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Quiescent *Saccharomyces cerevisiae* forms telomere hyperclusters at the nuclear membrane vicinity through a multifaceted mechanism involving Esc1, the Sir complex, and chromatin condensation

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**ABSTRACT** Like other eukaryotes, *Saccharomyces cerevisiae* spatially organizes its chromosomes within the nucleus. In G\textsubscript{1} phase, the yeast’s 32 telomeres are clustered into 6–10 foci that dynamically interact with the nuclear membrane. Here we show that, when cells leave the division cycle and enter quiescence, telomeres gather into two to three hyperclusters at the nuclear membrane vicinity. This localization depends on Esc1 but not on the Ku proteins. Telomere hypercluster formation requires the Sir complex but is independent of the nuclear microtubule bundle that specifically assembles in quiescent cells. Importantly, mutants deleted for the linker histone H1 Hho1 or defective in condensin activity or affected for histone H4 Lys-16 deacetylation are impaired, at least in part, for telomere hypercluster formation in quiescence, suggesting that this process involves chromosome condensation. Finally, we establish that telomere hypercluster formation is not necessary for quiescence establishment, maintenance, and exit, raising the question of the physiological raison d’être of this nuclear reorganization.

**INTRODUCTION**

In yeast, just as in other eukaryotes, chromosomes are spatially organized (Taddei et al., 2010; Albert et al., 2012). This organization is thought to influence gene expression but also DNA repair, replication, and recombination (Cavalli and Misteli, 2013). In the G\textsubscript{1} phase of the proliferating cycle, the yeast nucleus adopts a configuration called “Rabl,” in which centromeres are gathered together close to the spindle pole body (SPB), the yeast equivalent of the centrosome, through short nuclear microtubules (Guacci et al., 1997; Jin et al., 1998, 2000; Bystricky et al., 2004). The nucleolus containing the ribosomal DNA localizes at the SPB opposite pole (Yang et al., 1989). Constrained by the nuclear envelope, chromosomes undergo ATP-dependent undirected motion, each locus mobility being determined by its position along the arm (Marshall et al., 1997; Heun et al., 2001; Bystricky et al., 2004; Hajjoul et al., 2013). Finally, the 32 yeast telomeres cluster into 6–10 dynamic and mobile foci mostly found in the nuclear membrane proximity (Palladino et al., 1993; Gotta et al., 1996; Hediger et al., 2002; Bystricky et al., 2005).

Yeast telomeric sequences are short (~350 base pairs) and made of T(G\textsubscript{1-3}) repeats with a 3′ G-rich extension. Repeats of X elements, and eventually of Y′ elements, compose yeast subtelomeric regions. Many proteins interact with telomeric DNA, including various telomerase-associated complexes, but also Rap1, which binds double-stranded DNA, the Ku proteins, and the Sirtuin (Sir) complex. The Sir complex is composed of the NAD\textsuperscript{+}-dependent histone deacetylase Sir2 and the silencing proteins Sir3 and Sir4. This complex is...
recruited to telomeric DNA in part by Rap1, which interacts with Sir3 and Sir4. The Sir complex then associates with the hypoacetylated N-terminal tails of the histones H3 and H4 and spreads from telomeres into subtelomeric regions (for reviews, see Taddei et al., 2010; Wellinger and Zakian, 2012; Kupiec, 2014).

Telomere localization at the nuclear periphery is dependent on intricate and partially redundant pathways involving Esc1, the Ku complex, and Mps3, an essential integral nuclear membrane protein containing a SUN domain that is involved in SPB organization (Jaspersen et al., 2002; Nishikawa et al., 2003; Mps3, Esc1, and yKu80 interact with Sir4 (Taddei et al., 2010; Wellinger and Zakian, 2012). The nuclear pore proteins of the Nup84 subcomplex have also been involved in telomere tethering at the nuclear periphery (Therizols et al., 2006). In proliferating cells, telomere clustering into foci results from stochastic contacts generated by random motion, the chromosome arm length, and the nucleolus exclusion zone determining the telomere exploration radius (Schober et al., 2008; Therizols et al., 2010; Zimmer and Fabre, 2011; Wong et al., 2012). Telomere clusters are mobile and only transient (Schober et al., 2008; Therizols et al., 2010), their formation requiring intertelomere association through Sir3 oligomerization (Ruault et al., 2011). In fact, telomere clustering is thought to be generated via an aggregation/dissociation equilibrium, the association rate being dependent on geometrical parameters, and the dissociation rate being regulated by the Sir3 protein level (Hozé et al., 2013).

When cells cease to proliferate, they may enter a reversible non-dividing state called quiescence (Coller, 2011; O’Farrell, 2011; De Virgilio, 2012). Many years ago, it was shown that, in nondividing yeast cells, chromatin adopts a more compact arrangement compared with proliferating cells (Piñon, 1978; Lohr and Ide, 1979). More recently, Patterson and coworkers have shown that chromatin is more condensed in glucose-starved quiescent cells than in proliferating cells, this compaction being partly due to an increase in chromatin-bound Hho1, the yeast equivalent of the linker histone H1 (Schäfer et al., 2008). Chromatin compaction generally involves the N-terminal tail of the histone H4, and more precisely, deacetylation of the Lys-16 (Wilkins et al., 2014). Importantly, while acetylated H4K16 is predominant in proliferating cells (Smith et al., 2003), no acetylated H4K16 has been detected in quiescent cells by nano-electrospray tandem mass spectrometry (Ngubo et al., 2011). These data suggest that, upon quiescence establishment in yeast, chromatin adopts a compact conformation, maybe through an increased binding of Hho1 and a possible deacetylation of H4K16.

We have previously reported that, when quiescence is induced by glucose exhaustion, the yeast nucleus is drastically reshaped (Laporte and Sagot, 2014). Indeed yeast cells assemble a stable nuclear microtubule array that not only causes nucleolus relocation but also centromere declusterization and redistribution along the microtubule array (Laporte et al., 2013). Very recently, centromere declustering in quiescence was confirmed by chromosome conformation capture (3C) in an elegant study clearly demonstrating that the overall quiescent cells’ genome organization is significantly different from the one observed in proliferating cells (Rutledge et al., 2015). In fact, Broach and coworkers have established that, in quiescent yeast cells, intrachromosomal interactions increase at longer distance and subtelomeric regions interact with each other much more frequently (both intra- and interchromosomally), these rearrangements being dependent on the condensin complex (Rutledge et al., 2015). The topological reorganizations reported by Rutledge and coworkers were confirmed by Guidi and colleagues using 3C (Guidi et al., 2015). In the later study, fluorescence microscopy analyses demonstrated that, in a subpopulation of dense quiescent cells selected on a Percoll gradient (Allen et al., 2006), telomeres regrouped in one or two hyperclusters in the center of the cell nucleus in a Sir3-dependent manner (Guidi et al., 2015). Astonishingly, telomere hyperclusters were not observed by Rutledge and colleagues, who instead reported an increased number of Rap1-green fluorescent protein (GFP) foci in quiescence (Rutledge et al., 2015). These discrepancies hence question the existence of telomere hyperclusters in the nucleus of live yeast quiescent cells.

In this study, we have investigated the behavior of telomeres in a whole population of live yeast cells that entered quiescence following carbon source exhaustion. Using various telomere-associated proteins fused to fluorescent proteins and fluorescence in situ hybridization (FISH), we show that telomeres form two to three hyperclusters that localize close to the nuclear membrane. This nuclear membrane proximity depends on Esc1 but not on the Ku complex. Additionally, we demonstrate that telomere hypercluster formation not only depends on the Sir complex but also on the chromatin condensation machinery, the hyperclustering being affected in hho1Δ or condensin mutants. We further reveal that deacetylation of the histone H4K16 is critical for the quiescence-induced telomere hyperclustering process. Importantly, upon quiescence exit, telomere hyperclusters slowly disassemble independently of actin and microtubule dynamics. Finally, we unambiguously establish that telomere hyperclustering is not required for cell survival in early quiescence, raising the question of the physiological raison d’être of this specific nuclear reorganization.

RESULTS AND DISCUSSION

Telomeres do form hyperclusters upon quiescence establishment

On carbon source exhaustion, budding yeast cells leave the cell cycle and enter quiescence. In these conditions, we have analyzed by FISH the localization of subtelomeric regions (‘Y’ subtelomeric DNA sequences; Louis and Borts, 1995) in wild-type cells (WT). As previously described, 6–10 telomere clusters were detected in proliferating G1 cells (Palladino et al., 1993; Gotta et al., 1996). By contrast, in quiescent cells, only 2.3 ± 0.5 bright telomere clusters were observed (Figure 1A). These telomere hyperclusters contained Sir2 (Figure 1B), but also Sir3, Rap1, and Sir4 (Figure 1C, Supplementary Figure S1, A and B, and unpublished data), and form regardless of the culture temperature (for culture at 30°C, see Figure 1; at 25°C, Supplementary Figure S1F, and at 37°C, Figure 4D). Of note, in quiescent cells, one of the Sir2-GFP detected foci colocalized with the nucleolus (Supplemental Figure S1C). Kinetic analyses revealed that telomere hypercluster formation was a rather slow process that reached its plateau at –6–7 d of culture at 30°C (Figure 1D and Supplementary Figure S1D). A similar kinetic was observed when quiescence was induced by an abrupt carbon depletion, explaining why Rutledge and coworkers did not observe telomere hyperclusters after one day of carbon shortage when using this protocol (Rutledge et al., 2015; Supplemental Figure S1E). We also found that virgin and young mother cells were slightly more prone to form telomere hyperclusters than cells that have undergone more than four divisions (Supplemental Figure S1F). Our findings are consistent with chromosome conformation capture analyses demonstrating that intertelomere interactions increase in quiescent cells (Guidi et al., 2015; Rutledge et al., 2015). They are also in agreement with Guidi and colleagues, who found that subtelomeric regions form one to two clusters in 7-d-old W303 cells (Guidi et al., 2015). Taken together, these data clearly establish that telomeres do form hyperclusters upon quiescence establishment following carbon source exhaustion.
described telomere hyperclusters in the inner zone of the quiescent cells nucleus (Guidi et al., 2015), and will be discussed in the following sections.

To more precisely localize telomere hyperclusters, we took advantage of the nuclear microtubule bundle that emanates from the SPB in quiescent cell nuclei (Laporte et al., 2013). We coimaged Sir2-GFP together with the nuclear microtubule bundle (Figure 2C). The nuclear microtubule bundle was automatically detected and used as an axis of reference, the SPB being the origin of the axis (yellow arrow in Figure 2D) on two-dimensional (2D) projections of z-sections (maximum projection). The nuclei were consequently reduced to hemi-disks. The relative position of each telomere hypercluster was then determined using a dedicated MatLab script (Figure 2D and Materials and Methods). We found that, statistically, telomere hyperclusters were excluded from both the SPB region and an SPB opposite zone that corresponds to the tip of the nuclear microtubule bundle (Figure 2D). The exclusion of telomeres from this later zone could not be due to the presence of the nucleolus, as we have previously shown that the nuclear microtubule bundle assembly causes nucleolus displacement toward one side of the nucleus in more than 80% of the cells (Laporte et al., 2013). It is interesting to note that the telomere reorganization in quiescence is distinct from the bouquet arrangement adopted by meiotic cells that gather their telomeres together close to the SPB (Trelles-Sticken et al., 1999, and references therein). Consistently, we found that neither Csm4 nor Ndj1, two proteins involved in telomere rearrangement during bouquet formation, were required for telomere hypercluster formation upon quiescence entry (Supplemental Figure S1G).

Telomere hyperclusters localize close to the nuclear membrane

In quiescent cells, we found that telomere hypercluster movements were confined (Figure 2A, red line), contrasting with their mobility in proliferating G1 cells (Figure 2A, green line). In fact, in quiescent cells, as in proliferating G1 cells, we mostly observed telomere hyperclusters close to the nuclear membrane (<250 nm, Figures 2B and 3C). This is in striking contrast with Guidi and coworkers, who...
Telomere hyperclusters localize to nuclear membrane through Esc1 but not yKu

To get insight into the molecular relationships between telomere hyperclusters and the nuclear membrane in quiescent cells, we focused on yKu70/80 and Esc1, proteins known to play a role in telomere localization at the nuclear periphery in proliferating cells (Taddei et al., 2010; Welling and Zakian, 2012; Kupiec, 2014). In WT quiescent cells, we found that yKu80 colocalized with telomere hyperclusters (Figure 3A), while yKu70 could not be detected (unpublished data). Intriguingly, we observed that Esc1 localized as discrete zones all around the nuclear membrane of quiescent cells (Supplemental Figure S2A) with the exception of the SPB proximal zone and the region opposite to the SPB (Figure 3B), just as telomere hyperclusters do. Of note, in proliferating G1 cells, Esc1 was detected next to the SPB but was also excluded from the zone opposite to the SPB, probably because of the presence of the nucleolus in this region (Supplemental Figure S2B; Taddei et al., 2004).

Importantly, we found that telomeres were still hyperclustered in esc1Δ cells, but their localization close to the nuclear membrane was strongly impaired. Indeed, telomere hyperclusters randomly localized inside the nucleus (for Sir3-GFP, see Figure 3C; for Sir2-GFP, see Supplemental Figure S2C). Yet no significant difference in telomere hypercluster motility was measured between esc1Δ and WT quiescent cells (Supplemental Figure S2D). This suggests that the slow motion of telomere hyperclusters observed in quiescent cells was not a consequence of a tight interaction with the nuclear membrane. Additionally, deletion of yKu protein–encoding genes had no effect either on telomere hypercluster formation or localization to the nuclear membrane vicinity (Figure 3C and Supplemental Figure S2C), and no additional defect was observed when combining esc1Δ with yku deletions (Supplemental Figure S2, C and E). Taken together, our data demonstrate that quiescent cell telomere hyperclusters localize close to the nuclear membrane through Esc1.

Telomere hypercluster formation requires the Sir complex

In proliferating cells, the Sir complex has been involved in telomere clustering (Palladino et al., 1993; Gotta et al., 1996). Using FISH, we found that deletion of sir2Δ, sir3Δ, or sir4Δ affected telomere hypercluster formation in quiescent cells (Figure 4A). This is in agreement with the findings of Guidi and colleagues, who described the absence of Rap1-GFP hyperclusters in sir3Δ quiescent cells (Guidi et al., 2015). Additionally, we found that the Sir3 signal detected using red fluorescent protein (RFP) or GFP increased in quiescent cells compared with G1 proliferating cells (Supplemental Figure S3, A and B), and Western blotting indicated a slight augmentation of Sir3 steady-state level in quiescence (Supplemental Figure S3C). In proliferating cells, Ruault and coworkers have shown that overexpression of Sir3 leads to telomere hypercluster formation and that nonacetylable Sir3 can promote telomere clustering independent of its spreading in subtelomeric regions (Ruault et al., 2011). Because chromatin immunoprecipitation experiments did not detect any significant changes in the spreading of Sir3 between exponentially growing cells and quiescent cells (Guidi et al., 2015), we propose that, in quiescence, increased Sir3–Sir3 interactions promote trans interaction between Sir3-bound telomeres, leading to the formation of hyperclusters.
the recruitment of Sir3 and consequently participate in the Sir3-induced strengthening of telomere–telomere interaction in quiescent cells. Second, deacetylation of H4K16 has been associated with chromatin condensation, since it regulates the interaction between the histone H4 N-terminal tail and the H2A-H2B dimer within the nucleosomes (Wilkins et al., 2014). As mutations altering H4/H2A-H2B interaction such as H4K16A or H4K16Q also affected telomere hypercluster formation, we speculate that this process could be influenced by chromatin condensation.

In fact, in proliferating cells, telomere cluster formation is thought to be influenced by structural constraints (Hozé et al., 2013). The linker histone Hho1 has been involved in chromatin compaction during quiescence establishment (Schäfer et al., 2008). Interestingly, we found that telomere hypercluster formation was slightly impaired in quiescent hho1Δ cells (Figure 4C). Moreover, inactivating the condensin complex (smc2-8) drastically compromised telomere hypercluster formation (Figure 4D and Supplemental Figure S3D), in agreement with Rutledge and coworkers, who demonstrated that Smc2 is involved in chromosomes compaction in quiescent yeast cells (Rutledge et al., 2015). Therefore we could envision that, upon quiescence establishment, chromosome-arm condensation is strengthened, thereby reducing the telomere exploration area and indirectly triggering telomere hypercluster formation.

Telomere hyperclusters slowly disassemble upon quiescence exit

When quiescence exit was triggered by cell refeeding, the number of telomeric clusters per cell progressively increased, reaching 6.7 ± 1.7 upon entry into S phase (Figure 5A). As for other quiescent

This increased Sir3 recruitment in quiescence may rely on posttranslational modifications that were shown to modulate Sir3 interaction with chromatin in actively dividing cells (Arnaudo et al., 2013).

Ruault and colleagues have shown that, when Sir3 is highly over-expressed in proliferating cells, telomere hyperclusters localize inside the nucleus rather than at the nuclear periphery, possibly because of a competition between Sir3 and Esc1 for the binding of Sir4 (Ruault et al., 2011). We found that the Sir3 steady-state level in quiescent BY strains was slightly lower than the one we detected in quiescent W303 (Supplemental Figure S3C), a difference that may explain why telomere hyperclusters were found in the nucleoplasm of quiescent W303 cells (Guidi et al., 2015).

Telomere hypercluster formation requires H4K16 deacetylation and the chromatin condensation machinery

In quiescent cells, the histone H4 Lys-16 (H4K16) is solely found in its deacetylated form (Ngubo et al., 2011). To address the role of this histone H4 posttranslational modification, we analyzed telomere hypercluster formation in histone H4 mutants. No telomere hyperclusters were detected in H4 mutants bearing N-terminal tail deletion or H4K16A or H4K16Q point mutations. Conversely, a mutation preventing H4K16 acetylation (H4K16R) had no effect on telomere hypercluster formation (Figure 4B). Thus deacetylation of H4K16 seems to be a key event that is needed for telomere hypercluster formation. Two nonexclusive hypotheses can be envisioned regarding the influence of H4K16 deacetylation on telomere hypercluster formation. First, as Sir3 is known to preferentially bind unacetylated H4K16 (Carmen et al., 2002; Onishi et al., 2007; Oppikofer et al., 2011), the extensive deacetylation of H4K16 might increase the recruitment of Sir3 and consequently participate in the Sir3-induced strengthening of telomere–telomere interaction in quiescent cells. Second, deacetylation of H4K16 has been associated with chromatin condensation, since it regulates the interaction between the histone H4 N-terminal tail and the H2A-H2B dimer within the nucleosomes (Wilkins et al., 2014). As mutations altering H4/H2A-H2B interaction such as H4K16A or H4K16Q also affected telomere hypercluster formation, we speculate that this process could be influenced by chromatin condensation.

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cell-specific structures such as actin bodies, proteasome storage granules, or nuclear microtubule bundle (Sagot et al., 2006; Laporte et al., 2008, 2013), telomere hyperclusters disassembled even if the de novo protein synthesis was inhibited by cycloheximide (CHX; Figure 5B). During Saccharomyces cerevisiae meiosis, telomere movements were shown to be driven by actin cables (Koszul et al., 2008). Figure 5B shows that telomere hypercluster disassembly upon quiescence exit was clearly unaffected by latrunculin A, a drug causing the depolymerization of all yeast actin filament–containing structures (Ayscough et al., 1997), including the actin cables assembled within seconds upon quiescent cell refeeding (Sagot et al., 2006). At the present time, we do not know what causes telomere hypercluster disassembly upon quiescence exit. We can speculate that it may involve a sequence of events mirroring those occurring upon quiescence establishment, that is, the H4K16 reacetylation, the subsequent decrease in Sir3 interaction with subtelomeric regions, and chromosome arm decondensation upon entry into S phase.

**Telomeres hyperclusters are not required for cell survival in quiescence**

On quiescence establishment following glucose exhaustion, yeast cells assemble a nuclear bundle of microtubules that not only causes nucleolus relocalization but also centromere declusterization and redistribution (Laporte et al., 2013). Intriguingly, telomere hyperclustering was only slightly affected in some (kar3Δ and dyn1Δ) but not all (pnm1Δ) mutants in which the nuclear microtubule bundle formation is compromised (Supplemental Figure S3, E and F). Therefore, if the nuclear microtubule bundle formation clearly participates in several nuclear reorganizations occurring upon quiescence establishment (Laporte et al., 2013; Laporte and Sagot, 2014), it is not strictly required for telomere hypercluster formation. Accordingly, nocodazole treatment did not affect telomere hypercluster disassembly upon quiescence exit (Figure 5B), demonstrating that microtubule dynamics is not required for this process. Besides, the nuclear microtubule bundle assembly was not affected in hho1Δ, sir3Δ, sir4Δ, hho1Δ, yku70Δ, and esc1Δ cells (Figure 4 and Supplemental Figure S3G). These results demonstrate that microtubule remodeling in quiescence does not rely on telomere hypercluster formation or telomere interaction with the nuclear membrane. Therefore the reorganization of microtubules and telomeres can be dissociated and, as such, can be considered as independent quiescence-specific events.

Importantly, while the assembly of the nuclear bundle of microtubules is critical for cell survival in quiescence (Laporte et al., 2013), none of the mutations impeding telomere hypercluster formation or localization had a significant effect on cell viability in quiescence (Supplemental Figure S3H) or quiescent cells’ ability to give rise to a viable progeny upon quiescence exit (Figure 5B and Supplemental Figure S3I). The formation of other quiescent cell–specific structures (actin bodies or nuclear microtubule bundle) testify to accurate quiescence establishment in those mutants (Supplemental Figure S3G). A number of studies have proposed the existence of a correlation...
between gene subnuclear positioning and transcriptional activity in both yeast and mammals (Taddei et al., 2010; Albert et al., 2012; Nguyen and Bosco, 2015). Our data demonstrate that, upon glucose exhaustion, telomere hypercluster formation and localization are not required for survival in quiescence. Thus, if telomere hypercluster formation and localization modulate gene expression, this transcriptional regulation seems to not be mandatory for quiescence establishment, maintenance, and exit.

In conclusion, if the active assembly of quiescent cell–specific structures such as actin bodies or nuclear microtubule bundle are necessary for cell survival in quiescence (Sagot et al., 2006; Laporte et al., 2013), other reorganizations, like telomere hypercluster formation, are not strictly required for facing chronological aging (Figure 5). Of course, drugs or physical conditions to which mutants unable to assemble telomere hyperclusters in quiescence are sensitive could potentially be found. However, their specific effect on the quiescent state will be difficult to tackle. Nevertheless, at the present time, we cannot rule out that telomere hyperclustering may have a role in the long-term maintenance of yeast quiescence or could be involved in the fitness of quiescent exit.

Hence either telomere hypercluster formation in quiescence does have a physiological function, but we have not appreciated it yet, and an extensive amount of work is needed to shed light on the biological significance of this specific reorganization, or telomere hyperclusterization does not provide any beneficial trait. Why would cells actively embark on rearranging telomeres if this reorganization does not influence their survival in quiescence? In fact, telomere hypercluster formation could be just a passive consequence of other quiescence-induced modifications that may be vital for cells, such as chromatin hypercondensation (Piñon, 1978; Lohr and Ide, 1979; Schäfer et al., 2008; Rutledge et al., 2015) and/or the increase of nucleoplasm molecular crowding that alters the biophysical properties of the nuclear environment (Joyner et al., 2016). Therefore, to shed light on the obscure molecular mechanisms involved in quiescence survival, one of the key issues will be to establish whether each structure specifically assembled in quiescent cells results from a dedicated active process or whether it is just a passive consequence of another quiescence-induced phenomenon.

MATERIALS AND METHODS

Yeast strains and growth conditions

All the S. cerevisiae strains used in this study are isogenic to BY4741 or BY4742 available from GE Healthcare (Chicago, IL) and are listed in Supplemental Table S1. Yeast strains carrying GFP fusions were obtained from Invitrogen (Carlsbad, CA). Histone H4 mutants and the corresponding control strain were obtained from Invitrogen. The sir2Δ mutant (Strunnikov et al., 1995) was obtained from C. Boone (University of Toronto, Canada). The RFP (tdimer 2(12); Campbell al., 2002) sequence carried by plasmids p3695, p4589, p5041, p5043, and p5045 was integrated at the 3′ end of the NUP2, BIM1, RAP1, SIR2, and SIR3 endogenous loci, respectively. Three tandem copies of the GFP sequence carried by the plasmid p4587 were integrated at the 3′ end of the BIM1 endogenous locus (Laporte et al., 2013). Details of the constructions are available upon request. The Sir2-GFP fusion protein functionality was analyzed by measuring cell size using a Beckman Coulter Multi-sizer 4 (Beckman Coulter, Brea, CA). As expected sir2Δ cells were large, the median cell volume for sir2Δ being 53.4 μm³ (=40,000 cells counted). By contrast, the median cell volumes for WT, Sir2-GFP, and Sir2-RFP cells were 45.4, 45.5, and 43.8 μm³ (=25,000 cells counted), respectively. The functionality of the Sir3-GFP construct was attested by the viability of the Sir3-GFP pfd1Δ strain.

All experiments were done at least in duplicate and, unless specified, more than 200 cells were scored. Yeast cells were grown in liquid yeast–peptone–dextrose–adene (YPDA) medium at 30°C in flasks, as described previously (Sagot et al., 2006), except for Figure 4D and Supplemental Figure S3D, for which cells were grown at 37°C and 25°C, respectively.

For quiescence exit in the presence of different drugs (Figure 5B), cells were preincubated 25 min in the presence of the drug before quiescence exit. Drugs used were CHX (180 μM; Sigma-Aldrich, St. Louis, MO), nocodazole (7.5 μM; Sigma-Aldrich), and latrunculin A (200 μM; Enzo Life Sciences, Farmingdale, NY).

Cell staining

For identification of mother and daughter cells (Supplemental Figure S1F), 7-d-old WT cells expressing Sir2-GFP were incubated 5 min with Calcofluor white (20 μg/ml), then washed twice in phosphate-buffered saline (PBS) and imaged.

Cell viability (Supplemental Figure S3H) was scored after 5 min incubation in a solution containing 0.2% of methylene blue (Sigma-Aldrich) and 2% sodium citrate solution, pH 7 (Sigma-Aldrich).

For actin phalloidin staining (Supplemental Figure S3G), cells were fixed with freshly made paraformaldehyde (PFA) prepared as follows: PFA (3.8% final; Sigma-Aldrich) was suspended in PEM (0.1 M PIPES, 1 mM EGTA, 1 mM MgSO4, pH 6.9, with NaOH 5 N), vortexed every 20 min for 1 h at 70°C, and then centrifuged 5 min at 3000 rpm. After 1 h fixation at 30°C, cells were washed twice with PEM, resuspended in 1.5 ml of PEM + 1% Triton X-100, and incubated 3 min. Samples were then washed twice in 1.5 ml of PEM, resuspended in PEM containing 1/10 volume of Alexa Fluor phalloidin (Invitrogen) and incubated 24 h at 4°C. Finally, cells were washed twice, resuspended in a mounting solution containing 70% glycerol and 5 mg/ml para-phenylenediamine, and imaged.

The probe for FISH experiments was obtained by PCR with a template plasmid containing ~5 kb of Y′ element (pEL42H10; Louis and Borts, 1995) and using the primer pair GAAGATTGGCCCT-GCTCTTG/CCGTAAGCTGTCATTATT. PCR purification was followed by a nick translation labeling reaction using the Nick Translation kit (DIG-Nick Translation Mix; Roche, Mannheim, Germany; ref. 11745816910). After 5 min at 98°C, the probe was purified by ethanol precipitation and resuspended at the concentration of 25 ng/μl in a hybridization mix (50% formamide, 2×SSC, 10% dextran-sulfate, 0.5 mg/ml single-stranded salmon sperm DNA).

Spheroplasts were created as follows. Quiescent cells were fixed 15 min in 1.9% PFA (Sigma-Aldrich; diluted in PEM), centrifuged 1 min at room temperature, and washed twice in PEMS (PEM + 1.2 M sorbitol filter sterilized). Cells were then incubated 10 min in SH buffer (β-mercaptoethanol 0.5 M, Tris-HCl 0.1 M, pH 9.3) and washed twice in KCl Tris-HCl buffer (50 mM KCl, 2.5 M, Tris-HCl 50 mM, pH 7). Cells were washed twice in PEM, resuspended in 1 ml PEMS containing 27 mg/ml Zymo 20T (MP Biomedical, Santa Ana, CA; ref. 32092), and incubated at 37°C for 90 min (27 mg/ml Zymo 20T) and then vortexed every 20 min for 1 h at 70°C, and then centrifuged 5 min at 3000 rpm. After 1 h fixation at 30°C, cells were washed twice with PEM, resuspended in a mounting solution containing 70% glycerol and 5 mg/ml para-phenylenediamine, and imaged.

For FISH experiments (Figures 1A and 4, A and B, and Supplemental Figure S2E), 200 μl of the spheroplast preparation described above was centrifuged and washed sequentially with 150 μl 2×SSC, 150 μl 2×SSC 10% formamide, 150 μl 2×SSC 20% formamide, and 150 μl 2×SSC 40% formamide. Between each wash, cells were incubated 15 min. Then, 35 μl of the probe mix (150 ng of probe, 30 μl hybridization mix, adjusted to 38 μl with H2O) incubated at 75°C for
FISH protocol described above was then applied. Then washed once in PEM-BAL (300 μl) and twice in PEM (500 μl). After addition of GFP antibody (anti-GFP from mouse; Roche, ref. 11814460001, 1/50), cells were incubated overnight and then washed three times in PEM-BAL. The secondary antibody (sheep anti-mouse CY3 1/400; ThermoFisher Scientific, Waltham, MA) was added in PEM-BAL, and cells were incubated overnight and then washed twice in 100 μl PBS-BAG and once in 100 μl PBS. Cells were then mounted on poly-l-lysine–coated slides, with anti-fading reagent containing 4′,6-diamidino-2-phenylindole (DAPI) solution (Fluoroshield; Sigma-Aldrich, ref: F6057) before imaging.

For immuno-FISH experiments (Figure 1B), spheroplasts were washed twice for 30 min in 100 μl PEM-BAL (PEM + 1% BSA [Sigma, ref. B4287], 0.1% NaNO3, 100 mM lysine hydrochloride). After addition of GFP antibody (anti-GFP from mouse; Roche, ref. 118114460001, 1/50), cells were incubated overnight and then washed three times in PEM-BAL. The secondary antibody (sheep anti-mouse CY3 1/400; ThermoFisher Scientific, Waltham, MA) was added in PEM-BAL, and cells were incubated overnight and then washed once in PEM-BAL (300 μl) and twice in PEM (500 μl). For the last wash, 25 μl RNaseA (10 mg/ml) was added and incubated 3 h at 37°C. Finally, spheroplasted cells were fixed using 3% PFA for 30 min and then washed three times in 500 μl PEM. The FISH protocol described above was then applied.

Cell viability
Quiescent cells (n > 120 cells) were micromanipulated as described previously (Laporte et al., 2011). Plates were incubated 3 d at 30°C before colony scoring. Colony-forming capacity was addressed after 2, 7, and 14 d at 30°C by plating 200 cells, measured using a Beckman Coulter MultiSizer 4, on YPDA. Each strain was tested in duplicate, and each plat- ing was done in triplicate.

Fluorescence microscopy
Cells were observed in a fully automated Zeiss 200M inverted microscope (Carl Zeiss, Thornwood, NY) equipped with an MS-2000 stage (Applied Scientific Instrumentation, Eugene, OR), a Lambda LS 175-W xenon light source (Sutter, Novato, CA), a 100X 1.4 NA Plan-Apochromat objective, and a 5-position filter turret. For GFP imaging, we used a FITC filter (Ex: HQ487/25-Em: HQ535/40-BS: QS505l). For RFP imaging, we used a Cy3 filter (Ex: HQ535/50-Em: HQ610/75-BS: QS655l). For calcofluor imaging, we used a DAPI filter (Ex: 360/40-Em: 460/50-BS: 400). All the filters are from Chroma Technology. Images were acquired using a CoolSnap HQ camera (Roper Scientific, Tucson, AZ). The microscope, camera, and shutters (Uniblitz, Rochester, NY) were controlled by SlideBook software 5.0 (Intelligent Imaging Innovations, Denver, CO). Images are, unless specified, 2D maximal projection of z-stacks performed using a 0.25 μm step. For live-cell imaging, 2 μl of the cell culture was spotted onto a glass slide and immediately imaged at room temperature.

For fluorescence intensity measurement of Esc1-GFP (Figure 3B and Supplemental Figure S2B), a line scan (i1) of 5 pixel width was drawn along nuclear membrane using ImageJ software to simultaneously measure Esc1-GFP and Spc42-RFP fluorescence. Background was subtracted from these fluorescence intensities. The maximum intensity in Spc42 data (i.e., the SPB location) was used to align the two data sets. For Spc42 graph representation, the fluorescence intensity of the different measurements was summed. For Esc1 graph representation, data were processed as follows: Esc1 intensities <0 were set to zero, while intensities >0 were set to 1. This binary approach allows the construction of a probability map (sum of 0 and 1 divided by the number of measurements at a specific distance from the SPB) and avoids the overrepresentation of strong fluorescence signals.

For fluorescence intensity measurement of Sir3, Sir2, and Sir4 (Supplemental Figure S3, A and B), a circle (i1) containing both RFP or GFP signal and background was drawn around a telomeric focus (Supplemental Figure S3A) or around all telomeric signals (Supplemental Figure S3B) using ImageJ software. A circle two times larger at the same location was drawn to calculate the intensity of the surrounding background (i2). The real intensity (ir) was calculated as follows: ir = (ib i2) – (i1 i1) and ir = i1 – [ib(area i2 – area i1)]. Measures displayed on graphs were obtained by multiplying ir by area i1. Sir3-GFP/RFP signals measured in proliferation and in quiescence were multiplied by an arbitrary factor 2 to be on the same scale as Sir2-GFP/RFP signals.

For distance measurement (Figure 3C and Supplemental Figure S2C), distances between Sir3-GFP or Sir2-GFP foci and the nuclear membrane were determined using ImageJ. Given that the position of the nuclear membrane cannot be set precisely at the top or bottom of the cell, only telomere hyperclusters located in the equatorial plane of the nuclear membrane were scored.

Edge detection and heat-map construction
Individual cells were marked out manually directly from the raw data, and the corresponding z-stacks were exported as separate TIFF files using ImageJ. The detection and localization of telomere clusters and microtubule bundle were performed in each z-stacks using the Matlab (Mathworks) scripts summarized below. For each type of detection, the approaches used and the choice of analysis parameters were confirmed by comparison of the detected shapes with visual inspection. Individual telomere clusters were isolated and counted as follows: First, the averaged intensity (background) was subtracted from each plane of the z-stack independently and the resulting signal was smoothed with a 3 × 3 pixel Wiener filter and then normalized to its absolute maximum value over the entire stack. Then intensity thresholds were used to convert the z-stack to binary images. Matlab's Image Processing Toolbox shape-detection and morphological operation algorithms were next applied to detect and isolate 2D shapes in each plane of the z-stack, yielding a list of GFP spots with their three-dimensional (3D) coordinates and area. To prevent double counting of the same telomere clusters, we removed spots within a neighborhood of 7 pixels. This step eliminates the trace of a unique telomere cluster in adjacent planes of the z-stack (due to the blur along the z-axis), and ensures that the z coordinate of a cluster is attributed to the plane where its projection has the largest area (equatorial plane). Similarly, nuclear microtubule bundle signal (Bim1-RFP signal) was first normalized to its stack-wide maximum value and binarized using an intensity threshold. Then shape detection and morphological operations were performed to find the contour of the bundle in each plane. The extremity close to the brightest region of the bundle was identified as the SPB. For determination of the localization of the bundle extremities along the Z axis, the signal intensity was averaged over a 25-pixel window centered on the extremities of the trace of the bundle in each plane. The variations of these signals over the planes showed clear maxima in specific planes, indicating the most likely localization of each bundle's extremity along the z-axis. Finally, we calculated the distance of each telomere cluster to the two extremities of the bundle in 3D and normalized this distance to the...
length of the bundle to allow compilation of telomere cluster localization relative to nuclear microtubule bundle axis over multiple cells (Figure 2D, top right).

For heat-map construction, the nucleus was “sliced” along the nuclear microtubule bundle. The number of telomere clusters within each slice was normalized to the 3D volume of the slice, computed under the assumption of a spherical nucleus. The resulting heat map, shown in Figure 2D, bottom right, represents the probability density to find a telomere cluster in various regions of the nucleus. Note that the assumption of a spherical nucleus tends to underestimate the cluster density close to the equator. Indeed, for nuclei that tend to stretch along the microtubule bundle (as observed in vivo in some quiescent cells), the equatorial slices are smaller than for the spherical model, and the volumetric factor is overestimated.

**Mean square displacement (MSD) measurement**

For individual spot dynamics analysis (Figure 2A and Supplemental Figure S2D), Sir2-GFP spots were tracked over time in ImageJ to extract spot coordinates along 2D trajectories spanning 30–50 time points (4–7 min movies). For each trajectory, the MSD was computed as a function of the time frame between trajectory points (Michalet and Berglund, 2012). The presented MSD curve is averaged over multiple telomere trajectories (WT Gy cells: n = 54; WT quiescent cells: n = 127; escΔA quiescent cells: n = 62).

**Western blot**

Strains were grown overnight in YPDA medium and then diluted to OD_{600} = 0.1. Cells (10^8) were harvested in exponential growth (12 h) or after 7 d, and 200 μl of TCA 20%, 500 μl of acid-washed glass beads, and 200 μl of TCA buffer (20 mM Tris-HCl, pH 8, 50 mM ammonium acetate, 2 mM EDTA, 1 mM phenylmethylsulfonyl fluoride (PMSF), 1 μl of protease inhibitor cocktail [Sigma-Aldrich]) was added. Two 1-min rounds of vortexing were done to disrupt the cells. Samples were centrifuged at 14,000 rpm for 30 min at 4°C, and pellets were resuspended in 85 μl of SDS–PAGE sample buffer (120 mM Tris base, 3.5% SDS, 8 mM EDTA, 5% β-mercaptoethanol, 1 mM PMSF, 15% glycerol, 0.01% bromophenol blue). Extracts were boiled for 10 min and centrifuged at 14,000 rpm for 10 min. For immunoblotting, we used anti-Sir3 polyclonal antibody (1/200 dilution; yN-20; Santa Cruz Biotechnology, Dallas, TX). Loading was normalized according to anti-Ade13 antibodies (1:300,000; Escusa et al., 2006).

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