Expression of c-Kit receptor mRNA and protein in the developing, adult and irradiated rodent testis

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Abstract

Germ cell proliferation, migration and survival during all stages of spermatogenesis are affected by stem cell factor signalling through the c-Kit receptor, the expression and function of which are vital for normal male reproductive function. The present study comprehensively describes the c-Kit mRNA and protein cellular expression profiles in germ cells of the postnatal and adult rodent testis, revealing their significant elevation in synthesis at the onset of spermatogenesis. Real-time PCR analysis for both mice and rats matched the cellular mRNA expression profile where examined. Localization studies in normal mouse testes indicated that both c-Kit mRNA and protein are first detectable in differentiating spermatogonia. In addition, all spermatogonia isolated from 8-day-old mice displayed detectable c-Kit mRNA, but 30–50% of these lacked protein expression. The c-Kit mRNA and protein profile in normal rat testes indicated expression in gonocytes, in addition to differentiating spermatogonia. However, in the irradiated adult rat testes, in which undifferentiated spermatogonia are the only germ cell type, mRNA was also detected in the absence of protein. This persisted at 3 days and 1 and 2 weeks following treatment with gonadotrophin-releasing hormone (GnRH) antagonist to stimulate spermatogenesis recovery. By 4 weeks of GnRH antagonist treatment, accompanying the emergence of differentiating spermatogonia, both mRNA and protein were detected. Based on these observations, we propose that c-Kit mRNA and protein synthesis are regulated separately, possibly by influences linked to testis maturation and circulating hormone levels.

Introduction

Among the many regulatory factors that tightly control spermatogenesis, the interaction of Kit (known as c-Kit) receptor protein with its ligand, Kitl (also known as Kit ligand and stem cell factor, and referred to here as SCF), plays essential roles, including its specific influence on spermatogonial differentiation and survival at the onset of spermatogenesis (for review see Loveland & Schlatt 1997, de Rooij & Grootegoed 1998, Yan et al. 2000). The c-Kit gene encodes a transmembrane tyrosine kinase receptor for the SCF protein. SCF is produced as an integral membrane protein that may be cleaved to release a soluble, extracellular ligand (Loveland & Schlatt 1997). Expression and functional studies indicate that c-Kit is developmentally regulated in male germ cells during both fetal and postnatal development (for review see Mauduit et al. 1999) and localized in Leydig cells in both 6-day postnatal and adult mice (Manova et al. 1990, Rothschild et al. 2003) with the capacity to influence steroidogenesis.

In the mouse, c-Kit mRNA is detected at 7.5–12.0 days postcoitum in primordial germ cells, during their lineage specification and migration to the developing gonadal ridge (Manova & Bachvarova 1991), and animals with mutations in the genes encoding either SCF or c-Kit exhibit aberrant germ cell migration and proliferation during this period (for review see Mauduit et al. 1999, Zama et al. 2005). The c-Kit mRNA was also detected by in situ hybridization within clones grown in vitro from the germ cells of the newborn mouse testis, which are termed gonocytes, although the functional significance of this observation is uncertain, as no protein expression data were provided (Hasthorpe et al. 1999). In addition, lack of in situ protein expression has been previously reported in gonocytes.
(Yoshinaga et al. 1991). In contrast to the mouse, rat gonocytes have been shown to express both c-Kit mRNA and protein (Orth et al. 1996) which was indicated to be required for their normal migration from the centre to the perimeter of the seminiferous cord from day-1 to day-5 postpartum in vivo (Orth et al. 1997). Various studies employing in situ hybridization, immunohistochemistry and Northern and Western blotting at time-points concordant with spermatogonial differentiation have reported relatively high levels of c-Kit expression in the cytoplasm of the differentiating spermatogonia and persisting low levels in the meiotic pachytene spermatocytes (Manova et al. 1990, Sorrentino et al. 1991, Dym et al. 1995). The requirement for SCF/c-Kit interactions for progression to and survival in the meiotic pachytene stage of spermatogenesis has also been indicated (Packer et al. 1995, Vincent et al. 1998). However, amongst these studies persist discordant conclusions about the timing and expression pattern of c-Kit mRNA and protein at various stages in the juvenile and adult rodent testis, and no single study has addressed this issue in rats and mice simultaneously. An alternative c-Kit transcript, the tr-kit, lacking the extracellular, transmembrane and part of the intracellular kinase domain of the c-Kit receptor protein, is expressed in post-meiotic male germ cells (Albanesi et al. 1996).

More recently, the elevation of c-Kit expression has been linked with the commitment of spermatogonial stem cells to differentiate (Schrans-Stassen et al. 1999). It has also been reported that c-Kit-positive germ cells among the prospermatogonia population of the juvenile testis (termed undifferentiated spermatogonia by Schrans-Stassen et al. 1999) have less repopulating activity than the c-Kit-negative corresponding population (Ohbo et al. 1999). This is in general agreement with studies showing that only c-Kit-negative stem spermatogonia exhibit regenerative activity when transplanted into the adult mouse testis (Shinohara et al. 1999). Thus, knowledge of the timing of c-Kit expression and its regulation is essential for further studies regarding establishment and maintenance of the stem cell population during the first wave of spermatogenesis.

In this study we interrogated the expression profile of c-Kit mRNA and protein in two independent systems, purified spermatogonia derived from 8-day-old mice testes and irradiated adult rat testis. The irradiated testis model provides a powerful avenue for investigating the regulation of c-Kit expression spermatogonia, as irradiation destroys the radiosensitive differentiating spermatogonia, leaving the testis populated only by undifferentiated spermatogonia (Kangasniemi et al. 1996). Gonadotrophin-releasing hormone (GnRH) antagonists stimulate recovery of spermatogenesis when applied at 15 weeks after irradiation and recovery is assessed by scoring the percentage of tubules containing germ cells that have reached the B spermatogonial stage or later, termed the repopulation index. By 4 weeks of treatment, the repopulation index displays a significant increase relative to the 3-week treatment group, and this index increases steadily in subsequent weeks (Shuttlesworth et al. 2000). In order to examine whether the stimulated recovery of spermatogonial differentiation with GnRH antagonist treatment is linked to changes in the SCF/c-Kit signalling system, we studied the mRNA and protein expression profiles for c-Kit in this model. Surprisingly, in both of these systems we observed that c-Kit mRNA could be detected in the absence of protein expression.

This report is the first comprehensive comparison of the timing and expression of c-Kit mRNA and protein cellular expression in developing and adult mouse and rat testes. From examination of purified spermatogonial populations and adult irradiated rat testes we have provided evidence that, in purified juvenile mouse germ cells and recovering adult rat testes, c-Kit mRNA and protein are differentially regulated in spermatogonia. Our observations are in agreement with previous findings that, based on the age of the postnatal testes, spermatogonial differentiation is indeed concordant with the expression of c-Kit protein.

Materials and Methods

Animals

Sprague–Dawley male rats and C57BL and Swiss male mice ranging from juvenile to adulthood were obtained from Monash University Central Animal Services. Swiss mice were also obtained from the University of Newcastle Central Animal Facility. The animals were killed by decapitation (rats and mouse fetuses and juveniles) or cervical dislocation (adult mice) before tissue removal. Tissues were removed immediately after the animals had been killed. Tissue samples for RNA preparation were snap frozen immediately after collection and stored at −70°C until use. Tissue samples for in situ hybridization and immunohistochemistry were placed in Bouin’s fixative and processed as described below. All investigations of the developing rat and mouse tissues conformed with the NHMRC/CSIRO/AAC Code of Practice for the Care and Use of Animals for Experimental Purposes and were approved by the Monash University Standing Committee on Ethics in Animal Experimentation and by the University of Newcastle Animal Care and Ethics Committee.

For studies on irradiated testes, adult LBNF1 male rats (hybrids between Lewis and Brown–Norway) were purchased from Harlan Sprague–Dawley, Inc. (Indianapolis, IN, USA). Animals were allowed to acclimatize for 1 week before initiation of the experiment. All procedures were approved by the M D Anderson Cancer Institutional Animal Care and Use Committee.

Irradiation and hormone treatment

Animals were anaesthetized with 0.72 mg ketamine/kg and 0.022 mg acepromazine/kg (i.m.), and then a single 6 Gy dose was delivered to the lower part of the abdomen by a
60Co γ-ray unit (Eldorado 8; Atomic Energy of Canada, Ltd, Ottawa, Canada) as described previously (Shuttlesworth et al. 2000). GnRH antagonist treatment was also performed as previously described (Shuttlesworth et al. 2000). Briefly, 15 weeks after irradiation animals were given simultaneous injections of 1.5 mg Cetrorelix pamoate and 1.5 mg Cetrorelix acetate (ASTA Medica, AG Frankfurt, Germany), each at a different site in the upper portion of the dorsal region. A second dose of Cetrorelix pamoate injection was given 3.3 weeks after the first. This dose continuously suppresses intratesticular testosterone levels by at least 80% for 5 weeks (Shuttlesworth et al. 2000).

Preparation of testes sections

Tissue samples for in situ hybridization and immunohistochemistry were placed in Bouin’s fixative for 5 h immediately after collection, then dehydrated through a graded alcohol series and embedded in paraffin. Sections of 3–5 µm were dried onto Superfrost Plus II slides (Menzel-Glaser, Braunschweig, Germany).

Isolation of spermatogonia fractions

For isolation of spermatogonia, decapsulated testes from 20–30 8-day postnatal Swiss mice were pooled and incubated for 15 min each in 0.5 mg/ml collagenase (Sigma)/Dulbecco’s modified Eagles’ medium (DMEM; Invitrogen Australia Pty Ltd, Mount Waverley, Victoria, Australia) with agitation and then in 0.25% (v/v) trypsin/EDTA in DMEM. Tubules were dissociated manually by pipetting and washed in 0.5% (v/v) bovine serum albumin (BSA) in DMEM by centrifugation. Cell pellets were resuspended in 2 ml PBS, and then the dishes were treated with 2 ml 0.5 mg/ml BSA in PBS for 1 h at 37°C. The wells were rewash in PBS three times. Washed, isolated germ cells were resuspended in 2 ml DMEM/10% (v/v) FCS in wells at a final concentration of 4 × 10^5/well and incubated for 1 or 2 h at 32°C. Unbound cells were removed and centrifuged at 1500 r.p.m. for 5 min at 4°C, washed twice in PBS and then fixed in 4% paraformaldehyde for 30 min. Cells bound to laminin-coated wells were removed by incubation with 0.25% (v/v) trypsin/1 mM EDTA at room temperature for 5 min followed by vigorous pipetting. Cells were transferred to centrifuge tubes and washed and fixed as described above. Isolated laminin-bound and -unbound spermatogonia were placed onto slides and probed for expression of c-Kit mRNA and protein expression.

Quantitative mRNA analysis

For real-time PCR analysis of each postpartum time-point, total RNA from Swiss mice (two animals pooled per sample, with duplicate samples) and Sprague–Dawley rats (one individual animal per sample, three samples per time-point) were prepared using the Qiagen RNeasy (Qiagen, Victoria, Australia) following the manufacturer’s instructions for mouse samples and the acid–phenol extraction method for rat samples (Chomczynski & Sacchi 1987) Each total RNA sample was treated with DNA-free (Ambion, Austin, TX, USA) according to the manufacturer’s specifications. Five hundred nanograms of this DNA-free total RNA was used for each 20 µl reverse transcription reaction with 100 U Superscript III reverse transcriptase (Life Technologies, Grand Island, NY, USA) and oligo-dT primer, according to the enzyme manufacturer’s guidelines. Negative control reverse transcription samples were included and these did not have any Superscript III.

PCR samples were prepared in a final volume of 10 µl using Roche Diagnostics (Castle Hill, NSW, Australia) SYBR-Green PCR master mix containing 500 nM each forward (F) and reverse (R) primers for mouse (accession number: NM_021099 c-Kit; F, 5′-tcatcgagtgtgatgggaaa-3′; R, 5′-gggacctggttcagcaca-3′) and rat (accession number: NM_022264 c-Kit; F, 5′-ctggttgcagttccatagar-3′; R, 5′-tcaacagcccttcggagggac-3′) and 1 µl reverse-transcribed template. PCR was performed in the LightCycler 2.0 Instrument (Roche Molecular Biochemicals, Mannheim, Germany) using the following light cycler conditions: denaturation 95°C for 10 min, amplification 95°C for 15 s, 60°C for 5 s, 72°C for 10 s and 72°C for 7 min for 48 cycles. Melting curve analysis and agarose gel electrophoresis were used to monitor accumulation of the PCR products.

The PCR reaction for each sample was performed in duplicate. Negative controls, in which water was used in place of the reverse-transcribed template, were included for each primer pair to detect PCR amplification of any contaminating DNA. The amounts of Actb mRNA (accession number: NM_007393; F, 5′-aggctgtgctgtccctgtat-3′; R, 5′-aaggaagctggaagaacc-3′) for the mouse samples and GAPDH mRNA (accession number: BC013852; F, 5′-tggatcgagagactca-3′; R, 5′-tgtagatattggcagtt-3′) for the rat samples were measured in each sample template to enable normalization for sample loading of target gene values between samples. The selection of an appropriate housekeeping standard involved comparing abundant mRNAs in the testsis over the entire age series. When an analysis of both GAPDH and Actb was performed to validate those genes as standards, they each demonstrated
constant abundance over the different ages. Actb for the mouse and GAPDH for the rat was chosen to reflect the alignment of their amplification pattern with the number of cycles required for the c-Kit cDNA crossing points. To correlate the crossing points from the sample amplification plot with target mRNA copy number, a standard curve was produced for each product in every experiment using cDNA containing the target gene. The data were calculated as the means ± s.d.

In situ hybridization on tissue sections

In situ hybridization was used to localize c-Kit mRNAs in mouse and rat testis sections using digoxigenin (DIG)-labelled sense and anti-sense c-Kit probes. The probes were initially generated by producing a PCR product from mouse testis cDNA corresponding to base pairs (bp) 89 through 448 of the mouse clone accession number Y00864. The PCR product of 454 bp was cloned into the pGEM T-Easy plasmid (Promega, Madison, WI, USA) following the manufacturer’s instructions. The product was verified by sequencing, and it was shown by Northern blot analysis to hybridize to a single band of 5.5 kilobase pairs, as expected for c-Kit mRNA (Qu et al. 1988, data not shown).

The plasmid containing the c-Kit cDNA fragment was used to produce sense and antisense DIG-cRNAs using SP6 and T7 RNA polymerases in separate reactions containing DIG-UTP (Roche Molecular Biochemicals) and probe labelling was quantified according to the specifications of the DIG-UTP manufacturer. Hybridization was performed using 150 ng/µl of each DIG-cRNA, with sections incubated overnight at 55°C. Washes were performed in decreasing concentrations of SSC (at 2×, 1× and 0.1×) at the hybridization temperature. Single strength SSC (1×) is 0.15 M sodium chloride and 0.015 M sodium citrate.

Treatment with anti-DIG antibody (Roche Molecular Biochemicals; 1:1000 in Roche 1× blocking solution) and then final development with 5-bromo-4-chloro-3-indolyphosphate/nitro blue tetrazolium (BCIP/NBT; Astral Biochemicals; 1:1000 in Roche 1× blocking solution) £ C. The slides were incubated overnight at 52°C. The slides were then incubated in proteinase K for 5 min at 37°C before washing (2× SSC/50% (v/v) formamide) at 37°C for 5 min in PBS. Cells were then incubated in proteinase K for 5 min at 37°C (5 µg proteinase K/ml) and then incubated in 4% (v/v) paraformaldehyde at 4°C for 15 min. Cell were washed (2× 5 min in PBS) and hybridization was performed overnight at 52°C (25 ng probe/well) in a humid chamber. Slides were then incubated in 5 min in 2× SSC/50% (v/v) formamide at 55°C twice and then 5 min in 1× SSC twice. This was followed by treatment as described above with anti-DIG antibody and then final development with BCIP/NBT (Astral Scientific) at room temperature to visualize the cells containing c-Kit mRNA.

For immunohistochemical detection of c-Kit protein, following fixation and washing (2×) in PBS/0.02% (v/v) Triton X-100, spermatogonia were allowed to adhere to 12-well slides overnight at 4°C. The slides were washed once in PBS, permeabilized with 100% methanol for 5 min and washed again in PBS. The slides were then incubated with peroxidase (3% H2O2 in PBS) for 10 min at room temperature, washed twice in PBS and incubated with the C-19 primary antibody at 1:100 in 5% (v/v) normal rabbit serum in PBS overnight at 4°C. The slides were washed twice in PBS, incubated with a secondary antibody labelled with horseradish peroxidase (sheep anti-rabbit; DAKO) at 1:100 for 1 h at room temperature, and antibody binding was detected as a brown precipitate following development with DAB for 5 min, followed by treatment with Harris haematoxylin counterstain. The cells were mounted under glass coverslips in Depex.
Results

Expression of c-Kit mRNA and protein in the developing mouse testes

Quantitative measurements of mRNAs encoding c-Kit in mouse (0-, 6-, 10-, 16-, 24- and 42-day) testes were obtained using real-time PCR on total testis RNA samples. The results indicated that day-10 and day-16 time-points had the highest levels of c-Kit expression across the ages (Fig. 1A). In situ hybridization analysis of the mouse testis indicated no discernable staining at day 2 (Fig. 1B) and day 5 (data not shown). The c-Kit mRNA signal was first

Figure 1 Juvenile and adult mouse testes: c-Kit mRNA and protein expression profile. (A) Relative c-Kit mRNA expression profile in the mouse testis age series. Data were collected from four animals for each age. The relative c-Kit expression was quantified in duplicate for each group using real-time PCR. Results are normalized to ActB values and presented in relative units of concentration. Error bars represent S.E.M. Different letters above the bars represent the statistical significance between different ages (P < 0.05) (D = day). (B and B') Day-2 mouse testis. (C and C') Day-10 mouse testis. (D and D') Adult mouse testis. (B, C and D) Anti-sense cRNAs. (b,c and d) Sense cRNA controls. (B', C' and D') CD117 anti-c-Kit antibody. (b', c' and d') No primary antibody controls. Black arrowheads, gonocyte; thin black arrows, spermatogonium; red arrows, pachytene spermatocyte; asterisks, Sertoli cell cytoplasm; Int, interstitium; ES, elongating spermatid cytoplasm; white arrows, positively stained interstitial cells. Bars equal 50 μm.
evident at day 7 in the cytoplasm of some spermatogonia (data not shown). This timing is in accord with the appearance of differentiating spermatogonia (McCarrey 1993). The signals in the day-10 (Fig. 1C) testis samples were confined to the cytoplasm of the differentiating spermatogonia. In addition to differentiating spermatogonia cytoplasmic staining, there was also spermatocyte staining with a relatively lesser intensity and interstitial staining in the adult mouse testis (Fig. 1D).

Immunohistochemistry results identified c-Kit protein in the cell types and time-points at which c-Kit mRNA expression was observed in the mouse testis. Protein was not detected within cells of the seminiferous cords at day 2 (Fig. 1B') or at day 5 (data not shown); however, staining was evident in some interstitial cells associated with the persistence of some fetal-type Leydig cells at these ages (Fig. 1B'). Signals were first evident in some spermatogonia in the day-7 testis, consistent with the in situ hybridization results (data not shown), and these were distinct in spermatogonia in the day-10 (Fig. 1C'), day-15 (data not shown) and adult testis samples (Fig. 1D'). However, protein was observed in some spermatocytes from day 10 and higher ages at a lesser intensity. The adult testis also displayed protein expression of elongated spermatids (Fig. 1D'). This staining could be attributed to the expression of tr-kit in these specific germ cell types since the antibodies used recognize an intracellular sequence in the protein (Albanesi et al. 1996). tr-kit mRNA expression was, however, not detected because the probe was designed to correspond to a region encoding the extracellular domain of the c-Kit protein.

Expression of c-Kit mRNA and protein in isolated mouse spermatogonia

Both the laminin-bound and -unbound populations were probed for the presence of c-Kit mRNA and protein. This separation was intended to discriminate between spermatogonia with stem cell properties (laminin bound) and those that lack stem cell properties (laminin unbound) which can be expected to have differentiated based on the transplantation studies of Shinohara et al. (1999). We predicted that the latter population would be more likely to contain c-Kit mRNA and protein, while the former would not. In situ hybridization examination of isolated spermatogonial cells (>95% GCNA-1 positive) showed that c-Kit mRNA expression was detectable in the cytoplasm of 100% of spermatogonia from both the laminin-bound and -unbound populations (Fig. 2A). In contrast to the in situ hybridization data, protein detection in the cytoplasm of spermatogonia occurred in 54 ± 2% of laminin-bound and 69 ± 4% of the laminin-unbound cells (means ± S.E.M.) (Fig. 2B). These experiments were performed at least three times for both mRNA and protein expression. Cells with obvious red cytoplasmic staining in the presence of primary antibody were scored as positive for c-Kit protein.

Expression of c-Kit mRNA and protein in the developing rat testes

Quantitative measurements, using real-time PCR, of mRNAs encoding c-Kit in rat (1-, 2-, 3-, 5-, 7-, 9-, 15-, 20-, 30-day and adult) testes were obtained using real-time PCR on total testis RNA samples. The c-Kit mRNA level was highest in the rat testis on day 15 (Fig. 3A). To more precisely assess the pattern of change in mRNA levels, day-12 testis samples were included in a subsequent analysis (data not shown) with samples spanning day 9 through 26, and the day-12 sample had a higher relative level of c-Kit mRNA compared with either day 9 or day 15. Thus the peak in expression of this mRNA most likely occurs between day 9 and day 15. In situ hybridization analysis of the 4-day postnatal rat testes did not show any detectable signal within gonocytes or in any other cell type (Fig. 3B). At day 13 and day 15, a weak to moderate signal for c-Kit mRNA was observed in the cytoplasm of spermatogonia and spermatocytes (data not shown). The signal appeared to be more intense and distinct in spermatogonia of day 19, with staining in only

![Figure 2](https://www.reproduction-online.org)
some spermatocyte (Fig. 3C). Adult rat testes also exhibited similar staining patterns (Fig. 3D). Both day-19 and adult testes had detectable interstitial staining.

Immunohistochemical detection of c-Kit protein revealed faint staining in the cytoplasm of gonocytes of the day-4 (Fig. 3B') and day-6 rat testis (data not shown). Since no mRNA signal was detected in gonocytes it is possible that the c-kit protein is produced in male germ cells of the fetal testis, as previously documented (Manova & Bachvarova 1991) and the protein remains stable within these cells for several days. A similar observation has been made for the activin βA subunit in the fetal and
newborn rat testis (Meehan et al. 2000). A relatively weak signal was observed in differentiating spermatogonial subtypes at day 13 (data not shown), while at days 15 (data not shown) and 19 (Fig. 3C') most of the differentiating spermatogonia had readily detectable brown cytoplasmic staining, as did some pachytene spermatocytes. In the adult testis, c-Kit protein was observed in differentiated spermatogonia, pachytene spermatocytes and elongated spermatids (Fig. 3D'). As in the case of the in situ hybridization localization of c-Kit mRNA, the protein signal was not evident in all pachytene spermatocytes. The antibody staining of pachytene spermatocytes was more readily detectable in the day-15 and day-19 testis sections than in those from the adult testes, giving the impression that the protein was present in more of these cells and at a higher level. Staining of a distinct intracellular vesicle was also noted within some pachytene spermatocytes. Interstitial staining was present in the day-19 and adult testes sections (Fig. 3C' and D'). Tr-kit staining in the elongating spermatids was evident in the adult testis (Fig. 3D').

**Irradiated adult rat testis: a comparison of mRNA and protein expression**

*In situ* hybridization analysis of irradiated rat testis samples showed that c-Kit mRNA was detectable in the cytoplasm of spermatogonial cells from rats that received no GnRH antagonist treatment (Fig. 4A), 1 week of treatment (Fig. 4B), 2 weeks of treatment (Fig. 4C) and 4 weeks of treatment (Fig. 4D). Staining in interstitial cells, including Leydig and endothelial cells was particularly prominent in the samples from animals treated with GnRH antagonist for 4 weeks.

In contrast to the *in situ* hybridization data, protein was not detected in spermatogonia in the irradiated rat samples from animals receiving no treatment (Fig. 4A'), 3 days (data not shown), 1 week (Fig. 4B') and 2 weeks (Fig. 4C') of treatment. The signal was, however, readily evident in cells with the appearance of differentiating spermatogonia that were present in the 4-week GnRH antagonist treatment group (Fig. 4D'). Leydig cell staining was present in all samples, in agreement with the *in situ* hybridization data. These data were consistently observed using two different antibodies (CD117 and C-19 (data not shown)) that were verified as recognizing c-Kit protein by Western blot. These results were identical in triplicate experiments performed using testis sections from three individual animals.

**Discussion**

To provide for ease of comprehension, the findings have been summarized in Fig. 5. In essence, this study has illustrated the relationship between c-Kit mRNA and protein expression and highlights the key developmental time-points for the onset of differentiation of both rat and mouse germ cells during the first wave of spermatogen-esis. Published descriptions of c-Kit localization and expression do not come to a common conclusion about the expression of c-Kit mRNA and protein in pachytene spermatocytes (Manova et al. 1990, Sorrentino et al. 1991, Dym et al. 1995). In this study, both mRNA and protein expression data identified c-Kit in pachytene spermatocytes and showed that it is undetectable in round spermatids. The functional significance of expression in pachytene spermatocytes could relate to the survival of these cells during meiosis (Yan et al. 2000). The full-length c-Kit is, therefore, present both in the cytoplasm, and presumably the plasma membrane, of differentiating spermatogonia and pachytene spermatocytes in both rat and mouse. Comparison of mRNA levels measures in the total rat and mouse testis RNA by real-time PCR analysis with the appearance of c-Kit mRNA within specific cell types by *in situ* hybridization, in both rodents, revealed that expression levels increase at time-points concordant with the appearance of differentiating spermatogonia containing this transcript (McCarrey 1993). In comparison with the spermatogonial and spermatocyte staining, there was weak to no staining in the Leydig cells, a finding which suggests that Leydig cells contribute little to the total level of c-Kit mRNA in the testis at all ages tested. To complete this study, we were required to undertake a comparison of two commercially available antibodies that recognize the intracellular domain of the rodent c-Kit protein, and similarities and differences in their target recognition became apparent from this analysis. Though both antibodies (CD117 and C-19) gave similar outcomes when applied to Bouin-fixed sections of rat testis, only the CD117 antibody gave a clear signal on mouse testes sections.

In our analyses of two independent systems we have also uncovered evidence that c-Kit mRNA can be produced in the absence of detectable protein. Examination of spermatogonia isolated from the day-8 mouse testis indicates that c-Kit mRNA and protein are differentially regulated. Similarly, in the case of the irradiated adult rat testis, while mRNA was observed in spermatogonia in all sample groups, both mRNA and protein was detectable only in the differentiating spermatogonia present in the 4-week GnRH antagonist treatment group, suggesting that undifferentiated spermatogonia express c-Kit mRNA but not protein. Based on these observations, we hypothesize that distinct factors regulate c-Kit mRNA transcription and translation. These factors are presumably part of the microenvironment of the testis.

Our examination of the irradiated rat testis model provides a clue that factors regulating c-Kit protein synthesis in the testis are influenced by its exposure to hormonal stimulation. Coupling the present data with previous observations of spermatogonial morphology in these testes, we now understand that c-Kit protein expression occurs only when differentiating spermatogonia emerge in the testes of irradiated rats treated with the GnRH antagonist for 4 weeks and more (Shuttlesworth et al. 2000). In a study of W/Wv mutant mice, which produce
non-functional c-Kit receptor protein, spermatogonia expressing c-Kit are present in 2-week-old W/Wv mutant mice but are absent in 10-week-old animals (Ohta et al. 2003). Administration of a GNRH antagonist for 4 weeks to reduce intratesticular testosterone in 10-week-old W/Wv mice resulted in the emergence of c-Kit-positive spermatogonia. While production of c-Kit mRNA was not assessed in the latter study, it appears that reduction of the intratesticular testosterone concentration stimulates differentiation of spermatogonia in a manner independent of c-Kit function, but leading to the onset of c-Kit protein expression. Another known factor to stimulate the expression of surface c-Kit protein is BMP4, as shown by the expression of surface c-kit protein on 4-day postnatal spermatogonia after exposure to BMP4 (bone morphogenetic protein-4; Pellegrini et al. 2003). Microarray data for BMP4 expression in the irradiated rat testes indicate an elevation in this mRNA after 2 weeks of GnRH antagonist treatment group (O U Bolden-Tiller & M L Meistrich, unpublished results). These data coupled with our localization studies on c-Kit expression suggest that there could be translational regulation of c-Kit by BMP4. Current studies are underway to assess BMP4 protein expression at this stage.

In conclusion, the results of our study identify the key time-points and cell types for c-Kit mRNA production in the developing rodent testis and indicate that mRNA is...
present in the absence of detectable levels of protein. It also indicates that a spermatogonial stem cell could have c-Kit mRNA expression and exhibit regenerative capacity, while corresponding protein expression in the spermato- gonia results in lesser potential for regeneration. This observation is particularly intriguing, as it provides the opportunity to design appropriate studies for analysis of the regulation of both c-Kit mRNA transcription and its translation into protein, the latter of which is a key determinant of germ cell survival, proliferation and migration.

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