Spermatogonial Stem Cells: Implications for Genetic Disorders and Prevention

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Spermatogonial stem cells (SSCs) propagate mammalian spermatogenesis throughout male reproductive life by continuously self-renewing and differentiating, ultimately, into sperm. SSCs can be cultured for long periods and restore spermatogenesis upon transplantation back into the native microenvironment in vivo. Conventionally, SSC research has been focused mainly on male infertility and, to a lesser extent, on cell reprogramming. With the advent of genome-wide sequencing technology, however, human studies have uncovered a wide range of pathogenic alleles that arise in the male germ line. A subset of de novo point mutations was shown to originate in SSCs and cause congenital disorders in children. This review describes both monogenic diseases (eg, Apert syndrome) and complex disorders that are either known or suspected to be driven by mutations in SSCs. We propose that SSC culture is a suitable model for studying the origin and mechanisms of these diseases. Lastly, we discuss strategies for future clinical implementation of SSC-based technology, from detecting mutation burden by sperm screening to gene correction in vitro.

Introduction

S TEM CELLS PLAY a pivotal role in maintaining homeostasis in adult tissues; they self-renew, differentiate, and regenerate tissues upon injury. Unlike other stem cells, spermatogonial stem cells (SSCs) have the unique job of propagating genetic information to subsequent generations. In the following sections, we focus on the properties and dynamics of mammalian SSCs in vivo and in vitro, with special emphasis on implications for human diseases caused by germ line mutations. We conclude by considering strategies for future clinical implementation of SSCbased technology.

In Vivo Biology of Mammalian SSCs

SSCs reside along the basement membrane of the seminiferous tubules in adults and are unipotent (ie, lineage restricted). Developmentally, these cells originate from the epiblast and undergo fate specification as primordial germ cells (PGCs) [1]. PGCs enter the nascent endoderm, spread along the hindgut epithelium, and begin to migrate toward the genital ridge, the future testis [2,3].

After colonization of the gonad, PGCs give rise to prospermatogonia, a change that occurs in the fetal/early neonatal period in rodents. During this phase, prospermatogonia become mitotically quiescent until proliferation eventually resumes within the first week after birth [4,5]. Prospermatogonia migrate to the basement membrane and the spermatogenesis phase begins [6]. The differentiation of PGCs into prospermatogonia in humans is not well understood, but there seem to be strong parallels between the two species [7]. Postnatally, prospermatogonia mature into SSCs.

Various theoretical models have been developed to explain how SSCs concomitantly maintain the stem cell population through self-renewal while providing sufficient numbers of differentiating progeny. In rodents, the pool of undifferentiated spermatogonia (Spg), which includes SSCs, comprises A_s (single), A_{pr} (pair), and A_{al} (aligned) Spg. The definitions of A Spg subtypes were made according to their morphological arrangement on the basement membrane. The first model to describe stem cell renewal dates back to 1971 and is referred to as the A_s or static model [8]. It was proposed that A_s Spg comprise the only pool of true stem cells, whereas the balance of A Spg are irreversibly committed to differentiation.

An alternative model was presented recently by Yoshida's group, who demonstrated that these A_s , A_{pr} , and A_{al} cells are extremely interchangeable with significant cell fate plasticity [9–11]. The authors developed a pulse-labeling system, in which a pulse of tamoxifen was given to mice to activate a Cre recombinase (creER^{T2}) under control of the promoter either for Ngn3 (a marker of differentiating Spg [12]) or for *Gfr* α *1* (a marker of undifferentiated Spg [9]). Upon labeling,

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Compared with mice, primates show fewer spermatogonial subtypes, which can be divided into three distinct groups: A_{dark} (A_d), A_{pale} (A_p) (originally termed A1 and A2), and type B Spg [13,14]. A_d Spg, characterized by highly condensed chromatin, are thought to represent the reserve stem cells, whereas A_p Spg, which exhibit a more sparse chromatin distribution, may represent the active stem cells, even though definitive evidence is still lacking. A_d Spg, which are normally quiescent, became proliferative after irradiation but whether A_d Spg also contribute to steady-state spermatogenesis still remains to be addressed [15].

Some authors also proposed the existence of another type of A Spg, termed A transition Spg that display an intermediate phenotype between A_d and A_p [16–18]. Strikingly, this A_d/A_p model was proposed more than 50 years ago by Clermont, and, at present, very limited information is available about the kinetics of human Spg [19]. Nevertheless, substantial data exist for nonhuman primates, in which spermatogenesis has been extensively studied [14,17,20–23].

Derivation of SSC Cultures in Mice and Humans

Studies on SSCs have been hampered by the absence of specific markers, as well as the relative rarity of authentic stem cells present in the adult testis [11,24]. Yet, progress has been made in understanding the optimal culture conditions necessary to maintain SSCs in long-term culture, at least in rodents [25,26]. As with most other adult stem cells, SSCs require a specific niche to preserve their unique features.

In vivo, Sertoli and peritubular cells are the most prominent components of the niche, secreting factors that support SSC self-renewal and mitotic expansion [27,28], among which the most fundamental is glial cell line-derived neurotrophic factor (GDNF). The level of GDNF is critical for SSC homeostasis. Mouse *Gdnf* knock-out models exhibited impaired fertility [27,28], whereas *Gdnf* overexpression in the testis caused an initial abnormal expansion of undifferentiated Spg that resulted in sterility and tumor development [27]. As discussed hereunder, extensive efforts over the last 10 years have yielded a better understanding of how growth factors influence SSC renewal and differentiation.

A variety of strategies have enabled derivation of murine SSC lines with similar efficiency from both neonatal and adult mice [25,29]. Successful long-term culture conditions for SSCs were first established by the Shinohara group in 2003 [25]. The addition of GDNF, leukemia inhibitor factor, epithelial growth factor, and fibroblast growth factor-2 (FGF2) permitted maintenance of SSCs in vitro for up to five months. The colonizing potential of cultured SSCs was proven by injection into busulfan-treated mice and is discussed further hereunder. Another group reported the maintenance of SSCs in vitro for up to six months in serum-free conditions [26]. The outcome of SSC line derivation was also shown to be strain dependent [25,26].

In addition to SSC cultures from neonatal mice, long-term SSC cultures were derived from adult testis using modified culture conditions [29]. The studies already described have employed a variety of feeder cell types for maintenance of SSCs, including murine embryonic fibroblasts, testicular somatic cells (designated JK1), and SIM mouse embryoderived thioguanine and ouabain-resistant (STO) cells [30]. Thus, knowledge of factors supporting SSC growth has gradually enabled development of robust in vitro models.

The story of human SSC (hSSC) culture is completely different. As already described, innumerable differences exist between humans and rodents. As such, the dearth of knowledge in identifying hSSCs and their niche has obstructed progress in the field. Current obstacles include the difficulties in accessing healthy human tissue samples, as well as isolation and expansion of highly purified hSSCs in culture. Of note, somatic cell overgrowth has been encountered as a major challenge to assessing hSSCs in culture [31].

Various markers that were previously studied in other organisms have been reported to be useful in enriching hSSCs, although most are controversial. One of the first reported markers was THY1 (or CD90) [32,33]. An obvious problem for employing this antigen is its robust expression by the mesenchymal stem cells [34], and later studies suggested that SSCs, in fact, do not express CD90 [35]. Another marker widely employed is CD49f [32,36]. As with CD90, CD49f is expressed by contaminating mesenchymal cells, as well as by testicular somatic cells, limiting its utility [37]. As a consequence of this overlap between putative hSSC markers and mesenchymal markers, most of the earlier reports that claimed the successful isolation of SSCs have interpretive constraints [38,39].

Our group studied whether several novel markers identified previously in mouse models could be useful for hSSCs [39,40]. For example, ID4 expression was recently found to mark a small population of bona fide SSCs in mice, making it an attractive candidate marker for other species [41]. GPR125 is an orphan adhesion-type G-protein-coupled receptor expressed by undifferentiated Spg and it, too, was first described in mice [42]. We employed a validated, human-specific anti-ID4 monoclonal antibody and found that ID4⁺ cells coexpress GPR125 and reside on the basal membrane of the seminiferous tubules [39,40]. Although He et al. succeeded in isolating a pure GPR125⁺ population from healthy donor testis, they were unable to maintain the SSCs in culture for long [39,40].

In 2009, the successful achievement of long-term culture of hSSCs was reported [38]. The study described the colonization of testicular tubules of nu/nu male mice by the hSSCs maintained in vitro. However, hSSC xenotransplantation has significant interpretative limitations (see "SSC Transplantation" hereunder), and, in addition, certain important data were not provided in the study. For example, although they showed a large relative increase in stem cell activity over time, the absolute number of stem cells expanded and the ability to freeze-thaw the long-term SSCs was not addressed. Finally, the genetic stability of hSSCs in culture remains unknown.

Improved colonization has been achieved by sorting testicular cells using SSEA-4 before injection into the testis, although the analysis was performed only one month posttransplantation [32]. The same year, another group reported the successful isolation and maintenance of hSSCs in culture using an exclusion approach, by plating the cells on collagen and further sorting the unattached cells for CD9 [36]. The authors demonstrated the differentiation of the isolated SSCs into spermatids, albeit at low efficiency.

Substantial further efforts have been made by several groups recently to characterize and enrich the hSSC population by using various combinations of markers [31,43–46]. However, the unwanted expansion of testicular somatic cells that are propagated in culture, irrespective of the method of derivation employed, remains an impediment. In conclusion, hSSC culture technology is not nearly as mature as that for mice and other species.

SSC Transplantation

As already discussed, experimental investigation of SSCs is complicated by their paucity in the testis and the inability to efficiently separate them from other unwanted cell types. Nevertheless, good tools are available to quantify the success of SSC isolation by observing their biological ability to produce and maintain spermatogenesis in a transplant model, first developed in 1994 by Brinster and Avarbock [47]. They showed how cells isolated from a donor testis were able to restore spermatogenesis once directly injected into the seminiferous tubules of infertile mice. The transplanted cells could produce de novo spermatogenesis regardless of the nature of the infertility, whether genetic [48] or experimentally induced by busulfan.

Today, this transplantation technique is routinely performed in both mice and rats [49] as a readout for the efficiency of SSC derivation. Given that only a true stem cell can give rise to a clone that will subsequently yield spermatogenic cells [50,51], the transplantation protocol represents a useful functional assay and constitutes the best way to identify SSCs. As an extension of the seminal Brinster study, which used freshly isolated testis cell suspensions, many studies currently employ this method to verify the presence of stem cells in culture [25,26]. As discussed later, the transplantation model has been successfully used recently to obtain experimental support for the paradigm of paternal age effect (PAE)-associated diseases, such as Apert syndrome [52].

In 2002, a similar transplantation strategy was developed for the study of hSSCs [53]. In this xenotransplant system, the authors employed busulfan-treated nu/nu mice as recipients for transplantation of donor hSSCs. Intriguingly, they were able to detect the presence of single hSSCs on the basement membrane of seminiferous tubules up to six months post-transplantation. Thus, xenografted human Spg colonize and undergo limited proliferation but do not differentiate in the mouse testis and gradually disappear. This indicates that although the human and mouse niches are somewhat compatible, the mouse testis does not provide a sufficient microenvironment to support human spermatogenesis. As such, the rigorous assays employed for rodent SSC activity remain unavailable for hSSCs.

Fertility Preservation

Preservation of fertility has historically been the most straightforward potential clinical application of SSCs. This issue is particularly urgent in the case of young boys, often with childhood leukemia, who undergo gonadotoxic chemotherapy as part of a curative regimen. The survival rate of such individuals has greatly improved in recent years [54]. This has resulted in an increased number of long-term survivors who may ultimately want to have families, raising the question of how to proceed if infertility occurs in the absence of banked sperm [55,56]. Theoretically, a way to overcome this problem is to cryopreserve testicular tissue and/or SSCs before chemotherapy and to transplant them back into the testis once the patient has been cured from cancer.

As already discussed, a similar strategy is routinely performed in animal models but has not yet been formally tested in humans. One obstacle to advancing such a strategy is the inability to efficiently expand hSSCs in culture, as the number of SSCs recovered from a small tissue sample is not typically enough to restore spermatogenesis.

Although success in propagating SSCs in vitro may be a prerequisite for autologous transplant, there are other concerns that need to be considered. First, the genetic and epigenetic stability of SSCs during in vitro expansion needs to be determined. It has been demonstrated that murine SSCs are genetically stable even after long-term expansion in culture, suggesting that they may possess a mechanism that prevents them from accumulating deleterious mutations [57].

At the same time, SSCs, when in culture, are not immune to telomere shortening and may not be considered immortal. Some data indicate that this genetic stability is shared by hSSCs, providing support for the possibility of employing SSCs to restore male fertility [58]. To this end, Radford et al. undertook a clinical trial in which testicular cells from Hodgkin lymphoma patients were cryopreserved and autotransplanted back to the testis after completion of chemotherapy, but the results of follow-up analyses remain to be published [59]. Finally, removal of contaminating cancer cells also needs to be taken into account.

Use of SSCs for the Study of Pluripotency

The ability of antenatal germ cells to acquire pluripotency in vitro was discovered many years ago (see Matsui et al. [60]), yet it was only in 2004 that a landmark article demonstrated that postnatal SSCs, too, retain such characteristics [61]. We subsequently showed that long-term wild-type adult SSC lines in vitro preserve the latent plasticity that enables reprogramming, which was ultimately confirmed by others [42,62,63]. These features of SSCs raised the possibility of exploiting SSCs for the study of cellular reprogramming and potentially for therapeutic use. Thus, a number of studies have produced differentiated cell types from pluripotent SSC-derived cell lines [42,64,65]. However, paternal methylation imprints persist in SSCs and their derivatives, which could ultimately affect their functionality and safety in the context of therapeutic use [61].

In conjunction with availability of alternative sources of pluripotent stem cells (ie, induced pluripotent stem cells), the enthusiasm for pursuing derivatives of SSCs in this realm is currently limited. Nonetheless, significant effort has been exerted toward developing pluripotent lines from hSSCs, but long-term, self-renewing hSSC cultures remain elusive. An initial high-profile report of pluripotent stem cells derived from human Spg was ultimately retracted [33]. A number of groups have since provided evidence that postnatal human male germ cells, like their murine counterparts, have developmental plasticity, but most studies demonstrated neither robust evidence of pluripotency nor clonal origins for the precursor to the reprogrammed cells [66–68]. As such, the most important current application of reprogrammed cells originating from SSCs is confined to the domain of basic research studies.

SSCs and PAEs

The advent of high throughput DNA sequencing has recently enabled the direct interrogation of a unique category of human genetic variation, comprising de novo point mutations (DNMs), defined here as single nucleotide variants that are present in offspring but absent in constitutional (ie, somatic cell) genomes of the parents and vanishingly rare in the general human population [69]. For the purposes of the subsequent sections, we focus on DNMs and ignore other important sources of genetic variation (eg, copy number variants, insertion/deletions, and other structural aberrations) and common variants (eg, single nucleotide polymorphisms), due to the significance of DNMs in known disorders related to paternal age.

Consistent with historical predictions, recent sequencing studies of healthy individuals revealed that humans are born with dozens of DNMs, and most of the variation in this number is attributable to paternal age at conception [69,70]. On average, two to three additional DNMs are apparent in children with each additional year of paternal age at conception [69,71]. Furthermore, the location of DNMs is significantly influenced by a variety of factors, including sequence context (eg, CpG site) and regional timing of replication [69,70,72]. Notably, mutation rates are higher in functional regions of the genome, such as exons [70,72]. Finally, genic regions (as opposed to intergenic regions) are disproportionately affected by paternal age, suggesting that such age-related mutations have greater potential to manifest phenotypically [70].

As a tractable alternative to studying humans, researchers have assessed the mutation frequency of germ line cells in an experimental mouse model [73,74]. This transgenic system allowed them to directly measure the mutation frequency of an insert (ie, a *lacI* mutation reporter), which revealed lower mutation frequencies of germ line cells than somatic cells [74,75]. These studies also revealed that spermatogenic cells from old mice have increased mutation frequencies and an increased prevalence of transversion mutations than those from young or middle-aged mice [74,76]. Together with the human data, these findings suggest that an age-related increase in point mutations is a general feature of male gametogenesis in mammals.

Recent studies have found that a distinct subset of human genetic diseases originates from DNMs in the male germ line. Historically, these disorders, including Apert syndrome (*FGFR2* mutations), achondroplasia (*FGFR3*), Noonan syndrome (mutations in *PTPN11* and other genes), and Costello syndrome (mutations in *HRAS*), have been observed frequently in children from older fathers irrespective of mothers' age [77–79]. In fact, one of the earliest clues to the existence of such disorders was the observation by Weinberg in the early 20th century of a birth order effect on the incidence of achondroplasia [80]. The mutations in these disorders are paternally derived, yet the fathers do not harbor the

mutations in their somatic tissues, indicating the occurrence of de novo male germ line mutations [81–83]. These disorders are now commonly referred to as PAE disorders (see Goriely and Wilkie [84] for a detailed review).

The genes mutated in PAE disorders often belong to the receptor tyrosine kinase/RAS/MAPK signaling pathway and almost always comprise gain-of-function mutations. For example, in Apert syndrome, the mutation of FGFR2 at Ser252Trp manifests increased affinity of the mutant receptor for FGF ligand, which is likely to lead to activation of signaling even when availability of ligand is reduced [85]. Of note, PAE mutations generally exhibit milder gains-of-function than somatic mutations in cancer cells. Specific PAE mutations have been identified in sperm donors and increase with advanced donor age [86–89]. The increase in mutations is exponential, which is significant, because such an amplification could not be driven simply by the linear increase expected in the cumulative number of cell divisions in the normal aging testis.

Furthermore, testes obtained from normal older males have more mutations in PAE genes, and the cells bearing mutations are arranged in a spatially clustered manner [90– 93]. The clustering phenomenon is reminiscent of tumor expansion in somatic tissues and cannot be explained in statistical models by a local increase in hot-spot mutations. Thus, the simplest explanation is positive selection of PAE mutations in SSCs. That is, a PAE mutation may arise in a single SSC that acquires a selective growth advantage as compared with its wild-type neighbors (Fig. 1). Goriely and Wilkie termed this phenomenon "selfish spermatogonial selection" [84].

Although several attempts have been made to prove the mechanism of this phenomenon using computer modeling, selfish spermatogonial selection, in the absence of prospective, experimental data, is still largely hypothetical. One caveat of such statistical models is that many assumptions are required, including, for example, the mitotic rate of hSSCs, which is unknown and has only been inferred through a single, ethically dubious study in which a handful of prisoners were first administered H³-thymidine and then subjected to testicular biopsies [94].

De Novo Mutations and Complex Disorders

As discussed in the previous paragraphs, DNMs arise in the germ cells of older fathers, giving rise to various defects. In contrast to monogenic disorders, complex developmental disorders are genetically quite heterogeneous. Thus, it is typically more difficult to establish a direct association between specific DNMs and the disease phenotype. Also, unlike the monogenic disorders already described, the DNMs described to date in complex disorders tend to result in loss of function of the corresponding genes. Several studies detailed hereunder have highlighted a major contribution of DNMs in developmentally important genes. These mutations are believed to originate in parental germ cells before fertilization, and they tend to increase with the father's age [95].

Compared with whole-genome sequencing, which confers a greater sensitivity in detecting pathogenic mutations, whole-exome sequencing (WES) is more efficient in identifying novel disease genes [96]. One limitation of both approaches is the relatively small number of families

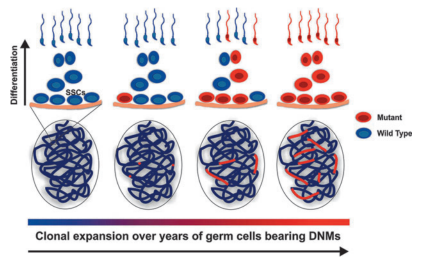


FIG. 1. Schematic representation of the paternal age effect hypothesis. De novo point mutations (DNMs) may occur in isolated spermatogonial stem cells (SSCs, *red dots* on the second testis) conferring an advantage to the mutant cells over the wild type cells. During the course of the years, the mutant cells subsequently take over the wild type cells leading to a clonal expansion and colonization of the testis tubules (*red fragment* in the third and fourth testes), increasing the possibility to pass those mutations to the offspring (*red spermatozoa*).

typically analyzed and the lack of a systematic analysis of the male germ lines, to determine the extent of accumulating mutations [71].

Neuropsychiatric Disorders, Congenital Heart Disease, and Cancer

The autism spectrum disorders (ASDs) comprise an array of developmental syndromes defined by a variety of symptoms and degrees of disability. Although a strong genetic component is well recognized, the underlying causes are still largely unknown, despite rapid progress. Recently, various independent WES studies demonstrated how disruptive (ie, nonsense, splice site, and frame shift) DNMs are highly recurrent in ASDs [91,97–99].

In particular, two of these studies carried out exome analyses not on family trios but rather on "quads" (two healthy parents, an affected child, and an unaffected sibling) [91,98]. Strikingly, the studies reached a common conclusion that the affected children exhibit an excess of potentially pathogenic mutations. These occurred in genes commonly expressed in the brain [91], were associated with the fragile X protein, FMRP [98], or involved β -catenin/chromatin remodeling protein networks, often associated with neuronal development [99]. Furthermore, both Neale et al. and O'Roak et al. observed a positive correlation between *CHD8* mutations and ASDs, suggesting the *CHD8* mutant phenotype as a novel syndromic form of autism [97,99]. Remarkably, a strong bias toward paternal rather than maternal origin for the culprit DNMs was also noted [98,99].

Similarly, a different study, using the same general strategy (ie, trio WES), showed a significant excess of DNMs in congenital heart disease with rare variants in genes expressed in the developing heart [100]. In particular, they found that DNMs in congenital heart disease tend to be associated with the methylation of lysine 4 on histone 3 (H3K4) [100]. In the same vein, earlier studies had indicated a strong effect of paternal age on the incidence of schizophrenia, another common genetically complex and heterogeneous disorder [101]. However, a very recent study employed five different models and concluded that, although age-related DNMs may account for 10%–20% of the increased risk, DNMs do not fully explain the predisposition to schizophrenia in the off-spring of older fathers [102].

It has also been proposed that DNMs in tumor-associated genes may have a role in cancer risk in children. Proving this assumption is again complicated by the heterogeneous, multistep nature of the disease. There are a number of studies that describe an association between early onset cancer and DNMs in father's gametes [103]. Recently a comprehensive study analyzed numerous genes known to be recurrently mutated in cancer, looking for evidence of germ line mosaicism [104]. Interestingly, they found that 8.5% of the patients analyzed harbored germ line mutations in genes related to cancer predisposition. Unfortunately, they did not assess the presence of these mutations in the parents of the affected children, which precludes addressing the origin of DNMs. Thus, further efforts are required to unravel a possible link between DNMs and cancer.

Development of Systems to Study SSC-Derived Mutations

Although de novo paternally derived mutations are likely to occur in many species, the disease phenotypes attributable to such mutations, thus far, appear to be a uniquely human phenomenon. As such, it would be ideal to employ hSSCs as a model system for studying paternal age-associated mutations. Unfortunately, the relatively primitive technology available for functional studies of hSSCs obviates substantial mechanistic studies. For example, if clonally derived, cryopreserved hSSC lines were readily available to investigators, significant in vitro studies could be combined with xenotransplantation assays to address the significance of key genes and pathways. However, as already noted, it would be important to first determine the genetic stability of hSSCs in culture.

Although such studies are not yet possible, important insights have come from static observations of archived human testis samples. The identification and marking of apparently clonal populations of mutant germ cells in the human testis were pioneered by the Wilkie and Goriely groups. In a series of unique studies, these investigators first predicted that certain immunohistochemical features would be unique to mutant clones present in the adult testis. Based on their earlier studies of spermatocytic seminoma [105], the investigators found two populations of tubules in the normal testis with differential levels of anti-FGFR3 staining: one with both normal signal intensity and morphology and a second with stronger FGFR3 expression and intense coexpression of MAGEA4 and phospho-AKT [106]. The latter (termed "immunopositive tubules") also frequently exhibited an unusual double layer of Spg.

In a recent subsequent study, the authors used laser capture microdissection, exon capture, and high-throughput sequencing to show that immunopositive tubules, as defined previously, were strongly enriched for mutations in growth factor signaling-related genes that could be implicated in PAE disorders [107]. Critically, adjacent nonimmunopositive regions selected as negative controls from the same testis did not have any of the tested mutations, indicating that their detection system was highly specific. Although the sample size was relatively small, it was somewhat unexpected that they detected only one of the common Apert syndrome-associated substitutions in FGFR2 (P253R) but not the other (S252W), nor any mutations in *RET*, another well-studied gene implicated in PAE.

Also of note, the individual immunopositive regions were surprisingly large (ie, up to millimeters in length of the tubule) but were relatively deficient in spermatogenesis compared with adjacent areas. These findings provide evidence for discrete clusters of mutant, clonal premeiotic germ cells, bearing DNMs in signaling-related genes, on a background of otherwise normal human germ cells, but suggest that many such putative clones do not actually contribute to disease in offspring.

In contrast to humans, animal models afford the opportunity to address SSC competition in a prospective, experimental manner. In fact, two historical studies incidentally revealed evidence of adult SSC selection. The first direct evidence for germ line selection in mammals may have come from Mintz. In the course of analyzing chimeric animals comprising cells from two different mouse strains, she cleverly noted a progressive shift in the proportion of offspring of each strain that appeared in subsequent litters from chimeric fathers [108]. Later, Jaenisch made a similar observation using a male mouse that was genetically mosaic for integrated Moloney leukemia provirus [109].

SSC competition has also been interrogated using transplantation assays in mice. The Shinohara group demonstrated that transplanted SSCs compete with endogenous stem cells for colonization [110]. In another series of relevant studies, the Brinster group assessed effects of aging and the niche microenvironment on SSC self-renewal [111,112]. They demonstrated that SSCs from old mice consecutively transplanted into young testes maintained self-renewal capacity and spermatogenesis for long past the normal life span of mice [111]. Although cell competition was not specifically interrogated, the data suggest that normal mouse SSCs do not acquire gross genetic or epigenetic abnormalities over time. Nonetheless, niche failure or deterioration could facilitate cell competition, particularly in humans.

To assess homeostatic behavior of SSCs in vivo, Nakagawa et al. first described life-long kinetics of mouse SSCs using genetic lineage tracing, as already mentioned [11]. In this study, it was demonstrated that stem cells are occasionally lost and compensated for by other stem cells that form larger sized clones over time. Indeed, using a similar lineage tracing setup in the mouse testis, we observed that certain SSC clones survive for more than 1 year, expanding their territories, but the remaining clones disappear (unpublished data) (Fig. 2). By analyzing SSC clonal dynamics using mathematical models, Klein and Simons proposed that neutral competition confers all stem cells with equal proliferation potential, but some stem cells are lost in a stochastic manner and are replaced by others [113].

Drosophila melanogaster has provided excellent models for cell competition, enabling comparisons of pairs of clones with different genotypes (ie, non-neutral competition; see Stine and Matunis [114] for a review). However, few nonneutral competition models in mammalian germ line systems have been described. One study used a competitive transplantation assay in mice, in which mutant stem cells were mixed with wild-type SSCs for transplantation into recipient testes. These data showed that most of offspring from transplanted SSCs, consisting of mixed populations of wild-type cells and those with loss-of-function for p21 cyclin-dependent kinase inhibitor, were derived from the p21-deficient SSCs and not from wild-type SSCs [115]. This suggests that the mutant SSCs successfully compete with wild-type cells to support long-term repopulation.

Using adult mouse SSCs, our group developed an experimental approach for interrogating PAE disorders [52]. In this study, SSCs with an ectopically expressed, Apert syndrome-associated FGFR2 mutation (S252W) exhibited significantly greater stem cell activity than those with ectopic wild-type FGFR2 in transplantation assays. Importantly, the S252W SSCs did not show any proliferative advantage when cultured alone in vitro in standard media, but exhibited increased competitiveness in the presence of wild-type SSCs. At the cellular level, S252W SSCs showed enhanced sensitivity to FGF, as evidenced by enhanced MAPK signaling with low dose FGF, suggesting an increased self-renewal capability of S252W SSCs, when growth factors are scarce. This experimental model supports the central hypothesis of positive selection propagation of SSCs in mammals.

Common genome editing techniques such as TALEN and CRISPR enable simple, rapid gene modification [116]. These new approaches will likely enhance the development of SSC culture systems designed to directly interrogate both the mechanisms of positive selection for PAE mutations and also the functions of other mutations arising in germ line stem cells. However, studies of SSC cultures are dependent on the post-transplantation reconstitution of spermatogenesis, a milieu that does not fully reflect the physiological behavior of SSC in vivo. To understand in vivo SSC competition in a normal environment, one would require a system to induce mutations in situ (ie, a genetic mosaic) and follow mutated cell fates over long intervals.

Significance of SSCs in the Setting of Disease Prevention and Diagnosis

The genetic contribution to PAE disorders is driven by DNMs. Although most of the best understood such disorders are individually rare and monogenic, it is likely that a substantial fraction of the risk also of common, genetically complex disorders (eg, autism and early onset cancers) is also driven by paternally derived DNMs. In fact, a recent preliminary report by the Developmental Disorders Study consortium estimates that the aggregate risk of a child being born with a developmental disorder (of any kind) attributable to DNMs is very large (\sim 1:450 to 1:200 live births)

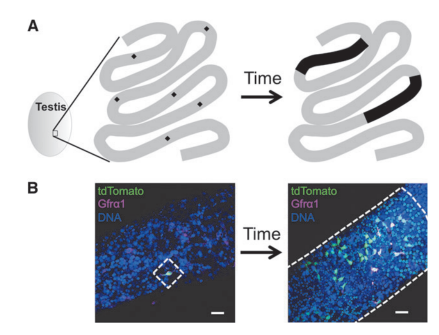


FIG. 2. Lineage tracing reveals the long-term kinetics of labeled SSCs. (**A**) An example of lineage tracing. In this segment of seminiferous tubules, a single SSC (*black diamond*; six SSCs are shown in the segment) is being labeled after induction. Certain clones (eg, only two clones out of six in this case) self-renew and expand to occupy the long segment (*black area*), but the rest disappear over time. By applying mathematical models to this phenomenon, Klein and Simons proposed a neutral competition model, in which all stem cells have equal proliferation potential, but some stem cells are lost in a stochastic manner [113]. (**B**) Actual images of clonal expansion in the lineage tracing system using the Sox2-creER^{T2} driver. Tamoxifen was administered to a Sox2-creER^{T2}; Rosa-tdTomato mouse at 7 weeks of age. *Right* after the induction, Sox2-creER^{T2} driver labeled A_{single} SSCs (*left*). After a long period of chase, SSC progeny clonally expand to form a large colony (*right*). Some of the undifferentiated spermatogonia retained Gfrα1 expression, which is known to mark SSCs. Scale bars: 40 µm.

[96]. Since the majority of such DNMs are of paternal origin, it is extremely important to understand the detailed pathogenic process that occurs upstream of transmission in the male germ line.

As techniques to efficiently derive and propagate hSSCs are being developed, new clinical potential applications of SSCs are coming into focus. With the advent of such lines, a number of strategies may emerge that could enable the prevention of genetic paternal age-related disorders. First, however, it will be necessary to determine which individually rare disorders are, in fact, exclusively of paternal origin. This information can sometimes be obtained using existing WES data by means of adjacent polymorphisms unique to the father or mother. However, informative polymorphisms are not always available, necessitating more in-depth genetic analysis.

For pathogenic DNMs that are exclusively of paternal origin, we can envision the development of genetic screening panels that could be applied to sperm samples of prospective older fathers, analogous to the diagnostic gene panels that are currently available for genotyping of somatic cancers. If evidence is found for a significant degree of risk, this would allow the prospective parents to make informed decisions about whether to proceed.

A series of further applications of hSSCs can be imagined, but each step in implementation entails serious concerns related to both safety and ethics. For men with insufficient healthy sperm (ie, those lacking known pathogenic DNMs), clonal SSC lines could be derived from a testis biopsy, expanded, and retested. If healthy lines are found, these could theoretically be employed in several ways. The first is autotransplantation back into the testis. Notably, proof-of-principle for autotransplantation has been already demonstrated in animal models [117– 120]. Second, given recent improvements in experimental derivation of gametes in vitro from stem cell precursors, it is likely that production of functional sperm from hSSCs will eventually become feasible. For example, functional male haploid gametes were recently generated in vitro from rodent stem cells and were successfully used to fertilize oocytes and produce offspring [121]. Also, chemical treatment (ie, retinoic acid exposure) has been used to efficiently induce meiosis in cultured mouse SSCs [121]. If such technologies could be adapted to hSSCs, these could be used for intracytoplasmic sperm injection to generate embryos for implantation.

Finally, given recent technical advances in genome editing, it may be possible to revert mutant alleles in SSCs back to the wild-type state before use of sperm. Each of the procedures already outlined would require extremely extensive, potentially prohibitive safety testing, first in primates and then in humans.

Although we do not attempt to delve into the ethical intricacies of assisted reproduction, several major issues bear brief a note. The following concerns are not entirely unique to the technical strategies already described, but are potentially more significant (eg, relative to simple in vitro fertilization [IVF]), given the greater extent of artificial manipulation of the cells involved. First, the genomes and epigenomes of gametes derived from long-term cultured cells could be altered in unforeseen ways. In contrast, shortterm culture of human embryos is already employed in standard IVF protocols, suggesting that at least some period of culture is compatible with normal development. Second, many future generations could be affected by genomic or epigenomic alterations due to long-term culture. From our current vantage point, these barriers are certainly formidable and may even appear prohibitive.

However, it is difficult to predict how novel technology may be able to successfully overcome such hurdles. Therefore, it is incumbent upon the research community to consider the risks and benefits far in advance of each stage of clinical testing and to proceed with extreme caution. In fact, recent guidelines from the International Society for Stem Cell Research, although supportive of research studies, prohibit clinical application of gametes that have undergone modification of the nuclear genome [122]. The growing interest around this new area is highlighted by the fact that human germ line editing is a major focus of a comprehensive study recently undertaken by the National Academies to address the clinical, ethical, legal, and social implications of the use of gene-editing technologies [123].

Conclusion

Myriad reasons exist to be optimistic about the future for using SSCs to understand and ameliorate human disease. The spectrum of human disorders for which the strategies already outlined are relevant is only likely to expand, as the ability to gather and interpret genomic sequencing data increases. Although impressive technological advances have already occurred in the field of SSC biology, many are limited to other organisms and have not been successfully applied to hSSCs. Yet, given the lag period for translating basic findings with other stem cell types (eg, embryonic stem cells) from model systems to humans, the question is not whether this will happen, but rather when will it happen. In the interim, the research and clinical communities will have ample time to consider the weighty issues that surround any manipulation of human gametes for therapeutic purposes.

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Author Disclosure Statement

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