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5	Clonal inactivation of telomerase
6	promotes accelerated stem cell differentiation
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1 Summary

2	Telomerase is intimately associated with stem cells and upregulated in cancer, where it serves essential
3	roles through its catalytic action in elongating telomeres, nucleoprotein caps that protect chromosome
4	ends ¹ . Overexpression of the telomerase reverse transcriptase (TERT) enhances cell proliferation
5	through telomere-independent means, yet definitive evidence for such a direct role in stem cell function
6	has yet to be revealed through loss-of-function studies. Here, we show that conditional deletion of TERT
7	in spermatogonial stem cells (SSCs) markedly impairs competitive clone formation. Using
8	lineage-tracing from the Tert locus, we find that TERT-expressing SSCs yield long-lived clones, but that
9	selective TERT-inactivation in SSCs causes accelerated stem cell differentiation thereby disrupting
10	clone formation. This requirement for TERT in clone formation is bypassed by expression of a
11	catalytically inactive TERT transgene and occurs independently of the canonical telomerase complex.
12	TERT inactivation induces a genome-wide reduction in open chromatin evident in purified SSCs, but not
13	in committed progenitor cells. Loss of TERT causes reduced activity of the MYC oncogene and
14	transgenic expression of MYC in TERT-deleted SSCs efficiently rescues clone formation. These data
15	reveal a required catalytic activity-independent role for TERT in preventing stem cell differentiation, forge
16	a genetic link between TERT and MYC and suggest new means by which TERT may promote
17	tumorigenesis.

1 Main Text

2 Telomerase is enriched in tissue stem cells and activated by somatic promoter mutations in many 3 cancers ²⁻⁴. The core of the telomerase enzyme is comprised of the catalytic subunit TERT and the 4 telomerase RNA component (Terc), a small non-coding RNA scaffold that encodes the template for 5 telomere addition¹. The critical requirement for telomerase in long-term cell viability is conserved from 6 single cell eukaryotes to humans. Cell proliferation in the absence of telomerase results in a lag phase 7 that is initially well tolerated while telomere reserves are ample. But, proliferation for extended periods in 8 the absence of telomerase culminates in senescence or cell death as telomeres progressively shorten 9 and eventually become dysfunctional. This is particularly evident in laboratory mice, which have very 10 long telomeres (40-80 kb vs 5-15 kb in humans). Telomerase knockout mice are initially viable, but 11 subsequent intergenerational breeding results in severe tissue defects in the advanced generations⁵⁻⁷. 12 These findings established the paradigm that TERT is required only for its role in synthesizing telomeres. 13 In contrast with these loss-of-function studies, overexpression studies have sugested that TERT 14 promotes cell proliferation independent of its enzyme function. Conditional transgenic TERT expression 15 caused proliferation of hair follicle stem cells⁸, skin basal layer keratinocytes⁹ and kidney podocytes¹⁰. 16 These effects of TERT were separable from telomere synthesis because they also occurred with a catalytically inactive TERT allele, or in mice lacking *Terc*⁸⁻¹⁰. In this context, TERT has been shown to 17 18 activate MYC, WNT and NFkB pathways⁸⁻¹⁴. However, the role of non-canonical functions of TERT in 19 tissue stem cells remains unclear due to the lack of definitive and immediate phenotype in TERT 20 knockout mice.

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Tissue homeostasis and carcinogenesis are shaped by cell competition, a mechanism to optimize cell composition in tissues by promoting replacement of damaged or unfit cells with more robust neighboring cells¹⁵. In renewing mammalian tissues including intestine, skin, and testis, competitive repopulation is a fundamental property of adult stem cells¹⁶. Stem cell competition results in a dominant expansion of

1 more fit "winner clones" and an elimination of less fit "loser clones" at the niche¹⁷. This competitive 2 behavior is also characteristic of carcinogenesis during which oncogenic mutations drive cells to clonally 3 expand their territory through a process of super-competition¹⁸. Among tissues where competitive 4 repopulation has been ovserved, testis shows high telomerase activity and exhibits unusual telomere 5 dynamics in that telomere lengths are preserved with aging, in contrast to the progressive shortening 6 seen in other human tissues. We previously found that spermatogonial stem cells (SSCs) express high 7 levels of TERT and TERT is downregulated with lineage commitment⁷. To understand the 8 telomere-independent role of TERT in stem cells, we developed a system to mark single SSCs 9 expressing TERT coupled with the ability to conditionally inactivate TERT using a lineage tracing 10 approach. Combining with transgenic rescue experiments and genomic assays, these studies establish 11 TERT as a mediator of stem cell competition in the testis, where it supports stem cell function through a 12 non-canonical mechanism independent of telomere synthesis.

1 Results

2 High TERT expression marks long-term SSCs that undergo stem cell competition

3 Spermatogenesis is a dynamic process to produce sperm, composed of mitosis, meiosis and 4 post-meiotic maturation (Extended Data Fig. 1a,b). In the testis, SSCs reside within a functionally and 5 morphologically heterogeneous population undifferentiated spermatogonia (US). Singly isolated Asingle 6 (A_s) US undergo incomplete cytokinesis, subsequently producing progressively elongating chains of 7 interconnected cells (A_{pr} - A₁₆) (Extended Data Fig. 1b)¹⁹. Maturation of US yields differentiating 8 spermatogonia (DS), which is accompanied by loss of stem cell potential. To measure Tert expression in 9 distinct spermatogonia subpopulations, we purified MCAM^{high} KIT⁻ US (US-h), MCAM^{med} KIT⁻ US (US-m) 10 and MCAM^{med} KIT⁺ DS (Extended Data Fig. 1a-c). Among these subpopulations, *Tert* mRNA expression 11 was high in both US-h and US-m cells, and sharply decreased in DS cells (Extended Data Fig. 1d). 12 Tert-Tdtomato reporter mice also showed high Tdtomato expression both in US-h and US-m (Extended 13 Data Fig. 1e). These data indicate that the entire US population exhibits high *Tert* expression.

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15 To functionally study TERT-expressing spermatogonia, we developed a lineage tracing assay using Tert^{CreER/+}; Rosa26^{lsl-Tdtomato/+} (Tert^{CreER/+}) mice, in which TERT-expressing cells are permanently labeled 16 upon tamoxifen-dependent activation of CreER to express Tdtomato (Fig. 1a,b)²⁰. Marking SSCs results 17 18 in long-lived Tdtomato⁺ clones, also known as patches, comprised of many daughter cells produced by the labeled SSC ²¹. If committed progenitor cells are labeled, only a small transient clone is generated 19 20 and these cells are lost through differentiation (Fig. 1a,b). Sparse labeling allows rare SSCs to be 21 marked, and in this context each patch derives from a single SSC. Thus, measuring the number of 22 Tdtomato⁺ patches allows a quantitative assessment of stem cell self-renewal activity.

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At two days after tamoxifen injection, TdTomato was detected similarly throughout the population of US
but not in KIT⁺ cells (Extended Data Fig. 2a-c). At three months, marking TERT⁺ cells yielded labeled
patches (Fig. 1c and Extended Data Fig. 2e). As we varied the dose of administered tamoxifen from 0.25

1 mg to 4 mg, patch number increased in a dose-dependent manner (Fig. 1d, and Extended Data Fig. 2d). 2 Patch length remained constant below 1mg but increased above 2mg, reflecting the fusion of two 3 independently labeled clones with high dose (Fig. 1e). In mice treated with 1mg tamoxifen and traced for 4 one year, patch length increased, clone number decreased, and the total aggregated Tdtomato⁺ patch 5 length (mean patch number times mean patch length) remained constant compared with those traced 6 for three months, consistent with previously described stochastic competition with unlabeled clones (Fig. 7 1d-f, and Extended Data Fig. 2d)²¹. At both three months and one year, patches were comprised of cells 8 throughout the spermatogenic lineage (Extended Data Fig. 2f). Altogether, these data show that 9 TERT-expressing SSCs generate long-lived clones and exhibit competition within the stem cell pool.

10

11 **TERT deletion impairs SSC-mediated clone formation**

12 To investigate a direct and immediate role for TERT in stem cells, we developed a competitive clone formation assay using Tert^{CreER/flox}:Rosa^{/sl-Tdtomato/+} (Tert^{CreER/flox}) mice (Fig. 1g, and Extended Data Fig. 13 14 3a-c). In this strain, activation of CreER induces Tdtomato labeling and concomitantly inactivates Tert in 15 the same TERT⁺ cell, enabling one to trace the fate of cell clones deriving from SSCs in which *Tert* has 16 been somatically deleted in an environment where most neighboring cells retain TERT expression. At 17 two days after 1mg tamoxifen treatment, GFRA1⁺ A_s and A_{pr} clones were marked indistinguishably in both Tert^{CreER/+} and Tert^{CreER/flox} mice (Fig. 1h,i). Pulse labeling efficiently eliminated Tert mRNA in 18 19 Tdtomato⁺ US (Fig. 1j,k). At three months and six months, deletion of TERT caused a marked reduction 20 in the number of Tdtomato⁺ clones and diminished mean patch length (Fig. 1I-n). Correspondingly, the 21 total aggregated Tdtomato⁺ patch length was sharply reduced (Fig. 1o). Together, these findings 22 indicate that TERT-deletion compromises long-term clone formation ability of SSCs under theses 23 competitive conditions.

24

TERT promotes SSC competition independent of its catalytic activity and the telomerase
 complex

1 To determine whether the effect of TERT in enhancing SSC competition depends on its catalytic activity. 2 we developed a system to rescue the defect in TERT-deleted cells using tetracycline-regulated TERT 3 transgenes: either one that is wild-type (TetO-TERT) or one that is catalytically inactive (TetO-TERTci) 4 due to a single amino acid substitution in the catalytic site ⁹. We produced compound mouse strains in 5 which activation of CreER simultaneously deletes the floxed Tert gene while inducing expression of 6 transgenic Tert by triggering expression of the tetracycline transactivator (tTA) that binds and activates 7 the TetO promoters (Fig. 2a). We analyzed how TERT loss affected clone formation in Tert^{CreER/flox}:Rosa^{lsI-Tdtomato/lsI-tTA} (Tert^{CreER/flox}) vs. Tert^{CreER/+}:Rosa^{lsI-Tdtomato/lsI-tTA} (Tert^{CreER/+}) controls and we 8 compared these results with restoration of Tert expression in Tert^{CreER/flox}:Rosa^{/sl-Tdtomato//sl-tTA}:TetO-Tert 9 (*Tert*^{CreER/flox} + *Tert*) mice or *Tert*^{CreER/flox}:*Rosa*^{IsI-Tdtomato/IsI-tTA}: *TetO-Tertci* (*Tert*^{CreER/flox} + *Tertci*) mice (Fig. 2a). 10 11 Transgenic Tert and Tertci expression was induced at a physiological range by treating mice with low 12 dose doxycycline, which suppresses binding of tTA to the TetO promoter (Extended Data Fig. 4a). 13 Transgenic expression of wild-type Tert, but not Tertci, restored telomerase activity (Extended Data Fig. 14 4b). Intriguingly, expression of either Tert or Tertci transgenes fully rescued patch number. average 15 patch length, and total patch length (Fig. 2b-e). Furthermore, expression of TERTci significantly increased total patch length compared to control *Tert^{CreER/+}* mice (Fig. 2e). These results indicate that 16 17 TERT promotes enhanced stem cell competition independent of its catalytic activity.

18

Terc encodes the RNA template for telomere addition and serves as the central scaffold for assembly of the telomerase complex. Therefore in the absence of *Terc*, TERT and the other components of telomerase fail to associate ^{1,22}. To determine whether formation of the telomerase complex is required for the TERT-dependent SSC clone formation, we produced first generation

23 *Terc^{-/-}: Tert^{CreER/flox} :Rosa^{ls/-Tdtomato/+}* (G1 *Terc^{-/-}: Tert^{CreER/flox}*) mice (Extended Data Fig. 4c. In this strain,

- telomerase was inactive in testes (Extended Data Fig. 4d). Three months after tamoxifen treatment,
- 25 conditional deletion of *Tert* in *Terc*-deficient mice impaired patch number, patch length, and total patch

length (Extended Data Fig. 4e -h). These findings uncouple the requirement for TERT in stem cell
 competition from engagement in the classical telomerase complex.

3

Somatic deletion of *Tert* in mice with very long telomeres is unlikely to cause telomere dysfunction ^{7,23,24}. 4 5 When telomeres become short, however, a DNA damage response activates the p53 tumor suppressor 6 protein, triggering either cell death or senescence ²⁵. To understand whether the impaired clone 7 formation in TERT-deleted SSCs is mediated by p53, we generated *Trp53^{flox/flox}:Tert^{CreER/flox}:Rosa^{ls-Tdtomato/+}* mice and examined competitive clone formation (Extended Data 8 9 Fig. 5a). At day five, the deletion efficiency of the *Trp53-flox* alleles in Tdtomato⁺ US was 44.4%10 (Extended Data Fig. 5b,c). We found that deletion of p53 failed to rescue the impaired clone formation 11 associated with Tert inactivation, as measured by patch number, patch length, and total patch length 12 (Extended Data Fig. 5d-g). Consistent with these findings, we found no accumulation of γ H2AX, a marker of DNA damage, in *Tert*^{CreER/flox} mice, whereas γ H2AX was elevated in G6 *Tert*^{Tdtomato/Tdtomato} mice 13 14 with dysfunctional telomeres (Extended Data Fig. 5h,i). Taken together, the effects of TERT in promoting 15 stem cell-derived clone formation is independent of catalytic activity, the DNA damage response and the 16 canonical telomerase complex.

17

18 Accelerated differentiation and impaired proliferation of SSCs lacking TERT

19 The preferential elimination of conditionally TERT-deleted SSCs could be caused by accelerated 20 differentiation, impaired proliferation, or increased apoptosis. To distinguish among these possibilities, 21 we investigated how clonal deletion of TERT influences SSC fate across at time points. At 14 days after tamoxifen administration, Tdtomato⁺ A_s and A_{pr} US were significantly decreased in *Tert*^{CreER/flox} mice, and 22 23 the reduction of those cells was rescued by transgenic expression of either TERT or TERTci (Fig. 2f and 24 Extended Data Fig. 6a). At six weeks when a comparable frequency of labeled patches were found in Tert^{CreER/+} and Tert^{CreER/flox} mice, labeled clones harbored significantly fewer US (Fig. 2g,h Extended Data 25 26 Fig. 6b.c). These results indicate that TERT deletion promotes differentiation of US. To understand how

TERT loss affects cell proliferation, we measured BrdU incorporation seven days after tamoxifen administration. The percentage of US incorporating BrdU was significantly reduced in *Tert^{CreER/flox}* animals and proliferation was rescued with transgenic expression of *Tert* or *Tertci* (Fig. 2i and Extended Data Fig. 6d). There was no increase in apoptosis by cleaved-PARP staining in *Tert^{CreER/flox}* mice, but apoptosis was elevated in testes from G6 *Tert^{Tdtomato/Tdtomato}* mice with critically short telomeres (Extended Data Fig. 6e,f). Taken together, impaired clone formation of TERT deleted SSCs is caused by accelerated differentiation and decreased proliferation.

8

9 TERT deletion compromises chromatin accessibility in SSCs but not in committed progenitors 10 To understand how TERT deletion affects global chromatin structure, we performed ATAC-seq, which allows an assessment of chromatin accessibility genome-wide ²⁶. To first define the patterns of 11 chromatin changes during normal spermatogenesis, we purified US-h, US-m, and DS from Tert^{CreER/+} 12 13 mice that had been injected with tamoxifen seven days prior to isolation, while spermatocytes (SP) and round spermatids (RS) were purified based on differential *Tert* promoter activity in *Tert*^{Tdtomato/+} mice⁷. 14 15 Principal component analysis (PCA) revealed that US-h and US-m clustered together in the lower left guadrant, consistent with their similar gene expression patterns (Extended Data Fig. 7a)²⁷. DS cells 16 17 localized in the upper left quadrant, while SP and RS populations were clustered together in the right 18 lower guadrant (Extended Data Fig. 7a), indicating that the PC1 axis captures the changes in global 19 chromatin state associated with differentiation. Similarly, Pearson correlation hierarchical clustering 20 showed a high correlation in open chromatin patterns among spermatogonia subpopulations, but abrupt 21 changes of chromatin accessibility globally upon entry to meiosis (Extended Data Fig. 7b). The number 22 of unique ATAC-seq peaks and promoter chromatin accessibility around transcription start sites were 23 highest in US-h and US-m and decreased significantly during differentiation into DS and SP (Fig. 3a,b, 24 and Extended Data Fig. 7c,d). The promoter region of *Tert* was accessible in US-h, US-m and DS, but 25 inaccessible in SP and RS, consistent with the expression pattern of TERT during spermatogenesis

(Extended Data Fig. 8a) ⁷. Altogether, these data reveal that SSCs show a marked increase in chromatin
 accessibility, indicating that a global reduction in chromatin accessibility occurs during spermatogenesis.
 3

4 To understand how conditional TERT-deletion influences chromatin accessibility in SSC populations, we performed ATAC-seq on US-h. US-m and DS isolated from *Tert*^{CreER/flox} mice. Pearson correlation 5 6 hierarchical clustering showed that the overall pattern of chromatin accessibility in TERT-deleted cells remained similar to that of *Tert*^{CreER/+} controls (Extended Data Fig. 7b). However, PCA revealed that 7 8 TERT deletion caused a shift in the US-h and US-m along the differentiation axis, whereas TERT loss 9 had no discernible effect on committed DS (Fig. 3c). Deletion of TERT in the US populations caused a 10 marked reduction in the number of open chromatin peaks and diminished chromatin accessibility 11 surrounding TSSs to a level resembling the control DS (Fig. 3a,b,d,e and Extended Data Fig. 7e). In 12 contrast, peak number was unaffected by TERT deletion in DS (Fig. 3 a,b,d,e and Extended Data Fig. 13 7e). This reduction in open chromatin peaks was evident in genes associated with stemness in SSCs, 14 including Ret, Gfra1, Cdh1, and Zbtb16, whereas those associated with differentiation including Kit, 15 Prm2, and Prm3 remained unchanged (Extended Data Fig. 8b-d). Pathway analysis revealed that genes in the MAPK signaling pathway, which promotes self-renewal of SSCs ²⁸, were particularly enriched 16 17 among those showing loss of open chromatin peaks in the TERT-deleted undifferentiated 18 spermatogonia (Extended Data Fig. 7f). Taken together, conditional inactivation of TERT caused a loss 19 of open chromatin selectively in the stem cell containing population but not in committed progenitors, 20 consistent with accelerated stem cell differentiation caused by TERT deletion.

21

22 TERT promotes competitive clone formation of SSCs through MYC

To understand how TERT promotes competitive clone formation in SSCs, we examined gene
 expression in TERT-deleted US-h cells by RNA-seq seven days after tamoxifen administration. PCA

25 showed that TERT-deleted US-h clustered separately from TERT⁺ controls (Fig. 4a). Using strict cutoffs

26 for significance, there were 23 genes downregulated and 116 genes upregulated in TERT-deleted US-h

1 (Extended Data Fig. 9a). Gene set enrichment analysis revealed that spermatogenesis-related genes 2 were upregulated in TERT-deleted US-h, consistent with their accelerated differentiation (Extended Data 3 Fig. 9b). Several gene sets were downregulated in TERT-deleted US-h cells, including E2F targets and 4 G2M checkpoints, reflecting the quantitative reduction in proliferation in these cells (Extended Data Fig. 5 9c). The most significantly downregulated gene set was 'MYC targets v1' and a second gene set 'MYC 6 targets v2' was also represented (Fig. 3b, and Extended Data Fig. 9c). Consistent with this, MYC protein 7 was significantly decreased in TERT-deleted US and MYC levels were restored by transgenic 8 expression of TERT or TERTci (Fig.4c, and Extended Data Fig.9d,e). Myc mRNA levels remained 9 unchanged, suggesting that TERT promotes MYC expression at the post-transcriptional level (Extended 10 Data Fig. 9f).

11

12 MYC is a transcription factor that promotes cell competitiveness by regulating cell proliferation, growth, 13 and metabolism ²⁹⁻³². MYC has been shown to promote SSC self-renewal and also an important 14 oncogene ³³⁻³⁵. Given the genetic links between TERT and MYC, we hypothesized that MYC 15 overexpression might rescue the failure of clone formation in TERT-deleted SSCs. To test this idea, we intercrossed Tert^{CreER/flox}: Rosa^{lsI-Tdtomato/IsI-tTA} mice with TetO-human MYC transgenic mice (Tert^{CreER/flox}) 16 17 +MYC) (Fig. 4d). This system allows simultaneous deletion of the residual TERT allele and activation of 18 transgenic MYC selectively in a lineage of TERT-expressing stem cells. To limit expression levels of 19 transgenic MYC, mice were treated with doxycycline, which reduced transgenic MYC mRNA levels by 20 5.6 fold (Extended Data Fig. 9g). Three months after tamoxifen treatment, the defects in clone formation 21 associated with TERT loss were dramatically rescued by MYC expression, as measured by patch 22 number, patch length and total patch length (Fig. 4e-h). MYC expression also restored the number of 23 GFRA1⁺ cells, proliferation and expression of MYC target genes (Fig. 4i,j and Extended Data Fig. 9h). 24 Taken together, these results indicate that the marked defect in stem cell competition associated with 25 TERT loss is due to impaired MYC function and establish an epistatic relationship between TERT and 26 MYC.

1

2 Discussion

3 TERT is expressed in many stem cell compartments governed by competitive repopulations and TERT 4 upregulation is selected for during human carcinogenesis. By tracing the fate of individual SSCs in vivo, 5 we found that loss of TERT severely compromises competitive clone formation and accelerates stem 6 cell differentiation. In this context, TERT-deleted stem cells represent "loser" clones outcompeted by 7 TERT-proficient "winner" clones. This requirement for TERT in competitive clone formation is 8 independent of telomeres and TERT catalytic function; instead, TERT maintains chromatin accesiblity 9 across many genes expressed in stem cells. These findings establish TERT as a key determinant 10 regulating competition between tissue stem cells by favoring self-renewal and disfavoring differentiation 11 (Extended Data Fig. 10). This result is surprising in that germline inactivation of TERT in mice is well 12 tolerated initially, although SSC function is ultimately compromised by telomere dysfunction after many 13 generations of telomerase-deficiency. Competitive behavior of SSCs may expain why germline 14 mutations in Tert do not cause an obvious phenotype in laboratory mice. Selection of winner clones and 15 elimination of loser clones in the testis may be determined in part based on competition for limiting niche 16 signals ³⁶. This niche-dependent competition between TERT-proficient and TERT-deficient cells may 17 amplify cell-autonomous defects caused by acute TERT deletion, while promoting elimination of 18 TERT-deleted SSCs. Homogeneous deletion of TERT throughout the organism does not create a 19 TERT-dependent competition, and in this context TERT-deficient cells are capable of self-renewal and 20 differentiation to produce sperm. Thus, germline gene inactivation identifies only one layer of activity, 21 whereas clonal, conditional gene deletion can reveal deeper aspects of gene function.

22

While TERT levels are important in maintaining stem cell function, they may play a similar role during carcinogenesis. TERT is upregulated in many human tumors through non-coding mutations in the proximal TERT promoter. These TERT promoter mutations show strong positive selection during tumor progression, as the prevalence of these mutations increases substantially from pre-invasive to invasive

1 stages. Our data establishing an epistatic relationship between TERT and MYC provide support for this 2 model. MYC activity is central in determining outcomes in cell competition; cells expressing higher MYC 3 outcompete those with lower MYC and MYC represents a key node in many human cancers ²⁹⁻³². TERT 4 upregulation in cancer likely increases telomerase activity and stabilizes telomeres, thereby preventing 5 the negative effects of dysfunctional telomeres on cell cycle and cell survival. At the same time, TERT 6 upregulation may enhance cell competition through MYC and its non-canonical effects in promoting 7 self-renewal and preventing differentiation. MYC can also control TERT levels by directly activating 8 transcription of *Tert* through promoter binding ³⁷⁻³⁹. Therefore TERT and MYC may comprise a positive 9 feedback loop that promotes cell competition in various tissues during development, homeostasis and 10 carcinogenesis. Our findings using clonal deletion in SSCs add important new complexity to our 11 understanding of TERT function. Taken together, this study establish a role for TERT in promoting clonal 12 competition in stem cells in vivo with important implications for understanding tissue homeostasis, cancer 13 development and telomere maintenance.

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13		

1 Methods

2 Animals. All animal protocols were approved by the Institutional Animal Care and Use Committee at 3 Stanford University. All experiments are in compliance with the ethical regulations of Stanford University. 7,10,20 4 Tert-Tdtomato mice were Tert-CreER. TetO-Tert. TetO-Tertci, previously reported Rosa-Is/Tdtomato⁴⁰, Rosa-Is/-tTA⁴¹, TetO-hMYC⁴², Terc-KO⁴³, Trp53-flox⁴⁴ mice were purchased from 5 6 The Jackson laboratory. Tamoxifen (Cayman) was dissolved in corn oil (Sigma-Aldrich) at 5 - 20mg/ml 7 by incubating at 50°C for 30min with mixing every 5min. Two- to three-month-old mice were 8 administrated with 0.25 - 4mg per 25g body weight tamoxifen by oral gavage or intra-peritoneal injection. 9 Doxycycline (Sigma-Aldrich) was dissolved in drinking water in light-protected bottles at 1 or 3 µg/ml and 10 changed every 4 days. BrdU (Sigma-Aldrich) was dissolved in PBS at 10mg/ml and intra-peritoneally 11 injected at 1.25mg per 25g body weight at 2 hours before sacrifice.

12

13 Generation of TERT-flox mice

14 9kb fragment of TERT locus was subcloned and Lox-Puro-lox cassette from pBS.DAT-LoxStop plasmid 15 (kindly gifted from David Tuveson) was inserted at the BsiWI site in the second intron. Another loxP 16 sequence and Ndel site were inserted at the Kasl site in the sixth intron. The targeting vector was 17 linearized and electroporated into J1 mouse ES cells. After positive selection with puromycin, correctly 18 targeted ES clones were selected by long-range PCR and Southern blotting, and then injected into 19 C57BL6 blastocysts to generate the knock-in line. To remove floxed Puro cassette, the knock-in line was 20 crossed with CMV-cre mice ⁴⁵ and puro-negative TERT-floxed mice were selected by PCR and Southern blotting using genomic DNA from tail tips. *TERT^{flox/+}* mice were born at normal Mendelian frequency. 21

22

23 Lineage tracing assay.

After tamoxifen injection, testes were detunicated, dissociated using fine forceps in PBS containing 1mg/ml collagenase IV (Worthington) for 10min to remove interstitial cells, and placed in cold PBS. Images were captured with a fluorescent dissection microscope and the patch number and length were

measured with ImageJ. Total patch length was calculated by multiplying the patch number by the
average patch length.

3

4 Whole-mount Immunofluorescence of seminiferous Tubules

5 Seminiferous tubules were dissociated using fine forceps in PBS containing 1mg/ml collagenase IV for 6 10min, fixed with 4%PFA at 4°C for 2h, cleared with 0.1% Igepal CA-630 (Sigma-Aldrich) in PBST, and 7 dehydrated and rehydrated by immersing in a gradient of methanol diluted with PBST (25%, 50%, 75%, 100%, 75%, 50%, 25%) at 4°C for 5min each. After washing in PBST, tubules were incubated in 8 9 blocking buffer (0.5% BSA PBST) (Sigma-Aldrich), followed by incubation with antibodies in Immuno 10 shot immunostaining Mild (Cosmo Bio) at 4°C for two days. After extensive wash with PBST, tubules 11 were incubated with secondary antibodies in blocking buffer at room temperature for 90min, washed 12 with PBST, and then mounted in Vectashield with DAPI (Vector laboratories). Images were captured on 13 a Leica SP5 confocal microscope and processed in Photoshop.

14

15 Section immunostaining

16 Testes were detunicated, fixed with 4% PFA at 4°C overnight, incubated in a gradient ethanol, xylen, 17 embedded in paraffin, and cut into 5-µm sections. After rehydration, antigen retrieval using Antigen 18 retrieval solution citric acid or tris-based (Vector) for 10min in a pressure cooker. Sections were blocked, 19 incubated with primary antibody at 4°C overnight. After washing with PBS, sections were incubated with 20 secondary antibodies at room temperature for 1h and mounted in Vectashield with DAPI. For co-staining 21 using rabbit anti-RFP antibodies and rabbit anti-PLZF antibodies, sections were antigen retrieved, 22 incubated with anti-RFP antibody, then with HRP-conjugated anti-rabbit secondary antibody as 23 described above, and signals were detected with TSA plus Cyanine 3 system (Perkin Elmer). Those 24 antibodies were stripped off by antigen retrieval, and sections were further stained with BrdU,

1 cleaved-PARP, or PLZF antibodies. For BrdU detection, those slides were treated with 2N HCl for 20min, 2 incubated with rabbit ant-PLZF and rat anti-BrdU antibodies at 4°C overnight, and signals were detected 3 by Alexa488-conjugated anti-rat IgG and Cy5-conjugated anti-rabbit IgG antibodies. For cleaved-PARP 4 detection, slides were incubated with rabbit ant-PLZF and mouse anti-cleaved PARP antibodies at 4°C 5 overnight, and signals were detected by Alexa488-conjugated anti-mouse IgG and Cy5-conjugated 6 anti-rabbit IgG antibodies. For triple staining using rabbit anti-RFP, rabbit anti-PLZF, and rabbit c-MYC 7 antibodies. sections were stained with anti-Tdtomato antibody using TSA plus Cyanine 3 system, and 8 antibodies were stripped off. Then those sections were stained with anti-MYC antibody with TSA plus 9 fluorescein system (Perkin Elmer) followed by antigen retrieval to remove antibodies. Finally, the 10 sections were further stained with anti-PLZF antibody and Cy5 conjugated anti-rabbit IgG. Slides were 11 mounted in Vectashield with DAPI. Images were captured on a fluorescent microscope and processed in 12 Photoshop. Signal intensity of c-MYC signal was quantified with Image J.

13

14 **TRAP** assays (Telomeric Repeat Amplification Protocol).

15 A two-step TRAP procedure was performed as previously reported⁴⁶. Extract fractions from whole testis 16 at 3 weeks or FACS-sorted undifferentiated spermatogonia were incubated with telomeric primers for a 17 30 min initial extension step at 30°C in a PCR machine, followed by 5 min inactivation at 72°C. Without 18 purification, 1µl of the extended reaction was PCR amplified (cycles of 30 seconds at 94°C, followed by 30 seconds at 59°C) in presence of ³²P end-labeled telomeric primers that has been purified using a 19 20 micro-spin G-25 column (GE healthcare). PCR reactions were resolved by 9% polyacrylamide gel 21 electrophoresis at room temperature, and the gel was exposed to a phosphor-imager and scanned by a 22 Typhoon scanner. The scanned image was quantitated using the TotalLab Quant software. 23 Representative gel images were presented among at least 2 repeats.

24

25 FACS analysis

Testes were detunicated, lightly dissociated in PBS, and incubated in PBS containing DNase I 1 2 (Worthington) and 1mg/ml collagenase I (Worthington) at 32°C for 10 min. Cells were centrifuged at 3 250g for 5min and supernatant was removed. After repeating collagenase I treatment, testicular cells 4 were further digested with TrypLE Express (GIBCO) at 32°C for 15 min. During enzymatic digestions, 5 seminiferous tubules were mechanically fragmented with vigorous pipetting every 5min. Cells were 6 sequentially filtered with 70 µm and 40 µm strainers, resuspended in cold FACS buffer (2% FBS, 1mM 7 EDTA in PBS), and incubated with antibodies on ice for 30min. After PBS wash, cells were resuspended 8 in cold FACS buffer containing DAPI, and analyzed and sorted with a BD Aria II (BD Biosciences). Data 9 was analyzed with FlowJo software. The list of antibodies is available in supplemental table.

10

11 **RNA in situ hybridization**

12 5 days post labeling, testes were collected and Tdtomato+ undifferentiated spermatogonia were 13 FACS-sorted, and cytospun at 300 r.p.m. for 5 min onto slides. Slides were fixed in 4% (v/v) PFA for 30 14 min at room temperature and processed for single-molecule RNA FISH using RNAscope 2.5 HD 15 Reagent KIT-RED (Advanced Cell Diagnostics) and probes against mouse *Tert* or *Trp53* (Advanced Cell 16 Diagnostics) according to the manufacturer's instructions.

17

18 Quantitative RT-PCR

For qRT-PCR, cells were directly sorted into Trizol LS (Thermo Fisher Scientific) by FACS and mixed with 100ng yeast tRNA as carrier. RNA was purified Direct-zol RNA Microprep (Zymo Research) and cDNA was synthesized using oligo-dT and SuperScript IV First-Strand Synthesis system (Thermo Fisher Scientific). For qRT-PCR of *Tert* and mouse *Myc*, Universal Probe Library Probe #066 and #72 (Roche) were used, respectively. For these reactions, TaqMan Fast Advanced Master Mix (Thermo Fisher Scientific) were used. For other qRT-PCR, PowerUp SYB Green Master Mix (Thermo Fisher Scientific)

- 1 was used according to the product manual. PCR analysis was done with a 7900HT Fast Real-Time PCR
 2 System machine (ABI). List of primers is available in supplemental table.
- 3

4 **RNA-sequencing**

5 US-h cells were directly sorted into Trizol LS by FACS and RNA was purified using Direct-zol RNA

6 Microprep. Genomic DNA was digested with on-column DNase treatment. RNA quality was checked by

7 Bioanalyzer 2100 (Agilent). RNA-seq libraries were constructed using SMARTer Stranded Total

8 RNA-seq Kit v2 – Pico Input Mammalian (Clontech), starting from 5ng total RNA. cDNA was synthesized

9 and amplified according to the manual. After the rRNA removal step, cDNA was amplified with 13 cycles

10 of PCR reactions. Quality of purified cDNA libraries was confirmed by Bioanalyzer 2100 and

11 concentration of cDNA. Libraries were sequenced on the Illumina NextSeq platform, generating about

12 16-24 million 75-bp paired-end reads per library. Four biological replicates per sample were analyzed.

13 Raw reads were trimmed by TrimGalore 0.4.0 (Babraham Bioinformatics), mapped to mm10 by tophat

14 2.0.13⁴⁷, analyzed by the DEseq2 packages ⁴⁸.

15

16 ATAC-sequencing

ATAC-seq libraries were made as described previously ⁴⁹ using the Omni-ATAC protocol. Adjustments 17 18 to the protocol were made to reflect two primary features of the cell types profiled in this work. First, the 19 amount of Tn5 transposase added to each reaction was modulated to maintain proportionality with the 20 number of cells assayed. For example, a normal reaction uses 50,000 cells and 2.5 ul of Tn5 21 transposase in a 50 ul reaction. In the case of rarer spermatogonial stem cells, only 5,000 cells could be 22 obtained so only 0.25 ul of Tn5 transposase was used in a 50 ul reaction. The difference in volume was 23 adjusted using water. Second, the ploidy of each cell type was taken into account and the amount of Tn5 24 was adjusted based on ploidy as well. For example, round spermatid cells are haploid, so transposition 25 of 50.000 cells would require 1.25 ul of Tn5 transposase in a 50 ul reaction. Similarly, spermatocytes are 26 4N meiotic cells so the amount of Tn5 transposase was increased proportionately and the amount of

water in the reaction was reduced. In all cases, regardless of cell number or ploidy, the reaction volume

1

2 of the transposition reaction was kept constant at 50 ul. All ATAC-seq reactions were performed using 3 homemade Tn5 transposase and Tagment DNA (TD) buffer ⁵⁰. Downstream amplification and 4 purification of libraries was performed as described previously ^{26,49,51}. 5 ATAC-seq data pre-processing was completed using the PEPATAC pipeline 6 (http://code.databio.org/PEPATAC/). The mm10 genome build (https://github.com/databio/refgenie) was 7 used for alignment. Briefly, all fastg files were first trimmed to remove Illumina Nextera adapter sequence using Skewer⁵² with "-f sanger -t 20 -m pe -x" options. FastQC 8 9 (http://www.bioinformatics.bbsrc.ac.uk/projects/fastqc) was used to validate proper trimming and check overall sequence data quality. Bowtie2⁵³ was then used for pre-alignments to remove reads that would 10 11 map to chrM (revised Cambridge Reference Sequence), alpha satellite repeats, Alu repeats, ribosomal DNA repeats, and other repeat regions with "-k 1 -D 20 -R 3 -N 1 -L 20 -i S,1,0.50 -X 2000 -rg-id" options. 12 13 Bowtie2 was then used to align to the mm10 reference genome using "--very-sensitive -X 2000 --rg-id" 14 options. Samtools ⁵⁴ was used to sort and isolate uniquely mapped reads using "-f 2 -q 10 -b -@ 20" 15 options. Picard (http://broadinstitute.github.io/picard/) was used to remove duplicates. Then the bam files were merged by conditions, and MAC2⁵⁵ was used to call peaks with parameter "-q 0.05 --nomodel 16 17 --shift 0". The narrow peaks were then filtered by the ENCODE 7 hg19 blacklist, as well as peaks that 18 extend beyond the ends of chromosomes. Bedtools ⁵⁶ was used to retrieve the reads of the called peaks 19 for each sample with multicov module. All the samples have similar sequencing depth, mitochondrial rate, 20 and duplication rate. The spermatocyte and the round spermatid samples have similar sequencing depth 21 compared to all other samples, but slightly higher mitochondrial rate and lower duplication rate, so have 22 more final reads after initial processing and filtering. To make all the samples comparable for the statistical analysis, we used final reads as the normalization factor. R package "DESeg2"⁴⁸ was used for 23 24 statistical analysis to identify significant peaks between different conditions. The differential peaks were 25 called between US-h CreER/+ and US-h CreER/flox samples. Peaks with FDR < 0.01 and fold change

1	larger than 2 or smaller than -2 were considered as significant. R packages "ChIPseeker" ⁵⁷ was used for
2	peak annotation. Package "ngsplot" ⁵⁸ was used for visualization of cumulated peak signal.

- 3
- 4 **Statistics:** No statistical methods were used to predetermine sample sizes. When comparing two
- 5 groups, *p* values were determined by two-sided unpaired *t*-test. When comparing more than two groups,
- 6 *p* values were determined by one-way ANOVA with Tukey's test. Values are presented as mean ± SEM.
- 7 The animals were randomly assigned to each experimental or control group. Graphs were generated by
- 8 the Prism 8.
- 9
- 10 **Data availability:** The source data for the RNA-seq study are available in the NCBI Gene Expression
- 11 Omnibus (GEO) repository under accession number GSE14659. All other data that support the finding
- 12 of this study are available from the corresponding author upon reasonable request.

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1 Figure legends

Fig. 1. TERT deletion impairs SSC-mediated clone formation. (a) Tert^{CreER/+}Rosa^{lsI-Tdtomato/+} mice. (b) 2 3 Lineage tracing using *Tert-CreER*. (c) Epifluorescence of Tdtomato and bright field image of untangled 4 seminiferous tubules in a single whole testis at 3 months after 1mg tamoxifen injection. Scale bars, 5mm. 5 (d-f) Mean patch number (d), mean patch length (e), and total patch length (f) after pulse labeling with 6 indicated tamoxifen dose and span (n=6, 6, 6, 4, 6, 5 mice from left to right). (g) Tert^{CreER/flox}:Rosa^{/sl-Tdtomato/+} mice. (h) Whole mount immunofluorescence of GFRA1 and Tdtomato. Scale 7 bars, 100 μ m. (i) Quantification of Tdtomato⁺ cells within GFRA1⁺ A_s and A_{pr} cells (n=4 mice per group). 8 9 (i) In situ hybridization against *Tert* mRNA in purified Tdtomato⁺ US at 5 days post-tamoxifen treatment. Undifferentiated spermatogonia from *Tert*^{CreER/CreER} mice was used as negative control. Scale bars, 10 11 40μm. (k) Quantification of foci of Tert mRNA (n=232, 243, 341 cells from left to right). (I) 12 Epifluorescence of Tdtomato in untangled seminiferous tubules at 3 months and 6 months 13 post-tamoxifen injection. Scale bars, 5mm. (m-o) Quantification of mean patch number (m), mean patch 14 length (n), and total patch length (o) (n=7, 7, 9, 8 mice from left to right). Data are represented as Mean \pm 15 SEM. 16 Fig. 2. TERT promotes SSC competition independent of its catalytic activity and the telomerase 17 complex. (a) Tert^{CreER/flox}:Rosa^{lsI-Tdtomato/lsI-tTA:} TetO-Tert or -Tertci mice. (b) Epifluorescence for Tdtomato 18

19 in untangled seminiferous tubules at 3 months after tamoxifen injection. Scale bars, 5mm. (c-e)

20 Quantification of mean patch number (c), mean patch length (d), and total patch length (e) (n=6 mice per

group). (f) Quantification of TdTomato⁺, GFRA1⁺ A_s and A_{pr} cells at 7 or 14 days post labeling (n=5, 5, 5, $\frac{1}{2}$)

5, 6, 6, 5, 5 mice from left to right). (g) Mean patch number at 6 weeks after tamoxifen injection (n=5

23 mice per group). (h) Quantification of percent Tdtomato⁺ clones containing PLZF⁺ cells (n=5,4 mice from

left to right). (i) Quantification of BrdU⁺ cells in Tdtomato⁺, PLZF⁺ undifferentiated spermatogonia (n=6, 5,

25 5, 6 mice from left to right). Data are represented as Mean ± SEM.

1 Fig. 3. TERT deletion compromises chromatin accessibility in SSCs. (a) Peak calls from ATAC-seq 2 data. Peak calls from each cell type are shown individually. Color indicates the type of genomic region 3 overlapped by the peak. (b) Average tag density of ATAC-seg reads around transcription start sites. (c) PCA of ATAC-seq data from US-h, US-m and DS cells purified from *Tert*^{CreER/+} versus *Tert*^{CreER/flox} mice. 4 5 (d) Heatmap representation of 11656 peaks that are significantly different between Tert^{CreER/+} and 6 Tert^{CreER/flox} in US-h, US-m, and DS or significantly more open either in SP or RS. Each row represents 7 one ATAC-seq peak. Color represents the relative ATAC-seq accessibility. (e) Venn diagrams of the peaks in US-h, US-m and DS cells from *Tert*^{CreER/+} and *Tert*^{CreER/flox} mice. 8

9

10 Fig. 4. MYC rescues competitive clone formation of TERT-deleted SSCs. (a) PCA of RNA-seq data from Tdtomato⁺ US-h. (b) Significant down-regulation of MYC target genes in *Tert*^{CreER/flox} US-h cells. 11 12 RNA-seg data was analyzed using gene set enrichment analysis. (c) Quantification of mean signal 13 intensity for MYC staining in PLZF⁺ undifferentiated spermatogonia using testicular cross sections at 7 days after labeling. (d) Tert^{CreER/flox}:Rosa^{IsI-Tdtomato/IsI-tTA}: TetO-hMYC mice. (e) Epifluorescence for 14 15 Tdtomato in untangled seminiferous tubules. Scale bars, 5mm. (f to h) Quantification of mean patch 16 number (f), mean patch length (g), and total patch length (h) (n=6, 8, 6 mice from left to right). (i) 17 Quantification of Tdtomato⁺ GFRA1⁺ A_s and A_{pr} clones at 14 days post labeling detected by whole mount 18 immunofluorescence (n=5 mice per group). (i) Quantitative analysis of Tdtomato⁺ BrdU⁺ PLZF⁺ 19 undifferentiated spermatogonia using testicular cross sections at 7 days post labeling (n=5 mice per 20 gropu). Data are represented as Mean \pm SEM.

21

22 Extended Data Figure 1. TERT expression in spermatogonia subpopulations. (a)

23 Spermatogenesis: seminiferous tubules are composed of four layers of male germ cells.

24 Undifferentiated and differentiating spermatogonia located in the basement layer undergo mitosis. They

translocate to the second layer when entering into meiosis and differentiate into spermatocytes. Haploid

26 spermatids produced by meiosis move toward luminal side during post-meiotic maturation. (b) A

1 functional, morphological, and gene expression heterogeneity in spermatogonia subpopulations. (c) 2 Flowcytometry analysis of wild-type testicular cells stained with α 6-Integrin, MCAM, and KIT. 3 α 6-Integrin-high cells were further separated into US-h, US-m, and DS based on KIT and MCAM 4 expression. (d) qRT-PCR of *Tert* and spermatogonia markers in FACS-sorted US-h, US-m, and DS. 5 Expression was normalized with Actb (n=3 mice per group). Data are represented as Mean \pm SEM. (e) Flowcvtometry analysis of testicular cells from $Tert^{Tdtomato/+}$ mouse. α 6-Integrin-high cells were gated and 6 7 further separated according to KIT and MCAM expression. Tert-Tdtomato expression was compared in 8 US-h, US-m, and DS.

9

10 Extended Data Figure 2. Marking and linage tracing of TERT⁺ cells. (a) Immunofluorescence of 11 Tdtomato and E-cadherin, a membrane marker of undifferentiated spermatogonia, in whole-mount 12 seminiferous tubules at day two. Scale bars, 100µm. (b) Quantification of Tdtomato⁺ cells in E-cadherin⁺ 13 undifferentiated spermatogonia (n=4 mice per group). Data are represented as Mean ± SEM. (c) Whole-mount immunofluorescence of KIT and Tdtomato in a *Tert^{CreER/+}:Rosa^{IsI-Tdtomato/+}* testis at two days 14 15 after high dose tamoxifen injection. Scale bars, 100µm. (d) Epifluorescence of Tdtomato in untangled 16 seminiferous tubules at 3-month (3m) or 1-year (1yr) after pulse labeling; injected tamoxifen dose shown 17 (0.25 - 4mg). Scale bars, 5mm. (e) Singly isolated Tdtomato⁺ patch. Scale bar, 1mm. (f) Cross sections 18 of testes immunostained with antibodies to Tdtomato at 3-month or 1-year after labeling. Scale bar, 19 50µm.

20

Extended Data Figure 3. Conditional deletion of TERT in SSCs using *Tert-flox* mice. (a) *Tert-flox*targeting strategy and Southern blot strategy. (b) Southern blot analysis of genomic DNA from mouse
tails. Locations of 5' and 3' probes are shown in A. (c) TRAP assay using mouse embryonic fibroblasts
from *Tert-flox* mice. Cells were transfected with adenovirus to express LacZ or Cre.

25

1 **Extended Data Figure 4. (a)** gRT-PCR analysis of *Tert* and *Tertci* in purified Tdtomato⁺ undifferentiated 2 spermatogonia at day 7. Effects of doxycycline addition to drinking water on Tert mRNA level. 3 Expression was normalized with Actb (n=3 mice per group). (b) Telomere repeat amplification protocol 4 (TRAP) assay using purified Tdtomato⁺ undifferentiated spermatogonia from indicated genotypes. (c) Tert^{CreER/flox}:Rosa^{IsI-Tdtomato/+}:Terc^{-/-} mice. (d) TRAP assay using whole testes from Tert^{CreER/+}:Terc^{+/+} and 5 6 Tert^{CreER/+}: Terc^{-/-} mouse. (e) Epifluorescence of Tdtomato in untangled seminiferous tubules. Scale bars, 7 5mm. (f-h) Quantification of mean patch number (d), mean patch length (e), and total patch length (f) 8 (n=7, 6, 8, 7). Data are represented as Mean \pm SEM. 9 10 Extended Data Figure 5. Persistent clone loss phenotype of TERT-deleted spermatogonial stem cells with conditional P53 deletion. (a) Tert^{CreER/flox}:Rosa^{lsI-Ttomato/+}:Trp53^{flox/flox} mice. (b) In situ 11 12 hybridization against *Trp53* mRNA. Tdtomato⁺ undifferentiated spermatogonia were sorted out at 5 days 13 post labeling and stained. (c) Quantification of foci number of Trp53 mRNA (n=174 and 135 cells from 14 left to right). (d) Epifluorescence of Tdtomato in untangled seminiferous tubules. Scale bars, 5mm. (e-g)

15 Quantitative data of patch number (e), average patch length (f), and total patch length (g) (n=6, 6, 7,7 16 mice from left to right). (h) Immunofluorescence of γ H2AX, PLZF, and Tdtomato in testicular cross 17 sections. *Tert^{Tdtomato/Tdtomato*} (G6) mice were used as positive control for γ H2AX staining. Scale bars, 30µm. 18 (i) Quantification of γ H2AX-positive unidiferentiated spermatogonia (n=4 mice per group). Data are 19 represented as Mean ± SEM.

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21 Extended Data Figure 6. Apoptosis in TERT-deleted undifferentiated spermatogonia. (a)

Immunofluorescence for GFRA1 and TdTomato at 14 days post labeling with tamoxifen. Scale bars,
100µm. (b) Epifluorescence for TdTomato in untangled seminiferous tubules at 6 weeks post labeling.
Scale bars, 5mm. (c) Immunofluorescence for Tdtomato and PLZF, a marker of undifferentiated
spermatogonia, using testis cross-sections at 6 weeks after tamoxifen injection. Scale bars, 50µm. (d)
Immunofluorescence for Tdtomato, PLZF, and BrdU using testis cross sections. Scale bars, 50µm. (e)

1	Immunofluorescence of PLZF, Tdtomato, and apoptosis marker, cleaved-PARP (cPARP), on testicular
2	cross-sections at 7 days after labeling. Tert ^{Tdtomato/Tdtomato} (G6) mice were used as positive control for
3	cleaved-PARP staining. Scale bars, 50µm. (f) Quantification of Tdtomato ⁺ cleaved-PARP ⁺
4	undifferentiated spermatogonia (n=4 mice per group). Data are represented as Mean \pm SEM.
5	
6	Extended Data Figure 7. Global changes of open chromatin structure in undifferentiated
7	spermatogonia after acute deletion of TERT. (a) Principal component analysis (PCA) of ATAC-seq
8	data. Colors represent indicated cell types from Tert-heterozygous mice. Arrow indicates direction of
9	differentiation. (b) Pearson correlation heatmaps of ATAC-seq samples. (c) Heatmap representation of
10	7597 peaks showing significant differences across US-h, US-m, DS, SP, and RS. Each row represents
11	one ATAC-seq peak. Color represents relative ATAC-seq accessibility. (d and e) Venn diagrams
12	showing the number of peaks found in spermatogonia subpopulations. (f) KEGG pathway analysis of
13	differential peaks between Tert ^{CreER/+} and Tert ^{CreER/flox} US-h.
14	
15	Extended Data Figure 8. Changes in open chromatin structure at specific loci after acute deletion
16	of TERT. (a-d) Read distribution of ATAC-seq data across different cell types. (a) Tert locus. (b) Ret and
17	Gfra1 loci. These genes are dominantly expressed in US-h. (c) Cdh1 and Zbtb16 loci. These genes are
18	dominantly expressed in both US-h and US-m. (d) Kit and Prm1/Prm2 loci. Kit is dominantly expressed
19	in DS and Prm1/Prm2 are dominantly expressed in RS.
20	
21	Extended Data Figure 9. Down-regulation of multiple signaling pathways in TERT-deleted US-h.
22	(a) Volcano plot of RNA-seq data of purified Tdtomato ⁺ US-h from <i>Tert^{CreER/+}</i> and <i>Tert^{CreER/flox}</i> mice.
23	Genes showing the significant changes (more than 2-fold change and P<0.0001) were colored with red.
24	(b and c) gene set enrichment analysis using RNA-seq data of US-h purified from Tert ^{CreER/+} and
25	Tert ^{CreER/flox} mice. nES: normalized enrichment score. (d) Immunofluorescence for MYC, PLZF, and
26	Tdtomato at day 7 after tamoxifen labeling. Scale bars, 50µm. (e) Quantification of mean signal

1 intensities for PLZF staining in undifferentiated spermatogonia (n=68, 71, 66, 78 cells from left to right). 2 (f) qRT-PCR analysis of Myc mRNA in purified Tdtomato⁺ US at day 7. Doxycycline was added to 3 drinking water. Expression was normalized with Actb (n=3 mice per group). (g) gRT-PCR analysis of 4 transgenic human MYC mRNA in purified Tdtomato⁺ US at day 7 after tamoxifen labeling. Doxycycline 5 was added to drinking water. Expression was normalized with Actb (n=3 mice per group). n.d.; not 6 detected. (h) gRT-PCR analysis of MYC pathway genes that were significantly down-regulated in gene 7 set enrichment analysis in figure 4b. Tdtomato⁺ US-h cells were purified at day 7 post labeling by FACS 8 and cDNA was synthesized. Expression level was normalized with Actb (n=6,6,5,5,5 mice from left to 9 right).

10

11 Extended Data Figure 10. Model for SSC competition driven by non-canonical functions of TERT.

12 Summary schematic of catalytic activity-independent functions of TERT in stem cell competition.

13 TERT-deleted SSCs are progressively eliminated from SSC pool through cell competition, reducing the

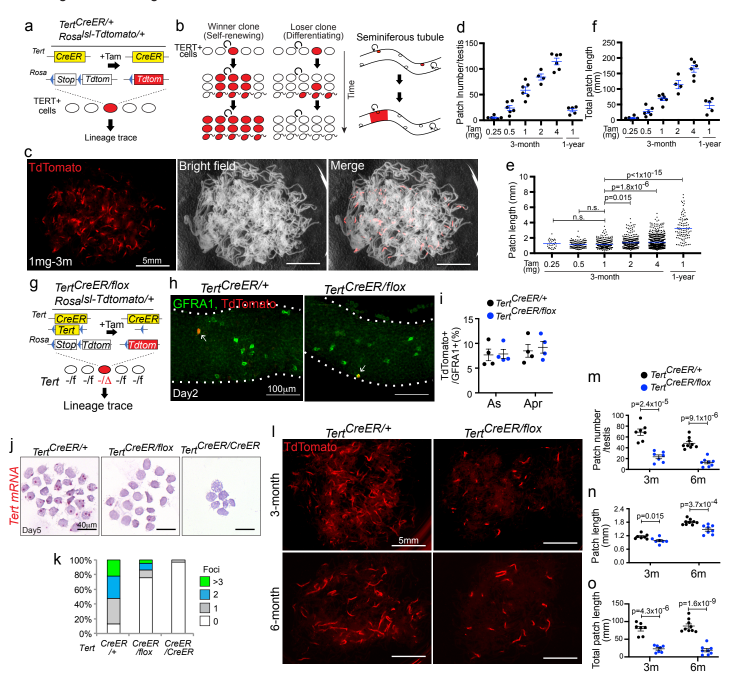
14 contribution of TERT-deleted SSCs to spermatogenesis over time (left). In wild-type SSCs, TERT

15 promotes competitive clone formation by up-regulating MYC protein. In SSCs lacking TERT,

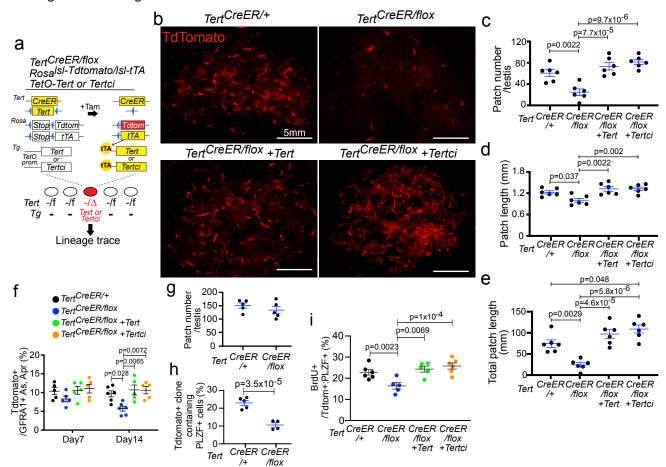
downregulation of MYC protein promotes rapid differentiation, impaired proliferation, and global loss of
 chromatin accessibility. MYC expression bypasses the requirement for non-canonical TERT, restoring

18 SSC-mediated clone formation.

Hasegawa et. al. Figure 1



Hasegawa et al. Figure 2



Hasegawa et al. Figure 3

