in response to its cognate ligand (androgen) through binding to an androgen response element (ARE).12,13

Similar to other members of the nuclear receptor superfamily, AR can be divided into four functional domains. They are: NH2-terminal transactivation domain, DNA-binding domain (DBD), hinge region, and ligand-binding domain (LBD). AR has two separate NH2-terminal transactivation domains which possibly interact with different coregulators or transcription factors in a promoter content-dependent manner.14 The DBD contains two zinc fingers that recognize specific DNA consensus sequences. AR homodimer binds to the inverted repeat ARE, GGTACAnnnTGTTCT.15-18 Apart from the formation of homodimer, it was reported that AR is capable of forming the heterodimers with TR4 (human testicular receptor 4, TR4, is an orphan member of the nuclear receptor superfamily) or ERα (estrogen receptor α), which results in a decrease in AR transcriptional activity.19,20 The LBD is responsible for the formation of the ligand-binding pocket, facilitating the interaction between AR and heat shock protein, and also interacting with AR NH2 terminus to stabilize the bound androgen.21,22

AR is of particular interest because of the observation that knockout of AR produced male mice displaying female-like appearance with arrested spermatogenesis.23 Although AR plays an indispensable role in spermatogenesis, only a few number of genes have been identified so far that are directly regulated by AR in the testis. The expression of X-linked Rhox5/PEM homeobox gene is a typical example of AR-mediated gene regulation in the testis.24-26 Barbulescu et al. have identified two functional AREs within 300-bp upstream of the Rhox5 transcription start site.27,28 The promoter region containing the regulatory sequences that directs AR-dependent expression specifically in Sertoli cells and confers AR stage-specific expression in adult testis.29,30 Recent studies from MacLean et al. have shown that another four Rhox genes (namely Rhox2, 3, 10 and 11) are dramatically upregulated in response to incubation with testosterone and cotransfection with an AR expression plasmid. Although the promoter sequences of the four Rhox genes have not yet been characterized, it is apparent that they all are androgen-dependent.26

Apart from AR knockout mice, a tissue-specific knockout mouse with the AR gene deleted in Sertoli cells [SC AR knockout mice] was generated to investigate how androgen/AR in Sertoli cells influence spermatogenesis.31,32 It was found that the SC AR knockout male exhibit similar phenotypes as that of AR-/- mice with more severe testis atrophy. SC AR knockout mice showed alterations in the expression of anti-Mullerian hormone (AMH), cyclin A1, Pem and sperm-1.31 The increase in the expression of AMH in mice leads to the reduction of testosterone production in Leydig cells. Significant reduction in germ cell number in SC AR knockout is associated with increased germ cell apoptosis and reduced expression of cyclin A1, Pem and sperm-1 genes that are important for late stage of germ cell development.31,32 Sertoli cell-specific AR knockout mice clearly demonstrated the functional significance of AR in Sertoli cells in maintaining spermatogenesis and steroidogenesis.

Using another SC AR knock-out model,32 it was shown that the loss of androgen receptors in Sertoli cells led to a disruption of the blood-testis barrier (BTB) integrity since biotin could
diffuse through the BTB. Using techniques of gene profiling, it was shown that the gene responsible for the "leaky" BTB in SC AR knock-out mice is likely to be claudin 3, which displays transient expression in newly formed tight junctions. However, it is noted that the SC AR knock-out mice used in this study were made with a floxed exon 1, yet the floxed animals had already displayed marked hypomorphic phenotype and the ultimate AR knockout was neither complete nor Sertoli cell selective, which may explain why did the SC AR knock-out mice had a serum testosterone level almost 40-fold of that of the wild type. Furthermore, the testicular claudin 3 level in adult rat testes was extremely low, and it is virtually undetectable beyond 45 days of age (Yan and Cheng, unpublished observations), making claudin 3 hardly an important structural component of the BTB in adult rats. Nonetheless, it is likely that testosterone and its receptor are important components that regulate BTB dynamics, much work is needed in the field to define the precise molecular target(s) of testosterone and AR at the BTB.

Retinoic Receptors

Retinoic acid receptors (RARs) and retinoid X receptors (RXRs) are two members of this family found in the testis. Ligand-dependent activation of RAR and RXR are essential to spermatogenesis based on the fact that infertility was observed in vitamin A-deficient rats and in RARα and RXRβ transgenic mice. In vitro binding studies have demonstrated that the natural metabolites all-trans-RA and 9-cis-RA are high-affinity ligands for RARs, whereas only 9-cis-RA has been shown to bind RXRs. Each family consists of three genes, namely α, β, and γ, and each of them exists as multiple isoforms. RXR is capable of forming homodimers (RXR/RXR), heterodimers with RAR (RXR/RAR) and with other types of nuclear receptors such as thyroid hormone receptor (RXR/TR), such characteristic enables this receptor family exerts combinatorial regulatory properties.

The homodimer and heterodimer function as RA-inducible transcriptional regulatory proteins through binding to DNA sequences called retinoic acid response element (RARE) or retinoid X response element (RXRE) located within the promoter of target genes. The consensus sequence of RARE is AGGTCAnnnnAGGTCA, whilst RXRE is direct repeats of AGGTCA with one nucleotide spacing (AGGTCAnAGGTCA). The RAR/RXR heterodimer binds to the RARE, with RXR occupying the 5' upstream half-site and RAR occupying the 3' downstream half-site.

Extensive studies using RARα and RXRβ transgenic mice have clearly demonstrated that retinoic acid-mediated gene regulation via RAR, and RXR play a crucial role in spermatogenesis. For instance, detailed morphological analysis in RARα knockout mouse testes showed that the typical characteristic of stage VIII tubule, where mature step 16 spermatids aligning along the tubular lumen, was not observed. Instead, a mixed population of germ cells was found in stage VIII tubule in RARα knockout male. For RXRβ knockout mice, failure of spermatid release occurred within the germinai epithelium and the epididymis contained very few spermatozoa. Although knockout of RARα and RXRβ resulted in male infertility, they displayed different seminiferous tubule morphology. These observations suggest that the downstream targets of RARα and RXRβ are not the same. Genes expressed in different testicular cells, namely Stra8 and bone morphogenetic protein 4 (BMP4) in germ cells, prostaglandin D2 synthetase in Sertoli cells, and fibronectin and laminin in myoid cells, were shown to be regulated by retinoic acid or retinol. However, the precise mechanisms of vitamin A-mediated gene regulations have yet to be elucidated. Whether the regulation of those genes are direct effects mediated through the interaction of retinoid receptors and their corresponding promoters, or whether other retinoid-regulated proteins mediate indirect regulatory effects remain to be determined. Identification of the regulatory mechanism on RA-RAR-mediated BMP4 expression in other cell lines has provided a blueprint to study the transcription regulation of BMP4 gene in germ cells.
Orphan Receptors

Germ Cell Nuclear Factor

Germ cell nuclear factor (GCNF), which is also known as retinoid receptor-related testis-associated receptor (RTR), is a novel member of the nuclear receptor superfamily of ligand-activated transcription factors. Since the natural ligand for GCNF has not been identified, GCNF is classified as an orphan receptor. GCNF binds as a homodimer either to direct repeat response elements (AGGTCA) without additional nucleotide or to extended half-site such as TCAAGGTCA (XRE). It does not form heterodimer with other nuclear receptors such as RXR. In vitro studies have revealed that GCNF is a sequence-specific repressor of transcription and it folds into a β-sheet that contributes to dimerization and the recruitment of corepressors. It can interact with other nuclear corepressors and with the repressor, RAP80, that is highly expressed in the testis. GCNF expression is restricted to the developing nervous system during embryogenesis, whereas the receptor is expressed during specific stages in maturing germ cells. Two transcripts of GCNF gene with sizes of 7.4 kb and 2.3 kb have been identified in spermatogenic cells. The 7.4 kb transcript is expressed during testicular development and is the predominant form in pachytene spermatocytes, whereas the 2.3 kb transcript is expressed predominantly in round spermatids. In situ hybridization studies have shown that the GCNF transcript levels remain low during the meiotic prophase in rats and mice, and increase substantially and reach maximal level in round spermatids at stages VI-VIII.

Up to now, several genes expressed in the testis were found to be regulated by GCNF. The temporal expression of protamine genes, prm-1 and prm-2, at stage I round spermatid is regulated reciprocally by GCNF and cAMP-response element modulator, CREM. Binding of GCNF to GCNF response elements of prm-1 and prm-2 promoters represses both basal and CREM-activated transcription, thus GCNF may play a role to shut down protamine gene expression in elongating spermatids. Apart from prm-1 and prm-2 genes, mitochondrial glycerol-3-phosphate dehydrogenase (mGPDH) and endozepine-like peptide (ELP) are two other testis-specific genes that are regulated by GCNF. Both promoters of mGPDH and ELP genes contain CRE/GCNF elements that can effectively bind to GCNF. The binding of GCNF to these motifs can interfere with CREM- transactivation. Apparently, GCNF is a crucial transcription regulator that regulates the temporal and spatial expression of several testicular genes during meiosis and the early haploid phase of spermatogenesis.

Testicular Orphan Receptors 2 and 4

Testicular orphan receptor 2 (TR2) and testicular orphan receptor 4 (TR4) constitute a subfamily of nuclear receptors. The TR2 and TR4 can modulate its target gene expression by forming homodimers and binding to the AGGTCA direct repeat (DR) sequences in its target genes. TR4 can modulate transactivation mediated by other steroid nuclear receptors through interaction with these steroid receptors. For instance, TR4 could interact with the androgen receptor (AR) and the estrogen receptor (ER) that suppress AR- and ER-mediated transactivation.

TR2 and TR4 have been shown to be expressed in mouse testes. TR2 is confined to meiotic and postmeiotic germ cells, whereas TR4 is predominantly expressed in primary spermatocytes, especially in late-stage pachytene spermatocytes. The expression of TR4 in round spermatid is stage-dependent and is confined to stage VII. Although the male knockout mice of TR2 and TR4 are fertile, the disruption of TR4 gene does affect spermatogenesis at the end of late meiotic prophase and subsequent meiotic divisions, thus delays the first wave of spermatogenesis in the TR4-/- mice. Gene disruption analyses indicated that TR4, but not TR2, is essential for normal spermatogenesis in mice.

Recent studies has demonstrated that TR4 can suppress the expression of 25-hydroxyvitamin D3 24-hydroxylase, Cyp24a1, through direct binding of TR4 to the vitamin D3 receptor response element (VDRE) in Chinese hamster ovary (CHO) cells. The VDRE shares similarity
with the hormone response element for the TR4, which contains two repeated half sites of AGGTCA; however, it is separated by a 3-nucleotide space. Using the TR4\(^-/-\) knockout mice model, Mu et al. showed that the expression level of Cyp24a1 increased in adult mice testes when TR4 gene was knocked out. Such observation indicates that testicular Cyp24a1 expression is also under the precise control of TR4.\(^{72}\) Cyp24a1 is the only gene identified so far that is regulated by TR in the testis, identification of the molecular targets, such as putative ligands of TR2 and TR4, and the mechanisms that affect meiosis may help in a better understanding of the role of TR in spermatogenesis.

**Transcription Factors Involved in Testicular Functions**

**Basic-Domain-Leucine-Zipper (b-zip) Family**

Members of the b-zip family that are known to be expressed in the testis include cAMP response element modulator (CREM), cAMP response element binding protein (CREB), and activating transcription factor 1 (ATF1).\(^{74,75}\) These proteins contain a basic DNA-binding domain with an adjacent leucine zipper that is required for dimerization and binding to a specific cis-acting element.\(^{76,77}\) CREM, CREB and ATF-1 are capable of forming homodimers and heterodimers in response to cAMP signaling pathway and bind to a regulatory DNA sequence, known as cAMP responsive element (CRE). A CRE is constituted by the palindromic consensus sequence, TGACGTCA.\(^{76-78}\)

**CREM**

Many isoforms of CREM are generated by alternative splicing. Among them, CREM\(_t\) is the isoform which has been extensively studied as its expression is restricted to the testis and is highly regulated during spermatogenesis.\(^{79}\) CREM\(_t\) mRNA transcript is found at high levels in pachytene spermatocytes and more advanced germ cells, while its protein is present only in post-meiotic spermatids, suggesting that CREM\(_t\) plays a role in late stages of spermatogenesis.\(^{80}\) The importance of CREM\(_t\) in spermatogenesis could be reflected in the gene knockout studies since spermatogenic arrest was observed at pachytene spermatocyte stage.\(^{81}\) A list of postmeiotic genes encoding structural proteins required for spermatid differentiation, including the transition proteins (TP1 and TP2), protamines (prm1 and prm2), RT7, testis angiotensin-converting enzyme (ACE), proacrosin and calspermin, were found to be the direct targets of CREM\(_o\).\(^{82-85}\) All of these genes contain the putative CREs for the binding of CREM\(_o\). It is apparent that CREM\(_t\) is a key transcription factor that controls postmeiotic germ cell differentiation.

Different from other CREM isoforms, the activation of CREM\(_t\) requires the association of a coactivator known as activator of CREM in testes (ACT).\(^{86}\) ACT is exclusively expressed in testes.\(^{86}\) ACT shows similar developmental expression pattern as CREM\(_t\) in testes and they are colocalized in spermatids.\(^{86-88}\) ACT displays intrinsic transactivation potential capable of converting CREM\(_t\) into a potent transcriptional activator, leading to the activation of CREM\(_t\) in a phosphorylation-independent manner.\(^{87,88}\) The presence of ACT in post-meiotic germ cells enables stage-specific activation of CREM\(_t\)-mediated gene transcription.

To elucidate the significance of ACT in CREM\(_t\)-mediated gene transcription in testes, gene targeting disruption in mice has been performed. It is surprising that male mice lacking ACT are fertile, which is different from the CREM knockout counterpart. Mice lacking ACT show some male reproductive defects including abnormalities in sperm heads and tails and reduced sperm motility.\(^{89}\) However, the expressions of CREM\(_t\)-dependent genes, such as TP1 and prm1, were not affected in ACT knockout mice. These results seemingly suggest that other yet-to-be identified coactivators exist in testes could compensate for the loss of ACT to modulate CREM\(_t\)-dependent gene transcription.

**CREB**

Similar to CREM, many CREB isoforms are generated by alternative splicing in the testis. Although the gene knockout analyses of CREB isoforms have been performed, the role of CREB
Transcription Regulation in Spermatogenesis

in spermatogenesis has not been fully elucidated. Since mice carrying mutations in all CREB isoforms exhibited severe developmental disorders and died shortly after birth. In situ hybridization analysis has shown that CREB mRNA is present in Sertoli cells in stages I-VIII tubules and the amount decreases to an undetectable level at stages IX-XIV. The cellular localization of CREB in the testis is quite different from CREM, whose protein is present only in post-meiotic spermatids. Several genes involved in spermatogenesis such as murine spermatogenesis-associated protein, claudin-II and nectin-2 have been found to be regulated directly by CREB via the CRE motif in the corresponding promoters. Interaction of CREB with other transcription factors, such as c-Jun, was found to be involved in regulating the nectin-2 gene transcription in Sertoli cells. In addition, overexpression of dominant-negative CREB in primary Sertoli cells could completely inhibit the FSH-induced c-Fos expression. Taken collectively, these data illustrate that CREB seems to play an intriguing role in regulating gene transcription in Sertoli cells.

Homeobox Family

Transcription factors belonging to this family contain the homeobox motif that is a highly conserved DNA-binding domain constituted by 61 amino acids. Transcription factors belonging to this family are grouped in subfamilies based on the homeodomain sequence as well as the gene structure. Our chapter does not attempt to cover all members of this family but highlights two subfamilies that show intimate relationships with spermatogenesis. They are the reproductive homeobox X-linked (Rhox) gene cluster and the POU-domain gene family.

Reproductive Homeobox X-Linked (Rhox) Gene Cluster

Rhox gene cluster presents a newly homeobox subfamily that contains 12 related homeobox genes. All 12 Rhox genes are organized into three subclusters, namely α (Rhox 1-4), β (Rhox 5-9) and γ (Rhox 10-12) on the X chromosome and are expressed in male and female reproductive tissues. All of them exhibit cell type-specific expression. In testes, all Rhox genes are restricted to Sertoli cells except Rhox4 which is predominantly expressed in Leydig cells. Apart from cell-type specificity, these 12 Rhox genes exhibit a colinear expression pattern in which an expression gradient is achieved spatially, temporally, or quantitatively, pertinent to their relative position within subclusters. For instance, the genes in subcluster α display both temporal and quantitative colinearity. Rhox1, the gene located at the distal 5' end of subcluster α express first (days 7-12 postpartum) followed by Rhox2 (day 12 postpartum), Rhox3 and Rhox4 (days 20-22 postpartum). Among them, Rhox1 is expressed at the highest level during testis development than other gene members in same subcluster and each subsequent gene in the same subcluster exhibits a stepwise decline in its expression level. It is believed that such colinear expression pattern observed in the Rhox cluster might provide Sertoli cells with a precise regulatory system to transduce temporally variable signals to germ cells at all stages of development. Clearly, future studies such as targeted disruption or knockdown approaches will be required to reveal the individual and overlapping function of these Rhox genes in spermatogenesis.

The importance of the Rhox gene cluster in spermatogenesis could be demonstrated at least by target disruption of Rhox5 gene in male mice. Ablation of Rhox5 gene by homologous recombination was subsequently found that mutant male are subfertile. Reduced sperm count and sperm motility along with increased germ cell apoptosis were observed in Rhox5-/- mice. Since the expression of Rhox5 is restricted to Sertoli cells, it is likely that Rhox5 plays a role in regulating the expression of Sertoli-cell genes that can modulate germ cell survival. Efforts should be made to elucidate the functional significance of each Rhox member in spermatogenesis and identify target genes that are regulated by the Rhox gene cluster.

POU Homedomain Proteins

Sperm-1

Sperm-1, belonging to the family of the POU (Pit, Oct, Unc) homeodomain proteins, is selectively expressed in male germ cells immediately preceding the first meiotic division and in
the haploid spermatids.\textsuperscript{96,97} Sperm-1 preferentially binds to an octamer DNA-response element with sequence of 5'-GCATATGTTATT-3' in which the optimal sequence differs from that preferred by other POU protein members.\textsuperscript{96}

Knockout studies of Sperm-1 in mice have been performed, null mice develop normal testis, apparently with normal spermatogenesis and produce normal number of motile sperms as those of normal mice, except that the Sperm-1 null male mice are subfertile.\textsuperscript{97} However, the molecular basis for this subfertile phenotype has not yet been elucidated. Thus, identification of the molecular targets and mechanism of action of sperm-1 may help in a better understanding its role in spermatogenesis.

\textbf{Oct-4}

Oct-4 is expressed in the postproliferative prospermatogonia until after birth in male embryos. Oct-4 expression continues in undifferentiated type A spermatogonia as spermatogenesis starts, and is downregulated when germ cells enter their differentiation pathway. There is no reexpression of Oct-4 in germ cells at any developmental stages of spermatogenesis.\textsuperscript{98,99} The downregulation of Oct-4 seems to be one of the molecular triggers in the commitment of meiosis in male germ cells, although the target gene(s) involved in such event has not been identified.

\textbf{C\textsubscript{2}H\textsubscript{2} Zinc Finger Family}

Transcription factors belong to this family must contain C\textsubscript{2}H\textsubscript{2} zinc finger motif (also known as Krüppel zinc finger motif), which is generally present in tandem arrays with the sequence of Y/F-X-C\textsubscript{2-4}-C-X\textsubscript{3}-H-X\textsubscript{2-4}-H, where X can be variable amino acids.\textsuperscript{100} These conserved cysteine and histidine residues are able to bond tetrahedrally to a zinc ion. Plzf and WT1 are two transcription factors that are known to possess a C\textsubscript{2}H\textsubscript{2} zinc finger and have been reported to have significant impact on spermatogenesis.\textsuperscript{101}

\textbf{Plzf}

Plzf is also known as zinc-finger protein 145 (zf145) that is expressed in the developing male gonad.\textsuperscript{102} In postnatal and adult testes, Plzf is restricted to spermatogonia that exhibit stem-cell like properties and is coexpressed with Oct-4, a transcription factor implicated in maintaining stem-cell population.\textsuperscript{102,103} The functional importance of Plzf has been revealed by two in vivo studies. Studies of naturally occurring Plzf-mutant (luxoid) mice and Plzf knockout mice have shown that both mutant mice exhibit a progressive loss of spermatogonia with age, associated with an increase in apoptosis, but without apparent defects in Sertoli cells.\textsuperscript{102,103} Spermatogonial transplantation experiments demonstrated that Plzf is a spermatogonia-specific transcription factor that is required to regulate self-renewal and maintenance of the stem cell pool as transplantation of spermatogonia isolated from Plzf-null mice failed to repopulate gonads that had been chemically depleted of germ cells.\textsuperscript{102} Up to now, no direct target gene of Plzf regulation has been identified. Apparently, it is an area that needs further investigation.

\textbf{Wilms' Tumor Protein (WT1)}

WT1 protein contains four COOH-terminal C\textsubscript{2}H\textsubscript{2} zinc fingers for DNA binding and one of each transcriptional repression and activation domains at its NH\textsubscript{2} terminus.\textsuperscript{104} WT1 plays a crucial role in the development of the genitourinary system.\textsuperscript{105} Conditional knockout of WT1 protein in Sertoli cells by embryonic day 14.5 could result in disruption of developing seminiferous tubules and progressive loss of Sertoli cells and germ cells.\textsuperscript{106} Using tissue-specific RNA interference (RNAi) approach that disrupts the expression of WT1 in mouse testes, studies have shown that increased germ cell apoptosis, loss of adherens junctions and impaired spermatogenesis were observed in siRNA-WT1 mice.\textsuperscript{107} Microarray analysis on siRNA-WT1 testes has found that a spectrum of genes encoding signaling molecules and structural proteins whose expressions were altered.\textsuperscript{107} For instance, integrin cytoplasmic domain associated protein 1α (Icap1-α) and epidermal growth factor receptor pathway substrate 8 (Eps8), which are signaling molecules that regulate actin-mediated cytoskeletal events, are altered in siRNA-WT1 testes.\textsuperscript{107} These results
suggest that Icap1-α and Ep8 are the target proteins of WT1 and WT1 is a crucial transcription factor in regulating spermatogenesis.

**GATA Family**

All GATA proteins contain a DNA-binding domain composed of two conserved multifunctional zinc fingers, C-X$_2$-C-X$_{17}$-C-X$_2$-C, where X represents variable amino acids.$^{108,109}$ GATA proteins recognize and bind to the DNA consensus motif, W|GATAR.$^{109}$ The N-terminal zinc finger is required for the specificity and stability of the DNA binding, whilst the C-terminal zinc finger is for the recognition and binding to the core GATA motif.$^{110-113}$ GATA interacts with cofactors such as Friend of GATA-1 and -2 (FOG-1 and FOG-2) and p300/CBP via the N-terminal or C-terminal zinc fingers, resulting in either activation or repression of gene transcription.$^{114-119}$

GATAs are essential transcription factors in mammalian reproductive development and function. Among six members of this family, GATA-1, -4 and -6 are found in testes. GATA-1 is expressed in mouse Sertoli cells from stages VII to IX of the seminiferous epithelial cycle.$^{120}$ GATA-4 is present in mouse testis throughout all developmental stages and localized to Sertoli cells and Leydig cells.$^{121-123}$ GATA-6 is expressed in neonatal, prepubertal, and adult testes and localized in Sertoli cells.$^{116,122}$ The GATA family members play equally important role in gonadal development, testosterone production and regulation of gene expression in testicular somatic cells such as Sertoli and Leydig cells.$^{118,119}$ For instance, GATA-4 is capable of activating the promoters of testicular genes including Mullerian-inhibiting substance (MIS), PII aromatase (Cyp19), SF-1, StAR and inhibin.$^{124}$ The examples mentioned herein are not intended to be exhaustive, readers are strongly encouraged to read earlier review to gain a more comprehensive view of this protein family.$^{118,119}$

**Nuclear Factor Kappa B (NF-κB) Family**

The NF-κB family of transcription factors regulates a wide variety of genes involved in spermatogenesis. The NF-κB family is composed of p50, p52, p65 (RelA), RelB and c-Rel.$^{125,126}$ which regulates transcription by binding as homo- or heterodimers to κB enhancer elements in the regulatory region of genes. Among five protein subunits, p50 and p65 have been shown to express in rat testes. Nuclear expression of p50 and p65 are cell-type and stage-specific. Nuclear p50 and p60 are highest at stages XIV-VII in Sertoli cells and stages VII-XI in spermatocytes.$^{127}$

Like another transcription factors, the NF-κB family of transcription factors can activate and repress testicular gene transcription. For example, TNF-α induces NF-κB binding to the cAMP-response element-binding protein (CREB) in AR promoters and elevates their promoter activities in Sertoli cells.$^{128,129}$ TNF-α has been reported to downregulate SF-1 transactivation of Mullerian inhibiting substance (MIS) gene in the testis by NF-κB. The SF-1-bound NF-κB could recruit histone deacetylases to inhibit the SF-1-mediated MIS gene activation.$^{130}$ Since TNF-α is a major cytokine secreted by germ cells, it is believed that the effect of TNF-α and its downstream regulators, NF-κB, may not be limited to those identified genes. Clearly, there is much remains to be investigated with regard to the function of NF-κB in spermatogenesis.

**Y-Box**

The family of Y-box proteins contains a conserved cold-shock domain (CSD) for DNA binding, a variable N-terminal domain thought for transactivation and a C-terminal tail for protein-protein interaction.$^{131,132}$ YB-1 was the first identified transcription factor that bound to the Y-box and the consensus DNA sequence was determined as CTGATTGGYYUU, a reverse sequence motif of the CCAAT box.$^{135}$

Mammalian germ cell homologues of *Xenopus* FRG Y1 and FRG Y2 have been identified in mouse testis, namely MSY1 and MSY2 respectively.$^{134,135}$ Similar to *Xenopus* homologues, MSY1 is ubiquitously expressed in somatic tissues; whereas MSY2 is expressed in meiotic and postmeiotic germ cells.$^{134,136}$ Several studies have revealed that Y-box proteins are needed to activate gene transcription in male germ cells, such as protamine 2 and cytochrome c genes.$^{137,138}$ Recent knockout studies further confirmed the functional significance of MSY2 in spermatogenesis.
Spermatogenesis is disrupted in postmeiotic null germ cells with many misshapen and multi-nucleated spermatids.\textsuperscript{139} Apart from MSY2, at least two other Y-box proteins, MSY1 and MSY4, are expressed in meiotic and postmeiotic germ cells.\textsuperscript{134,140} However, their roles on gene transcription pertinent to spermatogenesis remain entirely unknown.

**CAF1**
Chromatin assembly factor-1 (CAF1), also called as Cnot7, is the mammalian homolog of yeast CAF1.\textsuperscript{141} It is a component of the CCR4-NOT complex that has multiple roles in regulating transcription.\textsuperscript{142} CAF1-deficient male mice are sterile owing to oligo-astheno-teratozoospermia shown in two independent knockout studies.\textsuperscript{143,144} Maturation of spermatids is unsynchronized and impaired. Further studies have shown that the proper function of retinoid X receptor $\beta$ (RXR$\beta$)-mediated transcription in the testis requires the interaction of CAF1 through the AF-1 domain of RXR$\beta$, suggesting CAF1 functions as a coregulator of RXR$\beta$ in regulating transcription in testicular somatic cells as RXR$\beta$ is expressed in somatic Sertoli cells and Leydig cell.\textsuperscript{143}

**Testis-Specific Gene Expression**
Testis-specific gene expression could be in part achieved through the expression of testis-specific transcription factors, such as CREM$\alpha$, and cell type-specific components of the general or core transcription machinery as an increasing number of tissue or cell type-specific components of general transcription factors has been identified, such as TFIIA-$\tau$, a testis-specific isoform of TFIIA.\textsuperscript{145-147} An alternative approach to achieve tissue-specific gene expression is by permanent transcriptional repression of that particular promoter in nonexpressing cells via DNA methylation.\textsuperscript{148} A testis-specific expression of histone H1t is one of the examples belonging to this category. The repression of the histone H1t gene in nonexpressing cells is achieved by partial and full methylation of all seven CpG dinucleotides within the H1t proximal promoter, while these CpG dinucleotides are completely unmethylated in primary spermatocytes.\textsuperscript{149,150}

**Transcriptional Regulation of Cell Junction Dynamics**
The translocation of germ cells across the seminiferous epithelium during spermatogenesis requires extensive restructuring of cell junctions at the Sertoli-germ and Sertoli-Sertoli interface.\textsuperscript{151} It is believed that the transcriptional, post-transcriptional and post-translational regulations of cell junction proteins play crucial roles in controlling the assembly and disassembly of cell junctions, resulting in the progressive movement of germ cells to the adluminal from the basal compartment for the completion of spermatogenesis.\textsuperscript{152} Therefore, studies of the transcriptional regulation of junction proteins found at the ectoplasmic specialization (ES) and the blood-testis barrier (BTB) are crucial for the thorough understanding of spermatogenesis. In our laboratory, the transcriptional regulations of nectin-2 and claudin-11 in Sertoli cells have been studied.\textsuperscript{92,153} Nectin-2 is a junction protein localized at Sertoli cells and interacts at nectin-3 that is expressed in germ cells to form the heterotypic interlock between Sertoli and germ cells at the apical ES.\textsuperscript{154} Our recent studies have demonstrated that CREB and c-Jun are bound to the cAMP responsive element (CRE) motif of the nectin-2 promoter located between nucleotides -316 and -211 (relative to the translation start site), resulting in the upregulation of nectin-2 gene transcription. Apart from CREB and c-Jun, two members of Sp1 family, Sp1 and Sp3, are also positive regulators of the nectin-2 transcription.\textsuperscript{92} Analysis of the staged tubules has confirmed that the cyclic expressions of CREB and nectin-2 coincide with the event of apical ES restructuring between Sertoli cells and germ cells. It is believed that the tight regulation of the basal nectin-2 transcription by CREB, c-Jun and Sp1 are crucial to regulate the disassembly of adherens junctions between Sertoli cells and germ cells during spermiation (Fig. 1).
Apart from adherens junction proteins, we have also studied the transcriptional regulation of tight junction (TJ) proteins in Sertoli cells. Claudin-11 is a TJ integral protein found in testis and
Figure 1. A-B) A proposed model for the regulation of nectin-2 expression in testis. This model accounts for the functional cooperation of multiple transcription factors (Sp1 protein family, CREB and c-Jun) in regulating the basal nectin-2 gene transcription. It also illustrates how the cyclic expression of CREB in a spermatogenic cycle influences the nectin-2 gene transcription, which in turn regulates the assembly of SspJ (A) at stages II-VIII and disassembly at stages IX-I (B), resulting in spermiation.
CNS (central nervous system) myelin. In our study, we demonstrated that the overlapping GATA/NF-Y motif within the core promoter of claudin-11 gene is modulated by differential binding of various transcription factors, resulting in dual transcriptional control. We confirmed that GATA, nuclear factor YA (NF-YA), and cAMP response element-binding protein (CREB) form a complex in vivo and bind to the GATA/NF-Y region to promote claudin-11 gene transcription. GATA and CREB transactivation could be further modulated by the presence of Smad3 and Smad4 proteins. Binding of Smad proteins at the GATA/NF-Y motif could repress the GATA and CREB transactivation of claudin-11 gene. Such repression required the recruitment and physical interactions of histone deacetylase 1 and its corepressor, mSin3A, with Smad proteins. It is believed that cyclic changes in the ratio of positive regulators (GATA, NF-YA and CREB) to negative regulators (Smads) in the seminiferous epithelium during the spermatogenic cycle might provide the precise control in claudin-11 gene transcription.

Concluding Remarks and Future Perspectives

As we briefly reviewed and discussed herein, much work on the transcriptional regulation of spermatogenesis conducted in the past two decades was focused on individual transcription factor, and most of these studies relied solely on changes in phenotypes of the knock-out mice to assess the function of different transcription factors. However, the physiological linkage between different transcription factors during spermatogenesis remains unknown. Also, the molecular target genes of these transcription factors at different stages of the seminiferous epithelial cycle are largely unknown. Furthermore, how these genes and their proteins regulate different facets of spermatogenesis, such as germ cell cycle, meiosis, spermatogonial proliferation and renewal, germ cell apoptosis, cell adhesion and junction restructuring, germ cell migration, biochemical and morphological events pertinent to spermiogenesis, and others, remain unexplored. Nonetheless, with the recent advances in genomics and proteomics research, such as the use of gene profiling techniques coupled with mass spectrometry to identify target genes (proteins) important to transcriptional regulation in knock-out mice versus wild types, this shall provide an unprecedented opportunity for investigators in the field.

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Proteases and Their Cognate Inhibitors of the Serine and Metalloprotease Subclasses, in Testicular Physiology

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The testis is a highly dynamic organ not only in the fetal stage but also during postnatal development and in adult life. It is composed of two major compartments: the interstitium with the steroidogenic Leydig cells, and the seminiferous tubules. The seminiferous tubules are surrounded by peritubular cells. Tubules are composed of Sertoli cells and germ cells at different developmental stages. Sertoli cells play key roles in spermatogenesis. They are target cells for follicle stimulating hormone (FSH) and testosterone, responsible for the initiation and maintenance of spermatogenesis. They form the tubules and provide structural and nutritional support for the developing germ cells.1-4

The gonads emerge as an outgrowth and will develop either as a testis or an ovary, depending on the presence of the Sry gene located on the Y chromosome.5-6 In response to Sry, Sertoli cells differentiate. They synthesize the Mullerian Inhibiting substance, and they aggregate to form the cords together with peritubular cells originating from the mesonephros. Subsequently, Leydig cells differentiate in the interstitial milieu and start producing testosterone.7-10 At puberty, dynamic changes are associated with the transformation of the cords into tubules and the initiation of spermatogenesis. In adult life, germ cells migrate from the base to the apex of the tubule epithelium while differentiating further. Finally spermatids are released from the apex of the seminiferous epithelium into the tubular lumen, becoming spermatozoa. A new wave of spermatogenesis will initiate again.

Previous reports suggested that proteases or their inhibitors of the serine-, cysteine-, or metallo-protease family were involved in this spatiotemporal and highly orchestrated process, either during testis development11-13 or at specific stages of spermatogenesis.14-17 This chapter summarizes current knowledge about the occurrence and expression pattern of members of the metallo- and of the serine-family of proteases and inhibitors synthesized within the testis. We also report the various predicted functions for these molecules in the establishment and/or maintenance of the testicular architecture and in the process of spermatogenesis.

General Aspects of Proteases and Protease Inhibitors

A number of important processes that regulate the activity and fate of many proteins are strictly dependent on proteolytic events. For example, proteases are involved in the ectodomain shedding of cell surface proteins, the activation or inactivation of cytokines, hormones and growth factors, the exposure of cryptic neoproteins exhibiting functional roles distinct from...
the parent molecule, degradation of multiple extracellular matrix components facilitating cell migration and invasion. Accordingly, proteases are fundamental in nearly all complex processes of tissue maintenance, repair, growth and development, and alterations in the structure and expression patterns of proteases underlie many pathological processes including cancer, arthritis, osteoporosis, neurodegenerative disorders and cardiovascular diseases. The completion of the human genome sequence has allowed the determination of more than 2% of all human genes are proteases or protease inhibitors, reflecting the importance of proteolysis in human biology.\(^\text{18,19}\) The activity of proteases is regulated at multiple levels including the level of production, the activation of the protease which is generally synthesized in an inactive pro-form, and the production of specific inhibitors.

Proteases catalyse the hydrolysis of peptide bonds in proteins. They are of two types, the exopeptidases and the endopeptidases. The exopeptidases attack only peptide bonds localized at or near the amino or carboxy terminal portion of peptide chains. The endopeptidases, also named the proteinases, catalyse the hydrolysis of internal bonds in polypeptides. They are divided into 5 classes. Aspartic and metzincins proteases use an activated water molecule as a nucleophile to attack the peptide bond of the substrate. In the cysteine, serine and threonine classes the nucleophile is a catalytic amino-acid residue (Cys, Ser or Thr, respectively) that is located in the active site from which the class name derives. Analysis of the full repertoire of proteases present in the human, mouse and rat genome indicated that serine, metzincins and cysteine proteases are the most abundant proteolytic enzymes in rat, mouse and human (Table 1).\(^\text{20}\)

**The Metzincins**

Members of the metzincin superfamily are metalloproteinases that require zinc at their catalytic sites. Metzincins are distinguished by a conserved structural topology, a consensus motif containing three histidines that bind zinc at the catalytic site, and a conserved ‘Met-turn’ motif that sits below the proteinase active site zinc ion. The metzincins can be further subdivided into four distinct families, two of which i.e., the matrixins or matrix metalloproteinases (MMPs) and the adnalysin-related proteinases are abundantly expressed in the testis. The actions of these proteinases are inhibited by the tissue inhibitors of metalloproteinases (TIMPs).

**Matrix Metalloproteinases (MMPs)**

The MMPs are a family of extracellular matrix (ECM) degrading enzymes that share common functional domains and activation mechanism. These are Ca\(^{2+}\) and Zn\(^{2+}\)-dependent endopeptidases that are active at neutral pH. They are synthesized as secreted or transmembrane proenzymes and processed to an active form by the removal of an amino-terminal propeptide. MMPs can be activated by chaotrophic agents or by cleavage of the propeptide by members of the MMP family or other proteases such as the plasminogen activator of the urokinase-type. To date, more than 30 members of the MMP family have been identified. There are several distinct subgroups based on preferential substrates or similar structural domains: collagenases that are active against fibrillar collagen, gelatinases that have high activity against denatured collagens,

<table>
<thead>
<tr>
<th>Total</th>
<th>Aspartic</th>
<th>Cysteine</th>
<th>Metzincins</th>
<th>Serine</th>
<th>Threonine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human</td>
<td>561</td>
<td>21</td>
<td>148</td>
<td>186</td>
<td>178</td>
</tr>
<tr>
<td>Mouse</td>
<td>641</td>
<td>27</td>
<td>163</td>
<td>198</td>
<td>227</td>
</tr>
<tr>
<td>Rat</td>
<td>626</td>
<td>24</td>
<td>160</td>
<td>192</td>
<td>221</td>
</tr>
</tbody>
</table>

Adapted from references 19,20.
stromelysins that degrade noncollagen components of the ECM, membrane-type MMPs (MT-MMPs) that are transmembrane molecules and other less characterized members. Much of the functions attributed to MMPs to date are the results of the cleavage products of ECM components. These include the release of bioactive ECM fragments which may alter the ECM microenvironment, changing the cell from an adhesive to a migratory phenotype. MMPs may also activate chemokines, cytokines and growth factors synthesized as inactive pro-forms, inactivate the SERPINs (SERine Protease INhibitors) and generate soluble forms of a transmembrane receptor through shedding of the ectodomain. 18-21 MMPs are controlled at a transcriptional level depending on MMPs and on the tissue or cell type considered. Such a control is exerted by hormones, growth factors and cytokines as well as basigin or EMMPRIN (Extracellular Matrix metalloproteinase Inducer) which belongs to the immunoglobulin superfamily.22

The Adamalysin-Related Proteinases

This family includes the ADAMs which are cell-surface rather than secreted proteins that share a disintegrin and metalloproteinase domain. They are at least 32 ADAMs that have been cloned and sequenced, each containing a signal sequence followed in order by a pro-domain, a metalloproteinase or metalloproteinase-like domain, a disintegrin-like domain, a cysteine-rich domain, EGF-like repeats, a transmembrane domain, and a cytoplasmic tail. Accordingly, ADAMs potentially perform four distinct but complementary functions: proteolysis via the metalloproteinase domain, adhesion via the disintegrin domain, cell-cell fusion via a candidate hydrophobic fusion peptide in the cysteine-rich domain, and cell signaling via the intracellular domain. A large number of ADAMs show testis-specific expression and are mostly involved in sperm-egg recognition (Table 2).23-25 ADAMTS are the soluble counterparts of the ADAMs. They do not contain a transmembrane domain, but instead contain thrombospondin-1 motifs that permit ECM-association. To date, 19 ADAMTSs have been identified in human tissues, and two of them display testis expression (ADAMTS-2 and -20; Table 2).

### Table 2. General features of ADAMs and ADAMTS

<table>
<thead>
<tr>
<th>ADAMs: 32 Full Length cDNAs</th>
<th>ADAMTS: 19 Human Genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active metalloproteases: 17 ADAMs (1, 8-10, 12, 13, 15-17, 19-21, 24-26, 28, 30)</td>
<td>Exhibit a metalloproteinase catalytic domain. However, most of them have been cloned based on structural organization and have unknown substrates (i.e., the orphan ADAMTSs: 6, 7, 10, 12, 16-19)</td>
</tr>
<tr>
<td>Inactive metalloproteases: 13 ADAMs (2, 7, 11, 14, 16, 18, 22, 23, 28)</td>
<td>ADAMTS-1 and -8 are anti-angiogenic</td>
</tr>
<tr>
<td>Testis specific expression: 13 ADAMs (2, 3, 5, 16, 18, 20, 21, 24-26, 29, 30, 33)</td>
<td>ADAMTS (1, 4, 5, 8, 9, 15) are aggrecanases</td>
</tr>
<tr>
<td>Testis predominant expression: ADAMs (1, 4, 6, 31, 32)</td>
<td>ADAMTS (2, 3, 14) are pro-collagen N-proteinases</td>
</tr>
<tr>
<td>ADAMTS (9, 20) have a GON domain</td>
<td>ADAMTS-13 is a von Willebrand factor-cleaving protease</td>
</tr>
</tbody>
</table>

Note: When in testis, ADAMs are germinal cell products except ADAM 31 which is expressed in Leydig cells. Adapted from references 23-25.
The TIMPs

TIMPs are natural inhibitors of MMPs and inhibit the MMPs proteolytic activity by forming noncovalent 1:1 stoichiometric complexes resistant to heat denaturation and proteolytic degradation. Four TIMPs have been currently characterized and designated TIMP-1, -2, -3 and -4. They exhibit various N-glycosylation sites: two for TIMP-1, one for TIMP-3 and none for TIMPS -2 and -4. They are expressed in a variety of cell types and present in most tissues and body fluids. The TIMPs -1, 2 and 4 are secreted, whereas TIMP-3 is ECM-associated. TIMPs differ in many aspects including solubility, interaction with the proenzymes (proMMPs) and regulation of expression. TIMPs are 21-34 kDa proteins all possessing 12 conserved cysteine residues forming six disulfide bonds that fold the protein in two domains. The N-terminal domain of TIMP contains the MMP inhibitory domain. The C-terminal domain is involved in formation of complexes with the pro-enzymes, thereby regulating the MMP activation process.26-28 TIMPs are multifunctional proteins. In addition to inhibiting target proteinases, TIMPs participate in the MMP activation process through their ability to form complexes with proMMPs. Further, evidences accumulated that TIMP-1 and TIMP-2 transduce an intracellular signalling, although to date no specific TIMP receptors have been characterized.27-29 Originally, TIMP-1 was described for its erythroid-potentiating activity, and as such TIMP-1 plays a pivotal role in hematopoiesis.26 TIMPs -1 and -2 have also been demonstrated to display antiapoptotic and anti-angiogenic activity in various cell lines depending or not on MMP inhibition.29,30 The role of TIMP-3 has been deeply investigated mostly because inherited mutations in it lead to Sorbys fundus dystrophy, a degenerative eye disease.31,32 TIMP-4 has been less characterized but major functions for TIMP-4 have been described in implantation, heart function and ovulation.27,28

Serine Proteases and SERine Protease INhibitors (SERPINs)

The serine protease family is one of the earliest characterized and largest multigene proteolytic families, which has well characterized roles in diverse cellular activities including blood coagulation, platelet activation, fibrinolysis and thrombosis. The serine protease family can be further subdivided into 16 families including the plasminogen activators, the transmembrane-serine proteases and the kallikreins.20

Plasminogen Activators

In mammals, two major types of plasminogen activators have been identified, urokinase-type (uPA) and tissue-type (tPA). Even though both types of PAs catalyze the activation of plasminogen, the currently established functions of uPA-dependent plasminogen activation are mainly within physiological and pathological tissue remodeling processes involving degradation of matrix components and activation of latent proteinases or growth factors, whereas tPA is mainly involved in thrombolysis and neurobiology.33,34 However, it has been observed in gene deficient mice that PAs could substitute each other.34,35 Both PAs are released from cells as single chains with no (uPA) or low (tPA) activity, with cleavage of a polypeptide bond leading to the fully active two-chain forms. The most important feature of this system is the amplification loop achieved by the reciprocal activation of pro-PAs and plasminogen on the cell surface. Both plasmin-catalyzed conversion of pro-PA to active PA and the subsequent active PA-catalyzed conversion of plasminogen to plasmin are accelerated. Therefore, as long as pro-PAs and plasminogen are present, reciprocal proenzyme activation will maintain enzymatically active PAs and plasmin.33,34,36 Another consideration is that although tPA and uPA are secreted proteases, both can bind to cell surface via specific cell surface receptors, being thus protected from the inhibitory actions of the abundant plasma inhibitors.

At least eight apparently distinct plasin/plasminogen binding proteins have been proposed on various cell types, including α-enolase, amphoterin and annexin II.57,58 Annexin II is a 36 kDa, calcium-dependent, phospholipid-binding protein found on the surface of many cell types, which exhibits specific, saturable binding for both plasminogen and tPA. In addition, it has the interesting property of independently binding tPA but not uPA, anchoring both
tPA and plasminogen with high affinity in close proximity to each other on the cell surface, thus providing an environment in which plasmin production is greatly increased. The receptor for uPA is a cysteine-rich, highly glycosylated protein, which is attached to the cell surface by a COOH-terminal glycosylphosphatidylinositol (GPI) anchor. Both the inactive single-chain and the active two-chain uPA can bind to uPAR with high affinity. The receptor uPAR also binds the serum and extracellular matrix protein vitronectin, which is a ligand of αvβ3 integrin, an interaction that requires uPA. In contrast, plasminogen does not bind to uPAR. In addition to the membrane anchored form, cleavage of the GPI-anchor generates a soluble form of uPAR (suPAR). Although lacking a cytosolic domain, uPAR activates multiple intracellular signalling molecules through a connection with integrins, G-protein coupled receptors and caveolin. Signalling pathways induced by uPAR include cytosolic kinase pathways with the activation of intracellular tyrosine kinases, the Focal Adhesion Kinase (FAK) pathway leading to cytoskeletal reorganization, and intracellular calcium mobilization. It is also worth to note that both uPA (the aminoterminal fragment, ATF) and uPAR exhibit growth activities independant of their proteolytic activities.

Type II Transmembrane Serine Proteases (TTSPs)

TTSPs constitute a rapidly expanding family of serine proteases defined by the presence of an N-terminal signal anchor and a C-terminal serine protease domain, separated by a stem region containing an array of protein domains that varies widely between individual TTSPs. These enzymes are ideally positioned to interact with other proteins on the cell surface as well as soluble proteins, matrix components, and proteins on adjacent cells. In addition, TTSPs have cytoplasmic N-terminal domains, suggesting possible functions in intracellular signal transduction. TTSPs are synthesized as single chains zymogens and are likely activated by cleavage following an arginine or lysine present in a highly conserved activation motif. TTSPs are likely to remain membrane-bound following activation. Althought a few of the TTSPs are expressed across several tissues and cell types, in general theses enzymes demonstrate relatively restricted expression patterns, indicating that they may have tissue-specific functions.

Kallikreins

Kallikreins are represented by multigene families in humans and many animal species, especially in rat and mouse. Of particular interest are the glandular kallikreins, nowadays known as the tissue kallikreins. Kallikreins are expressed in a wide range of tissues including steroid-hormone producing or hormone-dependent tissues such as the prostate, breast, ovary and testis. Most, if not all, genes are under steroid hormone regulation, and there is a strong but circumstantial evidence linking kallikreins and cancer(s). Example is given with human kallikreins 2 and 3 (known as Prostate specific antigen) which are widely used tumor markers for prostate cancer. A total of 15 kallikrein genes is reported in the human genome versus at least 25 in the mouse species. Among them, 14 genes are presumed to encode serine proteases, the rest being pseudogenes. Interestingly, there are no homologs for human kallikreins 2 and 3 in the mouse or rat genomes. Tissue kallikreins are clustered at chromosome 19q13 in humans, 1q23 in rat and 7B2 in mouse. They share a similar genomic organization, being formed of five coding exons with very similar exon sizes. All kallikrein proteins are synthesized as prepro-peptides with a signal peptide at the N-terminus, followed by an activation peptide, and the mature protein. Certain ECM components such as fibronectin and laminin, IGFBPs (insulin growth factor binding proteins) and single chain tPA are substrates for kallikreins.

Serpins

The serpins are a superfamily of proteins with full-length coding sequences known or predicted to be about one-half of a total of 500, which fold into a conserved tertiary structural domain. The name serpin derives from the fact that most of the first identified serpins were inhibitors of serine proteinases. Today, this name is clearly inappropriate because a high number of serpins display no inhibitory action against serine proteinases while others inhibit...
cysteine proteinases. Nevertheless, the HUGO gene nomenclature committee recommended retention of the name with classification into clades based on phylogenetic relationships.\textsuperscript{45-48} For example, serpins in the A clade perform roles such as hormone transport i.e., thyroid-binding globulin (SERPINA6), corticosteroid-binding globulin (SERPINA7), and blood pressure regulation (angiotensinogen or SERPINA8) whereas serpins in the E clade are inhibitors of the plasminogen activators (SERPINE1 is plasminogen activator inhibitor 1 (PAI-1) and SERPINE2 is proteinase nexin 1 (PN-1). However, SERPINA5 also known as Protein C Inhibitor (PCI) or plasminogen activator inhibitor 3 (PAI-3) binds retinoic acid and targets activated Protein C and the plasminogen activators.\textsuperscript{49}

Serpins targeting serine proteinases have a unique suicide-substrate mechanism through an interaction with proteinases to form covalent complexes that are not dissociable when boiling in SDS but are sensitive to nucleophiles. Such a mechanism is based on a dramatic conformational change in the serpin. Thus the trapped complex is irreversible in nature. This feature is in marked contrast to what occurs with other classes of inhibitors, which instead used tight noncovalent association between the inhibitor and the proteinase, with little or no conformational change in either protein, to give a thermodynamically stable but reversible complex. Another specificity of serpins is that several of them including SERPINA5, SERPINE1 and SERPINE2 are activated by binding to heparin or other negatively charged glycosaminoglycans. The resulting enhancement in the rates of proteinase inhibition can be up to several 1000-fold suggesting that glycosaminoglycans are rate-limiting factors at sites of serpin action. In the case of the three serpins mentioned above, mechanism is a bridging mechanism in which glycosaminoglycans bind both serpin and proteinase to bring them in an appropriate interaction.\textsuperscript{45-48}

An Overview of the Repertoire in Testis

\textbf{MMPs and TIMPs}

The occurrence of these molecules is highly dependent on the species, the developmental age of the testis and its endocrine environment, as summarized in Tables 3 and 4.\textsuperscript{13,17,50-61} The MMP family has greatly expanded these last 20 years and a tissue distribution has generally been performed for each newly discovered MMP member using adult rat or mouse testes. Although the MMP-18, MMPs 23-26 and MMP-28 are present in the testis,\textsuperscript{50,62-65} there is no indication relative to their cellular localisation. In fact, most of the information available are on the gelatinases MMP-2 and MMP-9 probably because of the availability of a rapid and simple biological test i.e., the gelatin zymography. The human fetal testis is also the site of expression of MMPs and TIMPs.\textsuperscript{66} In the mouse, MMPs 2 and 9 are detected in fetal testes together with

\begin{table}[h]
\centering
\caption{The relative mRNA levels for all mouse MMPs and TIMPs and several ADAMs in testicular tissue from newborn mice}
\begin{tabular}{|c|c|c|c|c|}
\hline
\textbf{mRNA Levels in Newborn Mouse Testes} & \textbf{Very High Levels} & \textbf{High Levels} & \textbf{Moderate Levels} & \textbf{Weak/Very Weak Levels} \\
\hline
\textbf{MMPs, ADAMs} & MMPs: 1, 11, 14, 15, 19, 23 & MMPs: 2, 9, 28 & MMPs: 3, 7, 8, 12, 13, 16, 24, 27 & MMPs: 10, 17, 20, 21, 25 \\
\textbf{ADAMS} & ADAMS: 10, 12, 15, 17, 19 & ADAM-33 & ADAM-28 & \\
\textbf{TIMPs} & TIMPs 2, 3, 4 & TIMP-1 & & \\
\hline
\end{tabular}
\end{table}

Adapted from reference 50.
### Table 4. Summary of the findings reported for MMPs and TIMPs in the testis

<table>
<thead>
<tr>
<th>Cellular Localization</th>
<th>Local /Hormonal Regulation</th>
<th>Fetal Testis</th>
<th>Developing and Adult Testis</th>
</tr>
</thead>
<tbody>
<tr>
<td>TIMP-1</td>
<td>Sertoli cells, peritubular cells, but not in germ cells; adult Leydig cells</td>
<td>- FSH (J) via cAMP, IL1α(J), TNFα, (J) in Sertoli cell cultures</td>
<td>Sex-dimorphic pattern of expression; growth-promoting effect on the gonad</td>
</tr>
<tr>
<td>TIMP-2</td>
<td>20-day old rat Sertoli and peritubular cells, germ cells, adult Leydig cells</td>
<td>- FSH (J) via cAMP, Cytokines (J) in Sertoli cell cultures, TNFα opposes FSH-induced TIMP-2 expression</td>
<td>No sex-dimorphic pattern of expression</td>
</tr>
<tr>
<td>TIMP-3</td>
<td>Sertoli cells, peritubular cells</td>
<td>Role in the migration of the mesonephros towards the urogenital ridge</td>
<td>Enhanced expression of a 30 kDa MW band correlates with enhanced Sertoli-peritubular cell interactions</td>
</tr>
<tr>
<td>MMP-9</td>
<td>Sertoli cells, peritubular cells</td>
<td>- TNFα (J) in Sertoli cells, FGF2 (J) in cocultures of Sertoli and peritubular cells</td>
<td>No sex-dimorphic pattern of expression</td>
</tr>
<tr>
<td>MT1-MMP</td>
<td>Ubiquitously expressed</td>
<td>- Not regulated by FSH</td>
<td>No sex-dimorphic pattern of expression</td>
</tr>
<tr>
<td>MMP-2</td>
<td>Sertoli, peritubular and Leydig cells but not germ cells</td>
<td>- FSH (J) via cAMP, cytokine production inducers (J) in Sertoli cell cultures</td>
<td>No sex-dimorphic pattern of expression</td>
</tr>
</tbody>
</table>
the TIMPs 1-3 and EMMPRIN.\textsuperscript{13} FSH regulation has been explored using 20-day old rat Sertoli cells, and it was shown that FSH regulated MMP-2 but not MMP-9 or MMP-14, and TIMP-1 and TIMP-2.\textsuperscript{17,51,53-55,57-60,67,68} Cytokines largely involved in testicular physiology such as TNF\textsubscript{\alpha} (tumor necrosis factor \(\alpha\)), TGF\textbeta3 (transforming growth factor \(\beta3\)) and FGF2 (basic fibroblast growth factor) have also been shown to regulate various MMPs and TIMPs in a culture or coculture model of testicular cells.\textsuperscript{53-55,58,61} Leydig cells have been shown to express ADAM31,\textsuperscript{69} as well as TIMP-2 throughout development.\textsuperscript{70,71} It is yet unknown whether these proteins are under gonadotropin regulation in Leydig cells.

**Serine Proteases and Serpins**

PAs were the first serine proteases identified within the testis.\textsuperscript{72} Plasminogen is also synthesized within the testis.\textsuperscript{73} Originally, it was described that Sertoli cells were the site of synthesis of the two PAs, and FSH stimulates tPA while reducing uPA levels in the rat testis.\textsuperscript{74,75} Expression of uPA is also under a retinoic acid control.\textsuperscript{74,76} Pachytene and diakinetic spermatocytes exhibit immunoreactivity for tPA,\textsuperscript{77} indicating that a tPA proteolytic event may occur at the spermatocyte surface level. It would be interesting to determine whether the immunoreactivity corresponded to a tPA binding protein or a tPA receptor present on germ cell surface. Annexin II is a good candidate, because it acts as a receptor for tPA and its mRNA is represented in a testis cDNA library.\textsuperscript{78} By contrast, the receptor for urokinase has been identified on both Leydig cells and at Sertoli-germ cell contacts and/or germ cells,\textsuperscript{79} indicating that proteolysis involving plasminogen may occur in the vicinity of Sertoli and germ cells and at the Leydig cell membrane. The receptor for uPA has also been identified on sperm but in that case, uPAR would be involved in sperm-egg recognition.\textsuperscript{80} The binding of uPA to its receptor promotes cell adhesion by increasing the affinity of uPAR for vitronectin.\textsuperscript{81} It is thus of interest that vitronectin has been identified in the cytoplasm of Leydig cells\textsuperscript{82} and in germ cells,\textsuperscript{83} and that PAI-1 is a Sertoli cell product as well as a peritubular product.\textsuperscript{84,85} Indeed, PAI-1 might regulate cell adhesion or migration through competition with uPAR in binding to vitronectin.\textsuperscript{81} PAI-1 is downregulated by cAMP analogs and FSH in Sertoli cells, and up-regulated by locally produced cytokines (TGF\textbeta1, FGF2 and TNF\textsubscript{\alpha}).\textsuperscript{84,88} In contrast, PAI-3 or serpina5 is up-regulated by FSH and testosterone.\textsuperscript{89,91} Of interest is the recent finding that other serpins produced by Sertoli cells are also regulated by androgens, including eppin and the serpins a3n and a12n.\textsuperscript{91}

Germ cells are also a source of various serine proteases and inhibitors including the activated Protein C\textsuperscript{92} and its inhibitor (serpina5),\textsuperscript{79} the hepatocyte growth factor activator (HGFA) and its 2 specific inhibitors, the HAIIs.\textsuperscript{93} They also express the serpinb6b and testisin.\textsuperscript{94,95} Testisin (also named TESP5) is a GPI-anchored protein expressed by premeiotic testicular germ cells and is a candidate tumor suppressor for testicular cancer.\textsuperscript{95} Another TESStis Specific serine protease-1 TESSP-1 is a membrane-bound enzyme specifically expressed in type B spermatogonia and spermatocytes in the adult mouse.\textsuperscript{96} It is not known whether these proteases act within the seminiferous epithelium or later in sperm-egg recognition events as shown with most ADAMs.\textsuperscript{23}

Although few studies have explored the contribution of Leydig cells to the testicular protease repertoire, it is of interest that Leydig cells are known to express various serine proteases and serpins, and for some of these proteases, Leydig cells are the unique testicular site for their expression. For example, the serine protease originally named Leydin is in fact neutrotrypsin.\textsuperscript{18,97} Leydig cells are also the source of kallikreins 21, 24 and 27.\textsuperscript{98-100} Interestingly, LH-hCG was found to regulate several serine proteases and serpins identified in Leydig cells (including urokinase, matriptase-2, kallikrein-21, HAI-2 and PCI).\textsuperscript{93} indicative that common transcriptional signals may drive the expression of these molecules. Furthermore, kallikreins are regulated by testosterone and estradiol.\textsuperscript{98-100} Table 5 recapitulates most of the data available on serine proteases and SERPINs expressed in testis.
Proteases and Their Cognate Inhibitors of the Serine and Metalloprotease Subclasses

The α2-Macroglobulin

Sertoli cells synthesize and secrete α2-macroglobulin, a protease inhibitor with a large spectrum of inhibitory activities against proteinases of the thiol-, serine- metallo- and aspartic acid-families. Such unique inhibition of proteinases by α2-macroglobulin is based on a "trap mechanism" in which α2-macroglobulin is organized as a noncovalently associated dimer of disulfide-linked dimers, and physically sequesters the proteinase inducing conformational changes in the proteinase. Thus binding of proteinases to α2-macroglobulin is irreversible. In contrast to the hepatic protein, α2-macroglobulin is not an acute-phase protein in the rat testis, and it may bind to various cytokines and growth factors thus regulating their bioavailability.

What Potential Functions in Testicular Physiology?

Growth Factor and Receptor Activation and/or Receptor Shedding

Based on the described functions of proteases and inhibitors and considering testicular architecture and physiology, proteases and antiproteases may have a unique function in delivering growth factors trapped in the ECM, in activating growth factors or growth factor receptors, or in the shedding of transmembrane receptors generating soluble forms that would act as dominant negative and impede normal signal transducing pathway following ligand binding to its receptor. ECM is known to function as a reservoir of endogeneous growth factors, sequestering them in an inactive state and protecting them from proteolytic degradation. For example FGF2 which is deeply involved in testicular physiology does not contain a sequence signal for secretion, and it has been proposed that following environmental stimuli, FGF2 is released from the ECM through the action of proteases allowing it to bind to specific
transmembrane FGF receptors and transduce a signal. In addition to release growth factors stored in the ECM, proteases activate growth factors synthesised as inactive pro-factors. For example, uPA activates (at least in seminiferous tubules) pro-TGF-beta and pro-HGF, two decisive growth factors in testicular physiology. In addition, HGFA and hepsin are two serine proteinases recently identified in the mouse testis, in germinal cells and in peritubular or Sertoli cells, respectively. This is indicative that the pro-HGF produced by peritubular cells throughout development may be activated by hepsin whereas pro-HGF produced by adult Sertoli cells would be activated by either HGFA or hepsin, fueling the concept of paracrine between germ cells and Sertoli cells. Further, the testis is also the source for inhibitors of HGFA and hepsin, and one such inhibitor, HAI-2 (HGFA inhibitor type 2) is downregulated by LH-hCG in Leydig cells. Therefore, a proteolytic level of regulation probably exists together with a transcriptional level of regulation in the testis. However, its relative importance versus the transcriptional level of regulation is unknown. In this context, it should be mentioned that c-MET but also FGFR-1 may be specific targets for metalloproteases on the cell surface, yielding soluble receptors that may modulate the biological activities of their respective ligands.

**ECM Matrix Remodeling**

One of the most described roles for proteases and inhibitors relates to the degradation of extra-cellular matrix that forms a physical barrier for cells to invade. In a very comprehensive review on basement membrane and its testicular composition, ECM matrix remodeling is presented as a major event during organogenesis and growth whereas adulthood is characterized by a very low index in the turnover of extracellular matrix components. Furthermore, human pathological testes exhibit a hyalinisation of the seminiferous tubules that is accompanied by a lower sperm production ability and such a feature is also the hallmark of the testicular phenotype in the klinefelter syndrome. Thus, a physiological link is likely to exist between ECM and sperm production.

**Testis Cord Formation**

ECM constitutes a pathway along which cells may migrate, for instance during the migration of primordial germ cells. Indeed, PGCs express several integrins that may act as receptors for fibronectin and laminin that pave the PGC pathway toward the genital ridge. Another very important involvement of ECM proteins during testis organogenesis consists of the formation of a basement membrane between the epithelializing Sertoli cells and the mesenchymal peritubular cells. Originally, the genital ridge is composed of primordial germ cells and a thickened layer of coelomic epithelium. When the indifferent gonad has a XY genotype, SRY induces a cascade of gene expression which results initially in the migration of mesenchymal cells as well as endothelial cells from the adjacent mesonephros. No migration occurs in case of a XX gonad. Such a migration is accompanied by extensive restructuring which underlines a crucial need for balancing the proteinases/inhibitors ratio. Accordingly, major sex-related differences in the ECM components distribution and the expression of proteases and inhibitors have been reported. Knowledge in this area has greatly been enhanced with the development of microarray studies dedicated to the identification of genes expressed in a sex-dimorphic fashion during the initial phases of testicular differentiation. Table 6 summarizes data on the proteinases and inhibitors.

Investigations on the desert hedgehog (dhh) classified in the cysteine proteinases family, have also greatly increased our understanding on the role of the basement membrane in testis compartmentation. Sertoli cells are the source of Dhh whereas receptors are localized on peritubular cells and possibly Leydig cells. Interestingly, Dhh expression levels increase more than 45-fold at the time of testicular differentiation, and Dhh-null testes exhibit disorganized cords with occasional germ cells seen outside cords and abnormal Leydig cell development. It was concluded that these defects likely stemmed from abnormal peritubular stimulation due to the
lack of Dhh. Therefore, it would be of high interest to determine which proteinases and inhibitors may lie downstream of the Dhh/receptor complex.

**Testis Growth and Lumen Formation**

The prepubertal period is characterized by a rapid growth of the testis, the transformation of the seminiferous cords into tubules and the initiation of spermatogenesis. Specifically, tight junctions are formed between neighboring Sertoli cells thus creating the blood-testis barrier, and cords developed a lumen becoming tubules. Accordingly, Sertoli cells reorganize their cytoskeleton to support additional spermatogenic cell types as spermatogenesis is initiated, and tubules increase in diameter as well as in length. Several in vitro observations raise the possibility that proteases and inhibitors in response to hormonal (mainly FSH) and local stimuli such as HGF and FGF2 are critically involved in these substantial prepubertal changes.

For instance, using prepubertal rat Sertoli cells cultured in a two-chambered assembly to mimic the Sertoli cell barrier, it was demonstrated that proteases were implicated in the changes in the Sertoli cell cytoskeleton elicited by FSH and in modulation of the formation and maintenance of the Sertoli cell barrier. The nature of the protease(s) is not fully identified, but 2-macroglobulin opposed its action and increased integrity of the Sertoli cell barrier.67,136 MMP-2 and tPA are good candidates because they are Sertoli cell products and under FSH control.55,74,75 A second set of experiments was designed to examine cord and lumen formation by Sertoli cells cocultured with rat prepubertal peritubular cells or Sertoli cells cultured on a reconstituted ECM. These models have been extremely fruitful in evidencing cooperativity between Sertoli cells and testicular peritubular cells in the production and deposition of extracellular matrix components,137 and in highlighting the role of laminin in the morphogenetic cascade resulting in the formation of tubule-like structure.138,139 Other experiments also suggested that ECM components regulated the expression of tight junction proteins and the formation of a lumen.58,60 Inasmuch as MMPs and PAs degrade laminin, fibronectin and collagen

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**Table 6. Fold-increase in proteinases and inhibitors in fetal testes versus time-matched fetal ovaries**

<table>
<thead>
<tr>
<th>Name</th>
<th>1.5 &lt; Fold Change &lt;5</th>
<th>Fold Change &gt;5</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Serine proteinases</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Masp</td>
<td>E12.5</td>
<td>Neurotrypsin</td>
</tr>
<tr>
<td>Neurotrypsin</td>
<td>E12.5, E13.5</td>
<td></td>
</tr>
<tr>
<td><strong>Aspartic proteinases</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Caspase</td>
<td>7E12.5</td>
<td></td>
</tr>
<tr>
<td>Cathepsin</td>
<td>E12.5</td>
<td></td>
</tr>
<tr>
<td><strong>Cysteine proteinases</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ADAMTS2</td>
<td>E11.5</td>
<td></td>
</tr>
<tr>
<td>ADAMTS2, 5, 16</td>
<td>E12.5</td>
<td></td>
</tr>
<tr>
<td>ADAMTS2, 5, 7</td>
<td>E13.5</td>
<td></td>
</tr>
<tr>
<td>ADAM 12, 19</td>
<td>E13.5</td>
<td></td>
</tr>
<tr>
<td><strong>Metallo-proteinases</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Serpin B1a, serpin A3g,</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SPINT-2)</td>
<td>E12.5</td>
<td></td>
</tr>
<tr>
<td>(Serpin B1a, Serpin A3g)</td>
<td>E13.5</td>
<td></td>
</tr>
<tr>
<td><strong>Serpin</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Serpin B1a, serpin A3g,</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SPINT-2)</td>
<td>E12.5</td>
<td></td>
</tr>
<tr>
<td>(Serpin B1a, Serpin A3g)</td>
<td>E13.5</td>
<td></td>
</tr>
<tr>
<td><strong>Cystatin</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Cystatin 8, cystatin C)</td>
<td>E12.5, E13.5</td>
<td></td>
</tr>
<tr>
<td><strong>Tissue Inhibitor of MMPs</strong></td>
<td>TIMP-1 E13.5</td>
<td></td>
</tr>
</tbody>
</table>

E11.5, E12.5, E13.5, indicate the age at which the fold-change was registered. Data are from microarray studies of two laboratories.130,131
IV i.e., the major ECM components of the testicular basement membrane, any remodeling necessary to support the rapid and extensive growth of the prepubertal testis is thus expected to involve a delicate interplay between proteases and inhibitors, and certain growth factors. Indeed, various growth factors promote formation of cord-like structures by Sertoli cells in vitro. For instance, FGF2 which mediates mesenchymal-epithelial interactions of peritubular cells and Sertoli cells in the rat testis, promoted de novo testis cord formation and enhanced MMP-9, the 30 kDa glycosylated form of TIMP3 and PAI-1 in the cocultures.61 It would be of interest to investigate the expression pattern of proteases and inhibitors in the HGF-treated cultures because HGF is a powerful morphogen for Sertoli cells cultured on a reconstituted basement membrane. It not only promotes cords but also their further remodeling into tubules. 107,140 Additionally, the antisense strategy would constitute an elegant means to link morphogen cytokines, ECM components, restructuring events and involvement of proteases.

Few data are yet available in vivo to support these in vitro experiments. However, a recent study highlighted the role of laminin during the prepubertal period. Indeed, it was shown that male mice deficient for laminin alpha2 chain exhibited abnormal testicular basement membranes and displayed a defect in the timing of lumen formation, resulting in production of fewer spermatids. Furthermore, the authors demonstrated that laminin alpha1 chain corrected male infertility caused by absence of laminin alpha2 chain.141

Spermatogenesis and the Apical Migration of Germ Cells towards the Lumen

Different authors have been interested in understanding the dynamics of spermatogenesis which relies on the passage of the blood-testis barrier and the release of the elongated spermatids at the apex (spermiation). The description of the testis barrier is beyond the scope of this chapter and is treated in a recent and excellent review.16 However, it is noteworthy that the testis barrier is unique when compared to other blood-tissue barriers (e.g., blood-brain and blood-retina barriers) that it is composed of gap junctions, desmosomes, tight junctions and ectoplasmic specializations, precluding that the passage of germ cells requires a finely tuned process to allow its spatiotemporal opening and closing, not disturbing the integrity of the testis barrier which would provoke a pathological arrest of spermatogenesis. Because this situation is reminiscent of cell migration across ECM, different authors have concentrated their efforts in determining the composition of the junctions, most specifically those that are restricted to testis i.e., the ectoplasmic specialisations, and the way junctional proteins are transcriptionally and post-transcriptionally regulated. It was also reasonable to think that proteases which act like scissors would help germ cells in migrating along Sertoli cell membranes, and that specific inhibitors would restrict the activity of the proteases in a finely tuned regulatory fashion to preserve homeostasis. Therefore, a list of the cytokines, proteases and inhibitors present at the right time and at the right place has been tentatively established.142,143

First evidences came from the demonstration that the plasminogen activators were expressed as a function of the stages of the seminiferous epithelium, and an increased PA activity was found at the time of elongating spermatid translocation and spermiation at the stages VII-VIII.74,77,144,145 Interestingly, the expression of α2-macroglobulin fluctuates with the stages of the seminiferous epithelium, and immunostaining concentrated at stages I-VI, thus prior spermiation indicative that α2-macroglobulin may protect the integrity of the seminiferous epithelium against excessive proteolysis.146 Furthermore, it was shown that addition of germ cells to Sertoli cell cultures resulted in an enhancement of PA activity147 (and also of cathepsin L, a cysteine protease)148,149 and that this correlated in time with the dynamics of assembly/desassembly of the de novo adherens junctions formation between the cultured Sertoli cells. Furthermore, the expression of α2-macroglobulin but also of cystatin (a cathepsin L inhibitor) in the coculture model was consistent with the idea that proteases and their corresponding inhibitors were working synergistically supporting the evidence that they may be involved in the adherence of germ cells to Sertoli cells and the subsequent formation of intercellular junctions.150-152
Spermiation i.e., extrusion of elongated spermatids into the lumen, is the alternate major event occurring during stages VII-VIII. It is followed by the phagocytosis of the cytoplasm shed from the elongated spermatids, which are called the residual bodies.1,153,154 Interestingly, phagocytosis of residual bodies resulted in an increase PA activity155 in an in vitro model of coculture of Sertoli cells and residual bodies.156 Further, the addition of an anti-interleukin 1α antibody prevented the RB-induced enhancement of PA activity,155 thus emphasizing the role of this cytokine in the process of spermiation.157,158 Furthermore, given that the increased PA activity may facilitate the passage of germ cells across the testis barrier, it was suggested that a proteolysis-dependent message would participate in the synchronisation process of the spermatogenesis cycle,155 supporting the pioneering hypothesis of Regaud and Roosen-Runge.153,154 Two other cytokines have proven to be essential at least in the passage of the testis barrier by preleptotene spermatocytes. These are TGFβ3 and TNFα and the readers are encouraged to read recent reviews on the subject.16,143,152

Interestingly, stages VII-VIII are highly testosterone-dependent as demonstrated in models with testosterone deficiency in which a premature detachment of germ cells in the lumens of the tubules is described.159-162 In addition, androgens inhibit PA activity secreted by Sertoli cells in culture in a two-chambered assembly.163 Thus serpinA5 is of tremendous interest because it is upregulated by testosterone,90,91 it opposed PA activity and deficient mice develop male sterility.164 Specifically, lumens are filled with immature germ cells because of an unopposed proteolytic activity of the urokinase type.164 Such a testicular phenotype is reminiscent of the testicular phenotype described in mice deficient for claudin 11.165 Claudin 11 as well as claudin 1 and 3 are essential components of the testis barrier,165,166 and they are up-regulated by testosterone.166-168 In addition, claudins contribute together with MT1-MMP and TIMP-2 in activating MMP-2 secreted as a pro-form.169 MMPs may also be activated by urokinase. Thus, the germ cell enhancement of MMP-2 activity17 may in part, result from the increase in the activity of the PAs observed in Sertoli cell-germ cell cocultures and discussed above. In that context, it would be of interest to determine whether claudins are substrates for either PAs or MMPs, and to investigate claudin expression in the serpinA5-deficient testes and vice-versa.

Collectively, it appears that germ cells which do not bear classic characteristics of migrating cells regulate their own progression within the seminiferous epithelium, through a modulation of the expression pattern of the proteases and inhibitors produced by Sertoli cells, supporting the hypothesis that Sertoli cells act as facilitators of migration, and adding a new function to these nurse cells. Future experiments aiming at dissecting the kinetics of the reliant events of spermiation and translocation would be useful in deciphering such integrated system with hormones and local environment.

**Proteolysis and Steroidogenesis**

Because Leydig cells exhibit a specific repertoire of proteases and inhibitors and that several (if not all) of them are under gonadotropin regulation via cAMP,93 the question arises as to whether a link exists between steroidogenesis and proteolysis. Different arguments emphasize a role of ECM in the capacity of Leydig cells to respond to LH-hCG, and thus indirectly of proteases and inhibitors. For instance, it was shown that fibronectin and collagen IV induce downregulation of steroidogenic response to gonadotropins.170,171 Furthermore, TGFβ which is known to cause augmented fibronectin deposition172 and to elicit cytoskeletal changes in Leydig cells similar to those evidenced when these cells are cultured on plates precoated with fibronectin,173 inhibits DNA synthesis and antagonizes gonadotropin steroidogenic action in Leydig cells.173,174 However, direct evidences for involvement of proteases are still lacking.

**Lessons from Transgenic Mice**

Inasmuch as proteases and inhibitors are extremely abundant and redundant in their spectrum of actions, it is not surprising that very little knockout mouse models have to date contributed to our understanding on their roles in testicular function (Table 7). However, it should be
stated that most of the time no systematic analysis of the testes of the deficient mice had been undertaken unless the authors experienced reproductive difficulties. For example, male mice deficient in PAI-1, TIMP-1 or cathepsin L may still reproduce. However because several pieces of evidences supported a role for cathepsin L in germ cell movement during spermatogenesis, testes from deficient mice were carefully investigated and tubules were found to contain 32% fewer spermatids than the average tubule number of control mice. Such a study should be done on the MT1-MMP-deficient male mice which showed “no signs of sexual maturation”, as stated in the original publication. In other cases, the testis may not be the primary target as shown with male mice deficient in serpinE2 which are sterile because of altered semen protein composition. The different cases reported in Table 7 should be more informative with respect to the role of proteinases in testicular physiology, provided that an extensive study of the male reproductive system is done. Indeed, male mice deficient for EMMPRIN are azoospermic. Specifically, spermatogenesis is arrested at the metaphase of the first meiotic division, and the lumens are filled with round degenerated cells. Given that the expression of EMMPRIN correlates in time with the appearance of spermatocytes in the seminiferous epithelium, it is predicted that EMMPRIN is involved in the interactions between Sertoli cells and germ cells. However, no studies have yet reported the expression pattern of the MMPs known to be under EMMPRIN control in the deficient testes. Male sterility is also observed in transgenic mice with inactive alleles for ADAMTS-2, overexpression of MMP-7 and deficiency for serpina5, but the exact nature of the disorder remains to be fully characterized.

Conclusions and Future Directions

We herein provided a series of evidences highlighting that proteases may be active partners in establishing and maintaining testicular architecture, and in facilitating germ cell migration which constitutes a prerequisite for germ cell progression throughout the spermatogenic process. Therefore, it may be worth to revisit the phenotypes of transgenic male mice deficient for a proteinase

<table>
<thead>
<tr>
<th>Transgenic Mice</th>
<th>Phenotype of the Deficient Male Mice</th>
</tr>
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<tbody>
<tr>
<td>ADAMTS-2</td>
<td>Infertile with few sperms and large number of round spermatids but few elongate spermatids.</td>
</tr>
<tr>
<td>Cathepsin-L</td>
<td>Furless mice. Tubules contain 32% fewer spermatids per Sertoli cell than the average tubule number of control mice, as a result of reduced formation of preleptotene spermatocytes and their differentiation into pachytene spermatocytes.</td>
</tr>
<tr>
<td>EMMPRIN</td>
<td>Male mice are azoospermic because spermatogenesis is arrested at the metaphase of the first meiotic division, and the lumens are filled with round degenerated cells.</td>
</tr>
<tr>
<td>Matriallysin</td>
<td>Overproduction causes reduced sperm production beginning at 8 months of age. The architecture of the testes is altered.</td>
</tr>
<tr>
<td>MT1-MMP</td>
<td>Dwarfs with craniofacial and skeletal abnormalities; show no signs of sexual maturation.</td>
</tr>
<tr>
<td>Protease nexin-1 (Serpine2)</td>
<td>Sterility results from a defect in semen protein composition, which leads to inadequate semen coagulation and deficient vaginal plug formation.</td>
</tr>
<tr>
<td>Protease nexin-1 (Serpine5)</td>
<td>Sterility would be due to a destruction of the BHT because of increase urokinase activity within the seminiferous epithelium resulting in premature detachment of germ cells into the lumen.</td>
</tr>
</tbody>
</table>
or an inhibitor proven to be expressed at the time of translocation and/or spermiation. Furthermore inasmuch as various proteases, inhibitors, junctional components (e.g., claudins, occludins, JAMs) are under a complex hormonal regulation via gonadotropins and/or testosterone, and local regulatory control involving cytokines and growth factors, models with reduced testosterone bioavailability or with limited FSH or LH action coupled to microarray studies, as those recently published should be of tremendous benefit to fully understand the mechanisms that underpin the role of proteases and inhibitors in testis development and function.

Acknowledgements

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Antioxidant Systems and Oxidative Stress in the Testes

R. John Aitken* and Shaun D. Roman

Introduction

Spermatogenesis is an extremely active replicative process capable of generating approximately 1,000 sperm a second. The high rates of cell division inherent in this process imply correspondingly high rates of mitochondrial oxygen consumption by the germinal epithelium. However, the poor vascularization of the testes means that oxygen tensions in this tissue are low1 and that competition for this vital element within the testes is extremely intense. Since both spermatogenesis2 and Leydig cell steroidogenesis3,4 are vulnerable to oxidative stress, the low oxygen tension that characterizes this tissue may be an important component of the mechanisms by which the testes protects itself from free radical-mediated damage. In addition, the testes contain an elaborate array of antioxidant enzymes and free radical scavengers to ensure that the twin spermatogenic and steroidogenic functions of this organ are not impacted by oxidative stress. These antioxidant defence systems are of major importance because peroxidative damage is currently regarded as the single most important cause of impaired testicular function underpinning the pathological consequences of a wide range of conditions from testicular torsion to diabetes and xenobiotic exposure. This chapter sets out the specific nature of these antioxidant defence systems and also reviews the factors that have been found to impair their activity, precipitating a state of oxidative stress in the testes and impairing the latter’s ability to produce viable spermatozoa capable of initiating and supporting embryonic development.

Antioxidant Enzymes

Despite the low oxygen tensions that characterize the testicular micro-environment, this tissue remains vulnerable to oxidative stress due to the abundance of highly unsaturated fatty acids (particularly 20:4 and 22:6) and the presence of potential reactive oxygen species (ROS)-generating systems. ROS generation can be from the mitochondria and a variety of enzymes including the xanthine- and NADPH- oxidases,5,6 and the cytochrome P450s.7 These enzymes specialize in the professional generation of ROS or produce these toxic metabolites as an inadvertent consequence of their biochemical activity. In order to address this risk, the testes have developed a sophisticated array of antioxidant systems comprising both enzymatic and non-enzymatic constituents. Concerning the enzymatic constituents of this defence system, the induction of oxidative stress in the testes precipitates a response characterized by the NF-κB mediated induction of mRNA species for superoxide dismutase (SOD), glutathione peroxi-
Antioxidant Systems and Oxidative Stress in the Testes

The fundamental biochemistry of these antioxidant enzymes is summarized in Fig. 1 and involves the rapid conversion of superoxide anion \( \text{O}_2^- \) to hydrogen peroxide \( \text{H}_2\text{O}_2 \) in the presence of SOD in order to prevent the former from participating in the formation of highly pernicious hydroxyl radicals. The \( \text{H}_2\text{O}_2 \) generated in this manner is a powerful membrane permeant oxidant in its own right that has to be rapidly eliminated from the cell in order to prevent the induction of oxidative damage to lipids, proteins and DNA. The elimination of \( \text{H}_2\text{O}_2 \) is either effected by catalase or glutathione peroxidase, with the latter predominating in the case of the testes.\(^9,10\) GST on the other hand involves a large and complex family of proteins that catalyse the conjugation of reduced glutathione via the sulphydryl group to electrophilic centres on a wide variety of substrates in preparation for excretion from the cell. This activity is critical in the detoxification of peroxidised lipids as well as the metabolism of xenobiotics.
Given the importance of SOD in this defence strategy, it is not surprising that the testes contain not only the conventional cytosolic (Cu/Zn) and mitochondrial (Fe/Mn) forms of SOD but also feature an unusual form of extracellular SOD, (SOD-Ex) which is produced by both Sertoli and germ cells, particularly the former. There is also some evidence that the germ cells may stimulate the secretion of SOD-Ex by Sertoli cells through the actions of cytokines such as interleukin-1\(\alpha\). The importance of the cytosolic form of SOD (SOD1) was recently emphasised in studies of SOD1-knockout mice subjected to testicular heat stress. This treatment induced significantly enhanced levels of DNA strand breakage and cytochrome C leakage from the mitochondria of germ cells in these animals compared with the wild-type controls. Similarly, the importance of the mitochondrial form of SOD (SOD2) in controlling \(O_2^-\) leakage from testicular mitochondria has been emphasised by the finding that the mRNA for this enzyme is markedly higher in the testes than the liver, unlike GPx and catalase. Moreover, SOD-2 mRNA levels are developmentally and translationally regulated with maximal levels of expression in early post-meiotic germ cells.

Although catalase is of limited importance in the testes, there are several isoforms of GPx in this tissue that use glutathione (GSH) as a source of electrons to reduce H\(_2\)O\(_2\) to water. They are concentrated in the mitochondria, nucleus and acrosomal domain of differentiating spermatids. The phospholipid hydroperoxide GPx (PHGPx) is one of the most important GPx isoforms in a testicular context and is highly expressed in both spermatogenic and Leydig cells. Since most forms of GPx are selenium dependent it is possible to gauge the importance of these enzymes in the support of testicular function by examining the impact of selenium deficiency on male reproduction. Animals fed on a selenium deficient diet exhibit a significant reduction of testicular GPx activity and an accompanying loss of germ cells from the germinal epithelium of the testes. Moreover, selenium administration prior to the creation of oxidative stress in the testes using the torsion/detorsion model (see later) to create ischemia-reperfusion injury, has been found to suppress lipid peroxidation and improve the histopathological profile.

**Small Molecular Mass Antioxidants**

In addition to the major ROS processing enzymes, the testes rely heavily on small molecular weight antioxidant factors for protection against oxidative damage. These factors include ions and a wide variety of free radical scavengers, the nature of which are reviewed below.

**Zinc**

Zinc is an acknowledged antioxidant factor that as well as being a core constituent of free radical scavenging enzymes such as SOD and a recognized protector of sulfhydryl groups, is also thought to impair lipid peroxidation by displacing transition metals such as iron and copper from catalytic sites. In keeping with such a central antioxidant role, this element has a profound effect on the level of oxidative stress experienced by the testes. Thus rats fed a zinc deficient diet experience a decrease in testicular antioxidant potential and a concomitant increase of lipid peroxidation in this tissue. Conversely, zinc administration will counteract the oxidative stress created in the testes by exposure to lead. as well as the peroxidative damage induced by ischemia-reperfusion as a consequence testicular torsion-detorsion. Zinc administration has also been shown to attenuate the testicular oxidative DNA damage induced by cadmium as well as the decline in sperm production and testosterone secretion induced by this heavy metal.

**Vitamins C and E**

It has been recognized since the 1940s that vitamin E (\(\alpha\)-tocopherol) is a powerful lipophilic antioxidant that is absolutely vital for the maintenance of mammalian spermatogenesis. It is present in particularly high amounts in Sertoli cells and pachytenes spermatocytes and to a lesser extent round spermatids. Vitamin C (ascorbic acid) also contributes to the support of spermatogenesis at least in part through its capacity to reduce \(\alpha\)-tocopherol and
Antioxidant Systems and Oxidative Stress in the Testes

maintain this antioxidant in an active state. Vitamin C is itself maintained in a reduced state by a GSH-dependent dehydroascorbate reductase, which is abundant in the testes. Vitamin C or E leads to a state of oxidative stress in the testes that disrupts both spermatogenesis and the production of testosterone. Conversely, ascorbate administration to normal animals stimulates both sperm production and testosterone secretion. This vitamin also counteracts the testicular oxidative stress induced by exposure to pro-oxidants such as arsenic, PCBs (Arochlor 1254), cadmium, endosulfan and alcohol. Furthermore, endogenous ascorbate levels decrease dramatically when oxidative stress is induced in the testes by, for example, chronic exposure to lead, chromium, cadmium or aflatoxin. Vitamin E has also been shown to suppress lipid peroxidation in testicular microsomes and mitochondria and to reverse the detrimental effects of oxidative stress on testicular function mediated by exposure to such factors as ozone, iron overload, intensive exercise or exposure to aflatoxin, PCB, cyclophosphamide and formaldehyde. Furthermore testicular vitamin E levels have also been shown to fall significantly when oxidative stress is induced by exposure to pro-oxidant stimuli such as chromium.

Melatonin and Cytochrome C

The pineal hormone melatonin (N-acetyl, 5-methoxytryptamine) also plays a major role in protecting the testes from oxidative stress, given the significant stimulatory effect of pinealectomy on the oxidative damage recorded in the testes as a consequence of induced hyperthyroidism. Melatonin has two major attributes that set it apart from most other antioxidants. Firstly, it undergoes a two electron oxidation when acting as antioxidant, rather than the one electron oxidation favoured by many free radical scavengers. As a result, this compound cannot redox cycle and inadvertently generate free radicals. Secondly, melatonin is readily soluble in both lipid and aqueous environments and can readily cross the blood-testes barrier to protect the germinal epithelium. Melatonin levels in seminal plasma are depressed in infertile patients exhibiting poor motility, leukocytospermia, varicocele and non-obstructive azoospermia, all of which are conditions associated with oxidative stress in the male tract. Moreover, the intraperitoneal injection of melatonin has been shown to alleviate oxidative stress in the testes following the experimental induction of a left sided varicocele. Another small molecular mass free radical scavenger that has recently been shown to play a major role in reducing H$_2$O$_2$ is a testes-specific form of cytochrome C. This cytochrome C isoform is also a powerful activator of apoptosis, providing additional protection to the testes by virtue of its ability to facilitate the depletion of damaged germ cells.

Disruption of the Antioxidant Status of the Testes

Notwithstanding the antioxidant protection afforded to the testes in order to support its dual functions of steroidogenesis and sperm production, a wide variety of endogenous and exogenous factors are known to perturb these defences and generate a state of oxidative stress. In the following section, some of these factors are reviewed.

Cryptorchidism

The elevated temperatures associated with experimental cryptorchidism are associated with oxidative stress in the testes and a reduction in SOD and catalase activities. Consistent with these findings, direct exposure of spermatogenic cells to elevated temperatures was found to induce high rates of apoptosis via mechanisms that were associated with elevated levels of H$_2$O$_2$ generation and could be ameliorated by the addition of catalase. Moreover the consequence of heat stress on spermatogenic cells was exacerbated in SOD1-knock out mice via mechanisms that could be reversed by the addition of Tiron, a superoxide anion radical scavenger. The clinical significance of this finding can be seen in the high levels of DNA damage and ROS generation seen in the spermatozoa of patients with a history of cryptorchidism.
**Testicular Torsion**

Testicular torsion is a relatively common, painful condition that must be treated rapidly if the testes are not to suffer permanent damage. Prolonged torsion leads to testicular ischaemia and high levels of oxidative stress in the ipsilateral testes associated with NO and H$_2$O$_2$ production, increased lipid peroxide formation, isoprostane accumulation, antioxidant enzyme depletion and increased rates of mitochondria-mediated apoptosis in the germ line. Even short periods of ischaemia, for 3 hours or less, can lead to high levels of oxidative stress in the testes, depletion of testicular glutathione levels and the consequent disruption of spermatogenesis. Significantly, the level of peroxidative damage observed in testicular tissue increases following detorsion, indicating the induction of reperfusion injury. The biochemical basis for reperfusion injury is thought to involve a key metabolic enzyme, xanthine dehydrogenase, which becomes converted to a xanthine oxidase during ischaemia, due to oxidation of essential -SH groups and/or a limited proteolytic clip. As soon as the tissue is reperfused with blood, the xanthine oxidase is suddenly presented with oxidizable substrate in the form of xanthine/hypoxanthine and starts to generate copious amounts of ROS. The latter then induce high levels of peroxidative damage via mechanisms that are enhanced by the local release of transition metals. Although this scheme of events was developed to explain the tissue injury associated with conditions such as myocardial infarction, it also applies to the testicular injury associated with torsion-detorsion. The general notion that the testicular damage precipitated by temporary ischaemia is associated with oxidative stress is supported by the sudden induction of lipid peroxidation and the concomitant suppression of endogenous antioxidant activities including SOD, catalase and glutathione peroxidase. In addition, the tissue injury induced by testicular torsion/detorsion can be dramatically alleviated by pretreatment with exogenous antioxidants such as selenium, resveratrol, L-carnitine, caffeic acid phenethyl ester and garlic extract. Finally the enzyme purportedly associated with reperfusion injury, xanthine oxidase, can be inhibited by allopurinol and the latter is known to reduce the testicular damage associated with testicular torsion. Notwithstanding the importance of xanthine oxidase-mediated oxidative stress it should also be noted that neutrophil infiltration into the testes following torsion may represent yet another source of uncontrolled free radical generation responsible for mediating the pathophysiological consequences of temporary testicular ischaemia.

One of the major issues associated with the clinical management of unilateral testicular torsion is whether the ipsilateral testis should be removed in order to preserve the contralateral testis. In animal models prolonged testicular torsion results in excessive ROS generation, depletion of antioxidant enzymes and the appearance of oxidative damage in the contralateral testes. In light of these data, surgical removal of the ipsilateral testes would seem warranted if the period of ischaemia has been extensive.

**Varicocele**

The impaired venous drainage to the testes seen with varicocele is also associated with the disruption of spermatogenesis via mechanisms involving the induction of oxidative stress. In clinical studies, the presence of a varicocele has been shown to correlate with excess ROS generation by the spermatozoa, high rates of DNA damage in these cells and depleted antioxidant levels in the seminal plasma. In a recent study, surgical correction of left sided varicocele was shown to significantly improve sperm concentration, total count, morphology and motility in concert with significant improvements in the antioxidant status of the spermatozoa and seminal plasma. This reduction in oxidative stress secondary to varicocele excision was accompanied by a reduction in both protein carbonyl expression and DNA damage in the spermatozoa. Independent studies have also shown that the testicular expression of 4-hydroxy-2-nonenal modified proteins (another marker of oxidative stress) is significantly higher in patients that responded positively to varicocelectomy, suggesting that surgical treatments are capable of reducing oxidative stress in the testes. Immunocytochemical analyses of 8-hydroxy-2’-deoxyguanosine expression in the testes of varicocele patients also revealed particularly high levels of oxidative DNA damage in the
spermatogonia and spermatocytes that correlated well with the severity of the varicosity. The general concept that testicular pathologies associated with varicocele are linked with the induction of oxidative stress has been confirmed in animal models. Thus, creation of experimental bilateral varicocele in rats is associated with increases in lipid peroxidation and NO generation and a corresponding decrease in testicular antioxidant status. Moreover, the pathological consequences of experimental varicocele induction can be significantly reversed by the concomitant administration of an antioxidant, melatonin.

The site of free radical generation in varicocele patients is still open to conjecture. On the one hand, enhanced free radical generation by the spermatozoa and/or precursor germ cells has been repeatedly suggested, on the other, there is evidence to suggest that excess free radical generation may involve the spermatic vein itself. The excess generation of free radicals by the spermatozoa may be an indirect consequence of impaired spermatogenesis/epididymal function resulting in the retention of excess residual cytoplasm. The presence of excess cytoplasm has been positively correlated with the generation of ROS by human spermatozoa, via mechanisms involving the facilitated supply NADPH to oxidases in the sperm plasma membrane. These enzymes, including NOX5 and DUOX, both of which have been identified in human spermatozoa, are normally deprived of sufficient NADPH to drive free radical generation; what hexose monophosphate shunt activity there is, being largely devoted to the maintenance of glutathione reductase activity. However, when excess residual cytoplasm is present the limited substrate availability is no longer an issue and free radical generation can be initiated. The relevance of this model to the oxidative stress detected in cases of varicocele is clearly suggested by the effects of varicocelectomy. Thus, not only do the spermatozoa produced by such patients exhibit high levels of ROS in association with cytoplasmic retention but also surgical correction of this condition both prevents cytoplasmic retention and suppresses ROS generation. A causative association between these events therefore seems likely.

Hyperthyroidism

The induction of hyperthyroidism in rats is associated with oxidative stress in the testes as reflected by increased lipid peroxidation, elevated GSH levels and induction of antioxidant enzymes. The oxidative stress appears to be associated with a thyroxine dependent increase in mitochondrial activity and concomitant leakage of electrons from the mitochondrial electron transport chain. The oxidative stress precipitated by hyperthyroidism can be exacerbated by pinealectomy removing melatonin, an important testicular antioxidant, from the redox equation. These data resonate with clinical studies indicating that hyperthyroidism is associated with poor semen quality, particularly impaired motility, that normalize when the patients’ thyroid dysfunction is corrected and euthyroidism established. It should also be noted that hypothyroidism can induce oxidative stress in the testes as reflected by enhanced levels of H2O2 production and increased carbonyl generation. Clearly, normal testicular function is highly dependent on a functional thyroid system.

Diabetes

Experimental induction of diabetes in animal models has been shown to impair testicular function and decrease male fertility. Thus, diabetogens such as streptozotocin, enhance ROS generation and induce both lipid peroxidation and protein carbonyl expression in the testes. Moreover the oxidative stress associated with the diabetic condition is associated with DNA damage in the male germ line and high rates of embryonic loss in mated females (dominant lethal effect). These effects could be attenuated by the administration of antioxidants such as ascorbic acid, melatonin, taurine or an herbal mixture containing extracts from Musa paradisiaca, Tamarindus indica, Eugenia jambolana and Coccinia indica. In light of recent data showing an increased level of DNA damage in the spermatozoa of diabetic patients compared with non-diabetic controls, causative links between diabetes, oxidative stress in the male germ line and DNA damage appears both likely and clinically, extremely important.
Infection

Another factor that may cause oxidative stress in the testes is infection. Experimental models of infection, involving the intraperitoneal injection of bacterial lipopolysaccharide (LPS), induced lipid peroxidation in the testes and rapidly depleted this tissue of antioxidant enzyme activity in the form of SOD, catalase and the glutathione peroxidase-reductase couple. This oxidative stress was associated with the transient generation of pro-inflammatory mediators such as interleukin 1β, inducible nitric oxide synthase and cyclo-oxygenase-2. The same experimental infection model has also been used to demonstrate the particular sensitivity of Leydig cell steroidogenesis to oxidative stress induced by bacterial LPS. In these studies, the oxidative stress induced by LPS stimulated lipid peroxidation in Leydig cell membranes as well as significant reductions in steroidogenic acute regulatory protein (StAR) and 3β-hydroxysteroid dehydrogenase isomerase (3β-HSD) activity. Moreover, these effects were associated with the disruption of Leydig cell mitochondrial function and, specifically, the inhibition of StAR-mediated cholesterol transfer activity.

Physical Exertion

Physical exercise has been shown to up-regulate antioxidant activities in the testes of aging rats and may represent a practical way in which the detrimental effects of age on testicular function can be ameliorated. A similar case could be argued for the ability of moderate exercise to ameliorate the degree of oxidative damage inflicted on the testes by chronic ethanol ingestion. However, excess exercise can have the opposite effect, causing oxidative stress in the testes and generating high levels of lipid peroxidation in association with significant declines in the activities of key antioxidant enzymes including SOD, catalase, GST and GPx. Such stress has a significant inhibitory effect on both steroidogenesis and germ cell differentiation within the testes. The fact that these effects can be reversed by the administration of an antioxidant, α-tocopherol succinate, confirms the importance of oxidative stress in the aetiology of such exercise-dependent testicular dysfunction.

Reproductive Hormone Imbalance

The immediate endocrine environment of the testes has a major impact on the antioxidant status of this organ. Treatments including exposure to cyclophosphamide or dimethane sulfonate that diminish the intratesticular concentration of testosterone, inhibit the testicular expression of antioxidant enzymes such as GPx, SOD and catalase. Furthermore, these suppressive effects on antioxidant expression, as well as the disruption of spermatogenesis, can be reversed by the administration of exogenous gonadotrophin to artificially elevate intratesticular testosterone levels. Suppression of intratesticular testosterone with exogenous steroids, including both androgens and estrogens, similarly results in the suppression of antioxidant enzyme expression, a concomitant increase in peroxidative damage, the disruption of spermatogenesis and an increase in germ cell apoptosis. Intriguingly, the suppression of antioxidant activity in response to exogenous steroid treatment largely affects the Leydig cells that contain most of the catalase and GPx activities. Testicular SOD activities that are largely confined to the seminiferous tubules did not change dramatically under these circumstances. It is therefore possible that the site of free radical generation in response to gonadotrophin withdrawal involves electron leakage from the inhibited steroidogenic pathway of the Leydig cells. These free radicals then attack the germ cells within the seminiferous tubules leading to extensive apoptosis and the disruption of spermatogenesis. The fact that aminoglutethimide, an inhibitor of the P450 cholesterol side-chain cleavage, induces extensive lipid peroxidation in the testis supports this contention. Interestingly, over-stimulation of the Leydig cells by chronic exposure to hCG (100 IU/day for 30 days in rats) also stimulates high levels of ROS production from these cells, that in turn stimulate lipid peroxidation, reduction in antioxidant enzyme activities, germ cell apoptosis and the consequential disruption of spermatogenesis. Thus, as we saw with the involvement of the thyroid gland in the control of testicular function,
Table 1. Impact of antioxidants on testicular damage caused by testicular torsion/detorsion

<table>
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<tr>
<th>Antioxidant</th>
<th>Outcome</th>
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<th>Antioxidant</th>
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<td>Pentoxifylline</td>
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<td>Germ cell</td>
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<td>histopathology</td>
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<td></td>
<td>apoptosis</td>
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<tr>
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<td>Erdosteine</td>
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<td>Propofol</td>
<td>histopathology</td>
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</table>

MDA = malondialdehyde; XO = xanthine oxidase; GPx = glutathione peroxidase; MPO = myeloperoxidase; iNOS = inducible nitric oxide synthase.

a stable redox environment depends on an appropriately balanced gonadotrophic support, either hyper- or hypo- gonadotrophism will induce a state of oxidative stress in the testis.

Retinoids

While fluctuations in Leydig cell steroidogenesis may be one source of free radical generation in the testes, another is the Sertoli cell population. The latter has been shown to generate ROS following stimulation with all-trans-retinoic acid (RA), a vital cofactor for spermatogenesis. Exposure of rat Sertoli cells to RA led to activation of ROS generation, lipid peroxidation and, ultimately, a loss of cell viability. There is also some evidence to suggest that retinol might stimulate ROS generation in rat Sertoli cells and that this effect is accompanied by an up-regulation of testicular antioxidant enzymes including SOD, GPx and catalase. There may be nothing particularly specific about this effect since retinoids have been shown to stimulate ROS generation in a variety of other cellular systems. Nevertheless, the free radical generation triggered by retinoids in the testes may explain the testicular degeneration induced by hypervitaminosis A in the rat and the association between excess beta carotene intake and infertility in human males.

Impact of Xenobiotics

A wide variety of different xenobiotics have also been shown to induce oxidative stress in the testes in concert with the suppression of antioxidant mechanisms. A summary of these testicular toxicants is provided in Table 2. Heavy paternal smoking, for example, is known to
generate oxidative DNA damage in the male germ line in association with a 32% reduction in the α-tocopherol content of the seminal plasma. The role of oxidative stress in the genesis of this DNA damage is supported by the observation that in individuals subjected to an ascorbate depleted diet, the seminal plasma ascorbate levels decreased by a half, while DNA damage levels in the spermatozoa increased by 91%. Repletion of the ascorbate levels in the diet had the reverse effect and decreased DNA damage by 36%. Experimental exposure of rats to cigarette smoke also induces lipid peroxidation in the testes in association with disturbances in testicular antioxidant enzyme activity. The testicular damage induced by cigarette smoke exposure in rats is certainly oxidative in nature because it can be reversed by concomitant exposure to an antioxidant (caffeic acid phenethyl ester). In addition to smoking, excessive alcohol consumption also has a negative effect of testicular function through the induction of oxidative stress and the concomitant disruption of testicular antioxidant status. Furthermore, the ability of antioxidants such as vitamin C or lecithin to ameliorate this pathology, confirms the importance of oxidative stress in this context. In addition to inducing low sperm counts and poor sperm motility, it also appears that the oxidative stress created in the Leydig cells as a consequence of chronic alcohol exposure diminishes the steroidogenic capacity of the testes, lowering circulating testosterone levels.

Table 2 also highlights a number of metals that are known to induce oxidative stress in the testes and compromise male infertility. Chromium, for example, is a testicular toxicant that stimulates lipid peroxidation and suppresses antioxidant enzyme activities as well as ascorbate levels in the testes. Additional studies in monkeys have also shown that chromium administration decreases not only inhibit the classical array of antioxidant enzymes in the testes but also diminishes the testicular content of GSH as well as vitamins A,E and C, while H₂O₂ production and hydroxyl radical formation are increased. Additional transition metals such as iron also induce lipid peroxidation, protein carbonyl expression and lipid soluble antioxidant depletion in testicular tissue with the consequent disruption of spermatogenesis. Significantly, iron intoxication of male mice also induces a dominant lethal
effect characterized by high levels of embryonic loss in females mated to iron-exposed males. In this situation, the oxidative stress induced in the testes by acute iron overload must have so damaged the DNA in the spermatozoa that the resulting embryos were non-viable.

Heavy metals such as lead, cadmium and uranium have a similar effect on the testes disrupting spermatogenesis via mechanisms that involved the induction of lipid peroxidation, depletion of ROS scavengers and disruption of testicular antioxidant enzyme activity. Arsenic has also been shown to induce peroxidative damage in the testes elevating protein carbonyl expression and decreasing tissue GSH content and inhibiting $3\beta$- and $17\beta$- hydroxysteroid dehydrogenase activities. The importance of oxidative stress in the testicular toxicity associated with arsenic was emphasised by the ability of ascorbate to reverse these changes. Similarly, vanadate is a testicular toxicant that induces lipid peroxidation in the testes along with significant suppression of testicular SOD and catalase and the disruption of $3\beta$- and $17\beta$- hydroxysteroid dehydrogenase activities.

In addition to the above, Table 2 lists a wide variety of different industrial and environmental toxicants that are all capable of compromising male fertility by inducing a state of oxidative stress in the testes. These compounds include phthalate esters, sulfur dioxide, sodium fluoride, a range of environmental estrogens (e.g., PCBs, methoxychlor, bisphenol A, nonylphenol, chemotherapeutic agents (e.g., adriamycin, cisplatin, cyclophosphamide, hexachlorocyclohexane, 2,4,6-trinitrotoluene, aflatoxin, quinalphos, endosulfan, diethyl maleate, monensin, formaldehyde, alloxan, streptozotocin, acrylamide, and ozone. In addition to this list of xenobiotic chemicals that can induce oxidative stress in the testes, physical factors such as static magnetic fields and electromagnetic radiation in its various forms from heat to X-ray irradiation, can also trigger a state of oxidative stress in testicular tissue. Given the variety and prevalence of chemical and physical factors that can generate oxidative stress in the male gonad, there is an urgent need to identify antioxidants that can supplement the tissue’s own antioxidant strategies to rescue the testes from the consequences of ROS attack.

**Antioxidant Therapy**

In order to determine the relative potential of different antioxidants to address oxidative stress in the testes, the testicular torsion-detorsion model has been repeatedly used. Typically this model involves the application of antioxidant therapy prior to the creation of a brief period of oxidative stress and subsequent comparison of various testicular attributes (lipid peroxidation, histopathology, DNA damage or antioxidant enzyme status) with sham operated controls (Table 1). Such analyses have recorded significant protection against oxidative stress for factors as garlic extract, caffeic acid phenethyl ester (CAPE), N-acetyl cysteine, pentoxifylline, erdostein, resveratrol, dexpanthenol, L-carnitine, and propofol anaesthetic.

A variety of antioxidants have also been assessed for their ability to counteract oxidative stress in the testes created by alternative mechanisms. For example: (1) CAPE has been shown to protect the testes from the oxidative stress created by exposure to dizocilpine (MK-801), a drug that is commonly used to induce schizophrenia, and cigarette smoke (2) lycopene, the red plant antioxidant that is a major constituent of tomatoes, is capable of reversing the oxidative damage induced in rat testes following exposure to cyclosporin A or cisplatin (3) extracts from the herb *Lycium barbarum*, effectively protect the testes from the oxidative damage induced by heat stress and significantly suppress the oxidative DNA damage induced in mouse testicular cells by *H₂O₂* (4) MTEC (an aqueous-methanol extract of *Musa paradisiaca*, *Tamarindus indica*, *Eugenia jambolana* and *Coccinia indica*) protects against the oxidative testicular damage resulting from induced diabetes (5) lecithin administration protects the testes from the oxidative stress induced by chronic ethanol exposure (6) lipoic acid has been shown to inhibit the oxidative damage resulting from exposure to cyclophosphamide, adriamycin and X-irradiation (7) complex feruloyl oligosaccharides released from wheat bran have also been shown to protect the testes from the oxidative stress associated with alloxan-induced diabetes in rats (8) $\beta$-carotene
ameliates cadmium induced oxidative stress in the testes, suppressing lipid peroxidation and restoring SOD, GST and GSH to normal physiological levels. 166

One of the most effective antioxidants for the protection of testicular function is melatonin. This evolutionarily conserved compound has been shown to reduce oxidative stress in the testes induced by ethanol, 167 indomethacin, 168 X- irradiation, 169 streptozotocin-induced diabetes, 86 and cisplatin. 136 In vitro studies have also shown that melatonin and its immediate precursor N-acetyl-serotonin could inhibit ascorbate-Fe (II) induced lipid peroxidation in rat testicular microsomes and mitochondria. 170,171

The administration of antioxidants such as resveratrol, ascorbate or cocoa rich in flavanols to normal animals, not suffering from induced oxidative stress, also appears to improve testicular function, suggesting that oxidative stress is a consistent feature of testicular physiology. 172,173 In light of such results, antioxidants have frequently been administered to infertile men in the hope of improving the quality of the semen profile. Very few properly controlled double blind crossover trials have been conducted in this context. However where these conditions have been met, the results have been extremely promising. 174,175

Conclusions

Oxidative stress is a major factor in the aetiology of male infertility. At the level of the isolated spermatozoon, ROS attack can induce lipid peroxidation and DNA fragmentation disrupting both the motility of these cells and their ability to support normal embryonic development. 176-182 At the level of the testes, oxidative stress is capable of disrupting the steroidogenic capacity of Leydig cells 183 as well as the capacity of the germinal epithelium to differentiate normal spermatozoa. 184 A large number of independent clinical studies have demonstrated a correlative relationship between male infertility and evidence of oxidative stress in the ejaculate. 180,185 Moreover the literature reviewed in this chapter reveals an abundance of experimental data in animal models demonstrating a causal relationship between the induction of oxidative stress in the testes and the impairment of male reproductive function. However these two lines of evidence have not yet come together. Although oxidative stress is clearly a dominant feature in the aetiology of male infertility, the underlying causative mechanisms remain unresolved. The plethora of physical, chemical, and pathological factors that can apparently contribute to the induction of oxidative stress in the testes is impressive and suggests that the clinical picture will be extremely complex, with each individual being subject to a unique range of causative factors as a result of differences in occupational and environmental exposures, the presence of other pathological factors such as infection or diabetes, and genetic factors that could influence everything from the way in which specific xenobiotics are metabolised to the endocrine environment in which the testes have to function.

That there are so many factors capable of inducing oxidative stress in the testes strongly suggests that this is a vulnerable tissue that is both highly dependent on oxygen to drive spermatogenesis and yet highly susceptible to the toxic effects of reactive oxygen metabolites; in this context, the testis is very like the brain. While the testes clearly do possess highly specialized antioxidant defence enzymes such as extracellular SOD, PHGPx etc, there are clear benefits to be gained by treating susceptible individuals with exogenous antioxidants. Despite the evident clinical market for an antioxidant preparation specifically designed to support male reproductive health, it is remarkable how little effort has gone into the development of such a preparation and how poor most of the clinical trials in this area have been. In animal models an impressive range of antioxidant preparations has been examined and compounds identified that are clearly capable of crossing the blood testes barrier and protecting the germinal epithelium and Leydig cells from oxidative stress. The future imperatives for this area are to go beyond the superficial phenomenology that characterizes most of the clinical research in this area in an attempt to (i) gain insights into the underlying causes of oxidative stress in the male reproductive tract and (ii) develop optimized antioxidant preparations to treat pathologies arising from an imbalance in the redox status of these tissues. The journey will be long and difficult.
but ultimately more rewarding than the empirical approach that characterizes the current approach to treating the infertile male.

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Nitric Oxide and Cyclic Nucleotides: 
Their Roles in Junction Dynamics and Spermatogenesis

Nikki P.Y. Lee* and C. Yan Cheng

Abstract

Spermatogenesis is a highly complicated process in which functional spermatozoa (haploid, 1n) are generated from primitive mitotic spermatogonia (diploid, 2n). This process involves the differentiation and transformation of several types of germ cells as spermatocytes and spermatids undergo meiosis and differentiation. Due to its sophistication and complexity, testis possesses intrinsic mechanisms to modulate and regulate different stages of germ cell development under the intimate and indirect cooperation with Sertoli and Leydig cells, respectively. Furthermore, developing germ cells must translocate from the basal to the apical (adluminal) compartment of the seminiferous epithelium. Thus, extensive junction restructuring must occur to assist germ cell movement. Within the seminiferous tubules, three principal types of junctions are found namely anchoring junctions, tight junctions, and gap junctions. Other less studied junctions are desmosome-like junctions and hemidesmosome junctions. With these varieties of junction types, testes are using different regulators to monitor junction turnover. Among the uncountable junction modulators, nitric oxide (NO) is a prominent candidate due to its versatility and extensive downstream network. NO is synthesized by nitric oxide synthase (NOS). Three traditional NOS, specified as endothelial NOS (eNOS), inducible NOS (iNOS), and neuronal NOS (nNOS), and one testis-specific nNOS (TnNOS) are found in the testis. For these, eNOS and iNOS were recently shown to have putative junction regulation properties. More important, these two NOSs likely rely on the downstream soluble guanylyl cyclase/cGMP/protein kinase G signaling pathway to regulate the structural components at the tight junctions and adherens junctions in the testes. Apart from the involvement in junction regulation, NOS/NO also participates in controlling the levels of cytokines and hormones in the testes. On the other hand, NO is playing a unique role in modulating germ cell viability and development, and indirectly acting on some aspects of male infertility and testicular pathological conditions. Thus, NOS/NO bears an irreplaceable role in maintaining the homeostasis of the microenvironment in the seminiferous epithelium via its different downstream signaling pathways.

Introduction

Among the organs in the mammalian body, testis is one of the exceptional organs having complex cellular structures and organization. After puberty, testis functions as a sperm producing factory, generating up to millions of spermatozoa on a daily basis through the entire adulthood. In order to fulfill its reproductive function, testis is compartmentalized into two broad partitions, the seminiferous tubules and the inter-tubular areas (Fig. 1). In the seminiferous tubules, the
Nitric Oxide and Cyclic Nucleotides

The epithelium is physically divided into the adluminal compartment and basal compartment by the blood-testis barrier (BTB) which is constituted by adjacent Sertoli cells near the basement membrane. Different cell types situate in specialized testicular locations. Sertoli cells and assorted germ cell types namely spermatogonia, spermatocytes, and spermatids are found in the seminiferous epithelium. Myoid cells locate adjacent to the tubules and Leydig cells reside in the inter-tubular space known as the interstitium. Each cell type performs different function, however they are communicating with each other to share the core role in sperm production during spermatogenesis. Apart from that, the male sex hormone level namely testosterone in the systemic circulation is also produced and regulated by the Leydig cells in the testis via steroidogenesis. These processes cannot be fully executed, if they are not equipped with precisely regulated interactive mechanisms.
during spermatogenesis. In rodents, the germ-line lineage spermatogonia, initially residing on the basement membrane of the seminiferous epithelium must differentiate into preleptotene spermatocytes, which in turn, traverse the BTB at stages VII-VIII of the epithelial cycle to gain entry into the adluminal compartment for further development. In rodents, the germ-line lineage spermatogonia, initially residing on the basement membrane of the seminiferous epithelium must differentiate into preleptotene spermatocytes, which in turn, traverse the BTB at stages VII-VIII of the epithelial cycle to gain entry into the adluminal compartment for further development. During this event of germ cell movement, Sertoli cells also play a paramount role in determining the molecular events of germ cell development, including mitosis, meiosis, cellular differentiation and transformation. Sertoli cells accomplish this in part by monitoring the assembly and disassembly of inter-Sertoli junctions in the testes and partly by initiating cross-talk with germ cells. For instance, Sertoli cells are equipped with certain architectural machineries, such as microtubules, that interact with the movement-associated germ cell proteins (e.g., motor proteins) to provide this coordination. As such, premature germ cell release from the epithelium will be prohibitive. To sustain the optimal germ cell population in the testis, more than half (~75%) of the germ cells that are produced including spermatogonia, spermatocytes and spermatids undergo apoptosis, and are phagocytosed by Sertoli cells, thereby restricting the numbers of germ cells in the seminiferous epithelium. This spontaneous removal mechanism ensures that the limited resources from Sertoli cells (note: the number of Sertoli cells in adult rats remain the same throughout adulthood) are sufficient for germ cell development, and are within the capacity of the testes, by eliminating excessive germ cells to maintain germ cell quality.

Nitric oxide (NO) is a free radical synthesized by nitric oxide synthase (NOS). NOS is composed of two identical monomers with molecular weights ranging from 130 to 160 kDa. Three isoforms of NOS, NOS I, II, and III, are known to date, which are the alternate names for neuronal NOS (nNOS) (Mr, ~320 kDa), inducible NOS (iNOS) (Mr, ~260 kDa), and endothelial NOS (eNOS) (Mr, ~270 kDa), respectively. On the other hand, these NOS isoforms are functionally categorized into two groups, based on their intrinsic NO production efficacy. The constitutive group of NOS includes nNOS and eNOS, whereas iNOS belongs to the inducible form. Despite of these differences, all of them execute the same enzymatic reaction by converting L-arginine into NO and L-citrulline, using the cosubstrates of O2 and nicotinamide adenine dinucleotide phosphate (NADPH). In addition, tetrahydrobiopterin (BH4), flavin adenine dinucleotide (FAD), flavin mononucleotide (FMN), heme, calcium ions, and zinc ions are other necessary cofactors of this reaction. nNOS and eNOS usually synthesize NO in nanomole range, unlike the micromole range of NO generated by iNOS. Excessive level of NO, at >1 μM, has a direct detrimental effect on the physiological system, via the production of peroxynitrite, after interacting with oxygen or superoxide. In contrast, low concentration of NO, at <1 μM, works as an upstream regulatory molecule, goading the downstream signaling proteins, such as soluble guanylyl cyclase (sGC) that sequentially produces cGMP and activates protein kinase G (PKG). An inimitable form of nNOS is being documented as testis-specific nNOS (TnNOS). In short, the multifarious effects exhibited by NO are paramount in the general body metabolism. With the identification of a unique isoform of NOS, TnNOS, in the testis, an interest to examine the role of NOS and NO in spermatogenesis and steroidogenesis is expanding. As a complement to the earlier data regarding to the localization studies of NOS/NO and their related proteins and molecules, the functions of NOS/NO in spermatogenesis and steroidogenesis are being unfolded due to the establishment of several in vivo models or in clinical situations. Results derived from these studies advance the knowledge of NOS/NO in the testes and open the gate for the diagnosis and treatment of male reproductive dysfunctions.

NOS/NO in Junction Dynamics

Germ cells undergo various stages of differentiation during spermatogenesis, and these cells must also migrate from the basal compartment to adluminal compartment during maturation in the testis. To facilitate the course of germ cell movement, coordinated junction restructuring takes place between Sertoli and germ cells, resulting in the luminal release of spermatozoa. This phenomenon seems to be simple superficially; however, the mechanism(s) underneath is perhaps one
of the most complicated procedures in the mammalian body. Unlike other endothelia and epithelia with assorted junction zones, seminiferous epithelium contains hybrid junctions instead.\textsuperscript{18-20} The most imperative exposition of this heterogeneity is the BTB, which encompasses a minimal of three junction types: the tight junction (TJ), the gap junction (GJ), and the adherens junction (AJ).\textsuperscript{20} As such, the passage of preleptotene spermatocytes through the BTB depends on the dynamic changes of the junction structures at the BTB. In the testes, junctions are broadly classified into three types, the anchoring junctions, the TJ, and the GJ. The former junction is further divided into two sub-groups, the actin cytoskeleton-based AJ, known alternatively as the ectoplasmic specialization (ES),\textsuperscript{3,9} and the intermediate filament-based. Fundamental structure of junction complex consists of integral membrane or transmembrane proteins, which are indirectly linked to the underlying cytoskeletons via adaptors. In addition, peripheral regulatory proteins, such as kinases or phosphatases, are also found in the vicinity or in direct contact with these junction complexes. Thus, controlling the spatial and physical distributions of these junctions and their status are one of the important tasks for the successful generation of spermatozoa in a timely manner. Among numerous signaling molecules, NOS/NO is one of the important regulators of the junction integrity in the seminiferous epithelium.\textsuperscript{21} The roles of NOS/NO in the junction dynamics are also implicated in other endothelia and epithelia. Below are discussions of the pertinent studies of these structural components in the testes and how NOS/NO alters the junction stability and integrity by affecting integral membrane proteins, adaptors, cytoskeletons, and regulatory proteins.

**Integral Membrane Proteins**

Three types of transmembrane proteins in the testis have been extensively studied in recent years. First, occludins, claudins, and junctional adhesion molecules (JAM) are restricted to TJ at the BTB.\textsuperscript{4,21} Second, connexins are the known GJ proteins in the testes, with their distributions from the basal to the adluminal compartment in the seminiferous epithelium.\textsuperscript{20,22} Third, different assortment of proteins in anchoring junctions.\textsuperscript{9} Anchoring junctions are further divided according to their cytoplasmic associated cytoskeletons that serve as their attachment sites. Cadherins are the prominent proteins located between cells at the sites of actin-based AJ, such as basal ES at the BTB;\textsuperscript{9} whereas integrins are found between cells and extracellular matrix at the sites of focal adhesion as well as at the Sertoli-germ cell interface at the apical ES. The structures and the components of these testicular junctions are resemblance of other epithelia in mammals. However, their relative locations are unique in the testes. For instance, ES is a testis-specific AJ type, but it is also a hybrid anchoring junction having the properties of both AJ and focal adhesion.\textsuperscript{18,19} The physiological basis of these unusual features are currently unknown but recent studies have illustrated that there are cross-talk between different junction types in the seminiferous epithelium, which is mediated by the associated adaptors and/or the underlying cytoskeletons.\textsuperscript{15,20}

**Adaptors**

Adaptors are proteins that originally thought to have a restricted function by maintaining junction integrity in which they link the transmembrane proteins to the underlying cytoskeletons, such as actin, microtubule, or intermediate filament. Subsequent studies in recent years have shown that adaptors are functionally much more diversified than initially conceived, such as by recruiting signaling molecules (e.g., kinases, phosphatases) to the junction site and by mediating cross-talk between different junctional protein complexes in the testes.\textsuperscript{19} These expanded functionalities of adaptors are not entirely unexpected in view of their widespread occurrence in different junctional complexes. Amidst all the currently known adaptors, ZO-1 is known for its ability to conjoin occludin and connexin and it is found in both TJ and GJ.\textsuperscript{19} As such, one junction type (e.g., TJ) can impose indirect influence to other junctions (e.g., GJ) via a common adaptor (e.g., ZO-1). This view has been validated in at least two other studies in the testis. When rats were treated with a blocking connexin peptide, disruption of connexin functions using
pan-connexin peptide renders the disruption of occludin-based TJ, but not N-cadherin-based AJ in the testes. This proves the vulnerability of the occludin-associated complex following a dysfunction of connexin-based GJ, since pan-connexin peptide with sequence conserved among all connexins virtually blocks all connexin-associated functions in the testes. In addition, the structural association of γ-catenin, an AJ adaptor, and ZO-1, a TJ adaptor, is weakened after treatment of rats using adjudin, which is shown to have a role in disturbing the AJ without compromising the TJ in the testes. These findings substantiate the significance of adaptors that mediate cross-talk between different junction types in the testes. Apart from that, other functions are recently uncovered for adaptors, including their participation in cytokine signaling in the testes and immune-related activities. Based on this emerging evidence, adaptors are proteins with diversified physiological functions in maintaining spermatogenesis, in particular the events of junction restructuring.

Cytoskeletons

Actin filaments, intermediate filaments, and microtubules are the three cytoskeletons found in mammalian cells including testes. Actin filaments are assembled by the polymerization of the actin monomers, whereas microtubules are largely composed of α- and β-tubulins, which form the heterodimers as the basic constituents. In spite of the identification of at least five categories of intermediate filament elements in the testes, vimentin is one of the most studied intermediate filament components in the testes. Recently, the physiological roles ascribed to cytoskeletons are rapidly expanding. In addition to their roles that provide the cellular structural support and cell motility, they are proven to participate in other signaling mechanisms, such as in the transcriptional regulation of genes in the nucleus. Remarkably, nuclear actin and myosin are among two of the vital elements. Other functional activities include the organization of chromosomes, and their allocation and segregation during mitosis. In the seminiferous epithelium, actin is intimately involved in the development of acrosomes. On the other hand, tubulin-based microtubules are known to serve as the track, which works in concert with motor proteins to direct the trans-epithelial movement of spermatids from basal to the apical compartment or vice versa. Cytoskeletons have specific locations within the seminiferous epithelium. Due to the multiplexing nature of the junctions in the testes, the underlying junctions are intimately lying in the proximity of each other or even exhibiting an overlapping position. This further implicates the possible interaction among different cytoskeletons. Studies in other epithelia have illustrated that cytoskeletons mediate cross-talk between junctions. Interestingly, there are accumulating evidence regarding the signal transduction properties of different cytoskeletal elements, such as the intermediate filament. Due to the coherent complex nature of junctions in the testes, much work is needed in this area to decipher the functional roles of different cytoskeletons and their interactions during spermatogenesis.

Studies of NOS/NO in the Testes

Many proteins and/or molecules are known modulators of the junction integrity and functionality in the testes including NO. NO is highly efficient in these processes due to its small molecular size and diffusible nature, making its sites of action distance away from its production sites. The major source of NO in the testes derives from macrophages found in the microvessels and in the inter-tubular compartment in the interstitium, making Leydig cells as one of the most immediate responders to NO (Fig. 1). This effect inevitably alters the metabolic activities in Leydig cells, modulating their testosterone output during steroidogenesis (see below). Indeed, testicular macrophages are prominent cellular regulators in Leydig cell physiology, due to their proximal location with Leydig cells and that they are also the cellular source of growth factors and/or cytokines. Apart from that, NO also influences the activity of Sertoli and germ cells within the seminiferous tubules. However, the effects exhibited are not as remarkable as those observed in Leydig cells due to the intrinsic physical distance between the interstitium and the seminiferous epithelium. However, NO released from the different forms of NOS found in the
Nitric Oxide and Cyclic Nucleotides

Seminiferous epithelium is the major regulator of the spermatogenetic process. Several NOSs are found in different testicular cells. Three major forms of NOS and their defined locations in the testes are known to date (Table 1 and Fig. 1). Germ cells express eNOS, whereas Leydig cells produce iNOS abundantly. On the other hand, minor amounts of nNOS are produced by Sertoli cells. In spite of the presence of these classic NOSs, a testis-specific NOS variant has been identified and entitled as the truncated form of nNOS (TnNOS). Its expression is limited to Leydig cells, strongly implicating its role in steroidogenesis. This testicular forms of NOS seems to be an appropriate subject of an intensive investigation, however, its functions are not fully deciphered due to the lack of knockout mouse model.

In order to fully appreciate the physiological effects of NOS/NO, recent studies have extended to their effectors, which are the executioners of the NOS/NO signaling pathway. The components of the sGC/cGMP/PKG pathway are expressed in a cyclic manner during the seminiferous epithelial cycle in the testes (Table 1 and Fig. 1). Spatially, iNOS is depicted to exist in coherent locations with their associated proteins, such as sGC and PKG., as demonstrated by immunohistochemistry and immunofluorescent microscopy. These observations emphasize the close association of components of NOS/NO pathways in the seminiferous epithelium. A panel of junction proteins has been shown to be putative NOS-binding proteins using the techniques of coimmunoprecipitation and immunoblottings. For instance,

Table 1. Cellular localization of NOS signaling pathway components and their putative associated proteins in mammalian testes

<table>
<thead>
<tr>
<th>Proteins</th>
<th>SC</th>
<th>GC</th>
<th>AC</th>
<th>LC</th>
<th>MC</th>
<th>MF</th>
<th>EC</th>
<th>SP</th>
<th>Putative Associated Proteins†</th>
</tr>
</thead>
<tbody>
<tr>
<td>CNG 1</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>n.k.</td>
</tr>
<tr>
<td>eNOS</td>
<td>+</td>
<td>+</td>
<td>/</td>
<td>+</td>
<td>/</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>actin, β-catenin, eNOS, iNOS, N-cadherin, occludin, sGC, α-tubulin, vimentin</td>
</tr>
<tr>
<td>iNOS</td>
<td>+</td>
<td>+</td>
<td>/</td>
<td>+</td>
<td>+</td>
<td>/</td>
<td>-</td>
<td>+</td>
<td>actin, β-catenin, eNOS, iNOS, N-cadherin, occludin, sGC, α-tubulin, vimentin</td>
</tr>
<tr>
<td>nNOS</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>/</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>sGC, cGMP</td>
</tr>
<tr>
<td>PKG</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>β-catenin, sGC</td>
</tr>
<tr>
<td>sGC</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>TnNOS</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>n.k.</td>
</tr>
</tbody>
</table>

*Cellular localization was revealed by RT-PCR, IB, IHC, and/or IF. †Putative associated proteins were assessed by co-IP, IHC, and/or IF. This table was prepared based on the following research articles and reviews. Abbreviations used: +, presence; -, absence; /, not positively identified; AC, acrosome; CNG 1, cyclic nucleotide-gated channel 1; co-IP, coimmunoprecipitation; EC, endothelial cells in blood vessels; eNOS, endothelial NOS; IB, immunoblot; IF, immunofluorescent microscopy; IHC, immunohistochemistry; GC, germ cells; iNOS, inducible NOS; JAM, junctional adhesion molecule; LC, Leydig cells; MC, myoid cells; MF, myofibroblasts; n.k., not known; nNOS, neuronal NOS; NOS, nitric oxide synthase; PKG, protein kinase G; SC, Sertoli cells; sGC, soluble guanylyl cyclase; RT-PCR, reverse transcription polymerase chain reaction; SP, spermatozoa; TnNOS, testis-specific nNOS; ZO-1, zonula occludens-1.
iNOS and eNOS were shown to associate with occludin-based TJ complexes in the testes.\textsuperscript{44} Besides, these two NOS isoforms also link to the N-cadherin-based junction network.\textsuperscript{42} Additionally, sGC, the downstream effector of the NOS pathway, is structurally associated with both AJ and TJ protein complexes in the testes.\textsuperscript{43} In this context, NOS/NO is a putative regulator of junction restructuring in the testes as illustrated in other functional studies. Zinc (II) protoporphyrin-IX, a broad spectrum NOS inhibitor and a sGC inhibitor, was shown to perturb the TJ integrity in Sertoli cell cultures.\textsuperscript{44} This event also accompanied by a down-regulation in the amount of intracellular cGMP and the occludin protein steady-state levels.\textsuperscript{44} iNOS and eNOS were also shown to dissociate from the N-cadherin/β-catenin protein complexes during the Adjudin-induced AJ restructuring in the rat testes that led to germ cell depletion from the epithelium.\textsuperscript{42} Apart from that, the administration of KT-5823, a PKG inhibitor, delayed the actions of Adjudin in mediating germ cell loss from the seminiferous epithelium.\textsuperscript{42} Significantly, a tighter association of sGC with cadherin-based protein complex was observed during Adjudin-induced junction restructuring in vivo, whilst a weakened interaction was found between sGC and TJ protein complexes\textsuperscript{43} (Table 2 and Fig. 2). Based on these findings, the significance of the NOS/NO/sGC/PKG pathway in AJ dynamics in the testes is increasingly clear. In short, the NOS/NO pathway is one of the versatile mechanisms utilized by the testes to regulate junction dynamics.

**Studies of NOS/NO in Other Systems That Can Be the Basis of Future Studies in the Testes**

As described above, NOS/NO is a novel pathway to regulate AJ and TJ dynamics in the testes. However, information regarding the ability of NOS/NO in regulating other junction types, such as gap junctions and other anchoring junctions (e.g., desmosome-like junctions), are lacking. Recent studies have shown that NO regulates connexin 35-associated gap junction coupling in neurons.\textsuperscript{45} However, it remains to be determined if there is any interaction between NO and connexins in the testes. Also, NO has a negative role in modulating integrin-linked kinase, a protein kinase related to the integrin-associated protein complex, in rat kidneys\textsuperscript{46} (Table 2). However, these results still need to be validated in the testes.

**NOS/NO and Spermatogenesis**

**NOS/NO and the Hormone/Cytokine/Paracrine/Autocrine System in the Testis**

Hormones and cytokines are known to have versatile functions ranging from the modulation of endocrine systems to the fine-tuning of immune systems. Within the microenvironment in the testis, testosterone, the major male sex hormone, dominates the processes of spermatogenesis. The synthesis of testosterone by Leydig cells is under the control of luteinizing hormone (LH) released from the pituitary gland. For cytokines, they are mostly derived from the immune cells (e.g., macrophages) in the circulation and diffuse into the testicles via the blood vessels.\textsuperscript{39} Importantly, large part of the NO produced in the testes is derived from activated testicular macrophages, which are having high levels of iNOS.\textsuperscript{47} These exogenous and endogenous sources of hormones and cytokines are involved in constraining the expression of different forms of NOS in the testes, and as such, indirectly affecting the testicular NO steady-state level.\textsuperscript{21}

On the other side, NO coordinates the testicular production of hormones and cytokines. Exposure of Leydig cell cultures to an NO donor, S-nitrosoglutathione can elicit an inhibition of testosterone production in vitro.\textsuperscript{47} Other studies have demonstrated that the activated macrophage-produced NO is associated with a reduction in the testosterone producing activity in Leydig cells.\textsuperscript{48} Notably, this inhibition is at least in part involved with the blockade of the P450 steroidogenic enzymes.\textsuperscript{48}
Table 2. Participation of NOS/NO in the regulation of junctions in testes and other epithelia

<table>
<thead>
<tr>
<th>Organ or Tissue/Junction Type</th>
<th>NOS</th>
<th>Target Junction Proteins</th>
<th>Regulation</th>
<th>Selected References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Testis/TJ</td>
<td>eNOS, iNOS</td>
<td>n.k.</td>
<td>-</td>
<td>44</td>
</tr>
<tr>
<td>Testis/AJ</td>
<td>iNOS</td>
<td>N-Cadherin, β-Catenin</td>
<td>-</td>
<td>42</td>
</tr>
<tr>
<td>Hepatobiliary duct/GJ</td>
<td>iNOS</td>
<td>Occludin, ZO-1, ZO-2, ZO-3</td>
<td>-</td>
<td>76</td>
</tr>
<tr>
<td>Brain/TJ</td>
<td>gNOS</td>
<td>Occludin</td>
<td>+</td>
<td>77</td>
</tr>
<tr>
<td>Vascular/GJ</td>
<td>gNOS</td>
<td>Connexin 37</td>
<td>-</td>
<td>78</td>
</tr>
<tr>
<td>Muscular/GJ</td>
<td>gNOS</td>
<td>Connexin 43</td>
<td>-</td>
<td>79</td>
</tr>
<tr>
<td>Neuronal/GJ</td>
<td>gNOS</td>
<td>n.k.</td>
<td>+</td>
<td>80</td>
</tr>
</tbody>
</table>

This table only contains selected examples of NOS and the role of NO in junctions including the testes and other nongonadal tissues. Other important studies are not cited due to the page limit. Abbreviations and symbols used: +, positive regulation; -, negative regulation; nk, information not known; AJ, adherens junction; eNOS, endothelial NOS; iNOS, inducible NOS; GJ, gap junction; gNOS, studies involved the activation of general NOS activity in the systems; NO, nitric oxide; NOS, nitric oxide synthase; TJ, tight junction; ZO, zonula occludens.

In short, the bi-directional relationship manifested by NOS/NO and the hormone/cytokine level is crucial in maintaining the physiological function of the testes.

**NOS/NO in Germ Cell Development and Differentiation**

**Germ Cell Apoptosis and Germ Cell Output in the Seminiferous Epithelium**

The number of total germ cells produced by the testes is tightly regulated in order to secure the production of viable and fertile germ cells within the supporting capacity of the Sertoli cells in the testes. In the rat, Sertoli cells begin to proliferate at day 16 post-coitus, and at birth, the number of Sertoli cells is about 1 million per testis; and by day 15 post-natal, the number of Sertoli cells rises to ~40 million, however, proliferation ceases to occur thereafter. The Sertoli cell number remains relatively constant throughout adulthood. Thus, this fixed number of Sertoli cells cannot nurture unlimited number of germ cells generated from the primordial spermatogonia. To remove redundant and abnormal perhaps unhealthy germ cells, an elimination mechanism involving germ cell apoptosis is being utilized by the testes. More than half of the germ cells, perhaps ~75%, particular spermatogonia and spermatocytes, undergo spontaneous apoptosis during normal spermatogenesis. Besides estrogens (e.g., 17β-estradiol) that regulate the germ cell number via apoptosis (see below) (for a review, see ref. 51), another mechanism via phagocytosis is also used by the testis. In brief, apoptotic germ cells destined to be phagocytosed by Sertoli cells are recognized via an externalized phosphatidylserine on the apoptotic germ cell surface, which, in turn, coheres the class B scavenger receptor type I (SR-BI) on Sertoli cell surface since Sertoli cells also serve as the scavenger in the epithelium. If the SR-BI on the Sertoli cell surface is inactivated by a monoclonal anti-SR-BI antibody, this causes an increase in residual apoptotic germ cells in the epithelium. Using these mechanisms, the sperm output by the testis can be finely maintained without overwhelming the Sertoli cells.

Notwithstanding, uncontrolled apoptosis would disintegrate the harmonized microenvironment and the Sertoli:germ cell ratios in the seminiferous epithelium during spermatogenesis. As such, these apoptotic events can also be triggered by external stimuli or artificially induced, such
as testosterone deprivation and local testicular heating,\textsuperscript{53} which are detrimental to spermatogenesis. Based on these other models, several germ cell apoptotic pathways have also been identified. At the molecular levels, caspases, especially caspase 3,\textsuperscript{53} and Fas/Fas ligand (Fas L)\textsuperscript{54} are the main regulatory molecules. These pathways are probably inter-connected and mediated by junction proteins, such as connexins, during apoptosis in the testes.\textsuperscript{23} There are reports in the literature illustrating an excessive NO level can directly trigger massive germ cell apoptosis in the testes. These are clearly depicted in the artificial spermatic vessel ligation model\textsuperscript{55} and in the mouse model of congenital cryptorchidism,\textsuperscript{56} both of which were shown to elevate NO testicular levels and uncontrolled germ cell apoptosis. The role of NO in germ cell apoptosis has been further strengthened by the observation that treatment of Hoxa 11 knockout mice having congenital cryptorchidism using L-NAME, an NOS inhibitor, to block the NO-mediated effects can grossly attenuate germ cell apoptosis.\textsuperscript{56} Normal aging process also illustrates the association of NO and apoptosis.\textsuperscript{57} In aged testes, aggravated NOS activity and an induced iNOS level were shown to accompany with high incidence of germ cell apoptosis.\textsuperscript{57} iNOS and eNOS are also postulated to participate in germ cell apoptosis in the testes.\textsuperscript{21} Specifically, eNOS has been known to be highly expressed in degenerating germ cells in comparison with other germ cell types.\textsuperscript{58,59} In addition, iNOS plays a determining role in restricting the germ cell numbers in the testes, since testes from iNOS\textsuperscript{57} mice are heavier on average when compared to normal testes,\textsuperscript{60} and a marked reduction

![Figure 2. The NOS/NO signaling pathways that regulate AJ and TJ dynamics in the testes. This is a simplified schematic diagram summarizing recent studies regarding the role of the NOS/NO signaling pathway that regulates junction dynamics in the testis using the Adjudin model.\textsuperscript{42-44} Abbreviations used: AJ, adherens junction; ES, elongated spermatid; JAM, junctional adhesion molecule; MC, myoid cells; N, Sertoli cell nucleus; NO, nitric oxide; NOS, nitric oxide synthase; PKG, protein kinase G; PS, pachytene spermatocyte; RS, round spermatid; SC, Sertoli cell; sGC, soluble guanylyl cyclase; SPC, preleptotene/leptotene spermatocyte; SPG, spermatogonium; TJ, tight junction; ZO-1, zonula occludens-1.](image)
on germ cell apoptosis, in particular pachytene spermatocytes and round spermatids, is noted.60 These observations thus clearly illustrate the unequivocal role of NO in triggering germ cell apoptosis in the testes. Nonetheless, the downstream NO pathway(s) that regulates germ cell apoptosis is presently not known.

**NOS/NO in Fertility and Pathogenesis**

Male fertility has always been a primary concern in human reproduction. At the molecular level, fertility partly refers to the successful fusion of male and female gametes, which depends on several factors for its completion. The quality and quantity of sperm and sperm concentrations are several of the key issues directly affecting fertilization. As mentioned earlier, apoptosis is one of the key processes that maintains the quality and quantity of germ cells in the testes.61 Thus, if apoptosis fails to control these variables, a lack of viable and fertile germ cells will be the consequence. If testes fail to produce sufficient number of healthy spermatozoa without defects, infertility is unavoidable. Several etiologies are known to goad infertility, such as varicocele, tumors, and azoospermia. Also, DNA integrity in sperm is another major determinant.62 It is known that unbalanced levels of NO or dysregulation of NOS in the testes contribute to some of these defects (Table 3). For instance, a reduced testicular eNOS protein level was observed in patients with idiopathic azoospermia versus patients with obstructive azoospermia or varicocele and healthy individual.63 Besides, excess NO in human varicocele conditions was demonstrated to be harmful to the sperm motility.64 Despite all this, a physiological level of NO is required for successful fertilization.65 Apart from that, NO is also known to participate in sperm transport from the rete testis to the epididymis. The propelling force that transports spermatozoa to the epididymis via the rete testis is generated by the contraction and relaxation of the tunica albuginea in the testis, which, in turn, is regulated by PKG, a downstream effector of NOS/NO.66 In short, NOS/NO has diversified roles in testis pathogenesis.

**Concluding Remarks and Future Perspectives**

NO is a versatile molecule with diversified functions ranging from coordinating cell and blood vessel permeability to junction regulation. It is produced by four isoforms of NOS,
Molecular Mechanisms in Spermatogenesis

namely eNOS, iNOS, nNOS, and TnNOS. NOS/NO is depicted to regulate disparate junction dynamics, at AJ and TJ, in the testes. However, the significance of NOS/NO in the regulation of other junction types, such as gap junctions and desmosomes, are not fully deciphered and much effort is needed to uncover the associated mechanisms and identify the proteins that are involved. Apart from that, NOS/NO is illustrated to have roles in monitoring the levels of hormones and cytokines, indirectly controlling the processes of spermatogenesis and steroidogenesis. Importantly, NOS/NO also takes part in the regulation of germ cell development and differentiation, partially via the coordination of germ cell apoptosis and maintaining the correct ratio of Sertoli:germ cells in the epithelium, securing the efficacy in the production of viable and fertile spermatozoa. As such, NOS/NO is crucial to maintain male fertility and pathogenesis. An expanded research in this area can form the foundation of identifying candidate molecules for male contraception and to understand male reproductive pathogenesis.

Acknowledgements

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The Sertoli Cell Cytoskeleton

A. Wayne Vogl,* Kuljeet S. Vaid and Julian A. Guttman

Abstract

The cytoskeleton of terminally differentiated mammalian Sertoli cells is one of the most elaborate of those that have been described for cells in tissues. Actin filaments, intermediate filaments and microtubules have distinct patterns of distribution that change during the cyclic process of spermatogenesis. Each of the three major cytoskeletal elements is either concentrated at or related in part to intercellular junctions. Actin filaments are concentrated in unique structures termed ectoplasmic specializations that function in intercellular adhesion, and at tubulobulbar complexes that are thought to be involved with junction internalization during sperm release and movement of spermatocytes through basal junctions between neighboring Sertoli cells. Intermediate filaments occur in a perinuclear network which has peripheral extensions to desmosome-like junctions with adjacent cells and to small hemidesmosome-like attachments to the basal lamina. Unlike in most other epithelia where the intermediate filaments are of the keratin type, intermediate filaments in mature Sertoli cells are of the vimentin type. The function of intermediate filaments in Sertoli cells is not entirely clear; however, the pattern of filament distribution and the limited experimental data available are consistent with a role in maintaining tissue integrity when the epithelium is mechanically stressed. Microtubules are abundant in Sertoli cells and are predominantly oriented parallel to the long axis of the cell. Microtubules are involved with maintaining the columnar shape of Sertoli cells, with transporting and positioning organelles in the cytoplasm, and with secreting seminiferous tubule fluid. In addition, microtubule-based transport machinery is coupled to intercellular junctions to translocate and position adjacent spermatids in the epithelium. Although the cytoskeleton of Sertoli cells has structural and functional properties common to cells generally, there are a number of properties that are unique and that appear related to processes fundamental to spermatogenesis and to interfacing somatic cells both with similar neighboring somatic cells and with differentiating cells of the germ cell line.

Introduction

The cytoskeleton in animal cells consists of actin filaments, intermediate filaments and microtubules. These filamentous polymers are associated with a plethora of associated molecules that determine the balance between soluble subunits and polymers, the way in which the polymers are linked to others of the same and different types, and the way in which various molecules, protein complexes and organelles are transported and positioned in cells. The cytoskeleton not only is a determinant of cell shape, but also plays significant roles in cell division, intracellular transport and cell movement, and in establishing and maintaining tissue organization and integrity.

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The Sertoli Cell Cytoskeleton

In Sertoli cells, actin filaments, intermediate filaments and microtubules are abundant and are concentrated in specific regions (Fig. 1). They also are dynamic; that is, their patterns of organization change during spermatogenesis. Actin filaments are shown in red, intermediate filaments are in blue, and microtubules are in green. Actin filaments are concentrated in ectoplasmic specializations and tubulobulbar complexes. Intermediate filaments are concentrated around the nucleus and extend to desmosome-like attachments with adjacent Sertoli and spermatogenic cells and to hemidesmosome-like attachments with the basal lamina. Microtubules are predominantly oriented parallel to the long axis of the cell.

Figure 1. Schematic diagram showing the distribution of major elements of the Sertoli cell cytoskeleton at progressive stages of spermatogenesis in the rat seminiferous epithelium. Actin filaments are shown in red, intermediate filaments are in blue, and microtubules are in green. Actin filaments are concentrated in ectoplasmic specializations and tubulobulbar complexes. Intermediate filaments are concentrated around the nucleus and extend to desmosome-like attachments with adjacent Sertoli and spermatogenic cells and to hemidesmosome-like attachments with the basal lamina. Microtubules are predominantly oriented parallel to the long axis of the cell.

In Sertoli cells, actin filaments, intermediate filaments and microtubules are abundant and are concentrated in specific regions (Fig. 1). They also are dynamic; that is, their patterns of organization change during spermatogenesis. Interestingly, all three elements of the cytoskeleton are related, at least in part, to sites of intercellular attachment and may play significant roles in translocating, positioning, and anchoring spermatogenic cells in the seminiferous epithelium and in establishing and maintaining epithelial organization. This association with intercellular junctions and with specialized structures related to junction turnover indicates that the cytoskeleton also may participate in the mechanism of sperm release and in the process by which spermatocytes move through basal inter-Sertoli junction complexes as these cells move from basal to adluminal compartments of the epithelium.
Actin Filaments

**General**

Actin is the most abundant protein in eukaryotic cells and is associated with such fundamental processes as cytokinesis, cell movement, intracellular transport, cell polarity, and cell/cell and cell/substratum attachment. Actin monomers are globular proteins that under appropriate nucleation conditions polymerize into polar filaments that have a diameter of 5-8 nm. There are more than 60 classes of actin binding proteins that sequester monomers and nucleate, cap, sever, stabilize, cross-link and move along filaments. The actin-dependent motor proteins associated with cargo movement along actin filaments are the myosins.

There are two major models of generating actin filament structures in cells. One involves the Arp2/3 complex that results in three-dimensional networks of branched actin filaments. The other involves the formins and results in long unbranched filaments that can be cross-linked into bundles. The formins and Arp2/3 complex may work cooperatively at certain sites to generate actin structures.

**Actin Filaments in Sertoli Cells**

Actin filaments in Sertoli cells are concentrated in two specific regions: (1) ectoplasmic specializations, and (2) tubulobulbar complexes (Fig. 1). At both sites, the filaments are related to areas of the plasma membrane involved with intercellular attachment. Ectoplasmic specializations are primarily a form of actin-related intercellular adhesion junction. Tubulobulbar complexes are structures proposed to internalize junctions prior to sperm release and movement of spermatocytes through basal junction complexes. At ectoplasmic specializations, actin filaments occur in bundles in which the filaments are unipolar and close-packed into paracrystals (hexagonal arrays). At tubulobulbar complexes, the filaments form three-dimensional networks.

**Ectoplasmic Specializations**

Ectoplasmic specializations are tripartite structures consisting of a layer of actin filaments together with regions of the plasma membrane involved with intercellular attachment on one side of the filament layer and a cistern of endoplasmic reticulum on the other (Fig. 1). The intercellular gap between cells at ectoplasmic specializations narrows to about 90 angstroms.

Ectoplasmic specializations occur at two locations in Sertoli cells (Fig. 2): (1) At heterotypic sites of attachment between Sertoli cells and spermatids in apical regions of the seminiferous epithelium; (2) As part of the homotypic junction complex between neighboring Sertoli cells near the base of the seminiferous epithelium.

At apical sites, ectoplasmic specializations assemble as spermatids polarize and become situated in apical Sertoli cell crypts. Ectoplasmic specializations develop in Sertoli cell regions of the crypts juxtaposed to acrosomal regions of spermatid heads. No structures similar to ectoplasmic specializations develop in the spermatid heads. At certain stages of spermatogenesis, notably stage V in the rat, intermediate filament associated junctions (desmosome-like) appear to intercalate themselves into ectoplasmic specializations in regions adjacent to the dorsal curvature of spermatid heads, and then later disappear. When spermatids are mechanically separated from the seminiferous epithelium, intact ectoplasmic specializations remain attached to the surfaces of the cells, indicating that the three components of the junction (plasma membrane, actin filaments and endoplasmic reticulum of the Sertoli cell) are a structural unit.

At basal sites of attachment between Sertoli cells, ectoplasmic specializations occur in each of the adjacent cells and therefore the junctions are bilateral. Ectoplasmic specializations appear to overlap with and incorporate other junction types such as tight and gap junctions into their structure (Fig. 3). Together with desmosome-like junctions and the other junction types, ectoplasmic specializations form large junction complexes. Tight junctions in these complexes form the “blood-testis” or “blood-Sertoli” barrier.
Most available evidence, from studies where localization clearly occurs both at apical and at basal sites and/or where codistribution with actin filaments has been determined, is consistent with the conclusion that the integral membrane adhesion molecules present at mature ectoplasmic specializations in vivo are mainly α6β1 integrin,24,25 nectin-2,26 and JAMs.27
N-cadherin also is reported present at the sites. Presumably the ligand for N-cadherin is another N-cadherin in the adjacent membrane. The ligand for α6β1 integrin is not entirely clear, although laminin α3 recently has been identified as a candidate at apical sites. The ligand for nectin-2 is nectin-3 in spermatid membranes at apical sites and another nectin-2 on the neighboring Sertoli cell at basal sites. Nectin-2 knockout male mice are infertile and show loss of actin filaments and intercellular adhesion at ectoplasmic specializations—a result that highlights the importance of this type of adhesion junction to spermatogenesis.

In addition to integral membrane adhesion proteins and actin filaments, other elements present at ectoplasmic specializations include the adaptor proteins vinculin and afadin, that facilitate the relationship between integral membrane adhesion elements and the underlying actin cytoskeleton. The catenins also are reported as present. Actin binding proteins concentrated at the sites include α-actinin, fimbrin, espin, Keap 1, and cortactin. Although myosin VIIa is present in the structures, myosin II is not and the actin bundles are not considered to have contractile potential. Nonmuscle coflin is not localized at the sites. Paxillin is not concentrated at the sites but is generally expressed in the cytoplasm of Sertoli cells.
The regulation of ectoplasmic specializations recently has received a great deal of attention because they are thought to contain potential molecular targets for male contraceptives. It is proposed that intercellular attachment at ectoplasmic specializations is regulated directly by TGF-β3 through the Ras/MEK/ERK signaling pathway. Attachment also may be influenced at a higher order level by hormones.

The mechanisms of actin polymerization and depolymerization at ectoplasmic specializations have not been studied in detail, although changes in actin dynamics are suspected to be linked to changes in the status of adhesion elements in the plasma membrane. The unipolar, hexagonally packed and un-branched arrangement of actin filaments in ectoplasmic specializations indicates that predominantly a formin-based mechanism of polymerization rather than an Arp 2/3-based mechanism may be involved. Although not yet localized to ectoplasmic specializations, the formins have been implicated in actin polymerization at adherens junctions generally in epithelia and a model involving activation by Rho GTPases has been proposed. Once filaments are formed at ectoplasmic specializations, espin, fimbrin and α-actinin are likely involved with cross-linking the filaments into bundles and inter-relating the bundles to each other.

It has long been suspected that the endoplasmic reticulum component of ectoplasmic specializations may function to regulate calcium within the micro-environment of the junction plaque and thereby influence junction dynamics, possibly through effects on regulating calcium dependent actin-severing proteins such as gelsolin/scinderin. This class of proteins also binds to phosphatidylinositol 4,5-bisphosphate (PtdIns(4,5)P2) in the plasma membrane and is released when PtdIns(4,5)P2 is converted to inositol (1,4,5)-trisphosphate (Ins(1,4,5)P3) and diacylglycerol by phosphoinositide-specific phospholipase C. The Ins(1,4,5)P3, which is released from the plasma membrane, acts to release calcium from the endoplasmic reticulum, which in turn stimulates the actin severing activity of the free actin-binding proteins. The hypothesis that actin filament dynamics, particularly disassembly, is regulated in part by calcium dependent actin-binding proteins that also bind to PtdIns(4,5)P2 in the plasma membrane is attractive because it ties together the three structural elements of ectoplasmic specializations—the endoplasmic reticulum, the actin layer, and the plasma membrane.

We have reevaluated our original experiments in which we demonstrated the presence of gelsolin at the sites and feel our initial findings may largely have been due to secondary binding to the sites of exogenous gelsolin present both in serum used in our antibody blocking buffers and in serum contaminants in our spermatid/ectoplasmic specialization isolates. Although the calcium dependent actin-binding protein/phosphoinositide hypothesis generally is still viable, we now believe that gelsolin may not be a significant player. This conclusion is consistent with the observation that the reproductive capacity of gelsolin knock-out animals is not impaired. The roles of the endoplasmic reticulum and of calcium-dependent actin-binding proteins at ectoplasmic specializations remains to be clarified.

Function of Actin filaments in Ectoplasmic Specializations

The arrangement of actin bundles in mammalian ectoplasmic specializations, and the hexagonal packing of actin filaments within the bundles, may function structurally to reinforce and stabilize large intercellular adhesion domains in an epithelium where cells progressively change position and shape during spermatogenesis and in areas where the plasma membrane has an irregular contour. This is particularly true at apical sites where Sertoli cells are attached to spermatid heads. The observation that the patterns of actin bundles in ectoplasmic specializations change to conform to the maturing shape of spermatid heads during spermatogenesis is consistent with this function. The loss of adhesion between Sertoli cells and early elongate spermatids and the disruption of the structural and functional integrity of basal junction complexes after intratesticular injection of cytochalasin D are consistent with the argument that actin filaments stabilized adhesion domains in the plasma membrane.
Tubulobulbar Complexes

Tubulobulbar complexes are elaborate structures that develop at apical and basal sites of intercellular attachment in the seminiferous epithelium (Fig. 1). They are most pronounced at apical sites of adhesion between Sertoli cells and spermatids (Figs. 4 and 5) where the formation of multiple generations of these structures precedes sperm release. In this apical location, the structures consist of a number of tubular extensions from the plasma membrane of spermatid heads that protrude into corresponding plasma membrane invaginations of the adjacent Sertoli cell. The structures have swollen or “bulbar” ends that often have small tubular extensions with “coated pits” at their termini. A network of actin filaments cuffs the long tubular invaginations of the Sertoli cell and the bulbar regions are closely related to cisternae of endoplasmic reticulum. The terminal parts of the complexes are associated with numerous vesicles, some of which have been identified as lysosomes. The large bulbar regions of the complexes are thought to “bud” from the structures and be degraded.

A number of functions have been attributed to tubulobulbar complexes. They have been suggested to anchor spermatids to the epithelium based on the observation that they are the last structures to disengage at sperm release. They also are considered to function as a route by which Sertoli cells can eliminate cytoplasm from spermatids. Support for this argument comes from the observation that the development of tubulobulbar complexes correlates with the time during which cytoplasmic volume of spermatids dramatically decreases, and spermatids are much larger when tubulobulbar complexes do not develop in association with them. Another suggested function is that they facilitate the shaping of spermatid...
heads. An additional function, and the one that we favor, is that tubulobulbar complexes internalize integral membrane junction proteins during junction disassembly, a hypothesis initially put forward by Russell and later by Pelletier.

Figure 5. Apical tubulobulbar complexes (rat). A) Electron micrograph of a section through a late spermatid head and associated tubulobulbar complexes (TBCs). Multiple tubulobulbar complexes are shown in cross section. B) Electron micrograph of a longitudinal section through a tubulobulbar complex (TBC). C,D) Paired DIC and fluorescence images of tubulobulbar complexes (arrows) associated with the head of a late spermatid. The filamentous actin in (D) is labeled with fluorescent phalloidin and appears as an elongate cuff around the tubular membrane components of each complex. E) Electron micrograph a cross section through a single tubulobulbar complex. The actin filaments of the complex form a network surrounding the membrane components. This organization of actin filaments into a network is clearly different from the hexagonally packed bundles of filaments seen in ectoplasmic specializations (see Fig. 2D).
Generally in cells integral membrane junction proteins are synthesized in the rough endoplasmic reticulum where they are inserted into the membrane. They generally pass through the Golgi, and then “junction vesicles” are transported to and inserted into the basolateral plasma membrane. Here the proteins are mobile in the plane of the membrane and eventually may become incorporated into junctions. There is recent evidence that the trajectory of junction vesicles from Golgi to plasma membrane may not be as direct as initially thought. E-Cadherin, for example, may pass through a recycling endosome prior to delivery to the plasma membrane.65 During constitutive or induced junction turnover, the junction proteins in the membrane are internalized by endocytosis and may either be degraded or recycled. The emerging mechanism for the internalization of adherens and tight junction components appears to be one of disengagement from ligands on the adjacent cell and endocytosis of the proteins in each adjacent cell as part of either clathrin- or a caveolin-mediated pathways66-70 or by clathrin and caveolin independent mechanisms such as macropinocytosis71 depending on cell context and regulation.72

Interestingly, intact gap junctions, consisting of connexons in the plasma membranes of both adjacent cells, are typically removed from the surface and internalized by one of the cells and targeted to lysosomes.73,74 A similar mechanism recently has been observed for “moving” epithelial cells internalizing intercellular tight junctions.75 In this experimental system (MDCK and mouse Eph4 cells), adjacent junction membranes do not detach from one another. Rather, they are coendocytosed as double membrane vesicles into one of the adjacent cells and targeted for degradation. This type of mechanism also has been observed in colon and ovarian adenocarcinoma cell lines.76,77 We propose that this mechanism may be the normal method of junction internalization by tubulobulbar complexes in the seminiferous epithelium of the testis.

The model we propose is one in which Sertoli cells internalize junction proteins by the “budding-off” of large double membrane vesicles, from tubulobulbar complexes, that then either fuse with lysosomes and are degraded or are selectively recycled to the plasma membrane. In other words, junction proteins in one cell together with the attached junction proteins in the membrane of the adjacent cell are internalized together at tubulobulbar complexes. Although this mechanism would basically be a caveolin- and clathrin-independent pathway, our model also includes the possibility that some sorting by clathrin- and caveolin-mediated pathways of proteins may occur prior to and/or after budding of double membrane vesicles from the complexes, and that some of these junction molecules may eventually be recycled back to the plasma membrane. The observation that there are small vesicles, “coated pits” and multivesicular bodies associated with tubulobulbar complexes is consistent with this possibility.

Pertinent to the junction internalization hypothesis are a number of key observations made by Russell and coworkers57,59,60,78 in their now classic ultrastructural descriptions. First, the structures develop in areas occupied by ectoplasmic specializations or in areas where ectoplasmic specializations are pharmacologically perturbed,57 indicating that there may be a functional relationship between junctions and tubulobulbar complexes. Second, tubulobulbar complexes not only develop at apical junctions between Sertoli cells and spermatids, but also develop at basal junction complexes between adjacent Sertoli cells (Fig. 6), indicating that the primary function of tubulobulbar complexes is not related only to spermatid maturation. Finally, ultrastructurally identifiable gap and tight junctions occur in the bulbous regions of basal tubulobulbar complexes,78 indicating that junction elements do occur in these structures.

We recently have confirmed that tubulobulbar complexes at apical junctions with spermatids develop in regions previously occupied by ectoplasmic specializations. We also have found that double membrane vesicles occur at the ends of the complexes and that lysosomal and endosomal markers are associated with vesicles in similar locations.61 In addition, protein kinase Cα, a signaling molecule known to regulate endocytosis of junction proteins in other systems,66,79,80 is concentrated in regions containing tubulobulbar complexes. Significantly, immunological probes for nectin-2 (Sertoli cell) and nectin-3 (spermatid) react in a vesicular pattern with the ends of tubulobulbar complexes in fluorescence experiments, indicating that the two molecules each from different cells may be internalized together by Sertoli cells.
The junction internalization hypothesis of tubulobulbar function does not rule out the possibility that other processes may be involved in junction turnover, particularly at basal sites where junctions must disassemble above and assemble below spermatocytes as these cells translocate between basal and adluminal compartments of the epithelium (Fig. 7). At basal sites, tubulobulbar complexes are smaller than at apical sites and are much less numerous than might be predicted if they were the sole means for junction turnover. One alternative possibility is that junction components separate from those in the adjacent Sertoli cell and are endocytosed directly from the plasma membrane as occurs in other epithelia, and then are either degraded or recycled. Another is that some junction molecules dissociate from their ligands and move within the plane of the membrane around translocating spermatocytes to reattach to their partner ligands below.
At both basal and apical sites, the contributions from reused (recycled or moved in the plane of the membrane) and of newly synthesized junction proteins to the formation of junction complexes below spermatocytes and to newly formed ectoplasmic specializations at sites of attachment to early elongating spermatids needs to be investigated further.

Figure 7. Shown here are a number of alternatives for what may happen to integral membrane junction proteins during junction turnover in the seminiferous epithelium. Intact junctions may be internalized by tubulobulbar complexes and then either be recycled to newly assembling junctions or be degraded. Alternatively, junction proteins may disengage from their ligands in adjacent membranes and then be internalized by standard endocytic pathways in each cell, and then degraded or recycled. Another possibility is that junction proteins disengage from their ligands, move within the plane of the membrane, and then are recruited into newly assembling junctions elsewhere in the cell. Newly assembling junctions incorporate new protein from synthetic pathways and ‘old’ protein from recycling compartments. Although a basal junction complex between two Sertoli cells is illustrated here, a similar model could be proposed for junctions that disassemble during sperm release and that assemble in association with early elongate spermatids. It is likely that the Sertoli cell cytoskeleton, particularly actin filaments and microtubules, are involved in the proposed pathways.

At both basal and apical sites, the contributions from reused (recycled or moved in the plane of the membrane) and of newly synthesized junction proteins to the formation of junction complexes below spermatocytes and to newly formed ectoplasmic specializations at sites of attachment to early elongating spermatids needs to be investigated further.
Actin Filament Dynamics at Tubulobulbar Complexes

Little is known about the actin filaments associated with tubulobulbar complexes. What is known is that the actin filaments appear arranged into a three dimensional cuff that surrounds the length of each tubular component of the complexes. The arrangement of filaments into a network is consistent with the possibility that the Arp2/3 complex is involved in formation of the structures, although this remains to be verified.

The fact that tubulobulbar complexes originate from areas previously occupied by ectoplasmic specializations raises the possibility that the actin network might result partly from a remodeling of preexisting actin filaments in the junction plaques. Support for this possibility is that tubulobulbar complexes contain some of the same protein components that are found in ectoplasmic specializations.61

Among actin-related components known to be associated with tubulobulbar complexes are paxillin,42 espin, myosin VIIa, and Keap 1.61 One component found associated with tubulobulbar complexes that does not appear to be concentrated at ectoplasmic specializations is the calcium-independent actin-depolymerizing protein nonmuscle cofilin,41 a protein that is regulated by phosphorylation. The presence of cofilin at tubulobulbar complexes fits nicely with an Arp2/3 model of actin network formation at these sites.

The Function of Actin at Tubulobulbar Complexes

The function of the actin filament networks at tubulobulbar complexes is not known; however one obvious possibility is that the filaments facilitate the formation and maintenance of the elongate tubular shape of the complexes. This conclusion is consistent with the observation that tubular parts of the complexes do not form after intratesticular injection of cytochalasin D.81 During normal spermatogenesis, disassembly of the filament networks near the ends of the complexes may lead to the formation of the bulbs and to the eventual budding of large double membrane vesicles from the complexes. The biological function of having numerous elongate tubes associated with junction disassembly may be related to increasing the surface area for the internalization of the huge amounts of junctional membrane during sperm release and during the movement of spermatocytes through basal junction complexes.

Intermediate Filaments

General

Intermediate filaments generally are classified into five Types (I - V) on the basis of gene substructure and sequence homology in a specific region of the proteins.82 Type I to IV form 8-12 nm diameter cytoplasmic filaments while type V consists of the lamins that form part of a meshwork on the inner aspect of the nuclear membrane. Unlike actin filaments and microtubules, intermediate filaments are formed from filamentous rather than globular subunits and the resulting polymers lack polarity. There are no intermediate filament dependent motor proteins. Also, filament dynamics are not dependent on nucleotide binding and hydrolysis by subunits and there is only a small pool of soluble subunits in vivo.82 Importantly for their function, and unlike actin filaments and microtubules, there is growing evidence that cytoplasmic intermediate filaments are flexible, extensible and tough.83,84

Generally in epithelial cells, intermediate filaments form perinuclear networks that extend peripherally to attach to cell/cell and cell/substratum adhesion junctions.85 Through this relationship of strong yet elastic cytoskeletal elements with adhesion sites, a significant function of intermediate filaments is to provide tissues, particularly epithelia, with mechanical support and the ability to maintain tissue integrity and be resilient to mechanical deformation.85

Other functions of cytoplasmic intermediate filaments include establishing the positions of organelles, providing scaffolds for signaling and other protein complexes, facilitating resistance to metabolic stress, and determining cytoarchitecture.82,86-88 The nuclear lamins have been implicated in determining nuclear shape and providing structural support for the nuclear
membrane. They also may play roles in positioning nuclear pores and anchoring chromatin to the nuclear envelope, and in the biogenesis of nuclear membranes.89

**Intermediate Filaments in Sertoli Cells**

In Sertoli cells, cytoplasmic intermediate filaments are major components of the cytoskeleton. The filaments mainly consist of vimentin (Type III),90 although keratins (Type I and II) are expressed during development,91 and in certain other situations.92,93 Nestin (Type IV) also is expressed in Sertoli cells during development.94 Nuclear laminas are present in Sertoli cells,95,96 but have not been extensively studied.

Intermediate filaments in terminally differentiated Sertoli cells form a network around the nucleus with peripheral extensions to desmosome-like junctions with adjacent Sertoli and spermatogenic cells and to hemidesmosome-like structures that connect the cell to the basal lamina (Figs. 1, 8, 9 and 10).1,97 When spermatids are embedded deep within Sertoli cell apical crypts, transient desmosome-like junctions are intercalated into the ectoplasmic specializations in regions along the dorsal curvature of the spermatid heads (Figs. 1, 8 and 11).1 At these locations, intermediate filaments appear to pass through a defect in the ectoplasmic specialization and associate with a density on the plasma membrane. Desmosome-like junctions are dynamic during spermatogenesis11 so it is not surprising that the intermediate filament patterns also change during spermatogenesis.1

Generally in cells, desmosomes consist of the desmosomal cadherins (desmogleins and desmocollins) that are the integral membrane adhesion molecules, the armadillo proteins (plakoglobin and plakophilins) that link the desmosomal cadherins to the intermediate filament cytoskeleton and/or function as signaling or regulatory molecules, and the plakins (desmoplakin, plectin, envoplakin and periplakin) that regulate the linkage between the membrane complexes and underlying intermediate filaments.98 Hemidesmosomes consist of the integral membrane proteins integrin (α6β4) and type XVII collagen (also called bullous pemphigoid antigen-2) that links the membrane to the basal lamina, and the plakins (bullous pemphigoid antigen-1 and plectin) that link the integral membrane proteins to intermediate filaments.99,100

Remarkably little is known about desmosome-like and hemidesmosome-like junctions in the seminiferous epithelium. Desmosome-like junctions with spermatogenic cells are macular or spot-like in appearance. They are characterized by sub-plasma membrane densities on both the Sertoli and spermatogenic cell sides of the junctions (Fig. 10)101 and by a discontinuous and poorly defined intermediate dense line between adjacent plasma membranes. Intermediate filaments are related only to the Sertoli cell side of the junctions. Desmosome-like junctions between neighboring Sertoli cells are similar to those formed with spermatogenic cells except that intermediate filaments are related to both sides of the junctions.11 The junctions are most apparent in basal regions where they form part of, and are intercalated into, the junction complexes between Sertoli cells (Fig. 10A)15,102 and appear to form discontinuous belts around the cells. Desmosome-like junctions between Sertoli cells also are present in apical regions.11

The presence of desmosomal cadherins and desmosome related armadillo proteins at desmosome-like junctions in the seminiferous epithelium has not been confirmed; however, there exists the possibility that members of the classical cadherins and related catenins may be incorporated into these junctions.42,102,103 If true, the presence of classical cadherins at these sites, in addition to any desmosomal cadherins that occur at the sites, would enable these desmosome-like junctions to interact with actin filaments as well as with intermediate filaments. In Sertoli cells, the interaction would be mainly with intermediate filaments, although maybe not exclusively so. In spermatogenic cells, where intermediate filaments do not associate with the plaques, the interaction might instead be with the cortical actin network. One of the plakins (plectin) has been localized to the sites (Fig. 8)20 and presumably participates in linking the vimentin filaments to the junction plaques in Sertoli cells. Plectin also is associated with the Sertoli cell nuclear membrane where it may serve a similar function (Fig. 8).20
Virtually nothing is known about the molecular components of the small hemidesmosome-like junctions located at the base of Sertoli cells, which is surprising considering the important structural and signaling properties of these junctions in other systems.99
The function of intermediate filaments in Sertoli cells has not been clearly defined. The vimentin knock-out mouse is reproductively viable\textsuperscript{104} and the morphology of the adult seminiferous epithelium appears remarkably normal.\textsuperscript{105} The only phenotype observed in the Sertoli cells is a complete lack of cytoplasmic intermediate filaments. Even though desmosome-like membrane associated plaques are present, there are no intermediate filaments associated with them. It is possible that intermediate filaments in Sertoli cells play a mechanical strengthening role, but they do so only when the epithelium is stressed in a particular fashion or to sufficient levels. It has long been known that desmosome-like junctions maintain seminiferous epithelial integrity when exposed to hypertonic buffers.\textsuperscript{101} Intermediate filaments may facilitate this role in this and other\textsuperscript{106} mechanically stressful situations.

Figure 9. Electron micrographs of intermediate filaments (arrows) that surround the nucleus (A) and that also are associated with hemidesmosome-like structures at the base of the cell (arrowheads) (B).
Figure 10. Electron micrographs of desmosome-like junctions. A) Shown here is a section through parts of two Sertoli cells (SC) and part of a spermatogenic cell (GC) (rat). In the seminiferous epithelium, desmosome-like junctions occur both between adjacent Sertoli cells (examples are circled in this image) and between Sertoli cells and spermatogenic cells (GC) (examples are boxed in this image). In this image, desmosome-like junctions are clearly intercalated into the basal junction complex indicated by the ectoplasmic specializations (arrowheads). B) A desmosome-like junction between two Sertoli cells (ground squirrel). The cells have been fixed, extracted with detergent and treated with tannic acid. The arrows indicate intermediate filaments. C) A desmosome-like junction between a Sertoli cell and spermatogenic cell fixed using conventional techniques (rat). Intermediate filaments (arrows) are present only on the Sertoli cell side of the junction.
Microtubules

General

Microtubules are dynamic cytoskeletal components that influence cell shape and provide tracts for intracellular vesicular transport. Microtubules are long, 25 nm in diameter, tubular polymers formed by α and β tubulin heterodimers. These heterodimeric subunits confer an intrinsic polarity to microtubules providing distinct plus- (fast growing) and minus- (slow growing) ends, allowing these structures to be used as rails for directional cargo transport by the microtubule
dependent kinesin and dynein motor proteins. In most cells microtubules originate at nucleation sites called microtubule organizing centers (MTOCs), the classical center of which is the centriole containing centrosome. Microtubules then radiate out to the cell periphery with their minus ends, or slow growing ends, at the MTOC. Other general patterns of microtubule distribution are common, as is noncentrosomal nucleation of microtubules.

**Microtubules in Sertoli Cells**

Sertoli cells are tall columnar epithelial cells that maintain their elongate morphology largely with the microtubule cytoskeleton. Experiments using the microtubule disrupting drugs colchicine and vinblastine in the testis result in a dramatic loss of Sertoli cell architecture as well as the sloughing of germ cells.

Sertoli cell microtubules are abundant apical to the basally located nucleus where they are predominantly arranged parallel to the long axis of the cell (Fig. 12), although microtubules do occur in basal regions of the cell and form elaborate arrays in apical cytoplasm associated with late spermatids. Unlike what occurs in many other cells, microtubules are nucleated at the periphery of Sertoli cells and not in regions associated with the perinuclear centrioles. This conclusion is based on the observations that the majority of microtubules are oriented parallel to the long axis of the Sertoli cell.
with their minus-ends at the apex and their plus-ends at the base of Sertoli cells.\textsuperscript{118,119} Also, microtubules polymerize from the periphery or apex of Sertoli cells in the testis during recovery from nocodozole treatment.\textsuperscript{120} Additionally, the microtubule nucleating protein, \(\gamma\)-tubulin, is present at peripheral sites.\textsuperscript{121} Moreover, when \(\gamma\)-tubulin is over expressed in Sertoli cells it concentrates in peripheral regions associated with elongate spermatid heads.\textsuperscript{122}

During spermatogenesis, the microtubule patterns in Sertoli cells change; in other words, the patterns are stage specific and change according to the developmental stages of spermatogenic cells with which they are associated.\textsuperscript{1,115,117,123} In lateral processes associated with early spermatogenic cells, microtubules are scarce. However, a marked increase in microtubules is apparent in Sertoli cell projections associated with round spermatids. Later, microtubules concentrate in the cytoplasm around spermatid heads adjacent to the acrosomes, and form linear tracts of microtubules parallel to Sertoli cell crypts containing the attached elongate spermatids. Prior to sperm release, Sertoli cell microtubules often are concentrated in, or form specialized structures in, cytoplasm associated with spermatid heads. In addition, microtubules are present in projections associated with the lobe of residual spermatid cytoplasm. The findings that microtubules in Sertoli cells contain tyrosinated tubulin, which tend to be correlated with unstable populations of microtubules, and that the levels of tyrosinated tubulin change during spermatogenesis, are consistent with the observation that microtubule patterns in Sertoli cells are dynamic during spermatogenesis and with the possibility that the degree of microtubule stability may be stage specific.\textsuperscript{117,123}

In addition to a structural function, microtubules also are essential for the directional transport of intracellular cargo generally in cells. These cytoskeletal elements use the molecular motors, kinesin and cytoplasmic dynein to transport vesicles, proteins, and other cellular elements throughout the cell. These motors convert ATP as an energy source for their motility events. The kinesins together constitute a superfamily that has recently been reorganized to contain 14 families plus an additional group of orphan kinesins.\textsuperscript{124} Hirokawa's group\textsuperscript{125} has identified 45 kinesin genes in mouse and human databases. These protein machines transport cargo primarily towards the plus-ends of microtubules, although an increasing number of minus-end directed motors have been discovered. Cytoplasmic dynein is generally considered a minus-end directed molecular motor, although bidirectional movements of single dynein-dynactin complexes have been reported in vitro.\textsuperscript{126} Both of these groups of motor proteins have developed elaborate techniques for selecting cargoes, docking those cargoes to appropriate motors and targeting these cellular components to the appropriate locations.\textsuperscript{107,127-129}

In Sertoli cells, general housekeeping and other transport events such as those that occur between different membrane compartments and those involving the transport of organelles from one position to another likely involve microtubule-based motor proteins. This conclusion is based mainly on data from cells in general, and on the correlation of organelle position with microtubules,\textsuperscript{116,130,131} changes observed after microtubule perturbation\textsuperscript{81,114,152} and localization of motor proteins in Sertoli cells.\textsuperscript{133-135}

One transport event that occurs during spermiogenesis is the translocation of spermatids in the seminiferous epithelium.\textsuperscript{136-139} This event occurs as the result of a change in position of apical crypts that contain the differentiating germ cells. When the crypts initially form in the apices of Sertoli cells, they are shallow and, together with the attached spermatids, occur in apical regions of the epithelium. At certain stages, the crypts deepen (most dramatically at stages IV-V in rat) and the attached spermatids, specifically their heads, become located near the base of the epithelium. As spermatids mature, the crypts again become shallow and spermatids are returned to the apex of the epithelium for eventual release. The biological significance of transporting spermatids deep into the epithelium is not known. What is known is that the process occurs generally in vertebrates and appears fundamental to the process of spermatogenesis. Entrenchment within the epithelium may provide increased mechanical support for elongate shaped spermatogenic cells, facilitate development of the sperm tail, provide increased surface contact for exchange of materials or signals between Sertoli cells and the developing spermatids, or have other as yet unknown functions.
Based on observations that Sertoli cell microtubules are found in close opposition to ectoplasmic specializations attached to spermatid heads\cite{131,140} and that microtubules are oriented parallel to the direction of spermatid translocation, it has been proposed that the mechanism responsible for changes in position of elongate spermatids during spermatogenesis in mammals involves the transport of ectoplasmic specializations and the attached spermatids along microtubule tracts in Sertoli cells.\cite{115,140} It is proposed that microtubule-based motors attached to the cytoplasmic face of the endoplasmic reticulum component of ectoplasmic specializations transport the junction/spermatid complex first basally and then apically in the epithelium.\cite{115,118,141}

A number of lines of evidence support this hypothesis. First, pharmacological disruption (depolymerization) of microtubules in Sertoli cells alters the position of spermatids in the seminiferous epithelium.\cite{106,114} Second, microtubules bind to isolated ectoplasmic specializations in an ATP-dependent fashion. Moreover, the binding has a profile of sensitivity to pharmacological agents that is consistent with those of some known microtubule based motors.\cite{141} At the ultrastructural level, this nucleotide-dependent binding occurs on the cytoplasmic face of the endoplasmic reticulum component of the junction plaques.\cite{142} Third, isolated junction plaques transport microtubules. In an in vitro motility assay, exogenous microtubules move, in an ATP-dependent fashion, on isolated spermatid/junction complexes.\cite{143} Fourth, junction plaques transport microtubules in both directions. To account for the “down-and-up” movement in vivo, isolated junction plaques should move microtubules in plus-end and minus-end directions. This prediction has now been verified.\cite{144} Fifth, cytoplasmic dynein is associated with the junction plaques. Antibodies to the 74 kDa intermediate chain (IC74) of cytoplasmic dynein strongly react at the light microscopic levels with Sertoli cell regions known to contain the junction plaques,\cite{135} and react with the cytoplasmic face of the endoplasmic reticulum component of the junction plaque at the ultrastructural level.\cite{144} Significantly, this immunoreactivity occurs during the period of spermatid translocation. This result is the first evidence, from immunolocalization studies, that a microtubule-based motor occurs in plaque regions at the appropriate time during spermatogenesis. Further support for the association of cytoplasmic dynein with the junction plaques was obtained by using gelsolin (an actin severing enzyme) to disassemble the plaques and release the endoplasmic reticulum component of the plaques into solution. When assessed by immunoblots, GRP94 (a marker for endoplasmic reticulum) and cytoplasmic dynein (indicated by IC74) were enriched in supernatants collected from low speed centrifugation of gelsolin treated spermatid/junction complexes relative to controls.\cite{135}

Although cytoplasmic dynein has been localized to ectoplasmic specializations and is likely the motor responsible for the apical movement of spermatids in the mammalian seminiferous epithelium, the plus-end directed motor, likely a kinesin, responsible for the basal movement has yet to be identified.

**Concluding Remarks**

The association of the cytoskeleton with sites of intercellular attachment and with processes such as the release of sperm cells from the epithelium, the movement of spermatocytes through basal junction complexes between Sertoli cells and the translocation of spermatids in the seminiferous epithelium indicates that the cytoskeletal machinery of Sertoli cells is likely fundamental to spermatogenesis and may potentially contain molecular targets for contraception.

Although much has been learned about the organization and function of the Sertoli cell cytoskeleton over the last few years, there still is a great deal to be learned.

The mechanism of regulating actin dynamics in ectoplasmic specializations and at tubulobulbar complexes has yet to be defined. The role of the endoplasmic reticulum both at ectoplasmic specializations and at tubulobulbar complexes is completely unknown, except that at apical ectoplasmic specialization microtubule transport machinery has been coupled through the endoplasmic reticulum to the junction plaque in order to translocate and position spermatids in the epithelium. The precise role of actin at ectoplasmic specializations and tubulobulbar complexes remains to be determined.
The molecular composition of desmosome-like and hemidesmosome-like junctions in Sertoli cells, and how their linkages to the intermediate filament cytoskeleton is formed and controlled remains to be elucidated. Desmosome-like junctions are a major component of the intercellular contact machinery in the seminiferous epithelium and, like ectoplasmic specializations, their association with spermatogenic cells is cell specific and therefore dynamic. Desmosome-like junctions between neighboring Sertoli cells are part of, and are integrated into, the basal junction complex and are therefore presumably disassembled above and assembled below translocating spermatocytes. The hypothesis that intermediate filaments, together with their attachment to desmosome-like and hemidesmosome-like junctions, maintain epithelial integrity when the tissue is mechanically stressed has yet to be experimentally tested.

Although microtubule-based transport mechanisms are fundamental to intracellular motility generally in cells, the coupling of this machinery to a junction plaque to position adjacent cells in an epithelium is unique to Sertoli cells. Cytoplasmic dynein has been localized to the junction sites; however, the predicted plus-end directed motor, likely a kinesin, has not. The microtubule-based spermatid translocation hypothesis has yet to be tested in vivo, and the biological function of spermatid translocation itself remains to be determined.

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The Sertoli Cell Cytoskeleton


The Sertoli Cell Cytoskeleton

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CHAPTER 12

Blood-Testis Barrier, Junctional and Transport Proteins and Spermatogenesis

Brian P. Setchell*

Functional Evidence for a Blood-Testis Barrier

The term “blood-testis barrier” appears to have been first used by Chiquoine in an article on effects of cadmium on the testis, but evidence for such a barrier already existed, dating back to the early years of the twentieth century (see ref. 2 for early references). In a number of studies, it was shown that some dyes when injected into animals, stained most tissues, with the notable exceptions of the brain and the seminiferous tubules of the testis. The former observation was rapidly taken up and developed to form the basis for the concept of the blood-brain barrier, but it was only with the studies of Kormano that the true significance of the earlier observations on the testis was recognized. He showed that dyes which were excluded from the tubules of adult rats readily penetrated those of prepubertal animals. In addition, Kormano noticed that staining of interstitial cells with acriflavine also fell around the time of puberty, suggesting a change in the blood vessels as well. At about the same time as Kormano’s studies, Waites and I showed that testis blood flow measured by indicator dilution with rubidium gave much lower values that with iodoantipyrine, while similar values were obtained in most other organs except brain, suggesting that rubidium was also excluded to some extent from parts of the testis, as it was from the brain.

Also around this time, Waites and I devised techniques for collecting fluid from the rete testis (RTF) of sheep and from the rete testis and seminiferous tubules (STF) of rats, and we found that both RTF and STF differed appreciably in composition from either blood plasma or testicular lymph collected from a vessel in the spermatic cord. That such differences, especially those for small hydrophilic organic compounds such as inositol could be maintained provided further evidence that there was not free communication between the various fluid compartments inside the testis, and this was confirmed in studies on the rate of penetration of various radioactive markers from the bloodstream into RTF in rams or RTF and STF in rats.

There are three cell types between the fluid inside the blood vessels and that in the lumina of the seminiferous tubules, namely the endothelial cells lining the blood vessels, the peritubular tissue and the Sertoli cells. These last are the only cells to extend all the way from the peritubular tissue to the lumen of the tubule, with the developing germ cells lying either between the base of a Sertoli cell and the peritubular tissue, or in the intercellular space between a pair of Sertoli cells or in crypts in the luminal surface of a Sertoli cell. All three cell types could conceivably influence the rate of entry of substances into the tubules, although most attention has been directed to the Sertoli cells (see next section).

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Other techniques used to estimate the effectiveness of the blood-testis barrier include the measurement of the volume of distribution of a marker known to be excluded from STF, such as Cr-EDTA, inulin or sucrose, and relating this either to the volume of the interstitial tissue or to the value obtained when the efferent ducts had been ligated 24 h previously, so that the fluid secreted during that time had been retained in the lumina of the seminiferous tubules. Other studies used as a marker, hexamethonium iodide, which has been shown not to penetrate the blood-brain barrier, a zinc complex of carnosine labeled with C-14 and Zn-65, or a biotin tracer. Another approach is to relate the amount of a labeled compound to the amount of Tc-99 or I 125 labeled albumin appearing in the testis and brain of mice following an intravenous injection. From these data, an entry rate (Kₚ) for the marker can be calculated, but this value in the testis could be influenced by changes in vascular permeability as well as in permeability of the tubular barrier.

The latest development has been the use of magnetic resonance imaging of the testis, before and after intravenous injection of gadopentetate dimeglumine. Qualitative evidence for a barrier in young animals is provided by the development of a lumen and the secretion of fluid in the tubules.

While most evidence for the involvement of the Sertoli cells is morphological (see next section) it should be remembered that when isolated Sertoli cells are cultured at high density on Matrigel in a two-chamber system, they form a confluent layer, which exhibits barrier properties, as shown by an increase in electrical resistance and directional secretion of a number of substances. However, the transepithelial resistance (TER) obtained (usually about 100 ohm.cm²) was usually much less than that seen with MDCK cells or keratinocytes (100-2000 ohm.cm²). Nevertheless, treatment of Sertoli cell cultures with FSH and testosterone could raise TER to between 580 and 1200 ohm.cm², and the cells were usually obtained from prepubertal rats, in which the barrier would not be fully formed (see below).

Structural Evidence for a Barrier

A Sertoli Cells

The existence of specialized junctions between pairs of Sertoli cells was recognised in the 1960's. Their significance became apparent when it was shown that electron opaque markers which were injected into the interstitial tissue or reached there from the blood stream, were restricted from entering the tubules to some extent by the peritubular myoid cells, but almost completely by the specialized junctions between pairs of Sertoli cells. The markers used included colloidal carbon, ferritin, horseradish peroxidase, lanthanum salts, and more recently biotin.

Peritubular Myoid Cells

Peritubular myoid cells form a single layer in rodents and several layers in primates around the seminiferous tubules. As long ago as 1901, it was suggested that this cell layer formed “a sort of dialysing membrane which regulates the composition of the fluid contained in the space that it limits” (une sorte de membrane dialysante qui règle la composition du liquide contenu dans l’espace qu’elle limite). The cells change in shape, structure, marker expression and rate of cell division around the time of puberty and respond in culture to endothelin, which as the name implies, is usually produced by endothelial cells, but in the testis is formed mostly by the Sertoli cells. The myoid cells produce PmodS, a protein which has a powerful influence on several Sertoli cell functions, although an effect on the blood-testis barrier has not apparently been examined.

The peritubular myoid cells prevented the passage of larger electron-opaque markers like colloidal carbon or thorium, and lanthanum penetrated the myoid cell layer in only about 15% of the tubules in rodent testes. However, in primate testes, the peritubular cells have much less effect in restricting the penetration of markers.
Nevertheless, the myoid cells may have an important influence in restricting the entry of retinoic acid (RA) into the tubules. Less than 1% of the RA in the testis is derived from plasma RA, much less than in any other tissue studied.\textsuperscript{68} This may be due to the presence in the myoid cells of the RA-degrading enzymes Cyp 26 a1, Cyp 26 b1 and Cyp 26c1,\textsuperscript{69} while the first stage of the formation of RA from retinol occurs in the Sertoli cells. It has been known for many years that spermatogenesis is arrested in Vitamin A-deficient animals, and retinoic acid is effective in restoring sperm production only in pharmacological doses (10 mg/week compared with 0.1 to 0.2 mg/week for retinol).\textsuperscript{70} The restricted entry of retinoic acid may explain this difference. The myoid cells also contain high levels of cellular retinol binding protein,\textsuperscript{71,72} which is probably involved in the transport of retinol into the tubules (see below).

**Endothelial Cells**

The endothelial cells in the testis are unusual for an endocrine tissue in that they are unfenestrated,\textsuperscript{74-76} although in the human testis, some capillaries in the lamina propria do have fenestrations.\textsuperscript{77} Endothelial cells in the rat testis also have a much lower density of vesicles than vessels in other tissues, except brain,\textsuperscript{78} suggesting that vesicular transport is less important in these tissues than elsewhere in the body.

**Structural Constituents of the Sertoli Cell Junctions**

In recent times, a large amount of information has appeared about the constituent proteins of the Sertoli cell junctions which constitute that part of the blood-testis barrier (Fig. 1). The main components include occludin, one or more of the claudins, zonula occludens (ZO), and junctional adhesion molecules (JAM’s). Occludin, claudin-11 and JAM-1 are transmembrane proteins, the extracellular parts of which join with similar structure on an adjacent Sertoli cell to form a tight junction. In the cytoplasm of the cells, the intracellular tails of the occludin, claudin and JAM molecules are joined to ZO-1 and ZO-2 molecules, which in turn are linked to actin chains.\textsuperscript{48}

Occludin is a 60 to 65 kDa protein with four transmembrane domains, one intracellular and two extracellular loops, and is present in the tight junctions between Sertoli cells in rats and mice, but not guinea pig or human.\textsuperscript{79} In mice which carry a null mutation of the occludin gene, the testes initially develop normally, but by 40 to 60 weeks of age, the tubules become atrophic, with complete loss of germ cells.\textsuperscript{80} Therefore, it is rather surprising that occludin first appears in the fetal testis at about day 13 pc (post-coitus), long before spermatogenesis is initiated, suggesting that occludin has functions other than the establishment of the barrier. In postnatal rats, at about day 5, the reaction for occludin becomes more intense and is then located along the lateral plasma membrane of the Sertoli cells. Then at day 14, the reaction appears as intense focal bands close to the base of the epithelium, near the presumed sites of the tight junctions which are forming at about that time.\textsuperscript{81} Injection into rats of a 22-amino acid synthetic peptide corresponding to the second extracellular loop of occludin perturbs the blood-testis barrier and disrupts spermatogenesis.\textsuperscript{82}

Claudins are a family of more than 20 proteins, about 22 kDa in size,\textsuperscript{83} and claudin-11 is present at tight junctions between Sertoli cells in testes, but again appears first during fetal life. Its concentration in the testis reaches a peak at about 6 days of age, and then appears to decline, probably due to the appearance of claudin-negative germ cells.\textsuperscript{84} Claudin-11 null male mice are sterile, and tight junctions appear to be absent in these animals as judged by freeze-fracture.\textsuperscript{85} Claudin-5, which is found only in endothelial cells,\textsuperscript{86} is present in endothelial cells in the rat testes,\textsuperscript{87} but as mice null for this peptide die within a few days of birth,\textsuperscript{88} it has not been possible to study the effect of lack of this protein on spermatogenesis.

Integrins are thought to be involved in junctions between testicular cells and extracellular matrix,\textsuperscript{88} but there is evidence\textsuperscript{89} that integrin α6 β1 is also present in Sertoli-Sertoli cell junctions, especially at certain stages of spermatogenesis, but also in Sertoli cell-only testes.\textsuperscript{90} In testis explants, the development of this suprabasal integrin occurred only in the presence of FSH.\textsuperscript{90}
Figure 1. A diagram illustrating the molecular architecture of the three multiprotein complexes found at the Sertoli-Sertoli cell junctions of the blood-testis barrier. The three complexes are: (1) Occludin-ZO1/ZO2; (2) claudin-ZO1/ZO2; and (3) JAM-ZO1. Also shown are the peripheral membrane proteins known to regulate Sertoli cell tight junction dynamics. Reproduced with permission from: Mruk DD, Cheng CY. Endocrin Rev 25:747-806, ©2004 The Endocrine Society.
Transport Proteins and the Blood-Testis Barrier

**Transferrin**

Iron is transported into the germ cells inside the blood-testis barrier by a mechanism involving a specific transport protein, transferrin. In the blood, iron is carried bound to transferrin secreted by the liver, and on reaching the testis, this complex binds to transferrin receptors on the basal surface of the Sertoli cells. The iron-transferrin complex is then internalised and dissociated, the apo-transferrin returned to the interstitial extracellular fluid and the iron is complexed to transferrin produced inside the Sertoli cell and secreted into the space between the Sertoli cell and the germ cells (Fig. 2). How the iron is moved across the Sertoli cell is still uncertain, but may involve a ferritin-like molecule. Sertoli cells in a bicameral culture system synthesize and secrete transferrin, and iron from basally applied human transferrin is transported through rat Sertoli cells and appears in the apical compartment bound to rat transferrin. Nevertheless, the concentration of transferrin in seminiferous tubule fluid is less than one-twentieth of that in interstitial extracellular fluid or blood plasma.

Other elements besides iron are bound by transferrin, and this may be important in causing the accumulation inside the tubules of potentially mutagenic radioactive substances like indium and plutonium.

Transferrin production by Sertoli cells is greater if the cells are derived from 17 day old rather than 10 day old rats, is reduced following hypophysectomy and not restored by testosterone treatment. It is stimulated by FSH, cytokines, a factor PmodS produced by the peritubular myoid cells, and heregulins, which may also come from the same source. The presence of germ cells in the tubule may also have an effect on transferrin production by the Sertoli cells, although different results were obtained when the germ cells were depleted with methoxyacetic acid. Sertoli cells also secrete an copper-transporting protein, ceruloplasmin, but it is not known whether this substance is involved in copper transport into the tubules.

Figure 2. A diagram illustrating the role of transferrin in the transport of iron and other metals into the seminiferous tubules. Diferric serum transferrin (Fe-sTF-Fe) binds to a transferrin receptor on the basal surface of the Sertoli cell. The transferrin-ferric ion-transferrin receptor complex is internalized into special compartments in the cell, acidified and broken down. The apotransferrin and the transferrin receptor are recycled to the cell surface, and the iron is moved through the cell to newly synthesized testicular transferrin (tTf) or is incorporated into ferritin in the Sertoli cell. The testicular transferrin with the ferric ions is released into the intercellular space between the Sertoli and germ cell and then binds to transferrin receptors on the surface of the germ cells. The net result is transport of ferric ions from the basal surface of the Sertoli cell to the adluminal compartment of the tubule. Reproduced with permission from: Sylvester SR, Griswold MD. J Androl 1994; 15:381-385, ©1994 American Society of Andrology.
Another divalent metal transporter DMT1 (Slc11a2) is also present in the Sertoli cells of the rat testis, although it is not primarily responsible for translocating iron across the epithelium, but in intracellular handling of iron during spermatogenesis.\textsuperscript{109,110}

**P-Glycoprotein**

P-glycoprotein (Pgp) is the product of the multidrug resistance 1 gene (MDR1 or ABC B1 in humans, and mdr1a (sometimes called mdr3) and 1b in mice and rats). It was originally identified in cancer cells which had become resistant to chemotherapeutic drugs.\textsuperscript{111-114} Subsequently, it was found that this protein was present in a number of normal tissues, and especially in the endothelial cells of the brain and testis.\textsuperscript{115-118} It is also present in other cells in the testis, including Leydig cells, macrophages, peritubular cells, Sertoli cells and late spermatids, although not detectable in spermatogonia, spermatocytes or early spermatids.\textsuperscript{119,120} However, the relative concentrations in the various cell types has apparently not yet been determined and another group has detected mdr 1 in germ cells, probably spermatogonia, in rats, as well as in endothelial cells in the testis.\textsuperscript{121} In endothelial cells from brain, Pgp is expressed only on the luminal surface, consistent with a role in protecting the brain from circulating lipophilic molecules which would otherwise cross the blood-brain barrier. However, in endothelial cells in the testis, Pgp is expressed on both luminal and abluminal surfaces, which suggests that it acts to exclude substrates of the transporter from the endothelial cells themselves.\textsuperscript{122} A mRNA from a related gene mdr 2 is also present in Sertoli cells, but at a lower concentration than in liver.\textsuperscript{109}

The testes and brains of mice in which the gene for mdr-1a has been deleted accumulate more ivermectin, digoxin, cyclosporin A, ondansetron, loperamide and vinblastine than controls.\textsuperscript{123-126} In other studies,\textsuperscript{127,128} similar results were obtained with amitriptyline and some of its metabolites, but not with fluoxetine. In mice in which both mdr 1a and 1b have been knocked out, the entry of the anti-Parkinson drug budipine into the testes and brains was enhanced.\textsuperscript{129} In these double knockout mice, the penetration of the steroids, corticosterone, cortisol, aldosterone and progesterone into the testes was also enhanced,\textsuperscript{130} although cortisol\textsuperscript{131} or prednisolone\textsuperscript{132} entry into the testis was unaffected in mdr 1a single knockout mice. Pgp also transports HIV protease inhibitors (HPI) used in the treatment of AIDS\textsuperscript{133} and pharmacological inhibition of the transporter enhances the penetration of the HPI nelfinavir into the testes of mice treated with LY-335979, a potent Pgp inhibitor, as well as in mdr-1a knockout mice.\textsuperscript{134} The penetration of saquinavir, another HPI into the testes of mice was also enhanced by treatment of the animals with another inhibitor of Pgp, GF120918.\textsuperscript{135} However, treatment of mice with a variety of Pgp inhibitors failed to increase the penetration of vinblastine into either testis or brain,\textsuperscript{136} and vincristine enters seminiferous tubule fluid reasonably rapidly,\textsuperscript{137} although it is a substrate for both Pgp and MRP\textsuperscript{126} The closely related efflux pump, breast cancer resistance protein (BCRP) is also found in the endothelial cells and peritubular myoid cells in the testis,\textsuperscript{120} but the structurally related protein encoded by the cystic fibrosis gene is not found in endothelial cells, but is expressed in spermatids in a stage-specific fashion.\textsuperscript{121,138}

**Multidrug Resistance Protein**

Multidrug resistance proteins (MRP) are other members of the ATP-binding cassette superfamily distantly related to Pgp. MRP1 is present in high concentrations in testes\textsuperscript{139} and is localized to the Leydig and Sertoli cells in human and mice,\textsuperscript{120,140} but cannot be detected in endothelial cells in the rat testis.\textsuperscript{141} Mice lacking the gene for this protein are much more sensitive to the damaging effects of etoposide phosphate,\textsuperscript{142} and methoxycarbonyl\textsuperscript{142} than normal mice, suggesting that it acts to exclude these drugs from the seminiferous tubules. MRP1 is also involved in glutathione-mediated transport of sulfated estrogens, and it has been suggested that the high levels of MRP1 in the Leydig cells may be responsible for the efflux of the hydrophilic sulfated conjugates from the cell.\textsuperscript{143} The anticancer drug methotrexate, which is transported out of cells by MRP, but poorly by Pgp,\textsuperscript{144} is virtually excluded from seminiferous tubule fluid.\textsuperscript{145}
**Other Transport Mechanisms**

Endothelial cells in the testis contain high levels of \( \gamma \)-glutamyl transpeptidase, \( \alpha \)-glutamyl transpeptidase, an enzyme usually associated with amino acid transport, and it has also been shown that endothelial cells of the larger blood vessels in the rat testis transport leucine with transport kinetics similar to those of brain and much lower than for other tissues. There is also a large amino acid transporter present in rat testis as well as brain and heart, but not other tissues. Endothelial cells in the rat testis also contain an endothelial barrier antigen (EBA), previously thought to be confined to nervous tissue, and an isoform GLUT-1 of the glucose transporter family, usually associated with brain and retina.

The peritubular cells in the mouse testis contain a specialized transporter protein involved in urea movement across plasma membranes, UT-A5, the levels of which are not related to the stage of spermatogenesis in adults but are coordinated with the stage of testis development, increasing around 15 days post partum. In rat Sertoli cells, there are also 4 other urea transporters, UT-A 1, 2, 3 and 4 present at all stages of spermatogenesis, and UT-B is present at stages II and III. UT-A3 was also present in some interstitial cells. Flux of urea across the walls of isolated perfused seminiferous tubules is inhibited by phloretin. It is interesting that there is some evidence for the active accumulation of radioactively labeled urea inside the seminiferous tubules of rats.

Evidence has recently been presented for the presence of a family of saturable nucleoside transporters in isolated Sertoli cells, as primary cultures or as polarized layers on Matrigel, some of which is sodium-dependent and can be inhibited with nitrobenzylthioinosine.

Binding proteins may also be important in the regulating the entry of retinol into the tubules. Homogenates of rat testis bind more retinol and retinoic acid (RA) than any other tissue examined, but in vivo, very little RA enters the tubules from blood. Both myoid and Sertoli cells in the testis contain a cellular retinol-binding protein (CRBP). The Sertoli cells also contain a number of retinoic acid receptors. Retinol circulates in the plasma bound to a retinol-binding protein (RBP), a 21 kDa protein which normally is present as a 76kDa 1:1 complex with transthyretin. This complex in the testis is confined to the interstitial tissue. When retinol bound to RBP was injected into the testis or under the capsule, it appeared in the tubules only after at least 30 minutes, whereas tritiated retinol injected mixed with albumin, spread rapidly throughout the testis. Early studies could not detect any interaction of RBP with cells in or on the seminiferous tubules. Nevertheless, both peritubular myoid and Sertoli cells appear to be involved in the transport of retinoids to the germ cells. Both cell types in culture are able to accumulate retinol from serum RBP by a saturable and competeable process, which involves recognition of the retinol-RBP complex at the cell surface, with subsequent internalization of the retinol but not the RBP. The first step involves the myoid cells, which bind the retinol inside the cell to newly formed CRBP, and the new complex is released into the space between the myoid cells and the Sertoli cells. The latter then take up just the retinol and complex it with new CRBP, before releasing the complex again to reach the germ cells.

Sertoli cells also contain a prostaglandin D2 synthetase, which also binds retinoic acid but not retinol. This protein is secreted into rete testis fluid, but its role in the transport of retinoids into the tubules in not yet clear.

**Factors Affecting Blood-Testis Barrier Function**

**Age and Hormones**

As already mentioned, studies on the penetration of certain dyes into the seminiferous tubules showed that these dyes were excluded only from the tubules of rats older than about 20 days. Subsequently, it was shown that electron-opaque markers injected into the interstitial tissue of the testes of rats entered the tubules freely up to 16 days of age, but between 16 and 19 days, the occluding junctions between the Sertoli cells appear and the tracers are effectively...
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prevented from reaching the tubular lumen.\textsuperscript{163} In immature rat testes, occluding junctions, as demonstrated by freeze-fracture, are absent, although gap junctions are present. Furthermore, perfusion with hypertonic lithium chloride caused the cells outside the Sertoli cells junctions in adult testes to shrivel, with no effect on those inside the junctions, whereas in the testes of 13 day old rats, cellular shrinkage occurred throughout the tubules.\textsuperscript{164} Shrinkage of adluminal cells in response to exposure to a hypertonic solution decreased between 14 and 18 days of age.\textsuperscript{43} Similarly in guinea pigs, Sertoli cell junctional complexes appeared around 15 days after birth,\textsuperscript{165} and in mice at about 16 days.\textsuperscript{166} In rats around 15 days of age, the barrier appears only in those parts of the tubule where germ cells have reached pachytene.\textsuperscript{167}

In seasonal breeders such as mink,\textsuperscript{168,169} viscacha,\textsuperscript{170} and Djungarian hamsters\textsuperscript{171} electron opaque markers are excluded by the Sertoli cell junctions during the breeding season, but during testicular regression, the tracer penetrates throughout the tubules. In the study on mink, the exclusion of the marker from the tubule was associated with the presence of a tubular lumen, rather than any particular type of germ cells. At the other end of life, the barrier in 24 month old rats was grossly deficient, with associated failure in spermatogenesis.\textsuperscript{172} The development of the barrier in young rats and mice can be retarded by the neonatal administrations of diethylstilboestrol.\textsuperscript{173,174}

The development of a lumen in the tubules is more gradual, beginning at around 10 days after birth, and with the diameter continuing to increase slowly to day 30 and then more rapidly to around day 50.\textsuperscript{22,43} Fluid secretion per unit weight of testis also continued to increase until about 45 days of age,\textsuperscript{42,44} and the volume of distribution of Cr-EDTA, which is normally excluded from the tubules, continues to fall until after 30 days of age,\textsuperscript{22} so the functional barrier appears to develop more gradually than the anatomical one.

The development of transepithelial electrical resistance (TER) in two-compartment Sertoli cell cultures is delayed by FSH for several days, and once established is decreased and then returns to control levels or increases. Testosterone alone caused a rapid increase in TER, and testosterone and FSH together resulted in the highest TER levels. Dihydrotestosterone was more effective than testosterone, whereas estradiol was without effect.\textsuperscript{45} Dibutryl cyclic adenosine monophosphate (cAMP) in low concentrations stimulated TER development, whereas higher doses were inhibitory. Cholera toxin mimicked the FSH effects.\textsuperscript{175} The effect of cAMP on the Sertoli cell tight junctions is probably mediated by a proteasome-sensitive ubiquitination of occludin.\textsuperscript{47} TGF-\beta3 also regulates blood-testis barrier dynamics, probably by determining the steady-state levels of occludin and ZO-1 via the p38 MAP kinase signaling pathway.\textsuperscript{176} Tumour necrosis factor \(\alpha\) injected directly into rat testes caused a temporary disruption of the blood-testis barrier, by reducing the levels of occludin, zonula occludens-1 and N-cadherin.\textsuperscript{177}

Testosterone, acting through its receptor in the Sertoli cells, regulates the expression of claudin-3, which encodes a transient component of newly formed tight junctions. Sertoli cell-specific ablation of androgen receptor results in increased permeability of the barrier to biotin.\textsuperscript{30} The effect of androgen withdrawal on the Sertoli cell junctions was studied either by hypophysectomy or by treatment of rats with ethane dimethane sulfonate to destroy the Leydig cells. These treatments led to degeneration of germ cells and the formation of numerous basally-located vacuoles, formed by multiple focal dilations of the intercellular space associated with the junctional complexes. As this occurred also in Sertoli cell-only testes, produced by fetal irradiation, it cannot be explained by spaces left by degenerating germ cells.\textsuperscript{178} The expression of occludin is also reduced by treatment of rats with the anti-androgen, flutamide.\textsuperscript{179} In an intratesticular androgen suppression model, using subcutaneous implants of testosterone and estrogen to suppress LH secretion and hence endogenous androgen production, the adherens junctions between the Sertoli cells and spermatids can be disrupted, without affecting blood-testis barrier integrity.\textsuperscript{180}

The Sertoli cell barrier to lanthanum develops normally in rats treated in utero with busulfan but at a later age around 30 days of age, at the time of the appearance of the first zygotene and pachytene cells in these animals.\textsuperscript{181} However, in prenatally irradiated rats, tight junctions, as detected by freeze fracture, were extensive by 3 months of age, although their ability to block
the penetration of markers was not examined.\textsuperscript{167} It is probably relevant that the fluid inside the Sertoli cell only tubules of prenatally irradiated or busulfan-treated rats was plasma-like in its potassium content, in contrast to the high potassium of normal fluid.\textsuperscript{16,183}

**Vitamin A Deficiency**

In rats made Vitamin A-deficient from weaning (20 days old), Sertoli cell junctions were intact and complete spermatogenesis was maintained up to 80 days of age. However, by 90 days, lanthanum could penetrate through the junctions and by 100 days severe regression of spermatogenesis had occurred.\textsuperscript{184} Different results were obtained by Ismail and Morales,\textsuperscript{185} who found that the junctions remained impermeable to lanthanum, even when spermatogenesis had failed in rats 104 days old, deficient since 20 days old. In a later study, following long-term deprivation of Vitamin A, the Sertoli cell junctions became permeable to lanthanum when spermatogenesis was arrested and remained so even when spermatogenesis was first reinitiated. Spermatocytes normally found in the adluminal compartment were apoptotic, while spermatocytes normally found in the basal compartment remained normal.\textsuperscript{186}

**Tissue and Blood Pressures**

If the efferent ducts leading from the testis to the epididymis are ligated close to the testis, the fluid normally secreted by the Sertoli cells to transport the immotile spermatozoa is retained inside the seminiferous tubules. These become progressively distended for between 24 and 36 h in rats, so that the testis becomes enlarged and turgid. Then the testis weight falls again and eventually by 21 days, spermatogenesis is completely deranged.\textsuperscript{52} During this time the blood-testis barrier, judged by the ratio of the space of distribution of Cr-EDTA to the measured volume of the interstitial tissue remained normal during the phase of fluid accumulation, but increased sharply as testis weight begins to fall again, indicating breakdown of the barrier. Surprisingly, by the time testis weight had returned to control levels, the barrier appeared to be functioning again, and it remained functional even when spermatogenesis was completely disrupted up to 3 weeks later.\textsuperscript{24,187} One author\textsuperscript{188} found that lanthanum penetrated more readily through the Sertoli cell junctions as early as 24 h after efferent duct ligation. However, other studies with electron opaque markers gave contradictory results.\textsuperscript{188-191}

In chronically hypertensive rats, the penetration of sucrose and 2-methyl-4-chlorophenoxyacetic acid into the testis is reduced, while that of the highly permeable antipyrine is unaffected.\textsuperscript{192} In rats with testicular degeneration induced by epinephrine, the barrier remains able to exclude lanthanum.\textsuperscript{193}

**Cadmium and Other Toxic Substances**

The testes of most mammals are extremely sensitive to the effects of cadmium salts, in doses which have little effect on other tissues. Early observations\textsuperscript{1} concentrated attention on the blood vessels in the testis, and there is no doubt that testis blood flow is reduced in rats as a result of increases in vascular permeability as early as several hours after a single injection of cadmium chloride.\textsuperscript{6} Later studies showed that permeability of the blood-testis barrier to rubidium probably preceded the changes in vascular permeability.\textsuperscript{194} In guinea pigs on the other hand, increased staining of the interstitial tissue with acriflavine injected subcutaneously occurred before an increase in staining of the seminiferous tubules.\textsuperscript{195} However, lower doses of cadmium affect spermatogenesis without noticeable changes in the vascular system, and these effects can be reduced by coadministration of zinc salts.\textsuperscript{196}

Exposure of bicameral Sertoli cell cultures to cadmium salts caused a progressive and dose-dependent drop in TER.\textsuperscript{197,198} The expression of occludin is decreased and u-plasminogen activator is increased in the presence of cadmium.\textsuperscript{198} Treatment of rats with low doses of cadmium chloride caused changes in the tight junction-associated microfilaments in the Sertoli cells by 24 h after injection, although no changes were found after 4 h.\textsuperscript{199} The fall in TER in the presence of cadmium was reduced if testosterone and FSH were added.\textsuperscript{198} The disruption
of the barrier is associated with a transient increase in testicular TGF-β2 and 3 and the phospho-rylated p38 mitogen activated protein (MAP) kinase, concomitant with a loss of occludin and ZO-1 from the barrier site. There is also a surge in α2-macroglobulin at the Sertoli-Sertoli cell junctions at the time of disruption of the barrier.

It is interesting that there are some strains of mice whose testes are much more resistant to the effects of cadmium, and this is associated with reduced transport of cadmium into the testes. The cadmium transporter is saturable and can be competitively inhibited by zinc, but not calcium, and appears not to be associated with any tubular cells, but is probably located in the endothelial cells.

The integrity of the blood-testis barrier is altered by intratesticular treatment of rats with cytochalasin D, a known microfilament inhibitor. Evidence for this was obtained from studies on the penetration of electron-opaque markers, from the effects of perfusion with hypertonic solutions and from the entry of radioactive inulin into seminiferous tubular fluid.

Another substance which has been shown to disrupt the blood-testis barrier is glycerol when injected into the testes of rats. These animals showed increased entry of radioactive inulin and albumin into seminiferous tubular and rete testis fluids, and also disrupted tight junction-associated actin microfilaments, occludin and microtubules in the Sertoli cells.

Other substances which appear to affect the blood-testis barrier include hexanedione, cis-platinum, sarin, and DEET but stainless steel corrosion products affects spermatogenesis without apparently interfering with the blood-testis barrier. Other treatments such as bisphenol A or Adjudin (AF 2364) disrupt the junctions between Sertoli cells and spermatids without affecting the blood-testis barrier. Freunds complete adjuvant injected into guinea pigs 7 days previously increased the entry of horseradish peroxidase into the seminiferous tubules.

Temperature and Cryptorchidism

The entry of radioactive albumin into rete testis fluid of rats was unaffected during or following heating of the testes, but the entry of K, Rb, Na, lysine and some steroids was increased during heating. The entry of Cr-EDTA into the tubules was not affected when spermatogenesis had been disrupted in rats by local heating of their testes. In surgically-induced cryptorchidism in rats, the blood-testis barrier appears to remain intact, but in spontaneous cases in humans, the penetration of lanthanum between the Sertoli cells depended on the extent of the loss of germ cells. In other conditions of spermatogenic cycle breakdown in humans, lanthanum entry is increased in maturation arrest and in irregular hypospermatogenesis, but in germ cell aplasia the barrier remains efficient.

Mutants and Hybrids

The blood-testis barrier is less efficient in Tfm and Sxr mice, but normal in Mo+/Y and Gy/Y mutants. There are defects in both the germ cells and in the blood-testis barrier in as-mutant rats, as demonstrated by the distribution of cytochrome-c in the testis, as well as from studies involving spermatogonial transplantation. The blood-testis barrier is deficient in hybrids between blue and silver foxes, and spermatogenesis is arrested at early pachytene.

Significance of the Blood-Testis Barrier

As has already been discussed, there are several obvious consequences of the operation of the blood-testis barrier. The first is immunological. The barrier isolates the developing germ cells from circulating antibodies in the bloodstream. It also means that the body’s immunological system does not “see” the haploid germ cells, and therefore a male can be immunized against his own spermatozoa. However, the isolation is not complete and Tung has concluded that “tissue barriers and antigen sequestration are important but not sufficient to protect germ cell antigens and prevent experimental allergic orchitis”. Some germ cells outside the barrier can certainly provoke an immunological reaction, even
peritubular cells, leading to autoimmune orchitis. Furthermore, mice immunized with syngeneic testis antigen have IgG deposits surrounding cells at the periphery of about half the tubule cross-sections, particularly those at stage 7 to 12. Also sera from testis-immune orchidectomized donors are able to transfer IgG passively into the testes of normal syngeneic recipients in an antigen-specific manner, although there is evidence that the rete testis and tubuli recti are the sites of the earliest and most frequent lesions. Therefore, other factors must be involved in making rodent testes, but not those of sheep or monkeys, immunologically privileged sites. Possible factors have been discussed recently by Hedger.

The second effect of the barrier relates to the endocrine system. Peptide hormones such as FSH and LH do not instantaneously pass from the blood even into the extracellular interstitial fluid, so that the Leydig cells begin to respond to a rise in blood LH even before there is any change in the LH levels in the immediate vicinity of these cells. FSH on the other hand acts principally on the Sertoli cells, and therefore must penetrate both the endothelial cell and peritubular cell layers. This is probably less important as the concentration of FSH does not seem to show such pronounced peaks as LH does, and therefore changes in its concentrations in blood are more likely to be reflected in the concentrations at the basal surface of the Sertoli cells.

The situation with steroids is less clear cut. Because of their relatively high lipid solubility, they should pass more readily through the barrier than the hydrophilic peptides, but there is some evidence that the concentration of testosterone in RTF and STF does not change as much as that in blood. This may suggest that there is a transport system for steroids in the tubules, but no further evidence for this idea has been presented. It is clear from the relative concentrations inside and outside the tubules that the androgen-binding protein secreted by the Sertoli cells preferentially inside the barrier certainly does not produce a higher concentration of the total (free plus bound) steroid there. In fact the concentration of free testosterone may be appreciably lower in STF. Conjugated steroids, which are produced in large amounts in the testes of some species such as pig and horse tend to be less lipophilic than the free steroids and therefore remain in higher concentrations in the interstitial extracellular fluid than inside the barrier.

Glucose is transported across the barrier by a transport system the capacity of which appears to be less than the capacity of the Sertoli cells to convert the sugar to lactate. The consequence of this is that there is very little glucose in the fluid inside the tubules and the developing germ cells prefer to metabolize lactate even in vitro.

An interesting recent development has been the identification of a number of specific transport proteins for xenobiotics in various cells in the testis. These transporters, Pgp and MDR have important consequences in determining whether a particular toxicant will affect spermatogenesis, but in the case of transferrin, it may result in the accumulation of mutagenic substances in the environment of the germ cells.

One of the most interesting aspects of the function of the blood-testis barrier is the fact that it cannot remain closed all the time, but must open at different points along each tubule at specific times in the spermatogenic cycle to allow developing spermatocytes to pass from the basal to the adluminal compartment. How this is achieved is still a matter of debate. Four theories have been advanced to explain this phenomenon: zipper, intermediate compartment, repetitive removal of membrane segments and junction restructuring. However, junction disassembly and reassembly seems to be the most likely explanation. Opening of the Sertoli-Sertoli cell junctions in a limited part of the tubule must occur without affecting the Sertoli-Sertoli cell junctions elsewhere in that tubule or Sertoli cell-germ cell adherens junctions in the same and other parts of the tubule. It appears that cytokines may be involved and o2-macroglobulin also appears to play a part. One of the most intriguing questions which remains to be answered is how spermatogonia injected into the lumen of a single seminiferous tubule either directly or via the rete testis can pass between pairs of Sertoli cells to take up a position adjacent to the peritubular tissue and repopulate that area of the tubules with developing germ cells. The recipient animals...
have usually been treated with busulfan or irradiation to eliminate endogenous spermatogenesis, but nothing appears to be known about the efficacy of the barrier in these animals. It is interesting that transmissible leukemic cells, when injected together with testicular cells into the tubules through the rete in normal rats can reach the intertubular tissue where they resume their uncontrolled multiplication and make the recipient animals leukemic.236

It has also been repeatedly stated that the specialized environment created by the barrier may be necessary for the germ cells to proceed through meiosis. However, just what these conditions are has yet to be defined, but the fact that spermatogenesis can proceed, albeit to a limited extent, in aggregates of testicular cells encaised in alginate237 may indicate that as long as Sertoli and germ cells are in reasonably close association, that is sufficient.

One fascinating possibility is that retinoic acid (RA), not derived from blood but newly formed from retinol by a two-stage process, may be involved in the switch of the germ cells from mitosis to meiosis in the testis. The first stage of this conversion involves the Sertoli cells and the second the germ cells.69,238 RA has been shown to cause the germ cells in the fetal ovary to enter meiosis, while in the fetal testis, meiosis is inhibited by destruction of RA by Cyp26 b1.239,240 The same enzyme that in the myoid cells, prevents the entry of RA into the tubules.68

A premeiotic germ cell-specific cytoplasmic protein encoded by the RA-responsive gene Stra8 is present in only less than half the tubule cross-sections in a mouse testis,241 although unfortunately these authors did not identify the stage of spermatogenesis at which this protein was expressed. It is interesting that in mice in which the gene for p27kip1 is knocked out, spermatocytes were often arrested at preleptotene242 and this mitotic inhibitor can be induced in cultured Sertoli cells by RA.243 Furthermore, the expression of mRNA for CRBP is highest in spermatogenic stage IX to XIV, when most of the mitoses in the tubules occur. Preleptotene spermatocytes appear first at stage VII but CRBP mRNA rises significantly only in stage VIII,157 when the meiotic DNA synthesis is occurring. The expression of mRNA for the retinoic acid receptor RARα is also highest at stage VIII in the rat testis, and this receptor is present in preleptotene spermatocytes as well as in round spermatids.244 This receptor is required for synchronization of the spermatogenic cycle, and in its absence, preleptotene spermatocytes do not proceed to leptotene in the first, second and third waves.245,246 However, others have shown that in mice lacking plasma RBP, Vitamin A deficiency does not delay the entry of preleptotene spermatocytes into meiosis, while spermatogenesis is blocked by delayed or arrested differentiation of spermatogonia.247-250 This suggests that Vitamin A may have several functions in the testis, and furthermore, there may be important difference between mice and rats in the responses of their testes to Vitamin A deficiency.247

The observation that when spermatogenesis is restored in previously Vitamin A-deficient rats, spermatocytes progress to pachytene but then degenerate until the barrier is reformed,186 would add emphasis to the need for the barrier for complete meiosis. Likewise, the finding that the barrier is disrupted by the injection of a 22-amino acid peptide corresponding to the second loop of occludin, accompanied by a cessation of spermatogenesis82 would strongly emphasise the importance of the barrier for spermatogenesis.

However, as already mentioned, there are a number of conditions in which spermatogenesis is disrupted but the barrier function appears to be intact, suggesting that other factors are also important for normal sperm production. Nevertheless, the blood-testis barrier remains an important factor in the physiology of the testis, in particular in relation to spermatogenesis.

**Future Directions**

There are a number of lines of research on the blood-testis barrier which could yield important results in the future. First, possible roles of the endothelial and peritubular cells in regulating entry of substances into the testis or of influencing the Sertoli cell barrier need reevaluating. This is because of the many peculiarities of the testicular endothelial cells, many of which they share with brain endothelial cells, the site of the blood-brain barrier3,4 and the recent demonstration of transport systems for urea in the peritubular cells. Studies on endothelial cells should
now be possible following the recent demonstration that these cells can be isolated from rat testes, and that when cocultured with interstitial cells, the endothelial cells enhance the production of testosterone.\textsuperscript{251} Techniques for isolation and culture of peritubular cells have been available for some years.\textsuperscript{63,64} While cocultures of peritubular and Sertoli cells have been used\textsuperscript{252} to study basement membrane gene expression, and the effect of proteins from pachytene spermatocytes\textsuperscript{253} and spermatids\textsuperscript{254} have been used to study their effects on secretion by Sertoli cells, no-one appears to have used cocultures of peritubular or germ cells and Sertoli cells in bicameral chambers (as illustrated in Fig. 1B,C in ref. 46) to study the effects of other cells on barrier function.

However, probably the most interesting problem in this area is the mechanism by which the Sertoli cell barrier is opened and closed again to allow the passage of the developing germ cells. Various theories have been advanced\textsuperscript{48} but more evidence is needed on local factors controlling the distribution of this process in relation to spatially and temporally determined stages of spermatogenesis. Related to this problem is the need for an explanation of the occurrence in fetal testes of the structural proteins associated with the Sertoli-Sertoli cell junctions, occludin and claudin.

One new area of interest in relation to the blood-testis barrier is the involvement of specific transport proteins, such as Pgp and MDR. These may have important toxicological consequences in determining whether a particular compound disrupts spermatogenesis. It is conceivable that toxins could either be normally excluded or concentrated inside the tubules by these transporters, and further information on their distribution and specificity is needed. This may be particularly important for the disruptors of the barrier, cadmium salts and glycerol, and studies on the transport of these substances should be undertaken.

Finally, there is the old question of the role of the barrier in creating the conditions necessary for meiosis which needs further study. Recent progress in stem cell transplantation\textsuperscript{235} and in vitro spermatogenesis\textsuperscript{237} may provide the tools for further study of this fascinating problem.

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Cross-Talk between Tight and Anchoring Junctions—Lesson from the Testis

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Abstract

Spermatogenesis takes place in the seminiferous tubules in adult testes such as rats, in which developing germ cells must traverse the seminiferous epithelium while spermatogonia (2n, diploid) undergo mitotic and meiotic divisions, and differentiate into elongated spermatids (1n, haploid). It is conceivable that this event involves extensive junction restructuring particularly at the blood-testis barrier (BTB, a structure that segregates the seminiferous epithelium into the basal and the adluminal compartments) that occurs at stages VII-VIII of the seminiferous epithelial cycle. As such, cross-talk between tight (TJ) and anchoring junctions [e.g., basal ectoplasmic specialization (basal ES), adherens junction (AJ), desmosome-like junction (DJ)] at the BTB must occur to coordinate the transient opening of the BTB to facilitate preleptotene spermatocyte migration. Interestingly, while there are extensively restructuring at the BTB during the epithelial cycle, the immunological barrier function of the BTB must be maintained without disruption even transiently. Recent studies using the androgen suppression and Adjudin models have shown that anchoring junction restructuring that leads to germ cell loss from the seminiferous epithelium also promotes the production of AJ (e.g., basal ES) proteins (such as N-cadherins, catenins) at the BTB site. We postulate the testis is using a similar mechanism during spermatogenesis at stage VIII of the epithelial cycle that these induced basal ES proteins, likely form a “patch” surrounding the BTB, transiently maintain the BTB integrity while TJ is “opened”, such as induced by TGF-b3 or TNFa, to facilitate preleptotene spermatocyte migration. However, in other stages of the epithelial cycle other than VII and VIII when the BTB remains “closed” (for ~10 days), anchoring junctions (e.g., AJ, DJ, and apical ES) restructuring continues to facilitate germ cell movement. Interestingly, the mechanism(s) that governs this communication between TJ and anchoring junction (e.g., basal ES and AJ) in the testis has remained obscure until recently. Herein, we provide a critical review based on the recently available data regarding the cross-talk between TJ and anchoring junction to allow simultaneous maintenance of the BTB and germ cell movement across the seminiferous epithelium.

Introduction

In adult rats, spermatogenesis takes place in the seminiferous tubules in testes, in which spermatozoa (1n, haploid) differentiate from spermatogonia (type A, 2n, diploid) in the seminiferous epithelium in ~58 days via six mitotic and two meiotic divisions consecutively.1,2 During this time, germ cells also undergo different phases of cellular differentiation with changes
Figure 1. The physiological relationship along the hypothalamic-pituitary-testicular axis that regulates testicular function, and the intimate structural relationship between Sertoli and germ cells in the seminiferous epithelium during spermatogenesis. A) This is a schematic drawing that shows the hormonal regulation of spermatogenesis in the testis via the hypothalamic-pituitary-testicular axis. LHRH released from the hypothalamus stimulates the production of LH and FSH by the pituitary gland, which in turn regulates Leydig and Sertoli cell functions, residing in the interstitium and the seminiferous epithelium, respectively. Testosterone (T) and inhibin (and others, e.g., follistatin) released by Leydig and Sertoli cells, respectively, provide the necessary feedback loops to maintain the endogenous levels of LH and FSH to regulate spermatogenesis. Any changes in the homeostasis of these hormones can perturb spermatogenesis. For instance, a reduction of intra-testicular T level can suppress spermatid (step 8 and beyond) adhesion, leading to spermatid depletion from the epithelium without disrupting the blood-testis barrier (BTB) integrity. The right panel represents a stage II-VI seminiferous tubule dissected from the testis by using a transilluminating stereomicroscope as described. Legend continued on following page.
in morphology particularly at the head region associated with chromatin condensation, as well as tail elongation (see Fig. 1). Earlier studies using PAS (periodic acid-Schiff reaction) staining has divided these cellular changes in the seminiferous epithelium in adult rats into 14 stages, which, in turn, are comprised of one seminiferous epithelial cycle, from stages I to XIV. Each stage displays a unique pattern of association of germ cells at different stages of their development with Sertoli cells in the seminiferous epithelium.\textsuperscript{1-6} If an investigator is visualizing a specific section of the tubule at stage I in the epithelium under a transillumination stereomicroscope, this stage progresses to II through XIV and returns to stage I in ~12.9 days in rats. As such, a type A spermatogonium requires to complete the epithelial cycle 4.5 times in ~58 days utilizing much of these time for mitosis before it divides and matures into 256 spermatozoa.\textsuperscript{1,4} This process is precisely regulated under FSH, LH, and testosterone via coordination between the hypothalamus, pituitary gland, and the testis, which is known as the hypothalamic-pituitary-testicular axis (see Fig. 1).

Besides these well defined hormonal events that occur along the hypothalamic-pituitary-testicular axis, spermatogenesis cannot proceed unless preleptotene spermatocytes residing in the basal compartment can traverse the blood-testis barrier (BTB) at late stage VII through early stage VIII (these two cycles combined last for ~3.5 days in rats), entering into the adluminal compartment for further development\textsuperscript{7} (Fig. 1). The BTB is a functional term initially coined by Chiquoine\textsuperscript{8} and others\textsuperscript{9} to illustrate the presence of a physiological barrier in the testis. The tightness of the BTB is comparable to the blood-brain barrier. It prevented staining of the seminiferous tubules when dyes (e.g., acriflavine used by Korman\textsuperscript{9}) were injected into animals (for a review, see ref. 10). Subsequent studies by Setchell and colleagues have illustrated many unique features of the BTB versus other barriers in mammals (e.g., the blood-brain barrier or the blood-retina barrier).\textsuperscript{10} For instance, besides Sertoli cells, peritubular myoid cells in rodents contribute to the BTB function by limiting the diffusion of lanthanum and colloidal carbon,\textsuperscript{11,12} however, in other blood-tissue barriers, the barrier function is contributed almost exclusively by the epithelial or endothelial cells alone.

Several hypotheses regarding the movement of germ cells across the BTB have been suggested. For instance, the “zipper theory” proposes that during germ cell migration across the BTB, new occluding fibrils form below the preleptotene/leptotene spermatocytes, to be followed by the break down of occluding fibrils above these spermatocytes.\textsuperscript{5,13} The “intermediate cellular compartment theory” of Russell\textsuperscript{7,14} suggests the presence of a unique compartment inhabited by the preleptotene/leptotene spermatocytes in transit where these spermatocytes are trapped in between two occluding zonules as confirmed by electron microscopy. The “repetitive removal of membrane segments theory” of Pelletier\textsuperscript{15} suggests that the upward movement of the migrating germ cells create stresses on the Sertoli cell junctional complexes, which, in turn, induces the formation of intercellular pockets. Each developing germ cell (except elongating spermatid) is trapped in one of these intercellular pockets, sealed at both ends by tight junction (TJ), and anchoring junctions [e.g., adherens junction (AJ) and desmosome-like junctions (DJ)]. Note: while DJ is an intermediate filament-based anchoring junction that localizes to the BTB, virtually...
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no functional study is found in the literature regarding this junction type. Also, the molecular composition is virtually unknown, and as such, DJ is not included in our remaining discussion herein] and moves progressively across the seminiferous epithelium until it is internalized in autophagic vesicles. However, neither the “zipper”, the “intermediate cellular compartment”, nor the “repetitive removal of membrane segment” theories can perfectly describe the processes of preleptotene spermatocyte migration across the BTB during stage VIII of the epithelial cycle, taking into account the interactions between integral membrane proteins at the BTB, their peripheral adaptors, kinases, phosphatases, cytokines, proteases and protease inhibitors.3,5,6,14-17 This is not entirely unexpected since these postulates by and large were based on morphological studies without any biochemical and/or molecular data available at the time. A recent hypothesis known as the “junction restructuring theory” has provided a more solid basis in delineating the complicated biochemical and molecular cascade of events behind this phenomenon (for a review, see refs. 6,17). The essence of this hypothesis suggests that germ cell movement is composed of waves of junction disassembly and reassembly that occur under the influence of cytokines (e.g., TGF-β3 and TNF-α) that are released by Sertoli and/or germ cells into the microenvironment to facilitate cell movement. For instance, these cytokines regulate the steady-state levels of proteins at the BTB (e.g., occludin, ZO-1, cadherins and catenins) and the homeostasis of proteases and protease inhibitors in the epithelium.17 The net result of these interactions determines the “opening” or the “closing” state of the BTB to facilitate germ cell movement. This biochemical theory, if confirmed and adequately characterized, should provide new clues for developing novel male contraceptives, such as by “shutting-down” the BTB to prohibit the migration of preleptotene spermatocytes from the basal to the adluminal compartment. Recent studies have also illustrated that this hypothesis fits quite well with the “zipper theory” (see below for further discussion).

The Concept of Endocytosis in BTB Dynamics

In other epithelia, endocytosis has been shown to play a pivotal role in junction restructuring to facilitate cell migration.18-20 This mechanism also provides the efficient means for restructuring junctional complexes at the TJ-barrier by perturbing its permeability without requiring de novo protein synthesis.18 For instance, it is suggested that intercellular movement is achieved by coendocytosis of apposing TJ-integral membrane proteins into the adjacent cells in a study using mouse Eph4 cells in vitro.19 Besides, there is growing evidence that the “leaky” intestinal epithelia in chronic disorders is mediated by abnormal internalization of TJ-integral membrane proteins via endocytosis.21,22 Interferon-γ (IFNγ), a proinflammatory cytokine, was shown to increase the intestinal permeability through its effects on TJ proteins such as occludin, junctional adhesion molecule-A (JAM-A) and claudin-1 in T84 cells.22,23 For instance, TJ proteins were internalized into large vacuoles via the RhoA/ROCK signaling pathway.22

In the testis, endocytosis is also employed by Sertoli cells for house-keeping activities, such as phagocytosis of germ cells undergoing apoptosis or anoikis. This process is also used to eliminate residual bodies from elongating spermatids prior to spermiation when fully mature spermatids (spermatozoa) are emptied into the tubular lumen.17 At present, it is not clear if testes employ similar mechanism as other epithelia to regulate the rapid turnovers of TJ- and AJ-integral membrane proteins at the BTB to facilitate germ cell migration. Obviously, this can certainly be possible. Cytokines, such as TGF-β and TNF-α, have been shown to increase BTB permeability via different signaling pathways.24,25 However, the mechanism(s) that reduces the TJ protein levels following cytokine treatment is not known. Does this simply the result of a decline in de novo protein synthesis or involve internalization of TJ-integral membrane proteins? Recent findings have suggested that endocytosis does occur in the testis at the apical ES to facilitate spermiation. For instance, the formation of tubulobulbar complexes (TBC, another testis-specific AJ type) at the Sertoli cell-elongated spermatid interface at late stage VIII of the epithelial cycle prior to spermiation by replacing the apical ES has been speculated to assist the release of mature spermatids via internalization of TBC-junction molecules.16 This postulate has been confirmed in a recent study26 which showed that the TBC indeed
appeared at the concave surface of the head of spermatids that was previously occupied by apical ES. Furthermore, the adhesion domains of nectin-2 and 3 were found to be internalized as membrane vesicles near the TBC at spermiation.\textsuperscript{26} The fate of these internalized adhesion molecules at TBC remains to be determined; they can either be degraded or recycled back to the cell surface. In other epithelia, most of the endocytosed molecules (e.g., E-cadherin and occludin) enter the recycling pathway so that they can be rapidly recycled back to the cell surface to maintain junction integrity, especially in unstable cell-cell contacts. For instance, a significant increase in E-cadherin recycling was detected in MDCK cells during [Ca\textsuperscript{2+}]-depletion-induced loss of cell adhesion.\textsuperscript{27,28}

It is conceivable that internalization of TJ and/or AJ integral membrane proteins takes place at the BTB between adjacent Sertoli cells. However, this would be a more complicated event since TJ, AJ (e.g., basal ES and basal TBC), and desmosome-like junctions are coexisting at the BTB (see Figs. 1, 2). In other epithelia, it is believed that endocytic recycling of TJ proteins (e.g., occludin) is mediated by Rab GTPases.\textsuperscript{28} and Rab8B was shown to structurally associate with membranous structures that are involved in trafficking between the plasma membrane and the early and late TGN.

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Figure 2. Electron micrograph of normal rat testes illustrating the typical BTB ultrastructural features. The features shown herein for BTB from an adult rat are applicable to mice and most other mammals. The basement membrane (asterisk), which is part of the tunica propria, appears as a homogeneous substance. This is a modified form of extracellular matrix (ECM)\textsuperscript{42} and is clearly visible underneath the collagen (type 1) fibrils. The BTB is composed of coexisting tight junction (TJ, blue arrowheads), basal ES (bracketed in red), desmosome-like junction (DJ, bracketed in green, see electron dense substances on both sides of the apposing Sertoli cell plasma membranes typified of desmosome), and adherens junction (AJ, bracketed in yellow). Basal ES refers to the ultrastructure between two adjacent Sertoli cells typified by the presence of actin bundles (red arrowheads) sandwiched between the endoplasmic reticulum (ER) and the plasma membrane that are found on both sides of the two adjacent Sertoli cells (SC). Bar, 0.2 \textmu m.
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Figure 3. A schematic drawing depicting the currently known protein complexes at the apical ectoplasmic specialization (ES) and the BTB. Apical ES (top panel) is limited to the interface between Sertoli cells and spermatids (step 8 and beyond) in the apical compartment of the seminiferous epithelium in adult rat testes. As discussed in the text, the apical ES is a testis-specific hybrid cell-cell actin-based adherens junction (AJ), tight junction (TJ), and cell-matrix focal contact anchoring junction type since it contains protein complexes that are found in AJ (e.g., cadherins/catenins, nectins/afadin), TJ (e.g., JAM/CAR/ZO-1), and focal adhesion complex (FAC) (e.g., αβ1 integrin/laminin-333) of other epithelia. Some of the peripheral adaptors (e.g., α- and β-catenins, ZO-1), kinases (c-Src) and signaling molecules (e.g., p120ctn) that are known to associate with these complexes in adult rat testes are also shown. The BTB (bottom panel) is found between two adjacent Sertoli cells near the basement membrane that segregates the seminiferous epithelium into the adluminal and the basal compartments (see Figs. 1C, 2). In adult rat testes, the BTB is constituted by coexisting tight and anchoring junctions (see Figs. 1C, Fig. 2). The BTB is composed of protein complexes that are found in TJ (e.g., JAMs/ZO-1, claudins/ZO-1, occludin/ZO-1), AJ/basal ES/basal TBC (e.g., N-cadherin/catenin), desmosome-like junction (e.g., connexins, desmogleins, Dsg), desmocollins (Dsc)) of other epithelia. This drawing was prepared based on several recent papers and reviews.17,74,89,111-113 Because of space limitation, only the integral membrane proteins and some of the better studied adaptors, kinases and signaling molecules are shown.
with E-cadherin in the rat testis at both the BTB and the apical ES\(^2\) (see Fig. 3). Moreover, other than intracellular movement, endocytosis can also mediate the disassembly of cell-matrix focal adhesion to facilitate cell movement along the extracellular matrix.\(^3\) Interestingly, the basal ES is a hybrid cell-cell and cell-matrix anchoring junction type because it is composed of proteins that are usually restricted to focal adhesion complex at the cell-matrix interface, such as FAK and vinculin.\(^4\) The disassembly of focal adhesion at the fibroblast-matrix interface is independent of Rho GTPases, instead, another GTPase called dynamin is involved. Dynamin is known to play a pivotal role in endocytic process. Among the three isoforms of dynamin, dynamin-2 and 3 are highly expressed in the testis, particularly in Sertoli and germ cells\(^5\) while dynamin-1 is neuron-specific. It is likely that dynamin may be involved in the internalization of junction molecules at the BTB. Ubiquitination is another physiological process that has recently been shown to play a crucial role in junction protein turnover using Sertoli cells cultured in vitro.\(^6\) For instance, ubiquitination is used to facilitate the rapid turnover of occludin at the inter-Sertoli TJ-barrier by targeting the ubiquitin-conjugating enzymes, such as Itch and UBC4, to occludin to induce its degradation by proteasomes or lysosomes.\(^7\) Obviously, much research is needed in this area to investigate the role(s) of integral membrane protein recycling and ubiquitination at the BTB to facilitate germ cell movement. These studies will also identify new targets for male contraceptive research, such as by disrupting the events of protein recycling and/or internalization at the BTB to perturb spermatogenesis.

**Some Unique Physiological Phenomena at the BTB: Unidirectional and Bidirectional Cross-Talk between TJ and Anchoring Junction (e.g., Basal ES and AJ)**

In multi-cellular organisms, cell-cell adhesion in epithelia is mediated through junction complexes which are constituted by TJ, AJ, desmosomes and gap junctions (for a review, see ref. 36). Each of these junction types has a specific physiological role. For instance, the TJ functions as a barrier to limit the passage of ions, water and other molecules between cells, and maintains cell polarity. The AJ links neighboring cells together by forming a continuous adhesion belt, desmosomes serve as an anchoring site and unite cells together while the gap junction is for communication between cells. Interestingly, BTB is unique in its morphological layout versus other blood-tissue barriers found in other epithelia/endothelia (for reviews, see refs. 2,3,5,6). In brief, in other epithelia/endothelia excluding the seminiferous epithelium, junctions are organized typically from apical to basal, in which TJ is present at the most apical region, to be followed by AJ and then desmosomes (for reviews, see refs. 37,38-40). As such, these junctions, in particular TJ, are furthest away from ECM. However, the TJ in the testis is located at the basal compartment of the seminiferous epithelium, closest to the basement membrane, a modified form of ECM\(^6\) and coexisting with AJ, basal ES, basal TBC (both basal ES and basal TBC are testis-specific AJ types restricted to the Sertoli-Sertoli cells interface at the BTB), gap junctions, and desmosome-like junctions (see Figs. 1, 2).\(^3\,\,16\,\,17\,\,43-45\)

Apart from the unique morphological features of BTB, the testis apparently is using a specialized junction restructuring mechanism in assisting preleptotene spermatocytes to migrate across the BTB, such that the post-meiotic germ cell antigens can be sequestered from the immune system while permitting continuous restructuring of the TJ-barrier and/or anchoring junction (e.g., basal ES and AJ) at the BTB. This apparently is in contrast to other blood-tissue barriers (e.g., blood-brain and blood-retina barriers), even though they are suggested to be dynamic in nature.\(^8\) Unlike the BTB, these blood-tissue barriers do not undergo such drastic restructuring to facilitate the migration of preleptotene spermatocytes which are 10-15 μm in diameter. Besides, except for stage VIII of the epithelial cycle, the BTB remains closed to limit the entry of preleptotene spermatocytes into the adluminal compartment. This implicates that there are unique molecules and mechanisms in place to coordinate these events.

In the past decade, different in vivo models have been used to study BTB dynamics and regulation.\(^17\) There are a number of interesting observations stemming from this body of work.
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For instance, studies using CdCl$_2$,\textsuperscript{46-48} an environmental toxicant, and glycerol\textsuperscript{49-51} have shown that these toxicants can irreversibly perturb TJ which, in turn, lead to anchoring junction damage, dislodging germ cells of all classes from the epithelium. These studies thus illustrate the presence of cross-talk between tight and anchoring junctions that a disruption in TJ can spread to anchoring junction such as apical ES, which is located at the opposite end of the TJ in the seminiferous epithelium (see Fig. 1C). These findings are consistent with the cellular physiology in other epithelia, such as the skin, in which a disruption of TJ leads to underlying AJ damage and vice versa,\textsuperscript{52-54} illustrating bi-directional cross-talk between TJ and anchoring junction. However, in studies using antispermatogenic drugs such as Ad judin\textsuperscript{55} and londamide,\textsuperscript{56} it was shown that anchoring junction (e.g., apical ES) disruption could be limited to the interface between Sertoli cells and elongating spermatids/elongate spermatids/round spermatids/spermatocytes (but not spermatogonia) without compromising anchoring (e.g., basal ES) and tight junctions between Sertoli cells at the BTB,\textsuperscript{17,57,58} illustrating unidirectional cross-talk between TJ and anchoring junction in the testis. These findings were also supported in a recent study using the model of McLachlan and O’Donnell by suppressing the endogenous testosterone level in the testis using testosterone/estradiol (TE) implants.\textsuperscript{59,60} For instance, suppression of intratesticular androgen level by using TE implants in adult rats can lead to disruption of apical ES at the Sertoli-germ cell interface (step 8 spermatids and beyond) without perturbing the BTB nor anchoring junctions between Sertoli cells and round spermatids/spermatocytes/spermatogonia.\textsuperscript{60,61} Indeed, using the androgen suppression model, we have confirmed that a disruption of apical ES can be limited to the Sertoli cell-germ cell interface without compromising the BTB integrity nor the anchoring junction between Sertoli cells and round spermatids/spermatocytes in the seminiferous epithelium.\textsuperscript{62} In short, these findings are drastically different from other blood-tissue barriers (e.g., in the skin) since AJ disruption in these barriers always leads to TJ damage,\textsuperscript{53,54} and the assembly and integrity of TJ depend on AJ.\textsuperscript{63,64} For instance, when the reassembly of junctional complexes in intestinal cells is initiated by an incubation of high-calcium medium, AJ is reassembled within ~30 min prior to TJ assembly.\textsuperscript{65} As such, these results illustrated that this unidirectional cross-talk mechanism is operating at the BTB in which a disruption of Sertoli-germ cell adhesion can be restricted to anchoring junction (e.g., apical ES) without interfering with TJ nor anchoring junction (e.g., basal ES) at the BTB.

Changes in Protein-Protein Interactions between Integral Membrane Proteins and Their Adaptors, and Cross-Talk between Junctions at the BTB in the Regulation of BTB Dynamics

To address the observations regarding the cross-talk between TJ and AJ (e.g., basal ES) at the BTB that regulate its timely opening to facilitate preleptotene spermatocyte migration at stage VIII of the cycle, recent studies have shifted the focus on the role of protein-protein interactions in BTB dynamics. Studies based on two different in vivo models, namely the Ad judin\textsuperscript{32,55,57} and the androgen suppression model have yielded some interesting observations that leads to a hypothesis to be tested in future studies. First, in the testosterone suppression model of O’Donnell,\textsuperscript{60,61} in which subdermal testosterone/estradiol implants were placed in adult rats to suppress the intratesticular T level, it was shown that a loss of testosterone in the seminiferous epithelium selectively perturbed apical ES junction (from step 8 spermatids and beyond), leading to spermatid depletion from the seminiferous epithelium. While the BTB integrity and functionality were not compromised,\textsuperscript{62} there were significant changes at the BTB as well, reminiscence of the seminiferous epithelium during spermatogenesis, such as stage VIII of the epithelial cycle. For instance, at the time of apical ES restructuring that led to spermatid depletion from the epithelium, there were enhanced production of several basal ES proteins (e.g., cadherins, integrins, JAM-A).\textsuperscript{57,66,74} Second, these observations were consistent with other findings using the Ad judin model in which the Ad judin-induced anchoring junction restructuring in the seminiferous epithelium also led to progressive depletion of germ cells from the seminiferous epithelium, beginning with
elongating/elongate spermatids, to be followed by round spermatids and spermatocytes.\textsuperscript{6,75} However, the BTB apparently remained “intact”.\textsuperscript{17} Likewise, an induction of an array of proteins at the BTB was also detected, such as cadherins, catenins, integrins,\textsuperscript{32,74-76} illustrating restructuring indeed occurred at the BTB, but unlike the apical ES at the Sertoli cell-spermatid interface that led to spermatid loss from the epithelium, such BTB restructuring did not lead to functional damage which compromise the immunological barrier, reminiscent of the seminiferous epithelium at stage VIII of the epithelial cycle. We thus postulate that these transiently induced basal ES proteins that form a “patch” are being used to temporarily “reinforce” the TJ-barrier function at the BTB during androgen suppression- or Adjudin-induced junction restructuring in the seminiferous epithelium so that the BTB integrity can be maintained as observed in these models. However, this transient induction in basal ES proteins can also “supersede” the TJ-barrier function temporarily when TJ must open to accommodate preleptotene spermatocyte migration at stage VIII of the epithelial cycle. This speculation was also supported by a study using fluorescent microscopy. For instance, in studies using the androgen suppression model, it was shown that during the androgen depletion-induced spermatid (step 8 and beyond) loss, the increased basal ES proteins (e.g., N-cadherin, β-catenin) at the BTB had surrounded the entire basal compartment, forming a transient “patch”, and extended into the seminiferous epithelium well beyond the BTB site.\textsuperscript{57,62,66} As such, these basal ES proteins could “patch” the TJ-barrier, reinforcing the BTB integrity, which is also the mechanism being used to maintain the transiently disrupted TJ-barrier at the Sertoli-Sertoli interface at stage VIII of the cycle, facilitating preleptotene spermatocyte migration while maintaining the immunological barrier at the BTB. It is obvious that this possibility must be carefully evaluated and investigated in future studies. While functional data are not yet available to support this “patch” hypothesis, recent findings in the field seem to favor this postulate. For instance, several TJ proteins, such as JAM-C and coxsackievirus and adenovirus receptor (CAR) are putative integral membrane proteins at the TJ of multiple epithelia,\textsuperscript{67-70} yet recent studies have shown that they are also putative Sertoli and/or germ cell products that are restricted to the apical ES.\textsuperscript{71,72} A recent report has also identified the presence of JAM-A in germ cells,\textsuperscript{73} which is also detected at the BTB colocalizing with ZO-1 by fluorescent microscopy,\textsuperscript{62} suggesting that it is likely a component of the basal ES at the BTB. These findings are significant because they illustrate that ectoplasmic specialization (e.g., apical and basal ES), a testis-specific AJ type, is indeed a hybrid of AJ and TJ as it consists components of TJ-integral membrane proteins. Thus, these TJ component proteins at the basal ES can confer the necessary tight junction functionality, however, when the TJ fibrils are disrupted, the transiently induced basal ES proteins serve as a transient “patch”. Nonetheless, this concept must be vigorously investigated in future studies.

One may argue that since there was a surge in anchoring junction proteins in the seminiferous epithelium, and these proteins were being used to “reinforce” the TJ-barrier function at the BTB, why would such an increase in anchoring junction protein levels (e.g., cadherins and catenins) in the seminiferous epithelium fail to retain germ cells outside the BTB. Based on limited available data, it seems that this loss of cell adhesion is the result of a loss of protein-protein association between cadherins and catenins at the Sertoli-germ cell interface.\textsuperscript{66} This is likely mediated by a surge in tyrosine phosphorylation of β-catenin, resulting in a reduction in its adhesive activity.\textsuperscript{66} Other studies have shown that tyrosine phosphorylation of AJ proteins (e.g., cadherins) facilitates junction disassembly in various epithelia,\textsuperscript{77,78} as well as in pathological conditions, such as tumor metastasis,\textsuperscript{79} whereas Ser/Thr phosphorylation, on the contrary, promotes adhesion. In addition, other molecules are operating side-by-side to regulate BTB. For instance, the integrity of endothelial TJ-barrier was found to be partly regulated by reactive oxygen species, such as hydrogen peroxide. These reactive oxygen species up-regulate tyrosine phosphorylation of FAK and paxillin (note: these are cell-matrix focal adhesion proteins but have now been found at the apical and basal ES, such as the BTB), as well as β-catenin and VE-cadherin in the vascular endothelium.\textsuperscript{80} Likewise, recent studies have shown that NO, a reactive oxygen species produced by NOS, regulates Sertoli cell TJ-barrier and Sertoli-germ cell anchoring junction function via the cGMP/PKG
Cross-Talk between Tight and Anchoring Junctions at the BTB That Regulates BTB Dynamics

The coexistence of TJ and AJ at the BTB has been illustrated by studies using electron and freeze-fracture microscopy since the 1970s. Recent studies using fluorescent microscopy have also colocalized TJ and AJ to the same site at the BTB. Even though occludin and cadherin at the BTB have no direct protein-protein interaction as demonstrated by coimmunoprecipitation, they are linked via their peripheral adaptors, such as ZO-1 and catenins. This finding is not entirely unexpected since at the early stage of TJ assembly, ZO-1 was shown to associate with catenins in MDCK cells. Besides, ZO-1 also serves as a cross-linker between the cadherin/catenin protein complex and the actin-based cytoskeleton in nonepithelial cells. In a more current study, ZO-1 was found to use the same domain for its interaction signaling pathway which is also associated with a significant decline in cadherin-NOS and catenin-NOS interaction. It is likely that these changes can lead to an increase in phosphorylation of the cadherin/catenin complex via the action of PKC. These observations thus support the notion that as preleptotene spermatocytes traverse the BTB, there must be a coordination in the disintegration of TJ and anchoring junction. We speculate that when TJ is “opened” to accommodate cell migration, the transiently induced basal ES proteins form a “patch” to supersede the temporarily loss TJ-barrier function (see text above). Thereafter, cell adhesion complexes in the “patch” also transiently “open” to facilitate cell movement by altering their phosphorylation status so that preleptotene spermatocytes can continue their migration. However, one may also argue if the basal ES and perhaps AJ transiently loss their adhesion function at the BTB to facilitate preleptotene spermatocyte migration, how can BTB maintains its cell adhesion and TJ-barrier function at the time. Interestingly, in studies using the Adjudin and the androgen suppression model, it was shown that there was a transient induction in TJ-proteins (e.g., occludin, ZO-1, JAMs) at the BTB when spermatids were depleting from the epithelium. We speculate that these induced TJ-proteins are being used to supersede the temporal loss of AJ function at the BTB to confer adhesion between migration preleptotene spermatocytes and Sertoli cells. For instance, it is known that the first extracellular domain of occludin can confer cell adhesion function. In short, it is likely that TJ and basal AJ at the BTB are working in concert to facilitate preleptotene spermatocyte migration while maintaining the TJ-barrier function and cell adhesion in this microenvironment.

In the testis, apical ES is considered to be a hybrid cell-cell actin-based AJ and cell-matrix focal contact anchoring junction type restricted to the interface between Sertoli cells and developing spermatids in adult rat testes. This conclusion is reached since several focal contact proteins usually restricted to the cell-matrix interface to facilitate cell movement (e.g., macrophages and fibroblasts) in other epithelia are also found at the apical ES. For instance, FAK is localized at the basal ES near the basement membrane while its phosphorylated and activated form, p-FAK-Tyr, is detected almost exclusively at the apical ES and is structurally associated with a nonreceptor protein tyrosine kinase, c-Src. Furthermore, their protein levels are significantly induced when germ cells are depleting from the seminiferous epithelium, as illustrated in the Adjudin and testosterone-suppression models. Other protein kinases, including Csk, CK2 and Fer kinases have also been found to associate with N-cadherin and catenin at the basal and apical ES. Integrin-linked kinase (ILK) was also reported to be associated with β1 integrin at the apical ES. Collectively, these findings coupled with recent reports that TJ proteins are also integral components of the apical and basal ES (see discussion above) thus illustrate that ES is indeed a unique hybrid anchoring junction type of AJ, focal contact, and TJ, that can confer AJ functionality (i.e., cell adhesion) while facilitate cell movement (e.g., focal contact function) and transiently confer TJ functionality (e.g., during BTB restructuring in which the induced basal ES proteins form a “patch” at the BTB). Thus, it is not entirely unlikely that basal ES can supersede the function of TJ and vice versa as discussed herein.
with occludin and α-catenin. As such, ZO-1 is not only a TJ adaptor; in fact, it shuffles between TJ and AJ and links these two junction types to cytoskeleton.

Using an in vivo model to study junction dynamics involving Adjudin, rats were treated with a single dose of this compound at 50 mg/kg b.w. by gavage to induce germ cell depletion from the seminiferous epithelium. The drug is targeted at the apical ES with minimal effect to TJs and basal ES. Interestingly, at the time of germ cell loss, the association between the protein complexes at TJ (e.g., occludin/ZO-1) and AJ (e.g., cadherin/catenin) via their corresponding adaptors was temporarily abolished, namely ZO-1 and α- and γ-catenins. In essence, ZO-1 was no longer associated with catenins, and as such, occludin and N-cadherin as well as their adaptors, ZO-1 and γ-catenin were diffusing away from the BTB site. Yet, their association was reestablished by day 7 after treatment when germ cells (e.g., elongating/elongate spermatids and some round spermatids) were depleted from the epithelium. Based on such timely dissociation between TJ and AJ protein complexes in the seminiferous epithelium, an “engagement and disengagement” theory was proposed to describe the unique mechanism employed by the testis to facilitate germ cell movement pertinent to spermatogenesis (Fig. 4). It is likely that under physiological conditions, such as at stages other than VII and VIII of the epithelial cycle, TJ and AJ (e.g., basal ES) are structurally “engaged” via their peripheral adaptors at the BTB to reinforce barrier integrity (Fig. 4). At the time of spermatogenesis and when preleptotene spermatocytes must traverse the BTB, which occur concurrently at stage VIII of the epithelial cycle, a transient “disengagement” between adaptors (e.g., ZO-1 and catenins) of the corresponding TJ and AJ protein complexes takes place to facilitate germ cell movement across the barrier, avoiding the unnecessary damage to TJ during AJ restructuring. This novel mechanism thus preserves barrier integrity while facilitating germ cell movement (Fig. 4). After spermatogenesis and the movement of preleptotene spermatocytes across the BTB, the adaptors, namely ZO-1 and α/γ-catenins become “engaged” again to strengthen the barrier. This theory not only provides a solid biochemical basis regarding the mechanism of germ cell movement across the BTB, it also provides the rationale for the coexisting TJ and AJ at the BTB. However, the identity of the protein(s) that “pulls” ZO-1 away from catenins to maintain the “disengaged” state and facilitate germ cell movement remains to be determined. Recent studies have shown that dynamins (e.g., dynamin II), large GTPases that serve as “pinchase-like mechanoenzymes” to facilitate the formation of endocytic vesicles by severing nascent endocytic pits from the plasma membrane, are likely to maintain such a “disengaged” state at the BTB. This work must be vigorously validated and expanded in future studies.

In this context, it is of interest to note that this theory does not account for the temporal disruption of TJ at the BTB, which must occur to accommodate preleptotene spermatocyte migration. Recent studies have unequivocally demonstrated that TNFα and TGF-β3 produced and secreted by Sertoli and/or germ cells into the BTB microenvironment can induce reversible disruption of the barrier as receptors for these cytokines are mostly resided in Sertoli cells. This is illustrated in a functional BTB assay by monitoring the diffusion of a small molecular dye (e.g., fluorescein 5'-isothiocyanate), from the systemic circulation into the adluminal compartment behind the BTB following TNFα treatment. Thus, it is highly plausible that these cytokines at the BTB perturb TJs, transiently open the barrier to facilitate preleptotene spermatocyte migration at stage VIII of the epithelial cycle. Indeed, the expression levels of both TNFα and TGF-β3 are highest at stage VIII of the cycle. Other studies using Sertoli cell cultures have shown that TGF-β3 regulates TJ-barrier function via the p38 and ERK MAP kinase signaling pathways. Furthermore, studies using the in vivo CdCl2 model have unequivocally demonstrated that BTB dynamics are regulated by TGF-β3 via its effects, at least in part, on the steady-state protein levels of protease inhibitors (e.g., α2-macroglobulin), and proteases (e.g., cathepsin L), which, in turn, assist BTB restructuring. Other studies have shown that MMPs and TIMPs are both present at the ES and BTB, and whose actions are coordinated by TNFα. Collectively, these data suggest that at the time preleptotene spermatocytes traverse the BTB, it is initially mediated by a transient loss in...
Cross-Talk between Tight and Anchoring Junctions

Figure 4: A-C) A schematic drawing that illustrates the engagement and disengagement mechanism between TJ and AJ at the BTB to facilitate preleptotene spermatocyte migration. A) The restructuring of TJs and AJs (e.g., basal ES) at the BTB are regulated by the "engagement (close)" and "disengagement (open)" mechanism to facilitate preleptotene spermatocyte migration across the BTB with minimal damaging effects on the barrier integrity. The left top panel shows the spatial arrangement of the seminiferous epithelium in which the BTB separates the epithelium into the basal and adluminal compartments. TJs (e.g., occludin/ZO-1) and basal ES (e.g., N-cadherin/γ-catenin) that coexist at the BTB are structurally associated with each other via the peripheral adaptors (e.g., ZO-1 and catenins). At stage VIII of the epithelial cycle, spermiation occurs when elongated spermatids emptying into the tubule lumen, and when preleptotene spermatocytes traverse the BTB, a transient opening of BTB must occur. Figure legend continued on following page.
protein-protein associations between AJ-proteins at the basal and apical ES (e.g., cadherins/catenins) via a surge in tyrosine phosphorylation of catenins (and/or cadherins). Additionally, AJ-associated proteins complexes are transiently disengaged from TJs and the surge in TJ proteins can supercede the temporal loss of cell adhesion at the BTB. During the migration of preleptotene spermatocytes across the BTB, Sertoli and germ cells secrete TGF-β3 and/or TNF-α, into the BTB microenvironment at stage VIII of the epithelial cycle. When these cytokines bind onto the receptors on the Sertoli cell, the cascade of signaling molecules (e.g., p-p38) are activated, which, in turn, reduces the levels of TJ proteins (e.g., occludin) at the BTB to "open" the barrier to facilitate germ cell migration. Ser/Thr phosphorylation and the temporal increase in AJ-proteins can substitute the TJ-barrier function by forming a "patch" in the "opened" BTB. After germ cells pass through the BTB, further TJ disassembly can be limited by the production of protease inhibitors such as α2-MG and TIMPs. Finally, TJ and AJ become structurally linked and "engaged", which is mediated via their peripheral adaptors to reinforce the BTB.

Additionally, it is of interest to note that the disengagement between TJ and AJ at the time of Adjudin-induced germ cell depletion from the epithelium does not involve actin disruption, since the level of F to G-actin increases as AJ disengaged from TJ. This illustrates that this germ cell loss event is entirely a junction restructuring process. Perhaps the increase in F-actin content provides additional scaffolding function to the BTB during germ cell depletion. Indeed, an increase in F- to G-actin ratio is detected in human adenocarcinoma cells with an increase in invasiveness, illustrating active AJ restructuring pertinent to cellular migration is associated with an increase in F-actin, consistent with results obtained from the Adjudin model.

In short, the biochemical events that occur at the BTB during spermatogenesis as discussed above are in agreement with the "junction restructuring theory". Also, these recent biochemical findings do not negate the "zipper theory" or the "intermediate cellular compartment theory". Indeed, these three theories are not mutually exclusive. In essence, the "junction restructuring theory" is a biochemical version of the combined "zipper" and "intermediate cellular compartment" theories, taking into account the molecular players in the junction restructuring events that occur during spermatogenesis. For instance, in the "zipper theory", the "old" TJ fibrils above the migrating preleptotene spermatocytes must be broken down after the formation of the nascent TJ fibrils below these migrating cells as illustrated in the "intermediate cellular compartment theory" in which preleptotene spermatocytes were shown to be trapped between TJ fibrils in the seminiferous epithelium. The "disengagement" between TJs and AJ at the BTB as discussed above (see also ref. 89) thus offers the biochemical mechanism to facilitate preleptotene spermatocyte migration across the BTB.
Table 1. Junction types and their molecular components at the BTB of adult rodent testes

<table>
<thead>
<tr>
<th>Junction Type</th>
<th>Component Proteins</th>
<th>Mr (kDa)</th>
<th>Interacting Partners</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TJ</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TJ</td>
<td>- Integral membrane proteins</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Occludin, occludin IB</td>
<td>60-65</td>
<td>ZO-1, ZO-2, ZO-3, cingulin, NOS</td>
</tr>
<tr>
<td></td>
<td>Claudin-1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12-24</td>
<td>20-25</td>
<td>ZO-1, ZO-2, ZO-3</td>
</tr>
<tr>
<td></td>
<td>JAM-A, B</td>
<td>40</td>
<td>ZO-1, cingulin, AF-6/afadin, PAR3, PAR6, aPKC, PICK-1, CASK, MUPP-1</td>
</tr>
<tr>
<td></td>
<td>CAR</td>
<td>46</td>
<td>ZO-1, MUPP-1, MAGI-1</td>
</tr>
<tr>
<td></td>
<td>CLMP (CAR-like membrane protein)</td>
<td>44-48</td>
<td>Occludin, ZO-1</td>
</tr>
<tr>
<td></td>
<td>BT-IgSF (Brain- and testis-specific immunoglobulin superfamily)</td>
<td>43.77-52</td>
<td>n.k.</td>
</tr>
<tr>
<td></td>
<td>Tricellulin</td>
<td>66-72</td>
<td>n.k.</td>
</tr>
<tr>
<td>TJ</td>
<td>- Peripheral membrane proteins</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ZO-1, 2, 3</td>
<td>210-225</td>
<td>ZO-1, ZO-2, ZO-3, myosin,</td>
</tr>
<tr>
<td></td>
<td>Clingulin</td>
<td>140</td>
<td>JAM-A, claudin, JAM, α and γ-catenins, connexin 43, actin, afadin, 4.1R</td>
</tr>
<tr>
<td>Basal ES and/or AJ</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basal ES and/or AJ</td>
<td>- Integral membrane proteins</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>N- or E-Cadherin</td>
<td>127-135</td>
<td>α and γ-Catenins, p120ctn, c-Src, NOS-2,3, ponsin, α-actinin, actin, Rab8b</td>
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<tr>
<td></td>
<td>Celsr cadherins 1, 2</td>
<td>320</td>
<td>Protocadherin a, Rab7</td>
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<td>Nectin-1, 2, 4</td>
<td>70-85</td>
<td>Afadin, ponsin, α2-macroglobulin, ZO-1, PAR3, PAR6, aPKC, PICK-1</td>
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<tr>
<td>Basal ES and/or AJ</td>
<td>- Peripheral membrane proteins</td>
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<td></td>
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<tr>
<td></td>
<td>α-Catenin</td>
<td>100-104</td>
<td>N-Cadherin, β and γ-catenins, ZO-1, p120ctn, α-actinin, actin, afadin</td>
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<td></td>
<td>β-Catenin</td>
<td>92</td>
<td>N-Cadherin, β and γ-catenins, p120ctn α-actinin, actin, NOS-2,3, PRKG</td>
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<tr>
<td></td>
<td>γ-Catenin</td>
<td>82</td>
<td>N-Cadherin, α and β-catenins, ZO-1, p120ctn, α-actinin, actin, Rab8b</td>
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<tr>
<td></td>
<td>p120ctn</td>
<td>65-120</td>
<td>N-Cadherin, β and γ-catenins, α-actinin, actin, Rab8b</td>
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<tr>
<td></td>
<td>Afadin/AF-6</td>
<td>205</td>
<td>Nectin, ponsin, α2-macroglobulin, actin, P-cadherin</td>
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<tr>
<td>Basal TBC</td>
<td>MN7</td>
<td>90</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Actin</td>
<td>42</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cofilin</td>
<td>20</td>
<td></td>
</tr>
</tbody>
</table>

continued on next page
Table 1. Continued

<table>
<thead>
<tr>
<th>Junction Type</th>
<th>Component Proteins</th>
<th>Mr (kDa)</th>
<th>Interacting Partners</th>
</tr>
</thead>
<tbody>
<tr>
<td>Desmosome-like junction</td>
<td>Desmocollin-1, -2, -3, Desmoglein-1γ, -1α, -1γ, -2, -3, 4</td>
<td>107-115</td>
<td>Plakophilin, desmoplakin, vimentin</td>
</tr>
<tr>
<td></td>
<td>Connexins 26, 30.2, 30.3, 31.1, 32, 33, 36, 37, 40, 43, 45, 46, 50, 57</td>
<td>27-70</td>
<td>Occludin, ZO-1, ZO-2, N-cadherin, α and β-catenins, p120ctn</td>
</tr>
<tr>
<td></td>
<td>4.1B*</td>
<td>110</td>
<td>Plakoglobin, desmoplakin, vimentin</td>
</tr>
<tr>
<td></td>
<td>4.1G*</td>
<td>130-140</td>
<td>Cadherin</td>
</tr>
<tr>
<td></td>
<td>LYRIC**</td>
<td>80</td>
<td>Occludin, ZO-1</td>
</tr>
<tr>
<td></td>
<td>P-glycoprotein (P-gp)</td>
<td>n.k.</td>
<td>n.k.</td>
</tr>
<tr>
<td></td>
<td>Multidrug resistance-associated protein (MRP1)</td>
<td>n.k.</td>
<td>n.k.</td>
</tr>
</tbody>
</table>

This table was prepared based on current findings in the field as discussed in the text. References given here are not exhaustive due to the limited page space. As such, only selected references are given herein and additional references can be found in the text and earlier reviews, 6,17,30,39,40,46,108-110,113-124. In the column for “Component proteins”, proteins that have been positively identified in the testis are underlined. In the column for “Interacting partners”, underlined proteins that are bold and italic are those that were shown to interact with the corresponding integral membrane proteins at the BTB; proteins that are underlined only are found in testes but were shown to interact with the corresponding target proteins in other epithelia; proteins that are included in the table but not underlined are those that were found in similar blood-tissue barriers other than the testis. n.k. = not known. *These proteins are found at the Sertoli cell-spermatogonia interface near the basal compartment, and may not be a component of the BTB; **. Protein that has not been identified in the BTB but it is known to interact with TJ-associated proteins.
Cross-Talk between TJ, Anchoring Junction, and GJ in the Seminiferous Epithelium Is Crucial to Spermatogenesis

In other epithelia, cross-talk between different junctions has been the subject of active investigation in recent years. For instance, cross-talk between cadherins and integrins is likely mediated by Rap1 GTPase. It was shown that the disassembly of AJ through endocytosis activated Rap1, which, in turn, enhanced the integrin cell-matrix adhesion by redistributing integrins to new adhesion sites. This thus avoids uncontrolled cell dissemination.\textsuperscript{102} Direct association between AJ proteins (e.g., nectin-1 and 3) and PAR-3 (a signaling molecule usually restricted to TJ and crucial to cell polarization working in concert with aPKC and PAR-6\textsuperscript{103}) was also reported in neural epithelial cells in which the affinity of nectins toward afadin (an adaptor of nectin) and PAR-3 was found to be similar.\textsuperscript{104,105} Protein interacting with C-kinase-1 (PICK-1), a scaffolding protein, interacts with both AJ (e.g., nectin) and TJ proteins (e.g., JAM-A, B & C but not claudins) in CHO cells.\textsuperscript{106} It is believed that PICK-1 is an adapter which coordinates cross-talk between TJ and AJ in epithelial cells. However, the significance (and its presence in the testis) of PICK-1 in cross-talk between TJ and anchoring junction (e.g., basal ES) at the BTB and between apical ES and BTB remains to be determined. Gap junctions (GJ) (e.g., connexin-43) are recently shown to associate with TJ (e.g., ZO-1 & 2)\textsuperscript{107,108} and AJ proteins (e.g., N-cadherin, α-catenin, β-catenin and p120ctn).\textsuperscript{109} These findings thus demonstrate the potential inter-dependent relationship between junction types in various epithelia including the BTB, which is likely mediated via peripheral adaptors. In some cases, their roles can be temporarily superseded by each other, as illustrated in the engagement and disengagement theory in the BTB.\textsuperscript{89} In others, the formation of a junction type requires the coassembly of another type. For instance, when either connexin-43 (GJ) or N-cadherin (AJ) was disrupted by RNA silencing, gap junctional communication as well as the mobility of NIH3T3 cells was reduced,\textsuperscript{109} illustrating a functional AJ and GJ linkage.

Concluding Remarks—Lesson from the Testis

In the past decade, much work has been done in dissecting the role of TJ and anchoring junction dynamics in spermatogenesis, however, the crucial information is still lacking in the field as highlighted in this review. For instance, how is the cross-talk between TJ and anchoring junctions initiated and regulated? Is the cross-talk limited between TJ and AJ (e.g., basal ES and apical ES) or is it extended to the desmosome-like junctions? What are the signaling molecules and pathways that are involved in this cross-talk? Answers to some of these questions will also unravel potential targets for male contraception and may offer an explanation for unexplained infertility. Nonetheless, recent findings regarding the regulation of different junction types at the BTB as reviewed herein have illustrated that the testis has a unique mechanism in place in which anchoring junction (e.g., basal ES) and TJ can be functionally segregated so that preleptotene spermatocytes can traverse the BTB without compromising the TJ-barrier function. Additionally, cytokines (e.g., TGF-β3 and TNFα) produced locally by Sertoli and/or germ cells and secreted into the microenvironment of BTB are also being used to transiently “open” the TJ-barrier to facilitate germ cell movement. It will be of interest and physiologically important to determine if this mechanism is used to facilitate food absorption across the epithelial TJ-barrier in small intestine or to facilitate transepithelial migration of neutrophils across the endothelial TJ-barrier in microvessels at the inflammation site.

Acknowledgements

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The Role of the Leydig Cell in Spermatogenic Function

Renshan Ge, Guorong Chen and Matthew P. Hardy*

Introduction

Two anatomically distinct compartments, respectively, the interstitium where Leydig cells reside and the seminiferous tubules contribute the major functions of the testis, testosterone secretion and sperm production. This structural partitioning is achieved by peritubular myoid cells, which surround the seminiferous tubules. Despite the anatomic separation of the two compartments, they are functionally coupled by endocrine and paracrine inter-communication. For example, the testosterone secreted by Leydig cells under the stimulus of luteinizing hormone (LH), diffuses into seminiferous tubules and drives spermatogenesis together with another gonadotropic hormone, follicle-stimulating hormone (FSH). This dependency of the seminiferous epithelium on testosterone illustrates the significance of the Leydig cell in spermatogenesis.

Leydig cells are polygonal in shape and are the major cell type within the interstitium where they are often found adjacent to blood vessels and the seminiferous tubules. In addition to Leydig cells, other cell types such as fibroblasts, macrophages and small numbers of mast cells are also present in the interstitial space. The Leydig cell is the principal source for the testosterone in systemic circulation in males. In rodents, there are two generations of Leydig cells, fetal and adult, and in humans, there is a middle generation of neonatal Leydig cells. The first generation is designated as fetal not because of a functional immaturity but rather due to its embryonic origin (see ref. 4, for review). Fetal Leydig cells originate from stem cells in embryo that have yet to be identified, and go on to become fully competent in androgen synthesis prior to birth, leading to fetal masculinization (see ref. 5 for review). Fetal Leydig cells remain in the testis after birth but do not proliferate or contribute significantly to androgen levels thereafter, and are gradually lost from the testis by attrition. The next generation (in the rodent) referred to as the adult Leydig cells forms during pubertal development. Adult Leydig cells are not derived from preexisting fetal Leydig cells but from undifferentiated stem cells that are present in the interstitium. Conceptually, development of adult Leydig cells can be divided into four stages: stem, progenitor, immature and adult Leydig cell, according to their morphological and biochemical characteristics. Stem Leydig cells are spindle-shaped and do not express Leydig cell markers such as LH receptor and steroidogenic enzyme proteins. However, given their status as stem cells, they are capable of self-renewal and commitment to differentiation into progenitor Leydig cells. Progenitor Leydig cells, which are also spindle-shaped have a low capacity for steroidogenesis and mainly produce androsterone. Progenitor develop into
immature Leydig cells which are round and have a well-developed smooth endoplasmic reticulum and abundant cytoplasmic lipid droplets. Immature Leydig cells secrete 5α-reduced androgens and primarily 5α-androstane-3α, 17β-diol (3αDIOL) and 5α-androstane-3β, 17β-diol (3βDIOL). Immature Leydig cells ultimately develop into adult Leydig cells at the completion of puberty, whereupon testosterone is the predominant androgen end product. The pubertal rise in testosterone levels and initiation of spermatogenesis are coincident and interdependent, as will be discussed in this review.

Abbreviations

**Genes:** CYP11A1, cytochrome P450 cholesterol side-chain cleavage enzyme; CYP17A, cytochrome P450 17α-hydroxylase/C17-20-lyase; HSD3B2, 3β-hydroxysteroid dehydrogenase 2; HSD17B3, 17β-hydroxysteroid dehydrogenase 3; Gnhr, gonadotrophin releasing hormone; Lhgr, luteinizing hormone receptor; ESR1, estrogen receptor α; CYP19A1, cytochrome P450 aromatase; ESR2, estrogen receptor β.

**Proteins:** LH, luteinizing hormone; FSH, follicle-stimulating hormone; P450c17, cytochrome P450 17α-hydroxylase/C17-20-lyase; P450sc, cytochrome P450 cholesterol side-chain cleavage enzyme; StAR, Steroidogenic acute regulatory protein; PBR, peripheral-type benzodiazepine receptor; 3β-HSD, 3β-hydroxysteroid dehydrogenase; 17βHSD3, 17β-hydroxysteroid dehydrogenase 3; P450arom, cytochrome P450 aromatase; GnRH, gonadotrophin releasing hormone; 3α-HSD, 3α-hydroxysteroid dehydrogenase; hCG, human chorionic gonadotropin; INSL3, Insulin-like protein 3.

**Chemicals:** T, testosterone; 3αDIOL, 5α-androstane-3α, 17b-diol; 3βDIOL, 5α-androstane-3β, 17β-diol; E₂, estradiol.

Androgen and Spermatogenesis

**Testosterone Biosynthesis**

Leydig cells have a cyto-architecture that is typical for a steroid secreting cell, in that they possess an abundant smooth endoplasmic reticulum and numerous mitochondria, which have tubular cristae. They are able to synthesize cholesterol from acetate or to take up this substrate for steroidogenesis from lipoproteins present in circulation. The cytochrome P450 17α-hydroxylase/C17-20-lyase (P450c17) enzyme, which catalyzes later steps in testosterone biosynthesis may be active in cholesterol synthesis, because it has recently been shown to exhibit squalene epoxidase activity. The movement of cholesterol to the inner mitochondrial membrane, the location of the cholesterol side-chain cleavage enzyme (P450sc), is the rate-limiting step of androgen synthesis. Transport of cholesterol is acutely regulated by LH and occurs in two steps: mobilization of cholesterol from cellular stores such as lipid droplets or the plasma membrane to the outer mitochondrial membrane and then the transfer of cholesterol from the outer to the inner mitochondrial membranes. Steroidogenic acute regulatory (StAR) protein has been viewed as indispensable in the second phase of this process. StAR is expressed in steroidogenic tissues in response to agents that stimulate steroid production, and mutations in the StAR gene result in steroidogenic failure such as occur in the disease of congenital lipoid adrenal hyperplasia. The mechanism by which StAR mediates cholesterol transfer in the mitochondria has not been fully characterized. Another mitochondrial protein, peripheral-type benzodiazepine receptor (PBR), has been shown to be involved in intra-mitochondrial transport of cholesterol. Further research will be necessary to establish a definitive mechanism of cholesterol transport, but it may be that the function of StAR is convey cholesterol to PBR, which in turn forms a pore from the external face in the outer mitochondrial membrane for its internal conveyance.

P450sc is located on the matrix (or internal) side of the inner mitochondrial membrane. The side-chain cleavage reaction uses three molecules of oxygen and three molecules of
The Role of the Leydig Cell in Spermatogenic Function

NADPH and a mitochondrial electron transfer system resulting in the formation of pregnenolone. The enzymes required for subsequent steps of steroidogenesis are located in the smooth endoplasmic reticulum (SER) membranes in the cytoplasm. The bulk of the pregnenolone formed in the mitochondrion diffuses into the SER and serves as a substrate for progesterone synthesis by the $\Delta^\text{3\,17}$-hydroxysteroid dehydrogenase (3\,$\Delta^\text{3\,17}$-HSD) enzyme. However, recent evidence shows that once pregnenolone is formed, it may be converted within the mitochondria to progesterone by a mitochondrial form of the enzyme 3\,$\Delta^\text{3\,17}$-HSD. Although multiple isoforms of 3\,$\Delta^\text{3\,17}$-HSD have been identified, the testicular 3\,$\Delta^\text{3\,17}$-HSD is coded by type I and II 3\,$\Delta^\text{3\,17}$-HSD gene ($\text{HSD3b2}$). Once formed progesterone is converted by P450c17 into androstenedione by two steps: 17\,$\Delta^\text{17}$-hydroxylation and C17-C20 cleavage. In the human, the gene for P450c17, $\text{CYP17A}$, is located on chromosome 10. The final step of androgen synthesis, conversion of androstenedione into testosterone is catalyzed by 17\,$\Delta^\text{17}$-hydroxysteroid dehydrogenase (17\,$\Delta^\text{17}$-HSD). There are multiple isoforms of 17\,$\Delta^\text{17}$-HSD, and the testicular isoform is type III 17\,$\Delta^\text{17}$-HSD (17\,$\Delta^\text{17}$-HSD3), coded by 17\,$\Delta^\text{17}$-HSD3 gene ($\text{hsd17b3}$). Testosterone either associates with androgen binding protein and moves into the seminiferous tubule lumen to regulate spermatogenesis or is further metabolized into different steroids, some of which are more biologically active such as dihydrotestosterone in reactions catalyzed by types 1 and 2 5\,$\Delta^\text{5}$-reductase (Fig. 1).

Androgen and Spermatogenesis

Spermatogenic dysfunction is often associated with impaired Leydig cell dysfunction. Between 12 and 15\% of men with diminished spermatogenesis have lower testosterone levels.
Leydig cell dysfunction is clearly associated with spermatogenic damage in rodent models.\textsuperscript{28,29} Agents that cause Leydig cell dysfunction, such as irradiation, induction of cryptorchidism, and vitamin A deficiency, have also been found to disrupt the histology of the seminiferous epithelium,\textsuperscript{29} pointing to a clear linkage between Leydig cell function and spermatogenesis.

Androgens are essential for the maintenance of normal spermatogenesis. Suppression of testosterone biosynthesis inhibits spermatogenesis. For example, mutations in the gonadotrophin releasing hormone gene (\textit{Gnrhpg/hpg}) dramatically lower serum testosterone levels\textsuperscript{30} to a point where spermatogenesis is blocked.\textsuperscript{31} In these animals spermatogenesis can be qualitatively rescued by androgen replacement therapy.\textsuperscript{30} This rescue is independent of LH and FSH action, because FSH alone fails to promote spermatogenesis beyond the meiotic stages.\textsuperscript{32,33} In the LH receptor (\textit{Lhcr}) knockout mouse Leydig cells, which are developmentally arrested at the progenitor cell stage and consequently have barely detectable testosterone in circulation, exogenous testosterone administration restores spermatogenesis.\textsuperscript{34} In rats, suppression of testosterone levels by hypophysectomy leads to an acute, stage-specific regression of the seminiferous epithelium.\textsuperscript{35-37} If testosterone levels are not maintained in hypophysectomized animals, mid-stage round spermatids and mature, elongated spermatozoa are lost, suggesting that androgen action is involved in spermiogenesis and the transition from round to elongated spermatids. When hypophysectomized rats are treated further with ethane 1,2-dimethanesulphonate, a chemical that destroys the Leydig cells, the elimination of residual testosterone prevents spermatogenesis from proceeding beyond meiosis.\textsuperscript{36,38}

Intratesticular levels of testosterone are very high compared to blood and the question arises as to how much testosterone is required for maintenance of spermatogenesis. Spermatogenesis is maintained, at a reduced level, when testicular concentrations are only 5\% of control.\textsuperscript{39} Suppression of FSH release induced by high dose administration of testosterone or estradiol (E\textsubscript{2}), or by anti-LH or -GnRH immunization, lowers intratesticular testosterone levels to 20\% to 40\% of normal while allowing spermatogenesis to proceed in a quantitatively normal manner.\textsuperscript{40,41} Recently, it has been shown that intratesticular testosterone levels as low as 2\% of normal are compatible with the continuation of spermatogenesis in LH receptor knockout mice.\textsuperscript{42}

Stimulation by androgen is a prerequisite for the differentiation of germ cells. However, evidence also points to a negative regulation of spermatogonial stem cells by androgen. Testosterone inhibits spermatogonial proliferation and differentiation, and this can be reversed by the androgen receptor antagonist flutamide.\textsuperscript{43,44} A similar phenomenon has been observed in juvenile spermatogonial depletion (\textit{jsd}) mice, in which germ cells regress to a spermatogonia-only phenotype following the first wave of spermatogenesis. Treatments that oppose androgen action, such as GnRH antagonist or flutamide administration, stimulate proliferation and differentiation of spermatogonia.\textsuperscript{45} The inhibition of spermatogonial differentiation by testosterone in \textit{jsd} mutant mice is achieved only at the normal scrotal temperature.\textsuperscript{46}

\textbf{Androgen Receptor-Mediated Regulation of Spermatogenesis}

Testosterone binds to the androgen receptor thereby initiating nuclear translocation of the ligand-bound receptor complex and the regulatory functions of the androgen receptor in modulating gene transcription.\textsuperscript{47} Spermatogenesis is dependent on a constant level of androgen receptor mediated activity, seen by the fact that a null mutation in the X chromosome-linked gene that encodes the androgen receptor causes complete Androgen Insensitivity Syndrome (AIS) in both humans and mice, with consequent small, abdominally positioned testes and spermatogenic blockade early in meiosis.\textsuperscript{48,49} In the testis, the androgen receptor is expressed in somatic cells including Leydig, myoid, and Sertoli cells\textsuperscript{50,51} and, in human males, in spermatogonia.\textsuperscript{51-53} Immunocytochemical localization studies of rat testes had placed androgen receptor expression in spermatogonia, but direct androgen action on these cells is apparently not a requirement for fertility: androgen receptor null germ cells have the capacity to form sperm when transplanted into the testes of wild type males.\textsuperscript{54,55}
A role for androgen receptor-mediated activity is seen in Leydig cells where it is to promote maturation of the steroidogenic pathway. In rodents it has been observed that Leydig cells do complete differentiation both in the naturally occurring androgen receptor loss-of-function, testicular feminization mutant and in conditional knockout mice in which androgen receptor expression is selectively deleted in Leydig cells. Due to the reduction of androgen action on the Leydig cell, testosterone production is sharply reduced in these animals, which causes an arrest of spermatogenesis.

Sertoli cells are the only somatic cell type in the testis that is in direct contact with differentiating germ cells. Testosterone withdrawal causes retention of mature spermatids and premature release of round spermatids, and this led to the general postulate that Sertoli cells are the primary mediators of androgen receptor-mediated regulation of spermatogenesis. The androgen receptor is expressed in a stage-specific manner in Sertoli cells as a function of the seminiferous cycle, which can be maintained (in the absence of gonadotropin stimulation) by administration of exogenous androgen. Androgen receptor expression in Sertoli cells is highest at stages VII-VIII. Upstream of the androgen receptor gene is an androgen response element.

Results of the selective knockout of androgen receptor expression in Sertoli cells suggest the existence of multiple androgen receptor-dependent steps in spermatogenesis. While there is no apparent defect in meiosis, the germ cells are defective in making the transition from round to elongating spermatids. A requirement for androgen receptor activity in Sertoli cells exists at three points: (A) progression through meiosis I; (B) transition from the round to elongating spermatid; and (C) in the terminal phases of spermatogenesis (spermiogenesis). Androgen action is evidently not needed for passage of the germ cell through meiosis.

Other Leydig Cell-Derived Biologically Active Steroids and Spermatogenesis

**Estrogen**

Testosterone can be considered a prohormone that is converted in target tissues into the more potent androgen, dihydrotestosterone. In addition, however, testosterone can be metabolized into other active steroids including estrogens. It has long been known that estrogens affect spermatogenesis (see ref. 65 for review). More recently, it has been established that both estrogen receptor α (ESR1) and aromatase knockout mice are infertile. Moreover, the presence of both estrogen receptor (ER) α and β isotypes in the male reproductive tract indicates the importance of estrogen action in male reproduction. E2 is believed to be the endogenous ligand of both ER isotypes, and has the same affinity in each case. Other ligands exhibit selectivity in ER binding. For example, one of the endogenous metabolites of androgen, 3βDIOL, binds ERβ with higher affinity compared to E2, and must therefore be considered as an estrogen.

Testosterone is metabolized to E2 via the microsomal P450 aromatase (P450arom), which is encoded by the CYP19A1 gene. The P450arom reaction uses three molecules of oxygen and three molecules of NADPH for every molecule of testosterone catalyzed. CYP19A1 has an age-dependent pattern of expression in the testis. In prepubertal and immature testis, Sertoli cells are the major sources of estradiol, whereas localization in Leydig cells becomes predominant as animals mature. LH stimulates the formation of E2 form in Leydig cells. CYP19A1 mRNA and activity is also found in germ cells from the pachytene spermatocyte stage to round spermatids. P450arom also is higher in mature spermatids than in earlier stage germ cells.

Thirty years ago, it was noted that 3βDIOL is an estrogenic metabolite of dihydrotestosterone. The synthesis of 3βDIOL from dihydrotestosterone is catalyzed by 17β-hydroxysteroid dehydrogenase 7 (17β-HSD7) (Fig. 1), with the enzyme functioning in this instance as a 3β-HSD. During the puberty, when Leydig cells are still immature, they...
express higher levels of 5α reductase and 3α-hydroxysteroid dehydrogenase (3α-HSD) and 17β-HSD7, and expression of these enzymes declines significantly with the transition to adulthood.10 From postnatal days 10 to 20, 3αDIOL is the major metabolite released from the testes, and later on 3βDIOL secretion becomes predominant.81 In the human testis, metabolism of testosterone to 5α-reduced products is also notable prior to puberty and, in contrast to rats, 3βDIOL is the major end product.

**Estrogens and Spermatogenesis**

In instances where spermatogenesis is inhibited and testosterone levels are lowered, estrogen levels are typically elevated.82 A subset of male infertility patients has been described that have an endocrinopathic increase in serum E2 concentrations.27 The change in estradiol activity can often be detected as an abnormally low testosterone to estradiol (T: E2) ratio, which is considered indicative of increased aromatase activity. It appears that either LH or human chorionic gonadotropin (hCG) can stimulate increased P450arom activity levels in Leydig cells, implicating overexpression to gonadotropin as a factor in abnormally high T: E2 ratios. Local synthesis of E2 in the Leydig cell blocks steroidogenesis at the P450c17. Men that have spermatogetic failure in association with an increased T: E2 ratio can be treated with P450arom inhibitors thereby improving their fertility status. There is, however, limited direct evidence that estrogen directly disrupts spermatogenesis (reviewed in ref. 65). Indeed, at the low concentration at which it is normally present in the male, estrogen may be physiologically beneficial for spermatogenesis. Consistent with this hypothesis the knockout of P450arom in mice (ArKO), causes an arrest at the round spermatid stage.67 Similarly, mice lacking ESR166,83-86 and CYP19A167,87 expression all have fertility problems. ESR2 is widely expressed in the male reproductive tract but has yet to be assigned a functional role in spermatogenesis.88-92 Male mice with an ESR2 knockout are fertile, but this does not exclude the possibility that ERβ-mediated signaling is involved in spermatogenesis. In this regard it has been noted that E2 stimulates proliferation of spermatogonia.93 The positive and negative aspects of estradiol’s effects on the testis must be taken into consideration when considering the impact of environmental estrogens on male reproductive health. Exposures of laboratory animals and wildlife to high levels of estrogenic chemicals including industrial chemicals such as pesticides (e.g., DDT, methoxychor), industrial chemicals (e.g., DES, tamoxifen) and phytoestrogens (e.g., genisterin) resulted in a number of abnormalities, including reduced gonad size, feminization of genetic males, and low sperm count and quality.94-96 For example, estrogen administration during the neonatal period or adulthood can impair the production and maturation of sperm.97,98

**Estrogen Receptor-Mediated Regulation of Spermatogenesis**

Throughout postnatal development, ERα is mainly expressed in Leydig cells and is not present in the seminiferous tubules.99-101 In rodents, ERα is abundantly expressed in the rete testis,101 efferent ductules99,101 and epididymis.99-101 It remains unclear whether ERα is expressed in sperm. There are no reports of ERα localization in rodent and human germ cells with the exception of a single study reporting its presence in rat spermatocytes and spermatids.102 ERβ is present at relatively low levels in spermatogonia92,103 and immature Sertoli cells92,103 relative to ERα. By day 21, ERβ is abundantly expressed in pachytene spermatocytes92 but not in other germ cells.103 In the mouse, ERβ is also specifically immunolocalized to the spermatocytes by day 12, but decreases to the undetectable level by day 26.104 ERβ expression is also seen in the epididymis from day 21 onward.104,105 Localization runs from the efferent ducts to the epididymis in the mouse106,107 and rat.85,100 In adult males of certain species, mouse, primate and human102,107-109 Leydig cells are observed to express ERβ, although other reports put this in dispute.91,110 Expression of ERβ in rat, monkey and human spermatogonia103,108,109,111 appears to be established. It is unclear whether ERβ remains present at later stages of sperm differentiation and further exploration of differences between species will need additional investigation.92,103,108,110,112,113
E₂ is thought to be the natural ligand of both ERα and β. Receptor binding analyses have shown that 3βDIOL can activate both ERα and ERβ with a Kd of 1.9 × 10⁻⁸ M, and that it has a negligible affinity for the androgen receptor. Other studies have shown that 3βDIOL may have a much higher potency when bound to ERβ, as measured by transcriptional activation, relative to ERα.

**The Leydig Cell as Target for Hormonal Contraception**

The hypothalamus secretes gonadotropin-releasing hormone (GnRH), which is transported to the pituitary where it stimulates gonadotropins secretion (LH and FSH). LH and FSH are required to initiate and maintain spermatogenesis. Testosterone secreted by Leydig cells has a negative feedback effect on the hypothalamus and the pituitary inhibiting LH and FSH secretion (Fig. 2). In humans, the negative feedback of testosterone on the hypothalamic-pituitary axis serves to inhibit secretion of both LH and FSH, which is the basis for the current hormonal approach to male contraception (Fig. 2). In mice, selective immuno-vaccination against LH receptor leads to the suppression of fertility in about 60% of treated mice. Administration of an antiserum against LH (passive immunization) disrupts spermatogenesis in rats. However, it is increasingly clear that inactivation of Leydig cell testosterone production will not be sufficient to suppress spermatogenesis. The current

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Figure 2. Stimulatory (—) and inhibitory (---) pathways exploited in male hormonal contraception. Testosterone and its metabolite estradiol exert negative feedback inhibition on the hypothalamic-pituitary axis. Based on the negative feedback, androgens alone or in combination with other sex steroids have been developed to suppress spermatogenesis.
approach towards inhibition of testicular function relies on the suppression of both LH and FSH secretion and in parallel, often with supplementation of exogenous androgen to avoid the symptoms of androgen deficiency. Of the steroids that exert negative feedback on the secretion of gonadotropins, estrogen has received less attention from the standpoint of contraceptive development. Studies in humans have shown that administration of estradiol may further enhance suppression of gonadotropin levels when coadministered with an androgen (Fig. 2).

The ER\(_\alpha\) knockout is associated with Leydig cell hyperplasia and testosterone levels in mice, and this may be due to the higher LH levels that are seen in the mutant. However, the same knockout is known to disrupt spermatogenesis. Since ER\(_\alpha\) is present in Leydig cells but not in seminiferous tubules, the disruption of spermatogenesis is due to aberrant morphology observed in the efferent ductules.

**Leydig Cell Proteins and Spermatogenesis**

**Insulin-Like Protein 3 (INSL3)**

In addition to testosterone and other steroid end products, Leydig cells secrete proteins that may affect spermatogenesis either directly or indirectly. Among these proteins, insulin-like protein 3 (INSL3) has received particular attention because it has been shown to stimulate testicular descent during development. INSL3 is a member of the insulin-like hormone superfamily and is expressed exclusively in fetal and adult Leydig cells. A family of orphan G protein-coupled receptors that are homologous to gonadotropin receptors has recently been identified and named as LGR 4 through 8. Male mice with a mutation in the INSL3 gene exhibit defects in testicular descent, including cryptorchidism (abdominal testes) due to abnormal development the gubernaculum (the connective tissue cord that connects the testis to the scrotal sac). Conversely, overexpression of INSL3 induces ovarian descent in transgenic females. Transgenic mice missing the LGR8 gene are also cryptorchid, and INSL3 is now known to be the endogenous ligand of LGR8. A role for INSL3 and LGR8 in the development of the gubernaculum and testicular descent has been established in that disruption of either gene causes cryptorchidism and leads to spermatogenic failure (due to the lowering of intratesticular testosterone levels, in combination with the increased temperature of the testis). Fetal and adult Leydig cells are the chief source of circulating, and the serum concentration of INSL3 can even serve as an index of the functional status of Leydig cell steroidogenic capacity. The gubernaculum is the main site of LGR8 expression, but this protein is also localized to Leydig cells and germ cells. In germ cells the binding of INSL3 to LGR8 prevents apoptosis, but the function of the LGR8 pathway in Leydig cells awaits elucidation.

**Oxytocin**

Leydig cells produce and secrete oxytocin, as do (to a lesser extent) epididymis epithelial cells. Oxytocin produced in Leydig cells is regulated by LH. Oxytocin stimulates testosterone production. It has been suggested that the function of testicular oxytocin is to stimulate contraction of the peritubular myoid cells that surround the seminiferous tubules, moving spermiated sperm to efferent ducts. Oxytocin receptors are located in Leydig cells and Sertoli cells and peritubular myoid cells.

**Summary**

Testosterone is the most well known of the products synthesized and secreted by Leydig cells and that have a defined involvement in the regulation of spermatogenesis. However, the Leydig cell is the main source of testicular estrogen and estrogen action has both positive and negative effects on spermatogenesis that can no longer be ignored. In addition, two protein
The Role of the Leydig Cell in Spermatogenic Function

products of the Leydig cell—INSL3 and oxytocin—influence spermatogenic function by stimulating, respectively, testis descent and movement of mature sperm to the excurrent ducts (Fig. 3).

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