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Eroded telomeres are rearranged in quiescent fission yeast cells through duplications of subtelomeric sequences

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Abstract

While the mechanisms of telomere maintenance have been investigated in dividing cells, little is known about the stability of telomeres in quiescent cells and how dysfunctional telomeres are processed in non-proliferating cells. Here, we examine the stability of telomeres in quiescent cells using fission yeast. While wild-type telomeres are stable in quiescence, we observe that eroded telomeres were highly rearranged during quiescence in telomerase minus cells. These rearrangements depend on homologous recombination (HR) and correspond to duplications of subtelomeric regions. HR is initiated at newly identified subtelomeric homologous repeated sequences (HRS). We further show that TERRA (Telomeric Repeat-containing RNA) is increased in post-mitotic cells with short telomeres and correlates with telomere rearrangements. Finally, we demonstrate that rearranged telomeres prevent cells to exit properly from quiescence. Taken together, we describe in fission yeast a mode of telomere repair mechanism specific to post-mitotic cells that is likely promoted by transcription.

Introduction
Telomeres are nucleoprotein structures that protect chromosomes ends from degradation and ensure replication of terminal DNA. They are composed of long stretches of double-stranded TTAGGG repeats that end with a 3’ single-stranded overhang. Because conventional replication of linear ends by DNA polymerase leads to loss of telomeric sequences, telomeres shorten progressively at each cell cycle division and this shortening triggers replicative senescence. This phenomenon participates to cellular aging by limiting the proliferative capacity of most cells in an organism. By contrast embryonic stem cells and germinal cells have an unlimited capacity to divide to ensure tissue renewal, regeneration and repair. To counteract telomere attrition, these stem cells use a reverse transcriptase named telomerase that extends the 3’ end of chromosomes ends thanks to its RNA associated template. However, with the exception of embryonic stem cells and germinal cells, in the majority of stem cells the telomerase activity is low or absent. Thus telomere shortening occurs during replicative aging in stem cells, possibly at a slower rate than that in normal somatic cells and may alter stem cells function. Indeed, the capacity of stem cells to enter and exit quiescence is imperative to tissue homeostasis and to the response to life-threatening challenges.

Quiescence is a common life form for the cell. Indeed the majority of the cells in adult human body tissues and organs are non-dividing postmitotic cells. Although telomeres attrition is correlated with cell division, telomere shortening has been also observed in somatic cells of brain regions or skeletal muscle, regardless of their replicative activity. This suggests that other factors than chronological cell division may cause telomeres shortening. Because telomeres are G-rich, they might be extremely sensitive to oxidative stress. Indeed, guanines can be modified to 8-oxyguanosine by reactive oxygen species (ROS). Thus, base alteration by oxidative stress or other DNA damage may alter the binding of telomeric protein, trigger DNA repair and accelerate the telomere shortening. Whether it concerns postmitotic cells or quiescent stem cells, the observations above raise the question of how telomeres are maintained in quiescence and how the replicative senescence will impact on cell ability to enter and exit quiescence.

*Schizosaccharomyces pombe* is a key model organism since it has a high level of conservation of telomeric proteins with mammalian cells and cells can be easily maintained in quiescence state by nitrogen starvation. Moreover, genetic is
facilitated in *S. pombe* since it has only three large chromosomes (Chr I, II and III with the smallest one (Chr III) containing rDNA). Telomeric repeats consist in 300bp of the degenerated sequences G\textsubscript{2-6}TTAC[A] while subtelomeres contain a mosaic of multiple segments that span \textasciitilde50kb of the telomere proximal-site that define sub-telomeric-elements 1 (STE1), 2 (STE2) and 3 (STE3) \textsuperscript{15}. Subtelomeres are heterochromatinized regions in which the methylation of the lysine 9 of the histone 3 (H3K9me) serves as a binding site for the heterochromatin protein 1 (HP1\textsuperscript{Swi6}). In spite of their silenced heterochromatin status, telomeres are transcribed in long non-coding RNA named TERRA \textsuperscript{16}.Transcription of TERRA starts within the adjacent subtelomeric sequences and contains a variable number of telomeric G-rich sequences. In addition to TERRA, fission yeast chromosome ends produce a variety of telomeric transcripts \textsuperscript{16,17}. Although telomeric transcription functions are extensively investigated, the role of TERRA in the biology of telomeres is still elusive.

As in mammals, telomeres from yeast cells with reduced telomerase activity progressively shorten with each cell division. At late time-point of senescence, most of the cells die or remain arrested. This arrest (or crisis) is caused by a DNA damage checkpoint that is activated directly as a result of unprotected short telomeres being recognized as irreparable double-strand breaks (DSBs). In fission yeast, cells eventually recover from growth and establish survivors after 100 generations \textsuperscript{18}. Most cells survive by circularizing all chromosomes (type I survivors) to bypass the need of telomerase. Telomere fusions are dependent on Rad52, the Rad16\textsubscript{XPF}-Swi16\textsubscript{ERCC1} endonuclease, the single-stranded DNA-binding protein RPA, and the Srs2 and Werner/Bloom Rqh1 helicases \textsuperscript{19}. Cells can also survive by maintaining their telomeric repeated sequences, presumably through homologous recombination (HR), type II survivors related to ALT mechanism \textsuperscript{20}. Finally, cell can survive continually amplifying and rearranging heterochromatic sequences (type III) \textsuperscript{21}. These survivors are called HAATI (for heterochromatin amplification-mediated and telomerase-independent) and require the heterochromatin assembly machinery, the histone methyl transferase Clr4. Most frequently the rDNA spreads from the subtelomeric regions of Chr III to the termini of all three chromosomes. In very rare cases, subtelomeric sequences (STE) spread from the subtelomeric regions of Chr I and II to multiple sites dispersed in the genome.

Although telomeres maintenance have been extensively studied in cycling cell, rare studied have been undertaken in quiescent cells \textsuperscript{22}. It is established now that
DNA damage occurs in quiescent cells and that distinct modes of DNA repair are used by quiescent and vegetative cells. For example, oxidative DNA damage produced by respiration is thought to be a source of DNA damage. To tackle this issue, we examine the stability of telomeres in quiescent cells population containing variable number of telomeric DNA repeats using fission yeast as a model. To this purpose, we investigate the stability of telomeres in G0 cells in the presence and absence of telomerase. We show that eroded telomeres are rearranged during quiescence. This reorganization corresponds to duplications of subtelomeric sequences by homologous recombination, named STEEx which are promoted by transcription at telomeres. We also demonstrate that STEEx formation that occurs specifically during quiescence prevents proper re-entry into proliferative phase. Our results highlight how non-dividing cells that harbor eroded telomeres may circumvent the lack of a functional telomere protection in the absence of replication.

Results

Eroded telomeres are rearranged in quiescence

Because DNA damage accumulates in G0, we wondered whether telomeric repeats might represent a specific target for DNA damage and how are they maintained during quiescence. Genomic DNA of cells maintained in G0 by nitrogen starvation from 1 to 8 days were digested by EcoRI and subjected to Southern blot analysis (Fig. 1a). As shown in figure 1a, telomeric repeats from WT cells were stable during quiescence. To further study the fate of dysfunctional telomeres in G0, we deleted the RNA of the telomerase TER1 in dividing cells to trigger progressive telomere shortening and replicative senescence. We first performed daily dilution of ter1Δ spore colonies in cycling cells and monitor their growth and telomere shortening (Fig. 1b). In this representative experiment, crisis was reached after 90-100 population doublings (pds) concomitantly with the progressive disappearance of telomeric signal. After 1, 3, 5 and 7 days of senescence, cells with different setting of telomere length (corresponding to 41, 55, 68 and 81 pds) were starved from nitrogen and maintained in quiescence for a total of 8 days (schematized in Fig. 1c). Cell counting and microscopic observations confirmed the absence of cell division during quiescence. After 1, 4, 6 and 8 days in quiescence, cells were collected and genomic DNA was extracted, digested by EcoRI, and subjected to Southern blot analysis (Fig. 1d). The same membrane was first hybridized with a probe specific to telomeric
sequences (Telo), then with a subtelomeric probe (STE1) and finally with a chromosomal probe used as a loading control. At day 1 of the senescence (S1, 41 pds) telomere size was reduced by about 120bp compared to WT and telomeres were stable in G0 (Fig. 1d, Telo probe). However, by hybridizing the same Southern blot with the subtelomeric probe, we observed the appearance of three additional bands (900, 1400 and 1500 bp) specifically in quiescence (Fig. 1d, STE1 probe) while the control chromosomal signal was constant. The subtelomeric STE1 signal was barely visible at day 1 but intensified after 4, 6 and 8 days during quiescence (Fig. 1d). When initial telomere size decreased after 3, 5 and 7 days of senescence, the appearance of STE1 signal was exacerbated. Amplification of the STE1 signal during quiescence was also confirmed by spot assays (Fig. 1e). Supplementary Figure 1a shows another example of a ter1Δ clone for which replicative senescence and quiescence were monitored except that the Southern blot was revealed with a probe that hybridizes to both telomeric repeats and subtelomeric regions (TeloSTE1).

Finally, we wondered whether an inactive form of the telomerase could provoke the formation of these rearrangements. We took advantage of a strain expressing the catalytically dead telomerase. Subtelomeric signal amplification was observed in G0 in this strain, suggesting that it is the absence of telomerase activity per se induces rearrangement at eroded telomeres in quiescence (Supplementary Figure 1b).

Collectively these results indicate that eroded telomeres are subjected to subtelomeric rearrangements in quiescence. These rearrangements increase with telomere shortening and time in quiescence. Important, these rearrangements occur in the total absence of cell division, without selection pressure, and at high frequency. Thus, these events are not related to survivors that occur in replicative senescence during crisis at low frequency ($10^{-4}$-$10^{-5}$), but rather represent a new type of rearrangements that takes place globally at eroded telomeres of the quiescent cells population.

**Subtelomeric rearrangements correspond to STE1-extended amplification**

In the following experiments, ter1Δ cells were placed in G0 after approximately 60 pds, at a time-point of senescence where rearrangements were observed. To gain insight into the mechanism by which telomere rearrangements were produced during
quiescence, telomere structure from telomerase positive (8 days in G0) and \( \text{ter1}\Delta \) cells (1 and 8 days in G0) were analyzed by Southern blot analysis with Telo, STE1 and STE2 probes. Schematic illustration of fission yeast subtelomeric regions cloned into plasmids (pNSU70 and pNSU64) in which probes and restriction enzyme sites are annotated, is shown in figure 2a. Telomeres from telomerase positive cells, maintained for 8 days in G0 (indicated by + in the Southern blot of Fig. 2a), were detected as bands of approximately 0.3, 1 and 1.5 kb with Apal, EcoRI and Swal digestion, respectively. With \( NsI \) cut, four bands were detected that corresponds to sequence heterogeneity in subtelomeric regions as previously described \(^{26,27}\). In \( \text{ter1}\Delta \) cells, telomeres were substantially shortened but still detectable after 1 and 8 days in quiescence (indicated by 1 and 8 in Fig. 2a). Digestions with EcoRI and Swal revealed by the STE1 probe confirmed the appearance of quiescence specific rearrangements while \( NsI \) cut unveiled a smear at high molecular weight, which intensified after 8 days in G0. In contrast to EcoRI and Swal digestions, the Apal digestion showed a mixed pattern made of a smear signal and discrete subtelomeric bands. No change in profiles was observed with the STE2 probe showing that the upstream subtelomeric STE2 region was not affected by these rearrangements. Indeed, distinct bands at 5, 4 and 2.5 kb were revealed by the STE2 probe with EcoRI, Swal and \( NsI \) digestions, respectively (Fig 2a, lower panel).

Taken together, these results indicate first, that subtelomeric rearrangements occur within the terminal fragments downstream the \( NsI \) site and second, that the amplified STE1 region contains several repetitions of DNA fragments that contain the EcoRI and Swal restriction sites. The subtelomeric repetitions that are amplified may or not contain the Apal site. The resulting structure of the subtelomeric amplification is schematized in figure 2b, which shows a representation of Apal-Apal subtelomeric block duplication.

**STE1-extended rearrangements are initiated at homologous repeated sequences**

We further characterized the STE1-extended repetitions by identifying two 226 bp regions of strict homology (Homologous Repeated Sequence; HRS) flanking the subtelomeric region. One is located in the vicinity of the \( NsI \) site while the second is adjacent to the telomeric repeats (Fig. 2b, 3a and Supplementary Figure 2). Hence, we designed several sets of oligonucleotides indicated in figure 3a to map the
amplified region by qPCR in the STE1-extended region. The qPCR analysis showed that G0 subtelomeric rearrangements corresponded to the expansion of a STE1-extended DNA delineated by the Apal-HRS sequences (named STEEx) (Fig. 3b). The results suggested that the recombination events that we observed during quiescence in cells with dysfunctional telomeres are initiated at the HRS sites in a replication-independent manner.

Furthermore, we wondered if STEEx occurs internally only in cis or also in trans with another chromosome end. To investigate this point, we deleted ter1 in a strain that contains a transcriptionally inducible telomere (called tiTel) 17. In this strain, a nat':Pnmt1 cassette is inserted in the subtelomeric region of the left arm of chromosome II (Fig. 3c) 17. Importantly, this modified telomere can be specifically detected using a nmt1 probe. ter1Δ tiTel cells were grown in minimal medium before nitrogen starvation. Then, STEEx formation was monitored at the tiTel telomere by Southern blot with the nmt1 probe (Fig. 3d). The endogenous genomic nmt1+ sequence was visualized as a band migrating at 2.5 Kb, while the nat':Pnmt1 tiTel cassette was detected at 4.2 Kb. The tiTel signal intensified upon nitrogen starvation. As for STEEx, the amplification of tiTel signal corresponds to the duplication of HindIII-blocks that contain the nat':Pnmt1 cassette (tiSTEEEx) (Fig. 3d). At day 8 of G0, a smear was clearly detected with the nmt1 probe, reflecting rearrangements between tiTel and natural telomeres (ti/natural-STEEEx) (Fig. 3d-e). These data show that duplication of subtelomeric sequences primarily occurs internally in cis and then propagates to other chromosome extremities.

**Rad51 and Ctp1 are involved in STEEx formation**

The results above indicate that subtelomeric rearrangements were initiated at HRS sites through HR events. To strengthen this assumption we checked the impact of the deletion of genes involved in HR and DNA-end resection on STEEx formation (rad51, mre11, ctp1 and exo1). First, we deleted ter1 in a rad51Δ background. Replicative senescence was strongly impacted by the absence of Rad51 showing the importance of HR pathway in absence of telomerase 28. Indeed, crisis occurred shortly after the deletion of ter1. Nevertheless, we could starved from nitrogen these cells after 30 pds (telomere repeats were reduced by approximately 120-130 bp) and keep them for 8 days in quiescence. FACS analysis showed that cell death increased
significantly in the rad51Δ strain in the presence or absence of functional telomerase (our unpublished results). Telomere structure from rad51Δ and rad51Δ ter1Δ cells was analyzed by Southern blot (Fig. 4a). In rad51Δ cells telomeres displayed a WT length and remained stable in G0. The signal reduction observed after 8 days in quiescence was caused by cell death. In ter1Δ rad51Δ, telomeres were shortened but not rearranged indicating that STEEx events did not occur. NsiI digestion profile of the rad51Δ strain displayed a comparable profile to WT ruling out the possibility that the absence of STEEx was due to differences in the outline of the subtelomeric regions (Supplementary Figure 3). Next, we deleted ter1 in mre11Δ, ctp1Δ and exo1Δ backgrounds (Fig. 4b-d). Senescence was accelerated in mre11Δ and ctp1Δ mutants while it was delayed in exo1Δ strain (our unpublished data). STEEx formation was observed in exo1Δ and mre11Δ mutants. Noteworthy, in ter1Δ mre11Δ the lowered STEEx level was likely caused by a significant death of ter1Δ mre11Δ cells in G0 (our unpublished data) . In contrast to the situation in the mre11Δ mutant, STEEx did not form in the ctp1Δ mutant (Fig. 4c) suggesting that Ctp1 is involved in STEEx formation. From these data, we concluded that Rad51-dependant HR and Ctp1 are required to STEEx formation while Exo1 and Mre11 appear dispensable. Unexpectedly Ctp1 was shown to function in STEEx formation independently of the MRN complex.

STEEx are different from HAATI

Although very rare, the spreading and amplification of subtelomeric region was observed in absence of telomerase in the HAATI\textsuperscript{STE} survivors.\textsuperscript{21} Hence, we wondered whether STEEx are comparable to HAATI\textsuperscript{STE}. We investigated chromosome structure of ter1Δ quiescent cells by pulsed-field gel electrophoresis (PFGE) (Fig. 5a). While chromosomes of HAATI survivors fail to enter into the gel, the chromosomes of STEEx were visible after ethidium bromide staining although the signal became fuzzy after 8 days in G0. Because HAATI requires heterochromatin formation, we next examined the importance of the H3K9 methyl-transferase Clr4, Swi6 (HP1) and Mit1 (ATP-dependent DNA helicase of the SHREC complex) in STEEx formation. Senescence was accelerated in swi6Δ and clr4Δ mutants compared to WT while it was unchanged in mit1Δ strain (our unpublished data). However, the absence of Clr4, Swi6 or Mit1 did not prevent STEEx formation (Fig.
Noteworthy EcoRI digestion pattern was modified in swi6Δ and mit1Δ mutants (Fig. 5b and 5d). Accordingly, it was previously established that deletion of clr4 or swi6 increased recombination among subtelomeric regions. Indeed, NsiI digestion profiles in the swi6Δ, clr4Δ and mit1Δ strains were different compared to WT (Supplementary Fig. 3). This heterogeneity in subtelomeric regions likely explains the different STEEx patterns in G0. From these results, we concluded that STEEx are clearly different from HAATiSTE.

**TERRA transcription promotes STEEx formation**

In the absence of DNA replication, we wondered if transcription could initiate recombination in quiescence. Recently, it has been shown in fission yeast that among the various telomeric transcripts, TERRA is enhanced when telomere size shortens. Moreover, the observation that the HRS contains the transcription start site of TERRA (TSS) further links TERRA to STEEx. Thus, we assessed TERRA levels in quiescent cells. As an internal control we used Fcp1 mRNA that is transcribed in both cycling and quiescent cells. RNA was extracted from WT and ter1Δ quiescent cells and reverse transcribed using a random hexanucleotide primer. cDNA was then subjected to real-time quantitative PCR (qPCR) with primers spanning a subtelomeric fragment downstream of the TERRA TSS (Fig. 6a). Note that the TERRA primers also amplify the ARRET and αARRET transcripts although they are poorly represented during senescence. Figure 6b shows RNA levels of one representative experiment while figure 6c displays the ratio of global TERRA signal over Fcp1 signal obtained from three independent experiments. As previously reported, we confirmed that TERRA was up-regulated in cycling cells when telomeres shorten. Interestingly, we observed that quiescent WT cells express TERRA at higher level than cycling cells either in the presence or in the absence of telomerase. Most importantly, we detected a significant increase of TERRA level during quiescence in ter1Δ cells, which correlated with STEEx formation.

To determine if qPCR signal amplification reflected extensive accumulation of TERRA molecules or transcription of long subtelomeric RNA (long ARRET/αARRET) we ascertained the RNA species in WT and ter1Δ quiescent cells by Northern blot analysis. The rap1Δ mutant was used as a control because it is known to facilitate the detection of TERRA which is otherwise barely detectable in WT cells.
The Northern Blot was hybridized with a Telo probe and a STE1 probe allowing the detection of TERRA / ARIA species and global subtelomeric transcription (except ARIA), respectively (Fig. 6d). Figure 6e shows that in WT cells TERRA / ARIA accumulates in G0 (upper panel). In ter1Δ cells, TERRA / ARIA accumulation was comparable to WT. Hybridization with the STE1 probe revealed TERRA / ARRET / αARRET molecules (Fig. 6e, lower panel). Strikingly, we observed a strong accumulation of subtelomeric transcripts in ter1Δ cells while the signal moderately increased in WT. This accumulation of transcripts was likely caused by ARRET / αARRET molecules because the signal amplification was not detected with the Telo probe (Fig. 6f). We thus concluded that STEEx formation correlates with transcription of subtelomeric regions. Importantly, although subtelomeric regions are expanded in STEEx we could not detect longer transcripts than in WT or rap1Δ strains.

RNA-DNA hybrids promote STEEx formation
The results above suggest that transcription of subtelomeric regions correlates with STEEx formation. In budding yeast, it has been described that RNA-DNA hybrids promote recombination in pre-senescent cells at telomeres 32. These hybrids are known to be removed by the RNase H1 and H2 enzymes 33. We wondered if RNA-DNA hybrids generated by telomeric transcription could influence STEEx formation. We deleted ter1 in cells lacking the RNase H1 (rh1) and investigated STEEx formation in ter1Δ rh1Δ quiescent cells. Senescence kinetics were carried out with ter1Δ and rh1Δ ter1Δ clones by successive streaking on YES agar plates. After streaks 1, 2, and 3 cells were grown in minimal medium, starved from nitrogen source and kept for 8 days in quiescence. Then telomere structure was analyzed by Southern blot (Fig. 7). At streak 1 telomere size was comparable for both ter1Δ and rh1Δ ter1Δ clones but STEEx appeared prematurely in rh1Δ ter1Δ cells compared to ter1Δ∆∆-∆. At streak 2 and 3, STEEx dramatically increased with time in quiescence. These results indicate that RNA-DNA hybrids that are likely to accumulate in ter1Δ rh1Δ cells during quiescence stimulate STEEx formation, further suggesting that telomeric transcription initiates subtelomeric rearrangements.

STEEx are counter-selected when cells exit from quiescence
To analyze cell viability during quiescence we micro-manipulated cell after increasing period of quiescence onto rich media and monitored the proportion of cell forming colonies (CFC). The Figure 8a shows the percentage of wild-type and ter1Δ cells that are unable to form a visible colony after 1, 4, and 8 days in quiescence. This was performed after quiescence of the senescence experiment presented in figure 1. While almost all telomerase positive cells exit G0 (nearly 100% of CFC for the wild-type), telomerase negative cells dramatically failed to exit G0 in a way that was correlated with the telomere shortening and the time in quiescence. Strikingly, more than 70% of cells under conditions in which telomeres were highly rearranged were unable to exit G0 (at senescence day 7 and 8 days of G0). We further analyzed the cells that were not generating colonies and found that all of them were either enlarging or forming micro-colonies of two to twenty cells (Figure 8a). These observations indicated that those cells were not dead in quiescence but were not capable to grow. By micro-manipulating ter1Δ cells before nitrogen starvation, we verified that the inability of telomerase negative cells to form a colony at exit of G0 was not caused by the shortening of telomeres. Indeed, senescent cells were able to form colony with high efficiency while nitrogen starvation of the same cells affected return to growth (Supplementary Figure 4a-b). Taking together, this indicates that STEEx prevented cells to exit properly from quiescence.

To investigate this point, we monitored by Southern blot how stable were the STEEx when quiescent cells were returning to growth. We observed that STEEx were poorly maintained in proliferative cells as indicated by the reduction of the STEEx signal after 2 and 3 days of growth (Fig. 8b left panel, refer to Fig. 1c to visualize G0 profile). PFGE showed that chromosomes were still linear at exit of G0 (Fig. 8b, right panel). We analyzed the telomeric pattern of 15 individual clones that exit from 8 days of quiescence (Supplementary Figure 4c). The common STEEx profile was lost although some reminiscent STEEx bands were still detectable in some clones. We inferred from these results that cells with rearranged telomeres either died or were arrested while re-entering into the cell cycle, or that STEEx are lost when cells replicate their DNA.
Discussion

In this study we investigated the stability of eroded telomeres in quiescent fission yeast cells. While WT telomeres are stable, we observed that eroded telomeres in senescent cells are strongly rearranged during G0. We demonstrated that these rearrangements, named STEEx, correspond to the amplification of subtelomeric blocks delineated by a homologous repeated sequence (HRS) and are promoted by transcription. STEEx formation depends on Rad51-dependent HR and Ctp1 but does not seem to involve Mre11, Exo1 or the main heterochromatin assembly machinery thereby differentiating STEEx from HAATI STE. Noteworthy, Ctp1 CTIP is known to function as co-factor of MRN complex to process DSB repair in fission yeast and human 34,35, however we have revealed that Ctp1 may act distinctively from Mre11 in processing eroded telomeres in quiescent fission yeast cells, likely by controlling resection. Two possible mechanisms for subtelomeric amplification are depicted in figure 9 (see legend for details). These mechanisms are related to BIR (Break Induced Replication) and rolling circle, depending on the usage of the HRS sequences template located in trans or cis of the eroded chromosome end. In both models, the eroded telomeres may generate ssDNA up to telomere proximal HRS. The distal HRS (located in the vicinity of Nsil site) from another chromosome (BIR) or the same chromosome (Rolling circle) could be used as seed to promote Rad51-dependent strand invasion, D-loop formation and DNA synthesis through migration of the D-loop. The presence of two EcoRI fragments of 1533 bp and 1461 bp from two chromosome ends in the WT strain (Figure 2a) and the HindIII fragment of 4200 bp in the tiTel strain (Fig. 3c) supports the Rolling circle process (cis- amplification).

Comparable subtelomeric rearrangements have been previously described in fission yeast in the absence of Ku complex or in a mutant of Pola 29,36, however our study represents evidence for a massive and unique subtelomeric amplification in non-dividing conditions. Somehow, STEEx formation shares similarities with Type I telomere recombination that give rise to survivors in S. cerevisiae cells lacking telomerase activity. However, in contrast to budding yeast type I survivors, STEEx are not maintained when cells exit quiescence to re-enter into the cell cycle. These rearrangements probably activate the DNA damage response (as do to some extend type I telomeres in S. cerevisiae) and are counter-selected in dividing cells. We thus
described in fission yeast a pathway to repair eroded telomeres that operates in telomerase minus cells and is promoted in quiescence. Recently, it was established that RNA interference (RNAi) is essential for quiescence in fission yeast, controlling heterochromatin formation at centromeres and rDNA. Our work additionally shows that telomere repair in quiescence requires distinct mechanism than those that operate in cycling cells.

As schematized in the model of figure 9, STEEx are initiated at regions of homology (HRS) that contain the TERRA transcription site. Although quiescence is known to be characterized by global reduction of transcription, we have established that TERRA was induced in quiescent telomerase positive cells and that TERRA but also ARRET/αARRET transcripts was significantly increased during quiescence in telomerase negative cells (Figure 6). Strikingly, TERRA level correlated with STEEx accumulation in G0 and accumulation of RNA-DNA hybrids exacerbates STEEx formation. Thus we inferred that transcription and possibly R-loop formation at TERRA TSS could facilitate HR and STEEx formation at eroded telomeres in quiescence. On the other hand, STEEx formation could in turn exacerbate the accumulation of transcripts in quiescence. Recently, it was shown in human cells that the recruitment at telomeres of the lysine-specific demethylase 1 (LSD1) required MRE11 and was promoted by TERRA. Future investigations will be required to determine whether TERRA stimulates the resection at eroded telomeres during quiescence.

STEEx was characterized by a three-band pattern that corresponds to EcoRI-subtelomeric DNA fragments issued from digestion of different chromosome extremities. Some of these DNA fragments can be detected at very low level during replicative senescence prior nitrogen starvation and the signal remained constant for each time point of the senescence (not shown). This suggests that STEEx formation may occur from preexisting subtelomeric regions, but are counter-selected when cells proliferate. This highlights that genome stability and telomere maintenance mechanisms adapt to growth environmental conditions.
Methods

Strains and Growth Conditions

All yeast strains used in this study are prototrophic and listed in Supplemental Table S1. Telomerase was deleted by substituting the *ter1* gene by kanamycin or hygromycin cassette by one-step homologous insertion using FwTer1\textsuperscript{275} and RevTer1\textsuperscript{276} primers (FwTer1\textsuperscript{275} 5’-AACGCAACGCCCATGCTTAGAAGGTTGACAAGGAAAATTAATCAAACGGT-3’ and RevTer1\textsuperscript{276} 5’-TTATCTCTCTCTTAGTACGCAAATAATACATTAATTTATTTATTTATATAT-3’).\textsuperscript{40} Strain were grown in minimal medium MM \textsuperscript{41}, at 32°C to a density of 6x10\textsuperscript{6} cells/ml, washed twice in MM deprived from nitrogen (MM-N), and re-suspended in MM-N at a density of 2x10\textsuperscript{6} cells/ml at 32°C, reaching after two rounds of cell division 8x10\textsuperscript{6} cells/ml \textsuperscript{24}. This density of cell bodies is stable for several weeks and was used as a source of G0 cells. Thiamine was added in MM-N to repress *nmt1* promoter at a final concentration of 15 µM. The re-entry into the vegetative cycle was performed by replacing the MM-N by one volume of fresh rich medium (YES).

Cell Viability at exit of quiescence

Exit of quiescence was monitored by micro-manipulation of single quiescent cells on rich-medium YES plates. After 3 days of incubation at 32°C cells colonies were counted.

Telomere Analysis and Spotting Assay

Genomic DNA was prepared from 2x10\textsuperscript{8} cells according to standard protocols and digested with the indicated restriction enzymes (New England Biolabs). The digested DNA was resolved in a 1,2% agarose gel and blotted onto a Hybond-XL membrane (GE Helathcare). After transfer, the membrane was cross-linked with UV and hybridized with different probes listed in Supplemental table S2.\textsuperscript{32}P labeling of DNA probes was performed by random priming using Klenow fragment exonuclease-(New England Biolabs), in presence of [α-\textsuperscript{32}P]CTP and hybridizations were performed in Church buffer at 65°C for Telo and subtelomeric (STE1) probes and 55°C for chromosomal probes. For spotting assays, several dilutions of genomic DNA were
directly spotted onto a Hybond-XL membrane (GE Healthcare). Radioactive signal were detected using a Biorad molecular imager FX.

Quantitative PCR analysis
Oligonucleotides used for qPCR: Subtelomeric primers (Fw-pNsiI\(^{601}\) 5’-
GCACCCTAACGCACTCAAGCCCAC, Rev-pNsiI\(^{602}\) 5’-
GAAATGTGTGGAGATGTATGGAG, Fw-pSwaI\(^{599}\) 5’-
CGAATACCTCGCCTTTAGCATCGGC, Rev-pSwaI\(^{600}\) 5’-GAAGATAAAAAGCAGAGGACTATATTGG, Fw-pApaI\(^{119}\) 5’-TATTTCTTTATTCAACCTTTGGACTTC, Rev-pApaI\(^{252}\) 5’-
GTGTGGAAGATGATGGAAG and chromosomic primers (Fw\(^{214}\) 5’-
TACGCGAGCACCCTTGGCATAT, Rev\(^{215}\) 5’-TTATCGACACCTGGGAGCCCATT)

RNA isolation and Analysis
Total RNA was isolated from 2x10\(^8\) of vegetative cells and 10x10\(^8\) of quiescent cells, using the hot phenol method \(^{42}\). Total RNA was treated three times with RNase-free DNase I (Qiagen) and reverse transcribed using the Superscript III reverse transcriptase (Life technologies) with random hexanucleotide primers (Sigma-aldrich). No reverse transcriptase control is used to assess DNA contamination. Quantitative PCR amplification of cDNA was carried out using the SYBR Premix Ex Taq II (Ozyme) with these cycling parameters: 1 cycle at 95°C for 30 sec, followed by 45 cycles at 95°C for 10 sec, 60°C for 15 sec and 72°C for 20 sec and a final extension step at 72°C for 5 min. TERRA cDNA was quantified by using the primers oF1 (5’- GAAGTTCACTCAATTAATTTGGGTAACGGAG-3’) and oR1 (5’-
GGGCCCCATAGTGGGGGCTATATTGTG-3’) \(^{17}\). Fcp1 cDNA was quantified by using oFc1-F (5’- CGAGACCAGTGAACCTGGGACTATGGGACGAGGACTGGC-3’) and oFc1-R (5’-
CATCAACGCCCAGAGGATA-3’) primers. For Northern blot analysis, RNA was electrophoresed in 1,2% formaldehyde agarose gels, transferred to Hybond-XL membrane (GE Helathcare) and hybridized to radioactively \(^{32}\)P labeled probes at 42°C (table S2). Oligonucleotide U6 (5’-ATGTCGCTAGTGGCATC-3’) was 5’ end-labeled with T4 polynucleotide kinase (New England Biolabs) in presence of \([\gamma-\text{P}]\text{ATP}\) \(^{17}\). Membranes were washed twice in 0,1X Ssc, 0,1% SDS for 20 min at 42°C. Radioactive signal were detected using a Biorad molecular imager FX.
**Pulsed Field Gel Electrophoresis (PFGE)**

1x10^8 cells were resuspended in 50 µl of SEZ buffer (1 M Sorbitol, 50 mM EDTA, 0,5 mg/ml zymolase-100T) and 30 µl of agarose low gelling (Sigma-aldrich) were added to a 0,75% final concentration. Cell suspension was transferred in one plug mold. Plugs were incubated at 37°C during 2 h and 12 h for vegetative and quiescent cells, respectively. Plugs were rinsed 30 min in TE-1% SDS and incubated twice 12 h in Proteinase K buffer (0,5 M EDTA, 50 mM Tris-HCl, pH 9.5, 1 mg/ml Proteinase K). Plugs were rinsed three times in TE containing 1 mM PMSF for 1 hour. Finally, plugs were incubated 3 h at 37°C in 1XTAE containing 100 µg/ml of RNase. Plugs were loaded onto 0,8% agarose gel in 1XTAE buffer. Electrophoresis was performed in a CHEF DR III pulsed-field electrophoresis system (Biorad) in 1XTAE buffer with the following settings: 48 h at 2 V/cm with a 1,8T switch time at an included angle of 106 degrees. DNA was visualized by ethidium bromide staining (1 µg/ml) for 30 min.

The authors declare that the data supporting the findings of this study are available within the paper and its supplementary information files.

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**Author contributions**

L.M. and S.C. designed and performed the experiments and analyzed the data. J.A. and S.M. performed the experiments. S.C., B.A. and V.G. designed the study. S.C. and V.G. wrote the paper.
Competing financial interests

The authors declare no competing financial interests.

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Figure 1. Eroded telomeres are rearranged in quiescence
a) Top, Relative position of the restriction sites in the telomeric and subtelomeric regions of S. pombe chromosomes based on pNSU70. Telomeric (Telo) and subtelomeric (STE1) probes used for Southern blot are represented. Bottom, Genomic DNA from WT cells was digested with EcoRI and Southern blotted. The membrane was hybridized with Telo, STE1 and chromosomal probes. R, replicative sample; M, molecular weight marker; 1, 4, 8 correspond to the number of days in quiescence. A chromosomal probe was used as a loading control.
b) Top, freshly deleted ter1 colony was maintained in exponential phase in liquid EMM medium for about 120 generations by serial dilutions. Population doublings were monitored by cell counting. Bottom, genomic DNA was digested with EcoRI and Southern blotted. The membrane was hybridized with Telo and chromosomal probes. Asterisks indicate the time points of the replicative senescence kinetics that have been shifted to nitrogen-deprived medium.
c) Schematic representation of a typical senescence-quiescence coupled experiment. Cells harvested at different type points of the senescence were shifted in deprived-nitrogen medium and maintained in senescence as indicated.
d) Genomic DNA from quiescent ter1Δ cells was digested by EcoRI and Southern blotted. Membrane was hybridized with Telo, STE1, and chromosomal probes. 1, 4, 6, 8 correspond to number of days for which cells were in G0 before harvesting. S1, S3, S5 and S7 indicate the days during the senescence kinetics at which cells were starved from nitrogen. Subtelomeric rearrangements are shown by asterisks.
e) Genomic DNA from quiescent WT and ter1Δ cells at S3 and S5 was spotted onto Hybond-XL membrane and hybridized with the indicated probes. Signal quantification has been performed with “Quantity one” software and STE1/Control ratio was determined.

Figure 2. Subtelomeric rearrangements in quiescence correspond to duplication of a STE1-extended region
a) Top, relative position of the restriction sites in the telomeric and subtelomeric regions within pNSU70 and pNSU64. Both pNSU64 and pNSU70 are representative plasmids that share significant restriction pattern similarity with pNSU65 / pNSU21 and pNSU77 / pNSU71, respectively 29. The position of telomeric (Telo) and subtelomeric probes (STE1 and STE2) used for Southern blot are represented.
Bottom, genomic DNAs from quiescent cells (ter1+, after 8 days in G0 (+)) and ter1Δ cells (after 1 and 8 days in G0 (1 and 8)) were digested by Apal, EcoRI, Swal and Nsil and Southern blotted. The membrane was hybridized with either Telo, STE1 or STE2 probes, as indicated. Ethidium bromide staining DNA was used as a loading control. Asterisks represent the residual signal from STE1 probe.

b) Schematic representation of an Apal-Apal subtelomeric duplication. STE1-Extended region (STEEx) within subtelomeric region of pNSU70 and pNSU64 (pNSU70\textsuperscript{x2} and pNSU64\textsuperscript{x2}). According with this scheme, EcoRI digestions of pNSU64\textsuperscript{x2} and pNSU70\textsuperscript{x2} generate two DNA fragments of 902 and 1461 bp, and a single DNA fragment of 1533 bp, respectively. These three DNA fragments correspond to the bands that accumulated during quiescence at eroded telomeres. The presence of telomeric repeats at STEEx ends is not established.

Figure 3. Duplications of STE1-extended region are initiated at Homologous Repeated Sequences

a) Position of the restriction sites in the telomeric and subtelomeric regions within pNSU70 and pNSU64. Homologous Repeated Sequences (HRS) are marked in red. Primers within or flanking HRS that are used for quantitative PCR are represented: pApaI, pSwaI and pNsiI.

b) Quantitative PCR analysis of subtelomeric regions in ter1+ and ter1Δ cells. Cycling ter1+ (Repl) or ter1Δ cells that were collected during replicative senescence (Sen) were shifted in deprived-nitrogen medium for 1, 4 and 8 days (G0D1, G0D4, G0D8). Fold enrichment corresponds to the ratio of subtelomeric signal over genomic control signal (See methods). Error bars indicate the SEM from three independent quantitative PCR.

c) Schematic representation of a natural telomere and the transcription-inducible telomere (tiTel). The naf:Pnmt1 cassette was inserted in the left arm of chromosome II. The integrated cassette contains a new HindIII restriction site, while the natural Apal site is disrupted upon integration. Specific tiTel probe (nmt1) and HRS are indicated.

d) STEEX formation at the tiTel. tiTel ter1Δ cells were collected after 59 pds during replicative senescence and shifted to deprived-nitrogen medium. Cells were harvested after the indicated days (1, 4 and 8) in quiescence. Genomic DNA was digested with HindIII and analyzed by Southern blot with a nmt1 probe. R
corresponds to tiTel ter1+ cells collected during replicative growth. Asterisk marks the nmt1 endogenous locus.

e) Schematic representation of nat':Pnmt1 cassette duplication (tiSTEEEx). According with this scheme, HindIII-digestion of rearranged tiTel generates fragments of 4.2 Kb. The combination of tiSTEEEx and natural STEEx lead to accumulation of ti/natural STEEx of heterogeneous size visualized as a smear with the nmt1 probe.

**Figure 4. STEEx rearrangements in quiescent cells depend on Rad51 and Ctp1 but not on Mre11 or Exo1.**

a-d) Telomere structure during quiescence of the indicated mutants. Genomic DNAs from single or double mutants were digested with EcoRI and analyzed by Southern blot with a Telo/STE1 probe. ter1+ cells and ter1Δ cells that were collected at different days during replicative senescence (S1 to S8) and shifted to deprived-nitrogen medium. Cells were harvested after the indicated days (1, 4, 6, 8) in quiescence. R shows ter1+ cells collected during replicative growth and M correspond to molecular weight markers. STEEx are shown by asterisks.

**Figure 5. STEEx formation in quiescent cells does not depend on heterochromatin assembly machinery.**

a) Analysis of STEEx chromosomes by pulsed field gel electrophoresis. Gel was stained with ethidium bromide. b-d) Genomic DNA from quiescent telomerase positive cells and ter1Δ cells with the indicated genetic backgrounds was digested with EcoRI and Southern blotted. Membranes were hybridized with a Telo/STE1 probe. Cells were harvested after the indicated days (1, 4, 6, 8) in quiescence. R shows ter1+ cells collected during replicative growth. STEEx are shown by asterisks.

**Figure 6. TERRA accumulates in quiescence**

a) Schematic representation of the *S. pombe* telomeric transcripts. Subtelomeric region is shown as a grey line and telomeric repeats represented by dark gray triangles. Oligonucleotides used for RT-qPCR (oF1 and oR1, and random hexanucleotide NNNN), TERRA Transcription Start Site (TSS), and HRS are indicated.
b) RNA from telomerase positive (WT) and ter1Δ cells was extracted in proliferative (R) and in quiescent (D1 to D8) cells. RNA levels were determined by RT-qPCR using a random hexanucleotide primer followed by qPCR with specific oligonucleotides for TERRA and control Fcp1. A control experiment was performed without reverse transcription step (-RT).

c) TERRA fold enrichment was determined as the ratio of TERRA over Fcp1 RNA levels. Error bars indicate the SEM from three independent experiments; P values are calculated from two-tailed t-test (*P=0.0355; ** P= 0.001; **** P< 0.001).

d) Schematic representation of the S. pombe telomeric transcripts. The STE1 and Telo probes used for the Northern blots are indicated by back and grey lines, respectively. The TERRA / ARIA and TERRA / ARRET / αARRET transcripts are detected with Telo and STE1 probes, respectively.

e) Northern blot analysis in telomerase positive (WT), rap1Δ, and ter1Δ cells. RNA was extracted in proliferative (R) and in quiescent (D1, D4 and D8) cells. The Northern blot was hybridized with a Telo, STE1 and U6 (loading control) probes.

f) Schematic representation of transcription in ter1Δ quiescent cells. Duplication of TERRA TSS in STEEx may lead to accumulation of new transcripts.

Figure 7. RNA:DNA hybrids promote STEEx formation.
Replicative senescence (S1 to S3) was carried out with ter1Δ and rnh1Δ ter1Δ cells by successive streakings on YES agar plate. Then cells were grown in minimal medium and shifted to deprived-nitrogen medium. ter1Δ and rnh1Δ ter1Δ cells were harvested after the indicated days (1, 4, and 8) in quiescence. Genomic DNAs were digested with EcoRI and analyzed by Southern blot with a Telo and STE1 probe. STEEx are shown by asterisks.

Figure 8. STEEx are counter selected upon exit from quiescence.
a) Wild-type and ter1Δ single cells were maintained in quiescence for 1, 4, or 8 days, micromanipulated and plated onto YES plate allowing them to exit quiescence (refer to figure 1 for senescence and quiescence). The percentage of cells that were not able to form a colony was plotted. Microscopic observation was performed to determine how cells died when they re-enter into cell cycle. We distinguished single
round cell, enlarge cell, and micro-colony (1 to 20 cells). Round cell died in G0 while enlarged or micro-colony attempted to exit G0.

b) *Left panel*, at several time points of the senescence kinetics (S1, S3 and S5) a population of 8-days quiescent cells were allowed to exit G0 by changing medium to YES. Cells were grown for 1, 2 or 3 days in YES medium and harvested. Genomic DNAs were prepared from the harvested cultures, digested with *EcoRI* and analyzed by Southern blot with a Telo/STE1 and control probes. STEEx are indicated by asterisks. *Right panel*, Pulsed Field Gel Electrophoresis (PFGE) analysis was performed with the indicated samples (S1D8, 1 day in senescence, 8 days in G0, and 1 day after exit; S5D8, 5 days in senescence, 8 days in G0, and 3 days of growth after exit). Gel was stained with ethidium bromide.

**Figure 9: Possible mechanisms for STEEx formation in quiescence.**

Resection at unprotected telomeres might occur in quiescence generating a recombinogenic 3’-overhang within the telomere proximal HRS. Homology search and strand invasion by HR machinery might be facilitated by the presence of R-loop within the distal end HRS, thereby forming a D-loop structure. R-loop could be formed in subtelomeric regions by the annealing of TERRA molecules with the distal HRS. Another possibility to generate R-loop would be that RNAs are transcribed from the distal HRS that also contains the TSS sequence. Rad51-mediated HR might occur either internally within the same telomere through a rolling-circle like mechanism, or with another chromosome extremity followed by Break Induced Repair (BIR) related process. D-loop migration and lagging-strand synthesis might then ensure duplication of subtelomeric block. Amplification of this process might lead to STEEx formation. It is not established if STEEx extremities contain telomeric repeated sequences.
Figure 1

(a) Diagram showing the position of Centromere, Telomere, STE1, and Loading control bands.

(b) Graph showing population doublings over time.

(c) Diagram illustrating replicative senescence (MM+N) and quiescence (MM-N) with telomere length.

(d) Table showing telomere length and STE1 expression over days in G0.

(e) Table showing STE1 expression and telomere length over days in G0 for WT, ter1Δ S3, and ter1Δ S5.
Figure 2

(a) Schematic representation of the genetic map of the chromosomes of the yeast species. The map includes the positions of the telomeres, centromere, and the sites of enzyme digestion (SwaI, ApaI, EcoRI, NsiI). The distance between the telomeres is marked as 500 bp.

(b) Enzyme digestion analysis of the chromosomes. The gel electrophoresis showing the fragments obtained after digestion with the indicated enzymes. The sizes of the fragments are indicated in kilobases (kb). The blocks labeled as block 1 and block 2 represent the regions of interest. The sizes of the blocks are 1533 bp and 1461 bp, respectively.
Figure 4

a

|        | ter1Δ         | rad51Δ ter1Δ |
|--------|---------------|--------------|
| WT (ter1+) | S4 | rad51Δ | S1 |
| R | 1 | 8 | R | 1 | 4 | 6 | 8 | R | 1 | 4 | 6 | 8 | (kb) |
| * |   |   |   |   |   |   |   |   |   |   |   |   |
| * |   |   |   |   |   |   |   |   |   |   |   |   |
| * |   |   |   |   |   |   |   |   |   |   |   |   |

High exposure

b

| mre11Δ | mre11Δ ter1Δ |
|--------|--------------|
| R 1 4 6 8 | S1 S3 S5 |
| 1 4 6 8 | 1 4 6 8 | 1 4 6 8 | (kb) |
| * |   |   |   |   |   |   |   |   |   |   |   |   |
| * |   |   |   |   |   |   |   |   |   |   |   |   |
| * |   |   |   |   |   |   |   |   |   |   |   |   |

Mre11Δ


c

| ctp1Δ | ctp1Δ ter1Δ |
|--------|--------------|
| R 1 4 8 | S2 S3 S4 S5 S6 M |
| 1 4 8 | 1 4 8 | 1 4 8 | (kb) |
| * |   |   |   |   |   |   |   |   |   |   |   |   |
| * |   |   |   |   |   |   |   |   |   |   |   |   |
| * |   |   |   |   |   |   |   |   |   |   |   |   |

Ctp1Δ


d

| exo1Δ | exo1Δ ter1Δ |
|--------|--------------|
| R 1 4 6 8 | S2 S4 S6 |
| 1 4 6 8 | 1 4 6 8 | 1 4 6 8 | (kb) |
| * |   |   |   |   |   |   |   |   |   |   |   |   |
| * |   |   |   |   |   |   |   |   |   |   |   |   |
| * |   |   |   |   |   |   |   |   |   |   |   |   |

Exo1Δ
Figure 5

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a

|        | ter1Δ | WT |
|--------|-------|----|
| R      | 1     | 4  |
|        | 8     | R  |

b

|        | swi6Δ | swi6Δ ter1Δ |
|--------|-------|-------------|
|        |       |             |
| R      | 1     | 4  |
|        | 6     | 8  |
|        | M     | (kb)       |

---

c

|        | cir4Δ | cir4Δ ter1Δ |
|--------|-------|-------------|
| R      | 1     | 4  |
|        | 6     | 8  |
| S1     |       | (kb)       |
| S3     |       |             |

---

d

|        | mit1Δ | mit1Δ ter1Δ |
|--------|-------|-------------|
|        |       |             |
| R      | 1     | 4  |
|        | 6     | 8  |
|        | S1    | (kb)       |
| S3     |       |             |
| S5     |       |             |
Figure 6

(a) Schematic representation of TERRA fold enrichment with STE1 probe and Telo probe.

(b) Bar graph showing TERRA fold enrichment from WT, ter1Δ, WT (-RT), and ter1Δ (-RT) with R, D1, D4, D8 conditions.

(c) Histogram showing TERRA fold enrichment from WT and ter1Δ with R, D1, D4, D8 conditions.

(d) Schematic representation of TERRA fold enrichment with STE1 probe and Telo probe.

(e) Gel electrophoresis image showing TERRA fold enrichment with STE1 probe and Telo probe.

(f) Schematic representation of TERRA fold enrichment with STE1 probe and Telo probe.
Figure 7

|     | S1 | S2 | S3 |     | S1 | S2 | S3 |
|-----|----|----|----|-----|----|----|----|
| *   | 1  | 4  | 8  |     | 1  | 4  | 8  |
| ter1Δ |    |    |    | mhlΔ ter1Δ |    |    |    |
| STE1 |    |    |    |     |    |    |    |
| Telo |    |    |    |     |    |    |    |

* indicates a significant difference.
Figure 8

(a) Bar graph showing the percent of cell mortality at the exit of G0. The graph compares WT (wild type) and ter1Δ strains over different days (D1, D4, D8). The graph indicates that ter1Δ strains show higher cell mortality compared to WT strains.

(b) Western blot analysis showing the expression levels of the STE1 gene and a loading control. The blot indicates different expressions at days 1, 2, and 3 after G0 exit. The ter1Δ strain shows a different pattern of expression compared to the WT strain.
Distal HRS

Proximal HRS

Heterogenous eroded telomeres

Resection at unprotected telomere (3’ overhang formation)

BIR

Rolling circle

Strand invasion within HRS

D-Loop formation

R-Loop

TERRA TSS

D-Loop migration

Lagging-strand synthesis

Block duplication

STEx