Single Stem Cell Imaging and Analysis Reveals Telomere Length Differences in Diseased Human and Mouse Skeletal Muscles

Elisia D. Tichy,1 David K. Sidibe,1 Matthew T. Tierney,2 Michael J. Stec,2 Maryam Sharifi-Sanjani,1 Harish Hosalkar,4 Scott Mubarak,4 F. Brad Johnson,5 Alessandra Sacco,2 and Foteini Mourkioti1,6,*

1Department of Orthopaedic Surgery, Perelman School of Medicine, The University of Pennsylvania, 112A Stemmler Hall, 3450 Hamilton Walk, Philadelphia, PA 19104-6081, USA
2Development, Aging and Regeneration Program, Sanford Burnham Prebys Medical Discovery Institute, La Jolla, CA 92037, USA
3Joint Preservation Center, Tricity Medical Center, Joint Preservation & Deformity Correction Center & Traumatic Brain Injury Program, Paradise Valley Hospital, National City, CA 91950, USA
4Department of Orthopedic Surgery, Rady Children’s Hospital, 3030 Children’s Way, San Diego, CA 92123, USA
5Department of Pathology and Laboratory Medicine, Institute on Aging, Perelman School of Medicine, University of Pennsylvania, Philadelphia, PA 19104, USA
6Department of Cell and Developmental Biology, Perelman School of Medicine, The University of Pennsylvania, Philadelphia, PA 19104, USA

*Correspondence: fmour@mail.med.upenn.edu
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SUMMARY

Muscle stem cells (MuSCs) contribute to muscle regeneration following injury. In many muscle disorders, the repeated cycles of damage and repair lead to stem cell dysfunction. While telomere attrition may contribute to aberrant stem cell functions, methods to accurately measure telomere length in stem cells from skeletal muscles have not been demonstrated. Here, we have optimized and validated such a method, named MuQ-FISH, for analyzing telomere length in MuSCs from either mice or humans. Our analysis showed no differences in telomere length between young and aged MuSCs from uninjured wild-type mice, but MuSCs isolated from young dystrophic mice exhibited significantly shortened telomeres. In corroboration, we demonstrated that telomere attrition is present in human dystrophic MuSCs, which underscores its importance in diseased regenerative failure. The robust technique described herein provides analysis at a single-cell resolution and may be utilized for other cell types, especially rare populations of cells.

INTRODUCTION

Telomeres are long, repetitive DNA sequences (5’-TTAGGG-3’) that are present at chromosome ends (Collins, 2000). During each cycle of DNA replication, telomeres shorten, as DNA polymerases have no primers available to complex with and extend DNA (Ohki et al., 2001). Telomere shortening can also result from aberrant nuclease activity (Wu et al., 2012). Significantly eroded telomeres activate the DNA damage response, inducing cellular senescence and/or the activation of cell death processes (Shay and Wright, 2005). Cells have evolved mechanisms to combat such a dilemma. Classically, the action of telomerase (TERT), an RNA primer (TERC/TR), and accessory factors can extend telomere length in cells where these components are expressed and active (Sarek et al., 2015). The proper functioning of this pathway could play a crucial role in the regulation of stem cell aging and the prevention of the stem cell dysfunctional phenotype observed in degenerative disorders (Blasco, 2007b; Flores and Blasco, 2010).

Telomerase activity is most active during early development, after which the activity becomes reduced (Harley and Villepontage, 1995). In the setting of degenerative disease, stem cells may lack the ability to extend telomere length, thus making them susceptible to premature dysfunction. Indeed, telomere shortening in relation to loss of self-renewal capacity has been reported in hematopoietic stem cells, induced pluripotent stem cells, and embryonic stem cells (Batista et al., 2011; Morrison et al., 1996; Niida et al., 2000). While telomere defects have been extensively studied in other systems and stem cell compartments (Flores et al., 2008), studies investigating telomere length dynamics in muscle stem cells (MuSCs) are lacking.

MuSCs, also known as satellite cells, are adult stem cells that localize between the sarcolemma and the basal lamina (Campbell and Stull, 2003). In undamaged muscle of adults, MuSCs remain quiescent (Brack and Rando, 2012). However, upon muscle injury a major tissue remodeling process occurs, leading to the activation and proliferation of resident MuSCs (Shi and Garry, 2006). Environmental cues lead to transcriptional activation of pathways inducing proliferation, differentiation, and fusion of differentiated progeny, which will comprise repaired muscle fibers (Wang and Rudnicki, 2011). Many muscle diseases, including muscle dystrophies such as Duchenne muscular dystrophy (DMD), present with multiple rounds of muscle damage and repair (Mann et al., 2011). Over time muscle weakness develops, resulting from a lack of complete regeneration (Wallace and McNally, 2009). A recent hypothesis to explain such an outcome is that the MuSC pool responsible for muscle regeneration gradually becomes less efficient at responding to and repairing damage as a result of stem cell defects (Dumont et al., 2015; Sacco et al., 2010).
However, it has not been studied whether critical telomere shortening in diseased MuSCs contributes to the progressive dysfunction that compromises their regenerative potential, in part due to the inability to quantitatively estimate telomere length in these cells. An optimized technique that is able to measure telomere length in a muscle cell type-specific way would be an invaluable tool to study the involvement of stem cells in the onset and progression of DMD as well as other skeletal muscle diseases.

Many methods exist to measure telomere length, either directly or indirectly (Montpetit et al., 2014). Direct methods such as telomere restriction fragment analysis (TRF) (Kimura et al., 2010) have several inherent shortcomings, including the requirement of a large sample size. Such assays are hindered by the low abundance of MuSCs within skeletal muscles (Morgan and Partridge, 2003). At the same time, human tissue is limited, and methods for propagating undifferentiated MuSCs in sufficient numbers to conduct TRF do not yet exist. While qPCR-based methods (O’Callaghan and Fenech, 2011) do not require as much starting material to perform, such assays can only measure the mean telomere length within the entire population of cells. Over the last few years it became widely accepted that MuSCs are a heterogeneous population (Ono et al., 2010; Tierney and Sacco, 2016). Therefore, when traditional qPCR-based methods are used to measure telomere length, information regarding individual cells within the population cannot be easily attained. Analyzing the expression or activity of TERT/TERC (Skvortsov et al., 2010) or using TRAP (telomerase repeated amplification protocol) activity assays (Fajkus, 2006) does not provide telomere information in a single-cell resolution context but rather in whole tissues. A robust technique for telomere length analysis is fluorescence in situ hybridization (FISH), in which a fluorescently labeled probe that is complementary to the telomere sequence is hybridized and fluorescence intensity is measured by immunofluorescence microscopy (Hande et al., 1999; Henderson et al., 1996). Historically, cells are treated with microtubule inhibitors to promote metaphase arrest, after which metaphase spreads are stained and analyzed (Hande et al., 1999). The caveats to such an approach are that only cultures with a high mitotic index could be analyzed (Ourlac-Garnier and Londono-Vallejo, 2011), and a large number of cells are needed (Howe et al., 2014), making this method challenging for quiescent stem cells directly isolated from a tissue. Assessment of telomere length in interphase cells using in situ hybridization and digital fluorescence microscopy on tissue sections (de Pauw et al., 1998; O’Sullivan et al., 2004) is challenging due to the fact that MuSCs are sparse within skeletal muscles. In fact, our own attempts to measure telomere length by adapting FISH analysis on skeletal muscle sections coupled with the MuSC-specific marker, Pax7, was abandoned due to the sparse number of MuSCs per section. Flow-FISH is a version of FISH that can be combined with the detection of cell surface markers and examines mean values of telomere length in populations of non-dividing cells in interphase (Hultdin et al., 1998). However, most available flow-FISH protocols focus on analysis of hematopoietic cells and cultured cells (Baerlocher et al., 2006), and its use with other tissues and cell types has not been reported. A second drawback of flow-FISH is that it only provides mean telomere length values per cell, in contrast to the quantification of individual telomere foci signals in the case of microscopy-based methods (Meeker et al., 2002). Importantly, in our experience, the heat denaturation step, an absolute requirement of this method (Lauzon et al., 2000), compromises the cell surface antigens used for MuSC isolation.

Although all previous methods have been utilized for a variety of tissues and cells, we desired a better quantitative microscopy-based method that will be applicable for use with the rare population of stem cells present in skeletal muscle tissues. Here, we describe a protocol, designated henceforth MuQ-FISH (muscle quantitative fluorescence in situ hybridization), which first enriches MuSCs prior to the application of a modified FISH method that is optimized for MuSCs of mouse (mMuSCs) and human (hMuSCs) origin. The development of MuQ-FISH allows for quantification of both telomere length and number in prospectively isolated MuSCs, regardless of species origin or disease state. MuQ-FISH has many differences compared with other FISH methods, including MuSC isolation and processing, telomere staining, and analysis. After validating the MuQ-FISH method using mice lacking the RNA component of telomerase (TERT/TR), we showed that there were no telomere length differences in MuSCs isolated from uninjured young and old mice. Additionally, telomere shortening became evident in MuSCs isolated from dystrophic mice. Interestingly, telomere shortening was also discovered in hMuSCs from DMD patients compared with healthy aged-matched human cohorts. Thus, the method described herein could become a standard tool to measure telomere length in other stem cell diseases and disorders, especially muscle-based diseases, and can potentially be expanded for use in other resident muscle cell types.

**RESULTS**

**Validation of MuQ-FISH**

The effect of telomere length on maintaining tissue homeostasis and stem cell function has been studied in many tissues (Flores and Blasco, 2010; Flores et al., 2008; Morrison et al., 1996; Sekulovic et al., 2011; Shekhani
et al., 2016). However, in the skeletal muscle field, existing methods are not well suited for stem cell analysis due to technical and biological limitations of the muscles (rare population, fragile cells, no MuSC expansion, and small tissue sample availability). Notably, several of the steps used in other FISH protocols that work well for other tissue stem cells (Flores et al., 2008) are not favorable for MuSC staining. To measure telomere length in a MuSC-specific way and avoid previous limitations, we developed MuQ-FISH. More specifically, we first isolated MuSCs from uninjured wild-type mice by fluorescence-activated cell sorting (FACS) (Figure 1A) based on cell surface markers, as described by Sacco et al. (2008). FACS was followed by a short plating (8–10 hr) and fixation with 4% paraformaldehyde (Figure S1A), prior to staining for MuQ-FISH (Figure 1B). In our experience, the plating of cells prior to MuQ-FISH does not alter key criteria, including the differentiation state or viability of MuSCs (Figures S1B–S1F). Fixed cells were then stained using a Cy3-conjugated telomere sequence-specific probe labeled at the C terminus, followed by nuclear staining with DAPI. It was critically important that during image acquisition for each experiment, the exposure settings were kept the same for each color channel and that the signal intensities were not too low or too high (Figure S2). For mMuSCs, z stacks were taken for each color channel and combined and flattened using an
extended depth of focus/field (EDF) algorithm (Figure 1B). EDF is a digital method of processing a series of images (optical sections) focused at a range of depths. This EDF method results in a single image, with all areas in focus (Hovden et al., 2011). We found that the use of EDF processing in MuQ-FISH considerably improves the image quality and increases the accuracy of telomere foci measurements. Images were scaled to 16 bit and monochrome images for each channel were opened in the telomere analysis software, Telometer, and processed to subtract background noise. DAPI-stained nuclei were manually encompassed, and this region was used to determine a region of interest, in which telomere signal intensities were measured (Figure 1C). Telometer then compiles data from intensity measurements, which can be used for different analyses. Notably, the software is capable of discerning telomere length of cells in different stages of the cell cycle and normalizing the data, based on DNA content (Figure S3). The protocol for measuring telomere intensities is identical for hMuSC analysis (Figure 1C), with the exception that z stacks are not required for analysis due to the more planar morphology of hMuSCs in culture.

Data obtained from the telomere software of MuQ-FISH can be analyzed in different ways. Common approaches include measuring the total signal intensities of the telomere reads and dividing that value by the total signal intensity of the DAPI (referred to as sum intensity) or identifying the mean intensity of the telomere signal divided by the total signal intensity of the DAPI (referred to as mean intensity). To delineate rare populations of cell data that may be buried by the average telomere analysis described, MuQ-FISH data can also be broken down and plotted as histograms, using the same datasets for sum intensity or mean intensity. Finally, information regarding the number of telomeres, as defined by the individual signal reads, can be obtained, which can also be depicted as an average or as a histogram revealing the telomere length distribution within each cell population of a given genotype or condition.

Using our newly optimized methodology, we validated our approach by FACS-isolating MuSCs from wild-type mice, and from mTRG1 and mTRG3 mice, the latter of which should globally present with shorter telomere length (Blasco et al., 1997). After collecting similar numbers of MuSCs from all groups and processing cells for MuQ-FISH (Figures 2A and 2B), we interrogated the telomere length and number of telomere foci in all groups. Applying the analysis methods described above, we found significant shortening of telomere length when either the sum or the mean intensity of the telomere signal was measured between the different genotypes (Figures 2C and 2D). To examine differences in the number of foci with detectable telomere repeat tracks in mMuSCs, we queried the number of telomere foci per nucleus and found significantly fewer telomere signals only in the mTRG3-derived MuSCs (Figure 2E), in accordance with lack of detectable telomere repeats in late mTR−/− generations (Blasco et al., 1997). This analysis provides a biological validation of the MuQ-FISH method as a protocol that measures telomere length and number of telomere foci in stem cells derived from skeletal muscles.

**MuQ-FISH Analysis of Young and Old MuSCs**

Telomere shortening is reported to be a hallmark of aging in some organisms (Blackburn et al., 2015; Harley et al., 1990; Wong et al., 2003). To identify the effects of aging on the telomere length and number in MuSCs, young (~2 months old) and old (~25 months old) wild-type MuSCs were FACS-sorted from mice in the absence of any induced injury and processed for MuQ-FISH (Figures 3A and 3B). This analysis revealed no major differences in either the telomere length or the number of telomere foci between MuSCs derived from young and old mice (Figures 3C and 3D). To further assess the data, we presented telomere length values as histograms to delineate any rare cell populations that were lost when total data were analyzed. Using this approach, no significant shortening of telomeres was observed during aging (Figures 3C and 3D), suggesting that in the absence of muscle injury, telomere length is stable in aging murine MuSCs.

**MuSCs Isolated from a Dystrophic Mouse Model with a Severe Phenotype Have Shortened Telomeres**

In hematopoietic and lung disease states, telomere attrition in stem cells affects their regenerative capacity (Alder et al., 2015; Armanios, 2013). To query telomere length of MuSCs derived from a skeletal muscle disease condition, we examined telomere differences in young (3–4 months old) control and aged-matched dystrophic mice. Wild-type, mdx (carrying a dystrophin mutation), mTRG2, and mdx/mTRG2 (combined dystrophin and TERC mutation) mice were first queried for differences in cell number during FACS isolation. mdx/mTRG2 mice exhibit a more severe muscle phenotype than mdx/mTRG1 mice or mdx alone (Mourkioti et al., 2013), while mTRG3 mice do not exhibit muscle defects under non-injury conditions (Sacco et al., 2010). While similar numbers of MuSCs were isolated between wild-type, mdx, and mTRG2 mice (Figure 4A), a comparison of mTRG2 mice with mdx/mTRG2 mice did show a decrease in MuSC numbers in the dystrophic mice, consistent with the reported difference in stem cell reserves available for regeneration during the progression of the dystrophic disease (Sacco et al., 2010).

We then processed isolated mMuSCs from wild-type, mdx, mTRG2, and mdx/mTRG2 mice for MuQ-FISH (Figure 4B). We showed variable shortening of telomeres
between these MuSCs by plotting sum intensity values, with longer telomeres found in the wild-type MuSCs, with the mdx/mTRG2 MuSCs displaying shorter telomeres (wild-type > mdx > mTRG2 > mdx/mTRG2; Figure 4C). Furthermore, the distribution of telomere length (histogram) shows a more profound shift to shorter telomere length of the mdx/mTRG2 MuSC population (red in Figures 4C and S4A), suggesting that the telomere shortening observed in mdx/mTRG2 MuSCs is not solely the result of the mTRG2 genetic manipulation, but rather seems to be an additive effect of the combination of mdx and mTRG2 in these cells. Similar reductions in telomere length were found using confocal microscopy (Figure S4B). Further characterization of mMuSC telomere dynamics was carried out, whereby numbers of foci were examined between diseased and control mMuSCs. Interestingly, the number of telomere foci was very dynamic between all groups examined (Figure 4D). Overall, these data demonstrate that MuSCs from a murine model with severe muscular dystrophy exhibit extensive telomere shortening.

**Figure 2. Comparative Analysis of Telomere Length and Number in Different Generations of TERC/TR Knockout Mouse-Derived MuSCs**

(A) Number of MuSCs isolated from 20,000 total events. Data displayed are the percentage of CD34+/α7-integrin+ MuSCs that comprised the total number of cells in the parent gate. Data are displayed as the mean ± SEM from n = 2–3 female mice per group of similar age.

(B) Representative images of cells processed for MuQ-FISH. Scale bar, 5 μm.

(C) Left: sum intensity of telomere signals divided by DAPI sum intensity. n ≥ 90 cells were analyzed per mouse, with n = 2–3 mice per category. Bars represent mean ± SEM. *p ≤ 0.05, **p ≤ 0.01. Right: histogram depiction of sum intensity measurements (mean ± SEM).

(D) Left: data representation of telomere mean intensity divided by DAPI sum intensity. Right: histogram representation of data in left panel. *p ≤ 0.05, **p ≤ 0.01.

(E) Left: relative number of telomere foci in wild-type, mTRG1, and mTRG3 mice. Right: histogram representation of number of telomere foci. *p ≤ 0.05. WT, wild-type.

WT, wild-type.
Dystrophic hMuSCs Exhibit Shortened Telomeres

To determine whether the telomere length observations in diseased mMuSCs recapitulate what happens in the human dystrophic condition, we collected hMuSCs from the biopsies of three healthy and three DMD-diseased teenage individuals (Figure 5A), and FACS isolation revealed that adequate hMuSC numbers were procured (Figure 5B). Following a short culture that allowed attachment of cells on slides (Figures 5C and S5), hMuSCs were processed for MuQ-FISH (Figure 6A) and telomere lengths and number were calculated. This analysis revealed that telomere length was significantly shorter in DMD-diseased hMuSCs compared with healthy MuSCs, with a clear shift toward shorter telomeres in the distribution representation (Figure 6B) without a significant loss in telomere foci (Figure 6C). These data support the notion that accelerated telomere shortening in the setting of muscular dystrophy is also a characteristic of hMuSCs, extending this finding to the human disease.

To address the specificity of hMuSC-telomere shortening, we conducted MuQ-FISH on patient muscle biopsies, followed by staining for the common leukocyte antigen, CD45 (Thomas and Lefrancois, 1988) as well as for α-smooth muscle actin, which labels smooth muscle cells (Skalli et al., 1987). In both cases telomere lengths, as assayed by MuQ-FISH, are not significantly different between healthy and DMD patient samples that stained positively for these markers (Figures S6 and S7). This result argues against ubiquitous telomere shortening in other skeletal muscle-derived cells other than MuSCs and suggests that the telomere shortening, which we observed in DMD patient-derived MuSCs, is likely the result of the disease process.

DISCUSSION

Telomere shortening can occur in actively proliferating cells, cells with aberrant nuclease activity, or cells experiencing

**Figure 3. Comparative Analysis of Telomere Length and Number in Young and Old Murine MuSCs by MuQ-FISH**

(A) Number of MuSCs isolated from 50,000 total events. Data are displayed as the mean ± SEM from n = 3 male mice per group of identical age.

(B) Representative images of cells processed for MuQ-FISH. Scale bar, 5 μm.

(C) Left: sum intensity of telomere signals divided by DAPI sum intensity. n ≥ 90 cells were analyzed per mouse, with n = 3 mice per category. Bars represent mean ± SEM. Right: histogram depiction of sum intensity measurements.

(D) Left: relative number of telomere foci in wild-type young and old mice. Right: histogram representation of number of telomere foci.
Figure 4. Muscle Stem Cells from Dystrophic Mice Exhibit Significant Telomere Shortening Compared with Control Mice

(A) FACS analysis of the number of mMuSCs isolated from mice of different genotypes, represented as percentage of the parent gate. Data are displayed as mean ± SEM. *p ≤ 0.05.

(B) Representative images of MuQ-FISH from mMuSCs of different genotypes. Scale bar, 5 μm.

(C) Left: plotted data of telomere sum intensity calculation. Bars represent mean ± SEM. *p ≤ 0.05, **p ≤ 0.01. Right: data plotted as histograms. Note the shift toward the shortest telomeres in the mdx/mTRG2 MuSCs. n = 3 mice per genotype and n ≥ 70 cells per mouse. See also Figure S4.

(D) Left: dot plot showing number of MuSC telomere foci between genotypes. Bars represent mean ± SEM. *p ≤ 0.05, **p ≤ 0.01. Right: histogram depiction of data shown on the left.
DNA damage, particularly in those lacking functional telomerase (d’Adda di Fagagna et al., 2003; Herbert et al., 1999). Significant telomere attrition leads to the activation of DNA damage signaling pathways (Viscardi et al., 2007), which induces cell cycle arrest, senescence, and/or cell death. Other consequences of telomere shortening include the alteration of gene expression (Ye et al., 2014), induction of epigenetic modifications and promoter methylation (Blasco, 2007a) and altered gene silencing near the telomere (Guan et al., 2013), as well as the alteration of gene expression megabases away from telomeres (Robin et al., 2014). These effects, either individually or in combination, contribute to stem cell dysfunction.

Stem cell functions decrease in many tissue types during aging, which often correlates with telomere shortening (reviewed in Sharpless and DePinho, 2007). Interestingly, short telomere lengths limit stem cell numbers and function in hematopoietic cells (Armanios, 2013), while short telomeres can contribute to stem cell failure in the lung (Alder et al., 2015). Moreover, telomere shortening is suggested to act as a biological clock that counts cell divisions to regulate the number of self-renewing divisions within each differentiation state in hematopoietic cells (Morrison et al., 1996). In the skeletal muscle system, the progressive loss of the mMuSC reserve plays a major role in determining the severity of the dystrophic phenotype (Dumont et al., 2015; Sacco et al., 2010). However, due to the lack of a proper method to measure telomere length in MuSCs, all previous analyses were not performed in MuSCs but rather in their differentiated myoblast derivatives (Ramunas et al., 2015; Sacco et al., 2010) or in whole muscle extracts (Decary et al., 2000). The technique described here is specific for the skeletal muscles and has many differences among previously reported FISH methods in other tissues, including modifications of the probe employed, the cell isolation and fixation method, the optimized staining procedure, and the incorporation of telomere software for unbiased analysis. MuQ-FISH allows for telomere foci number and length assessment of individual cells within the population and is applicable to both muscle cells cultured shortly in vitro and/or freshly isolated from in vivo sources. More specifically, MuQ-FISH has several advantages compared with other methods: (1) the analysis of telomere length can occur at the single-cell level, such that data from scarce populations are not diluted and lost; (2) no radioactive compounds are required; (3) the assay is optimal for analyzing cells where only a small amount of material can be obtained, as is generally the case for cells derived from human biopsies; (4) this protocol uses common and/or inexpensive equipment to complete; and (5) analysis of both proliferating and non-proliferating or quiescent cells is possible. Thus, factors influencing proliferation and/or cell homeostasis and their effects on telomere attrition can all be tested including, but not limited to, genetic manipulation, aging, and culture conditions such as oxygen tension, substrate rigidity, and the alteration of growth factors.

The current study established a new approach in a translationally relevant biological system, by measuring...
telomere length in MuSCs derived from young and old mice as well as in healthy and dystrophic mMuSCs and hMuSCs. The rationale for such a study is that in many stem cell systems, telomere shortening often occurs during aging (Armanios et al., 2009), as well as in many degenerative diseases (Calado and Young, 2009; Harley, 2005). Our analysis revealed that telomere length of mMuSCs deriving from old wild-type mice did not shorten when compared with young wild-type mMuSCs. Although telomere shortening is major determinant of aging in other systems (Blackburn et al., 2015; Harley et al., 1990; Wong et al., 2003), our finding of stable MuSC telomere length is in agreement with other reports measuring telomere length in whole muscle fibers from young and aged mice (O’Connor et al., 2009) as well as telomere length findings in skeletal muscles of non-human primates (Gardner et al., 2007).

Figure 6. MuQ-FISH Analysis Reveals Significantly Shortened Telomeres in hMuSCs Isolated from DMD Patients Compared with Age-Matched Healthy Individuals

(A) Representative images of telomere MuQ-FISH in healthy and dystrophic hMuSCs. Scale bar, 10 μm.

(B) Quantification of the average telomere length per nucleus (left) and the sum intensity distribution presented as a histogram (right). Note the clear shift toward shorter telomere length in the diseased population. n = 3 human biopsies per group and n ≥ 100 cells per patient were analyzed. Bar represents mean ± SEM. *p ≤ 0.05. See also Figures S6 and S7.

(C) Left: measurements of relative number of telomere foci in healthy and dystrophic hMuSCs. Bars represent mean ± SEM. Right: distribution of number of telomere foci shown as histograms.
dynamics in age-matched MuSCs derived from dystrophic mice undergoing chronic regeneration, as well as in genotype control MuSCs. Although mTRG2 mice have no reported skeletal muscle phenotype, their MuSCs exhibited significantly shorter telomeres compared with wild-type controls. This result validates the MuQ-FISH method and further demonstrates the sensitivity of the assay in evaluating premature telomere length in a stem cell-specific way. Intriguingly, we show that telomere shortening is present in mdx/mTRG2 MuSCs and to a lesser extent in mdx MuSCs. The order of telomere shortening in these groups correlates with the severe penetrance of the dystrophic phenotype observed in mdx/mTRG2 compared with mdx mice (Sacco et al., 2010). These findings support the notion that telomere shortening contributes to the stem cell dysfunction and impaired regeneration observed in muscular dystrophy. Our results of dissimilar numbers of foci between dystrophic and wild-type MuSCs are important to consider in light of previous studies that show an increased percentage of chromosomes with telomere signal free ends in dystrophic myoblasts (Sacco et al., 2010). In contrast to the extensively cultured myoblasts utilized in the previous study, MuQ-FISH is able to detect telomere foci in MuSCs soon after their isolation from dystrophic muscles, which may reconcile the signal-intensity differences observed between the two designs.

Although much is known about MuSCs and the tissue-regenerative abilities in these murine models, very little information has been obtained regarding humans. Current findings of regeneration processes in human muscle diseases are limited, due to the fact that hMuSC analysis has lagged behind. Using MuQ-FISH, we have successfully isolated dystrophic and aged-matched healthy MuSCs from human muscle biopsies and demonstrated premature telomere shortening in MuSCs DMD patients, while the numbers of telomere foci were not different, consistent with our mouse findings. Extension of MuQ-FISH to other resident cell types in human muscle sections demonstrated that telomere attrition occurs in MuSCs but that it is not an inherent process in non-skeletal muscle cell types. Thus, the premature telomere shortening observed in DMD implies an intrinsically limited functional potential of human dystrophic MuSCs, and the expansion of dystrophic MuSCs as a therapeutic strategy may have a limited window of effectivness before critical telomere shortening triggers replicative senescence and regenerative failure in DMD muscles.

In summary, we provide an optimized and validated assay to detect telomere length in stem cells isolated from skeletal muscles of mice and humans. Analyzing the role of telomere length in hMuSCs using the MuQ-FISH method described here will allow for a better understanding of the molecular genetic events that, in relation to telomere dysfunction, may lead to the delineation of muscle regeneration limitations in different human skeletal MuSC disease etiologies. Importantly, utilizing MuQ-FISH will allow for further investigation of the developmental and self-renewal processes and pathways involved in stem cell dysfunction in various skeletal muscle disorders. Diseases of particular interest include other muscular dystrophies, muscle atrophy, sarcopenia, myosarcomas, and chronic muscle injuries. Moreover, using this method we can extend the analysis to other cells isolated from the same tissues, with particular interest in cell types that influence MuSC function and muscle regeneration, such as fibroadipogenic progenitor cells (Joe et al., 2010). A better understanding of the molecular players and the exact mechanisms that regulate stem cell function, telomere length, and skeletal muscle homeostasis in humans should constitute an important advance in the field of regenerative medicine.

EXPERIMENTAL PROCEDURES

Mice
Mice were housed and bred in accordance with Institutional Animal Care and Use Committee guidelines outlined by the University of Pennsylvania. Additional information can be found in Supplemental Experimental Procedures.

Human Muscle Biopsy Procurement
Human muscle biopsies from healthy and DMD patients were obtained from the lower extremity muscles during surgical procedures as part of the patient’s clinical care plan at Rady’s Children’s Hospital, San Diego. Written informed consent from the parent or guardian was obtained for all subjects. The protocol was approved by the University of California, San Diego Human Research Protections Program and Institutional Review Board in accordance with the requirements of the Code of Federal Regulations on the Protection of Human Subjects. Three healthy and three DMD-affected biopsies were collected for this study.

Murine-Derived MuSC Quantitative Fluorescence In Situ Hybridization
After sorting, a minimum of 2,000 mMuSCs were plated onto laminin-coated (Sigma-Aldrich) 8-well chamber slides (Nunc; Labtek II, Thermo) in 50 μL of myoblast medium (Figure S1) and left at room temperature overnight, uncovered to dry onto the slides. This step results in the retention of more MuSCs on the slide following the staining procedure. The next day, cells were fixed in 4% paraformaldehyde/PBS for 15 min and rinsed once with PBS. Cells were permeabilized with 0.1% Tween 20/PBS for 5 min and washed with PBS twice for 3 min each. Cells were treated with 100 μg/mL pre-boiled RNase A for 20 min at 37 °C, followed by three PBS washes of 5 min each. Slides were dried. The Cy3-conjugated PNA telomere probe (1/300 dilution of 50 μM stock of TelC probe; F1002; PNAbio; 5’-CCC TAA CCC TAA CCC TAA-3’) was prepared in MuQ-FISH buffer (60% formamide, 5%
of 10× blocking agent for nucleic acid hybridization [not diluted to 1×; Roche], 2% 1 M Tris [pH 7.5], all in water). The probe and slide were preheated to 86°C for 10 min before the probe was added to the slides. After probe addition, slides were covered and incubated for an additional 10 min at 86°C, prior to cooling overnight at room temperature in the dark. The following day, slides were washed twice in a stepwise fashion with pre-warmed (55°C) 2× saline sodium citrate (SSC)/0.1% Tween 20, 1× SSC/0.1% Tween 20, and 0.5× SSC/0.1% Tween 20. Chamber slides were rinsed with PBS, disassembled, and coverslips were mounted with Fluoromount G with DAPI (SouthernBiotech).

Human-Derived MuSC Quantitative Fluorescence In Situ Hybridization

hMuSCs were cultured in DMEM-F12 with 15% fetal bovine serum and 1× antibiotic-antimycotic. For human MuQ-FISH staining, hMuSCs were plated and grown overnight in a 37°C/5% CO2 humidified incubator on collagen-coated 8-well chamber slides (Nunc; Labtek II, Thermo) and processed as stated in the mMuSC protocol.

Image Acquisition

Images of mMuSCs were taken using a Nikon eclipse 90i wide-field epifluorescence microscope equipped with a Prior Proscan III motorized stage, a Photometrics CoolSnap HQ2 14-bit digital camera, and a Nikon 100×/1.40 Plan Apo VC objective. Five to eight z stacks, which corresponded to the lower and upper boundaries of DAPI signal per nucleus, were taken for both the DAPI and Cy3 channels, at a 1-μm step. Samples are then imaged to determine each channel’s exposure time in a way that situates the intensities for all samples in the mid-intensity range (Figure S3). This control prevents overexposure or loss of signal detection with the selected exposure times during sample imaging (Figure S3A). Once an optimal exposure time is defined (Figure S3B), image acquisition utilizes the same exposure settings for the Cy3 (telomere) and DAPI signals for imaging of individual experiments. For each experiment, the intensity settings are not changed between sample groups. Images are taken without binning. Z stacks are combined using the extended depth of focus/field (EDF) option in the Nikon elements software. Images of hMuSCs were taken with a Nikon eclipse Ni-U wide-field epifluorescence microscope equipped with a Nikon Q1M1c 14-bit camera and a Nikon 100×/1.24 Plan Apo objective. A minimum of 30 cells per sample set is required for analysis; however, the more cells imaged, the more accurate the resultant data are representative of the source. Regardless, imaging must take place in the same time window for each experiment, and combination of multiple experiments performed on different days is not recommended, due to the inherent caveat of fluorescent intensity measurements differing between experiments.

Telomere Analysis

Telomeres were analyzed with the investigators blinded to genotypes and/or conditions using open-source software (Telometer; http://demarzolab.pathology.jhmi.edu/telometer/), as previously described (Meeker et al., 2002; Mourkioti et al., 2013). Other telomere-measuring software could also be used for such analysis, including but not limited to Telomap (Flores et al., 2008), TFL-Telo (Poon and Lansdorf, 2001), Tissue-Telo (Aida, 2014), TeloView (Chuang et al., 2004) and the commercially available program, Isis (MetaSystems). All pictures were converted to 16-bit individual channel monochrome images. Channels were normalized to remove background and the DAPI-positive region was manually encircled. Cy3 signals within this region are subjected to a rolling ball algorithm (mask), and halos and conjoined particles are manually eliminated/separated by the investigator. Notably, this type of editing should only be used for high-resolution images. For images captured at a lower magnification, this editing should not be used due to the small size of foci compared with the thickness of the eraser, as attempts to separate conjoined foci will result in severe signal losses. The program generates statistics on the entire region of the nucleus. Statistics returned include the intensity sum of all Cy3 telomere pixels for a given nucleus (proportional to the cell’s total telomere length) and the intensity sum of all DAPI pixels for the nucleus (proportional to total cellular nuclear DNA content). Additional measurements taking into account mean intensities of both telomere signals and DAPI signals are also generated. By taking the ratio of the telomere intensity measurements to the corresponding DAPI intensity measurements, one is able to compensate for ploidy differences.

Statistical Analysis

Values are presented as mean ± SEM. Analyses were subjected to unpaired Student’s t tests with Welch’s correction using GraphPad Prism 6 software. Significance is indicated in the figures by *p ≤ 0.05 and **p ≤ 0.01. Additional information can be found in Supplemental Experimental Procedures.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, seven figures, and five tables and can be found with this article online at http://dx.doi.org/10.1016/j.stemcr.2017.08.003.

AUTHOR CONTRIBUTIONS

E.D.T. designed and performed experiments, analyzed and interpreted data, and drafted the manuscript. D.K.S. performed experiments, analyzed data, and edited the manuscript. M.T.T. carried out hMuSC isolation, analyzed data, and edited the manuscript. M.J.S. carried out mMuSC isolation and edited the manuscript. M.S.-S. performed experiments and analyzed the data. H.H and S.M. procured human biopsies. F.B.J. provided mTRG3 mice and edited the manuscript. A.S. designed and supervised hMuSC experiments, interpreted data, and drafted the manuscript. M.T.T. carried out hMuSC isolation, analyzed data, and edited the manuscript. E.D.T. designed and performed experiments, interpreted and inter-
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