The age of heterozygous telomerase mutant parents influences the adult phenotype of their offspring irrespective of genotype in zebrafish [version 2; peer review: 2 approved]

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Abstract

Background: Mutations in proteins involved in telomere maintenance lead to a range of human diseases, including dyskeratosis congenita, idiopathic pulmonary fibrosis and cancer. Telomerase functions to add telomeric repeats back onto the ends of chromosomes, however non-canonical roles of components of telomerase have recently been suggested.

Methods: Here we use a zebrafish telomerase mutant which harbours a nonsense mutation in tert to investigate the adult phenotypes of fish derived from heterozygous parents of different ages. Furthermore we use whole genome sequencing data to estimate average telomere lengths.

Results: We show that homozygous offspring from older heterozygotes exhibit signs of body wasting at a younger age than those of younger parents, and that offspring of older heterozygous parents weigh less irrespective of genotype. We also demonstrate that tert homozygous mutant fish have a male sex bias, and that clutches from older parents also have a male sex bias in the heterozygous and wild-type populations. Telomere length analysis reveals that the telomeres of younger heterozygous parents are shorter than those of older heterozygous parents.

Conclusions: These data indicate that the phenotypes observed in offspring from older parents cannot be explained by telomere length. Instead we propose that Tert functions outside of telomere length maintenance in an age-dependent manner to influence the adult phenotypes of the next generation.

Keywords
telomerase, telomere length, zebrafish, age, sex, Tert

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Amendments from Version 1

We thank the reviewers for their insightful comments and suggestions. Following this we have made the following amendments to the manuscript:

1) We have added a sentence with references to the results to clarify that measuring telomere lengths using whole genome sequencing is an established methodology.

2) We have added a section to the discussion which details that sampling from a distribution of telomere lengths by descent from two different parents may lead to overall longer telomeres in one line of descendants compared to another.

3) We have removed an erroneous reference to DNA damage from the discussion.

4) We have changed the labelling in Figure 1F to clarify that the phenotypic embryo is maternal-zygotic mutant offspring from homozygous mutant parents.

5) We have added a supplementary figure and text in the results with new data on egg quality and quantity from heterozygous tert intercrosses. The raw data have been deposited on figshare.

6) We emphasise in the methods section that only non-phenotypic embryos with filled swim bladders were raised to adulthood for this study.

7) We have also added two new authors: Neha Wali: NW conducted the experiments for the revision. Derek L. Stemple: Due to initial delay of the authorship confirmation response DLS was removed from the authors list in the first submission. We have now received confirmation. All authors have agreed to the updated author list of this version 2.

See referee reports

Introduction

Telomeres are composed of non-coding repetitive sequences at the termini of each chromosome, protected by a complex of proteins called the shelterin complex. With each cell division the telomeres shorten due to the inability of DNA polymerases to replicate the ends of linear DNA, a phenomenon known as the ‘end replication problem’ (reviewed in 1). To circumvent this problem certain cell types express the ribonucleoprotein Telomerase, which adds telomeric repeats back onto the ends of chromosomes (reviewed in 2). When telomeres lose their protective cap, the linear DNA ends are recognised by DNA damage response proteins resulting in telomere fusions and ultimately leading to replicative senescence and cell death.

Telomerase is composed of an RNA molecule (hTR in humans, tert in zebrafish) containing a complementary sequence to the telomeric repeat hexamer which serves as a template for DNA synthesis and a reverse transcriptase enzyme (TERT in humans, Tert in zebrafish) which harbours an RNA binding domain and a reverse transcriptase domain.

Mutations in components of telomerase or other telomere-associated proteins lead to a class of human diseases collectively known as telomeropathies (reviewed in 14,15). These include dyskeratosis congenita (DC), idiopathic pulmonary fibrosis (IPF) and cancer. Telomerase is not expressed in most somatic cells but its expression is activated in germline tissue and tumour cells enabling these cell populations to divide potentially indefinitely, and thus activation of telomerase is a hallmark of many cancers. DC results from germline mutation in a number of telomerase-associated genes including DKC, TINF2, TERC and TERT, and its clinical features combine IPF, cancer predisposition, skin abnormalities and bone marrow failure. Interestingly, patients with autosomal dominant DC show disease anticipation, due to haploinsufficiency of either TERT or TERC, whereby the age of onset and severity of symptoms worsens in successive generations.

Mouse models were traditionally used to study telomere dysfunction, however laboratory mouse strains have very long telomeres and consequently mTR and mTERT homozygous mutants can be incrossed for several generations before phenotypes become apparent, although in the case of mTERT null mutants, genetic background contributes significantly to the generation in which phenotypes arise. More recently zebrafish models have been used to interrogate the function of telomerase in development and disease. Fish homozygous for the null mutation ter have shorter telomeres and show premature lethality and tissue degeneration. In addition, a p53-mediated reduction in cell proliferation and increased apoptosis are observed in these mutants.

Here we have characterised a new nonsense mutation in the zebrafish tert gene, tert, and show that the age of heterozygous parents influenced the adult phenotypes of their offspring. Offspring of all genotypes from older parents weighed less and showed a male sex bias. We used whole genome sequencing to estimate average telomere lengths and surprisingly found that the phenotypes observed were independent of telomere length. This highlights the need for further investigation into the additional roles of telomerase aside from maintaining telomere length.

Methods

Husbandry

This study was conducted in line with the ARRIVE guidelines. Zebrafish were maintained in accordance with UK Home Office regulations, UK Animals (Scientific Procedures) Act 1986, under project licence 70/7606, which was reviewed by the Wellcome Trust Sanger Institute Ethical Review Committee. The mutant alleles tert and tert were obtained from the Zebrafish Mutation Project and recovered from frozen sperm samples at the Wellcome Trust Sanger Institute. Embryos were produced...
through natural matings and maintained in an incubator at 28.5°C up to 5 days post fertilisation (d.p.f.). All efforts were made to ameliorate any suffering: fish underwent careful health checks twice a day and any fish that showed signs of distress, such as lethargy, failure to feed or abnormal swimming behaviour, were culled following the standard Schedule 1 method of anaesthetic overdose with subsequent destruction of the brain. Only morphologically normal larvae with inflated swim bladders at 5 d.p.f. entered the nursery to be raised to adulthood for this study.

Genotyping
DNA from embryos or fin biopsies was extracted and genotyped for tert<sup>sa6541</sup> or tert<sup>hu3430</sup> using KASP genotyping as previously described<sup>33</sup>.

Sperm extraction
Males were anaesthetised and sperm extracted by gentle abdominal massage as described previously<sup>3</sup>.

Weight measurements
Anaesthetised males were patted dry and weighed in a dish of system water on electronic scales.

Telomere length analysis
DNA was extracted from caudal fin clips and used to make Illumina libraries which were sequenced on four lanes of HiSeq X in 151 bp paired-end mode. Sequence data were deposited in ENA under accession ERP016250. FASTQ files were analysed using Computel<sup>36</sup> (v0.3, options: -proc 4 -nchr 25 -lgenome 1371719383 -pattern TTAGGG), FASTQ files were also aligned to the Zv9 reference genome using BWA (v0.7.15, options: mem -t 14 -p -Y -K 10000000) and analysed using TelomereHunter<sup>38</sup> (v1.0.4, options: -d -r TTAGGG -nf).

Results

**tert<sup>sa6541</sup>/+** fish age prematurity
Fish homozygous for the point mutation tert<sup>sa6541</sup> age prematurely, evidenced by spinal curvature, tissue degeneration and premature infertility and lethality<sup>33,34</sup>. We have characterised a new tert allele, tert<sup>sa6541</sup>, produced by the Zebrafish Mutation Project<sup>36</sup>, which carries a point mutation resulting in a premature stop codon and a predicted protein lacking the reverse transcriptase domain and most of the RNA-binding domain (Figure 1A). We compared the phenotype of tert<sup>sa6541</sup> fish to the published phenotype for tert<sup>hu3430</sup> to assess whether tert<sup>sa6541</sup> recapitulates the published zebrafish tert knockout phenotype<sup>13,34</sup>. Fish homozygous for tert<sup>sa6541</sup> show visible signs of ageing earlier than their siblings, including a wasting phenotype (Figure 1B), and homozygotes die prematurely (Figure 1C). As previously shown for tert<sup>hu3430</sup>, tert<sup>sa6541</sup> homozygous males become prematurely infertile; by 9–10 months of age, most (16/17) homozygous males failed to produce any sperm by gentle abdominal massage, whereas all (27/27) wild-type siblings produced sperm (Figure 1D and E). Furthermore maternal zygotic tert<sup>sa6541</sup> fish displayed a variety of embryonic phenotypes (Figure 1F) in accordance with the previously published phenotypes. Intercrosses from heterozygous fish (965 embryos from 3 separate clutches) showed no significant difference (Welch two sample t-test, p-val=0.49) in the number of phenotypic embryos compared to wild-type sibling intercrosses (614 embryos from 4 separate clutches).

To further confirm that the phenotypes observed in tert<sup>sa6541</sup> fish were due to the mutation in tert, we generated compound heterozygous fish using another uncharacterised tert allele. tert<sup>sa25076</sup> carries a point mutation resulting in a premature stop codon and a predicted protein lacking both the reverse transcriptase and RNA binding domains (Figure 1A). Embryos from compound heterozygous intercrosses displayed a range of phenotypes similar to those observed in tert<sup>sa6541</sup> intercrosses (Supplementary Figure S1A) and by 5 days post fertilisation (d.p.f.) compound heterozygous intercrosses had fewer non-phenotypic embryos compared to wild-type siblings (Supplementary Figure S1B).

The age of heterozygous tert<sup>sa6541</sup> parents influences the adult phenotype of their offspring
We observed when intercrossing heterozygous fish over several months (Figure 2A), that lines produced from older parents showed signs, at a younger age, of body wasting reminiscent of fish aged 18 months or older (Figure 2B). Homozygous fish from older parents were less plump (note sunken abdomen in 8 month old homozygote) and paler than older homozygotes derived from earlier matings. This led us to investigate the effect of tert<sup>sa6541</sup> parental age on the phenotype of their offspring.

We raised three tert<sup>sa6541/+</sup> lines; MR2430 and MR2431, which were the same age, and MR2884, which was 12 months younger and obtained from an outcross of MR2430 (see Supplementary Figure S2 for full family tree). We intercrossed MR2431 and MR2884 at 16 and 4 months of age respectively and raised their offspring alongside each other in the nursery to minimise the effect of environmental influences on survival. This revealed that homozygous fish from older parents had reduced survival compared to homozygous fish from younger parents. By contrast, survival of heterozygous and wild-type siblings was unaffected by parental age (Figure 2D). However comparing survival rates across multiple lines raised at different time points showed that this was not a consistent phenotype (Supplementary Figure S3). This may reflect the many different factors that contribute to survival.

Furthermore we observed an effect of heterozygous parental age on the sex ratios of their offspring. Irrespective of the age of the parents, there was a strong male sex bias among tert<sup>sa6541/+</sup> fish (Figure 2E). However as the age of the parents increased, this male sex bias became more pronounced. While 23/93 (24.7%) homozygotes were male in fish conceived at 12 months or younger, only 5/68 (7.4%) homozygous fish conceived at 16 months or older were female. Table 1 provides a list of the sex ratios compared to the expected ratio of 0.5 with significant deviations marked with an asterisk. The mechanisms of sex determination in zebrafish are poorly characterised and
Figure 1. \textit{tert}^{sa6541/sa6541} fish age prematurely. (A) Protein domain structure of zebrafish Tert. Depicted in yellow is the telomerase ribonucleoprotein complex RNA-binding domain (RBD), and in purple the reverse transcriptase domain (RTD). The position of three known alleles are shown. (B) \textit{tert}^{sa6541/sa6541} fish age prematurely. Homozygous fish display a wasting phenotype. Scale bar: 10mm. (C) \textit{tert}^{sa6541/sa6541} fish die prematurely compared to their siblings (n=92 homozygotes, n=92 heterozygotes, n=92 wild-types). (D) Photograph of capillaries containing sperm from wild-type sibling males (left) and clear fluid containing no sperm from \textit{tert}^{sa6541/sa6541} males (right). (E) Homozygous \textit{tert}^{sa6541} fish become prematurely infertile. All wild-type males but only 5.9\% of homozygous males tested, aged 9 or 10 months, produced sperm. (F) Maternal-zygotic mutant embryos derived from \textit{tert}^{sa6541/sa6541} intercrosses display a range of phenotypes by 24 h.p.f. including a reduction in head tissue and a shorter tail.
Figure 2. The age of heterozygous tert<sup>mut</sup> parents affects the phenotype of their offspring. (A) Family tree depicting relationship between lines shown in (B). Lines numbers are indicated according to our fish stock database with date of birth underneath. (B) Homozygous fish from older heterozygous parents show signs of body wasting at a younger age than those from younger heterozygous parents. 8 month old fish from 7 month old heterozygous parents appear older than 10 month old fish from 5 month old heterozygous parents. Scale bar: 10mm. (C) Family tree depicting relationship of lines shown in (D) and (F). (D) Homozygous fish from 16 month old heterozygous parents have reduced survival compared to homozygotes from 4 month old heterozygous parents which were raised at the same time. (E) Sex ratios of intercrosses from tert<sup>mut</sup> parents of different ages. Patterned boxes around line numbers indicate which lines were raised simultaneously in the nursery. Absolute numbers are indicated above each bar. (F) Box plot with scatter of weights of male fish from 4 or 16 month old heterozygous parents showing that fish from older parents weigh less than those from younger parents. (G) Box plot with scatter showing no significant difference between the weights of wild-type fish from wild-type parents aged 5 or 30 months.
are influenced by environmental factors. Nevertheless, the sex bias in tert
homzygous fish is specific to mutation in tert since lines produced from younger parents (12 months and below) have heterozygous and wild-type sibling populations that do not differ significantly from a 0.5 sex ratio with 120/208 (57.7%) and 63/108 (58.3%) females, respectively. This indicates that the male skew in homzygous fish is not a result of environmental conditions during the raising of the line. However, with advanced age of the heterozygous parents, a male sex bias also became apparent in heterozygous and wild-type offspring populations (Figure 2E). In heterozygous and wild-type fish from parents aged 16 months and older the female proportions dropped to 25/128 (19.5%) and 36/98 (36.7%) respectively. These fish were raised alongside lines from younger parents to eliminate environmental influence (indicated as different rectangles in Figure 2E). It is conceivable that heterozygous females have an oogenesis defect with downstream consequences for development. However, we did not find a statistically significant difference in clutch size (Welch two sample t-test, p-val=0.53) or fertilisation rates (Welch two sample t-test, p-val=0.14) between wild-type and heterozygous intercrosses (Supplementary Figure S4A). In addition, we investigated whether there was a general influence of parental age on sex ratios across lines raised in our facility. There was no measurable trend to either sex correlating with the age of either parent (Supplementary Figure S4B). In conclusion these data indicate that the age of heterozygous parents affects sex determination in their offspring.

We observed that tert homozygous fish from older heterozygous parents appeared smaller than those from younger parents when raised alongside each other. Given the sex bias in clutches produced from older parents we quantified the weights of 8 month old adult males from MR3039 and MR3040, which had parents aged 4 months at the time of conception, and 8 month old MR3041 males, which had 16 month old parents at the time of conception (Figure 2C). This revealed that all fish from 16 month old parents, irrespective of genotype, were significantly smaller than those from 4 month old parents (linear model formula: weight ~ genotype * parental age; p-value: <10^{-16}) (Figure 2F). Additionally, tertsa6541/tertsa6541 fish produced from intercrosses of older parents were also significantly smaller than their wild-type siblings (linear model formula: weight ~ genotype * parental age; p-value: 0.0001) (Figure 2F). This indicates that older tertsa6541/tertsa6541 parents produce smaller offspring irrespective of genotype, and that the effect of being homozygous is only significant when these fish have older parents. To assess whether smaller offspring was a general phenotype of older fish, we intercrossed wild-type fish aged 5 and 30 months, but found no significant difference between the weights of their male offspring (Figure 2G) suggesting that the size difference observed between the two tertsa6541 lines is a consequence of the interaction of parental tert heterozygosity with age.

### Phenotypes observed in fish from older heterozygous parents are independent of telomere length
To assess whether the reduced body weight and more pronounced male sex bias observed in adult fish from older heterozygous parents were a result of shorter telomere lengths, we used whole genome sequencing to estimate telomere lengths; a method which has been shown to produce comparable results to other methods of measuring telomere lengths such as Southern blot and qPCR. We extracted DNA from caudal fin clips, performed whole genome sequencing (mean coverage 6x) and determined average telomere length per sample using Computel. We fin clipped 8 fish from each of: an old tertsa6541 heterozygous stock (MR2430) which was 30 months old at the time of clipping; adult offspring of each genotype from MR3310, which were 7 months old at the time of clipping and were the progeny of an intercross of MR2430 at 22 months; a younger tertsa6541 heterozygous stock (MR2884) which was 17 months old at the time of clipping; and adult offspring of each genotype from MR3309, which were 7 months old when clipped and were the result of an intercross of MR2884 at 10 months (Figure 3A). Both intercrosses (MR3309 and MR3310) were raised at the same time to minimise the effect of environment on telomere length.

Telomere length data were analysed using one-way ANOVA, followed by Tukey’s test. For both parent-offspring families,

**Table 1. Sex ratios of fish from old or young heterozygous parents.** The sex ratio of fish with each genotype (GT) from young parents aged 12 months or younger, and old parents aged 16 months or older was compared to 0.5 using a Chi-squared test. The p-values were adjusted for multiple testing using Bonferroni correction. (*) next to the adjusted p-value indicates significance at p<0.05.

<table>
<thead>
<tr>
<th>Parental age at conception (months)</th>
<th>GT</th>
<th># fish</th>
<th># females</th>
<th>% female</th>
<th>p-val</th>
<th>Adj. p-val</th>
</tr>
</thead>
<tbody>
<tr>
<td>12 or younger</td>
<td>wt</td>
<td>108</td>
<td>63</td>
<td>58.3</td>
<td>0.083</td>
<td>0.5</td>
</tr>
<tr>
<td></td>
<td>het</td>
<td>208</td>
<td>120</td>
<td>57.7</td>
<td>0.027</td>
<td>0.159</td>
</tr>
<tr>
<td></td>
<td>hom</td>
<td>93</td>
<td>23</td>
<td>24.7</td>
<td>1.10E-06</td>
<td>6.57E-06*</td>
</tr>
<tr>
<td>16 or older</td>
<td>wt</td>
<td>98</td>
<td>36</td>
<td>36.7</td>
<td>0.009</td>
<td>0.052</td>
</tr>
<tr>
<td></td>
<td>het</td>
<td>128</td>
<td>25</td>
<td>19.5</td>
<td>5.41E-12</td>
<td>3.25E-11*</td>
</tr>
<tr>
<td></td>
<td>hom</td>
<td>68</td>
<td>5</td>
<td>7.4</td>
<td>2.01E-12</td>
<td>1.21E-11*</td>
</tr>
</tbody>
</table>
Figure 3. Telomere lengths of families with old or young parents. (A) Family tree depicting the relationship of the lines used in the telomere length analysis. (B) Box plot of average telomere lengths, estimated using Computel, of fish from two families, one with heterozygous parents aged 22 months and one with heterozygous parents aged 10 months.
the homozygous fish had significantly shorter telomeres than their parents (MR3309 p=0.01, MR3310 p=0.001) and both their wild-type (MR3309 p=0.0001, MR3310 p=0.03) and heterozygous (MR3309 p=0.02, MR3310 p=0.006) siblings (Figure 3B). Similarly tertmutated fish have been shown to have shorter telomeres than their wild-type siblings.33

Surprisingly, the older parental heterozygous stock had longer telomeres than the younger parental heterozygous stock (p=0.002) (Figure 3B). This is contrary to our expectations since MR2430 was 30 months old at the time of fin clipping whereas MR2884 was only 17 months old. Furthermore, within each parent-offspring family the average telomere length of both the wild-type and the heterozygous offspring did not differ significantly from that of their parents. This indicates both that the heterozygous and wild-type offspring were able to maintain their inherited telomere length and that the phenotypes observed in the heterozygous and wild-type populations from older heterozygous parents are not the consequence of shorter telomeres, considering either the absolute telomere length or the telomere length relative to their parents. The same pattern of telomere lengths was also found when using TelomereHunter, a tool that determines telomere lengths from sequencing data using a different method to Computel (Supplementary Figure S5A).

The offspring lines MR3309 and MR3310 used in this telomere length analysis were also used in the sex ratio analysis (Figure 2E). This enabled us to show specifically that the sex ratios are independent of telomere length. Despite having shorter average telomere lengths, MR3309 wild-type and heterozygous fish did not have a male sex bias. By contrast even though both had longer telomeres, MR3310 heterozygous fish had a strong male sex bias, whereas wild-type fish did not (Figure 2E).

Discussion
We have characterised a new mutation in zebrafish tert, tertmutated, and shown that it recapitulates the phenotype of the previously published tert knockout tertmutated. We focussed our analysis on the adult phenotypes resulting from tert haploinsufficiency in heterozygous parents, and showed that the age of heterozygous parents affected the phenotype of their adult offspring. Importantly, we have demonstrated that these adult phenotypes are not simply a consequence of shorter telomeres in offspring from older parents as one might expect, but instead our data point to a role for Tert independent of telomere lengthening.

There have been a number of studies showing that TERT function outside of telomere maintenance. TERT is involved in Wnt signalling, regulation of transcription and epithelial cell proliferation and haematopoiesis. Confirmation that TERT acts independently of telomere lengthening in these situations came from studying TERT mutants lacking reverse transcriptase function or the ability of a tert construct lacking the trec binding domain to rescue blood cell numbers in Tert-deficient zebrafish, or the effects of TERT in a TERC background. However the mechanism of TERT function, other than in telomere maintenance, remains poorly understood and which aspects of a TERT-deficient phenotype are due to defects in telomere maintenance and which to non-canonical roles of TERT is an area that requires further investigation.

When two lines from parents of different ages were raised simultaneously we observed reduced survival in homozygous offspring from older tertmutated heterozygous parents but no difference in the survival of wild-type or heterozygous siblings between the two lines. This suggests that tert genotype interacting with parental age underlies the reduced survival. However this pattern of reduced survival with increasing parental age cannot be robustly observed when comparing survival across many of the lines raised. This may be due to changes in environmental factors when lines were raised at different times.

We demonstrated that offspring of older tertmutated heterozygous parents weighed less than those from younger parents, and that this weight difference was more pronounced when specifically comparing homozygous offspring to wild-type siblings. This could reflect a slower rate of proliferation or increased apoptosis in fish from older parents. Mouse embryos homozygous mutant for mtTERT and zebrafish embryos resulting from homozygous tertmutated intercrosses are smaller than their wild-type counterparts, indicating that lack of TERT can influence size, and increased p53-dependent apoptosis has been shown in both zebrafish tert knockouts and mouse mTERT mutants. However a parental age-dependent effect on body weight has not previously been demonstrated to result from haploinsufficiency in the parental generation. Thus a combination of reduced proliferation and increased apoptosis may explain the reduced body weight of fish from older tertmutated heterozygous parents. It is surprising though that the effects of tert haploinsufficiency can affect even wild-type offspring in the next generation which are able to express zygotic Tert.

The strong male sex bias in tertmutated homozygous fish suggests they have accumulated DNA damage. Mutations in genes involved in DNA damage repair such as brcaca result in all fish developing as male. This is due to the developing oocytes dying, presumably as a result of an accumulation of DNA damage resulting from recombination during meiosis, and, without the presence of oocytes, juveniles develop as phenotypically male. The shorter telomeres of homozygous fish, as revealed by Computel and TelomereHunter analysis of whole genome sequencing data, cannot by themselves explain the accumulation of DNA damage since heterozygous offspring from young parents had a similar average telomere length to homozygous offspring from older parents, but failed to show a male sex bias.

We have used an average telomere length estimate to compare telomere lengths of fin biopsies, but this does not give us the resolution to be able to assess the lengths of individual telomeres. Whilst our data suggest that the phenotypes we observed are independent of telomere length, we cannot rule out the possibility that fish from older heterozygous parents contain one or two chromosomes.
with critically short telomeres that elicit genomic instability. It has been proposed that the shortest telomere length, not the average, is the most relevant measure for telomere dysfunction\textsuperscript{12}, however average telomere length remains a common measure and has been shown to correlate with health markers and survival in both humans\textsuperscript{33-39} and animals\textsuperscript{60,61}. Furthermore, we only measured telomere lengths of caudal fin biopsies and it is possible that other tissues will behave differently.

The average telomere lengths of older parental heterozygous \textit{tert}\textsuperscript{m541} fish was longer than for younger parental heterozygotes. This was in contrast to what we expected, as the general consensus is that telomeres shorten with age\textsuperscript{30-54}, and there was a 12 month age difference between the two lines. This age difference resulted from an additional outcross to wild type for the younger line. Successive intercrosses of \textit{mTERT} heterozygous mice have progressively shorter telomeres due to \textit{TERT} haploinsufficiency\textsuperscript{45}, but this is unlikely to explain the shorter telomeres in younger \textit{tert}\textsuperscript{m541} heterozygous fish. Firstly, we introduced a wild-type copy of each chromosome from our wild-type stock rather than intercrossing carriers for the additional generation, and, secondly, our telomere length measurements showed that wild-type and heterozygous offspring maintained their inherited telomere length, indicating that a shortening of telomeres is not inevitable with each generation of \textit{tert} heterozygous fish. Indeed it has been shown in humans that paternal age at birth is positively correlated with offspring telomere length\textsuperscript{46}. Instead the difference in telomere length suggests that the genetic background of each line plays a significant role in determining telomere length, as has been shown for mice\textsuperscript{42}. More specifically, it is possible that from the distribution of mean telomere lengths in the grandparents, individuals with longer telomeres were crossed to produce MR3310 and an individual with by chance shorter telomeres was crossed to produce MR2884. This notion is supported by the finding that telomere length in the wild-type and heterozygous offspring lies in the range of their respective parents, but is different between the families despite being of the same age.

Haploinsufficiency of either \textit{TERT} or \textit{TERC} causes dyskeratosis congenita (DC) in human patients. Our zebrafish loss of function model recapitulates haploinsufficiency of \textit{tert}, but this haploinsufficiency is not evident in the heterozygous parents themselves, and, instead, heterozygosity of \textit{tert} interacts with parental age to produce phenotypes in the next generation. This suggests that \textit{tert} haploinsufficiency manifests in gametogenesis and progressively worsens as parents age. It is tempting to speculate that offspring from older parents inherit shorter telomeres, leading to phenotypes in offspring from older parents, but our telomere length analysis indicates that this is not the case. How heterozygosity of \textit{tert} in the parental generation can affect the adult phenotype of their offspring in a parental age-dependent manner remains to be elucidated. A detailed analysis of the structure and stability of chromosomes in general, not just focussing on the telomeres, in gametes from parents of different ages will be needed to address this question.

Human diseases resulting from loss of \textit{TERT}, such as DC, comprise several different phenotypes. In the case of DC, patients suffer from idiopathic pulmonary fibrosis, increased cancer susceptibility, skin pigmentation defects, nail dystrophy and bone marrow failure. Given the emerging evidence for telomere-independent roles of \textit{TERT}, including the data presented in this study\textsuperscript{40-42}, dissecting the mechanisms of pathogenesis of different components of such diseases may reveal new areas that could be the focus of disease interventions.

Data availability
Whole exome sequencing data are available from ENA under Study Accession Number: ERP016250. All other raw data are available via Figshare at https://doi.org/10.6084/m9.figshare.c.3866671.

Competing interests
No competing interests were disclosed.

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Supplementary material
Supplementary Figure S1. \textit{tert}\textsuperscript{m6541/a23076} compound heterozygous fish display the same phenotype as \textit{tert}\textsuperscript{m6541/m6541} homozygous fish. (A) Embryos produced from intercrosses of compound heterozygous fish display a range of phenotypes similar to those resulting from \textit{tert}\textsuperscript{m6541/a23076} homozygous intercrosses. (B) Compound heterozygous intercrosses produce clutches with fewer non-phenotypic embryos at 5 d.p.f. than wild-type sibling intercrosses.

Click here to access the data.

Supplementary Figure S2. Family tree of \textit{tert}\textsuperscript{m541} lines used in this study.
The relationship between all the \textit{tert}\textsuperscript{m541} lines is depicted as a family tree. Line numbers are given according to our fish database with date of birth below.

Click here to access the data.
Supplementary Figure S3. Survival graphs of progeny from intercrosses of different aged tert<sup>−/−</sup> heterozygous parents. Each graph shows the survival of homozygous, heterozygous and wild-type offspring from individual intercrosses. Parental age is indicated for each intercross.

Click here to access the data.

Supplementary Figure S4. Clutch characteristics of tert<sup>−/−</sup> intercrosses and scatterplot of sex ratios in relation to parental age of 74 different lines.

Click here to access the data.

Supplementary Figure S5. Telomere lengths of fish from Figure 3A estimated using TelomereHunter.

Click here to access the data.

References


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Version 2

Reviewer Report 06 March 2018
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Noriyoshi Sakai
Genetic Strains Research Center, National Institute of Genetics, Mishima, Japan

I agree with the authors' comments that they did not find a statistically significant difference between heterozygous and wild-type fish in fertilization rates and clutch size. It may be difficult to find differences in gonad histology between heterozygous and wild-type in this condition. I think authors properly responded to my comments.

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 05 March 2018
https://doi.org/10.21956/wellcomeopenres.15254.r31126

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Karl-Lenhard Rudolph
Leibniz Institute For Age Research, Jena, Germany

The authors addressed my comments, I am in favor of indexing.

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
Scahill et al. investigate the influence of the parental age of heterozygous telomerase mutation (Tert<sup>+/−</sup>) carriers on the phenotype of the offsprings in zebrafish. Tert<sup>+/−</sup> zebrafish exhibit a shortened lifespan, reduced body mass, male infertility, and sex bias towards males in the offsprings. These genotypes were aggravated by the parental age of Tert<sup>+/−</sup> fish but did not correlate with shorter telomeres. The authors conclude that Tert mutation heterozygosity influences phenotypes in the next generation in a telomere length independent manner.

This is a very interesting and well-designed study. I have a few points of criticism that should be addressed:

1. The study investigates the phenotype of the offspring in one tree of families and the telomere length in another tree following the same mating scheme from the grandparent generation. The telomere length is measured by sequencing – a new method. The Box plot analysis of mean telomere length shows high spreads. Is it possible that by chance difference in telomere length appear in the offspring generation by selection of germcells with longer or shorter telomeres in the grandparent and parent generation? Can the authors provide some control that the sequencing analysis of telomere DNA content correlates well with telomere length analysis by classical methods such as Southern blot on fin clips?

2. The authors conclude that parental age of Tert mutation carriers influence phenotypes in offspring in a telomere length independent manner. Parental age effects mutations have been described in humans. Is it possible that short telomeres select for genetically or epigenetically altered germline stem cell clones that influence phenotypes in the offspring generation. This selection process may also select for genetically/epigenetically altered germline stem cells with longer telomeres. Even if the telomeres in the offspring would not show a correlation of telomere shortening with the occurrence of phenotypes, it would still be a telomere shortening induced phenotype as the process of germline stem cell selection would triggered by short telomere in the parental germline in an age dependent manner.

3. The authors discuss that the “accumulation of DNA damage could however be independent of telomere length as it has been shown telomere associated DNA-damage response proteins do not differ in length from those that are negative for DNA damage markers (51).” This paper investigates the chromatin response (de-compaction) to telomere uncapping induced by Trf1 and Trf2 deletion. This paper does not investigate the formation of DNA damage foci in response to telomere shortening – the most relevant process in call aging. There is instead good evidence that the
induction of DNA damage responses is indeed telomere shortening dependent and occurs at the critically short telomeres, as the authors discuss in the following paragraph of the discussion. The cited part of the discussion should be corrected as it is misleading.

Is the work clearly and accurately presented and does it cite the current literature?  
Yes

Is the study design appropriate and is the work technically sound?  
Yes

Are sufficient details of methods and analysis provided to allow replication by others?  
Yes

If applicable, is the statistical analysis and its interpretation appropriate?  
Yes

Are all the source data underlying the results available to ensure full reproducibility?  
Yes

Are the conclusions drawn adequately supported by the results?  
Partly

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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**Author Response 12 Feb 2018**

**Elisabeth Busch-Nentwich,** Wellcome Trust Sanger Institute, Wellcome Genome Campus, Hinxton, UK

Thank you for reviewing our manuscript and your insightful comments. Please find our point-by-point response below.

Scahill et al. investigate the influence of the parental age of heterozygous telomerase mutation (Tert+/−) carriers on the phenotype of the offsprings in zebrafish. Tert+/− zebrafish exhibit a shortened lifespan, reduced body mass, male infertility, and sex bias towards males in the offsprings. These genotypes were aggravated by the parental age of Tert+/− fish but did not correlate with shorter telomeres. The authors conclude that Tert mutation heterozygosity influences phenotypes in the next generation in a telomere length independent manner. This is a very interesting and well-designed study.

Response: Thank you for this positive assessment of our work.

I have a few points of criticism that should be addressed:

1. The study investigates the phenotype of the offspring in one tree of families and the telomere
length in another tree following the same mating scheme from the grandparent generation.

**Response:** The telomere length measurements in Figure 3 were conducted on two families (MR3309 and MR3310) that were also investigated for their sex ratios in Figure 2E.

The telomere length is measured by sequencing – a new method. The Box plot analysis of mean telomere length shows high spreads. Is it possible that by chance difference in telomere length appear in the offspring generation by selection of germcells with longer or shorter telomeres in the grandparent and parent generation? Can the authors provide some control that the sequencing analysis of telomere DNA content correlates well with telomere length analysis by classical methods such as Southern blot on fin clips?

**Response:** Telomere length measurement by Whole Genome Sequencing (WGS) is an established method and has been benchmarked against non-sequencing based methods such as southern blot ([https://doi.org/10.1093/nar/gku181](https://doi.org/10.1093/nar/gku181)) and qPCR ([https://doi.org/10.1371/journal.pone.0125201](https://doi.org/10.1371/journal.pone.0125201)) previously. We apologise for not making this more explicit before, and have added a sentence to introduce the method with references in the results section. The spread of telomere lengths is the same (at 4-6 kb) for MR2884 +/- parents as that found in human samples ([https://doi.org/10.1093/nar/gku181](https://doi.org/10.1093/nar/gku181)) and smaller for the rest. However, we absolutely agree that it is perfectly possible (and in fact likely) that the overall mean telomere length of the investigated families is determined through sampling from a distribution by descent from two parents. This is supported by our finding that telomere length in the wild-type and heterozygous offspring lies in the range of their parental generation. Therefore our assertions still stand, that 1) absolute mean telomere length is independent from the age of the parents and 2) mean telomere length cannot explain the morphological phenotypes in offspring from old parents. We have added the following sentence to the discussion to acknowledge the role of sampling from a distribution of telomere lengths: “More specifically, it is possible that from the distribution of mean telomere lengths in the grandparents, individuals with longer telomeres were crossed to produce MR3310 and an individual with by chance shorter telomeres was crossed to produce MR2884. This notion is supported by the finding that telomere length in the wild-type and heterozygous offspring lies in the range of their respective parents, but is different between the families despite being of the same age.”

2. The authors conclude that parental age of Tert mutation carriers influence phenotypes in offspring in a telomere length independent manner. Parental age effects mutations have been described in humans. Is it possible that short telomeres select for genetically or epigenetically altered germline stem cell clones that influence phenotypes in the offspring generation. This selection process may also select for genetically/epigenetically altered germline stem cells with longer telomeres. Even if the telomeres in the offspring would not show a correlation of telomere shortening with the occurrence of phenotypes, it would still be a telomere shortening induced phenotype as the process of germline stem cell selection would triggered by short telomere in the parental germline in an age dependent manner.

**Response:** This is a very interesting hypothesis, however currently we do not have evidence to support it. Although it is possible that short telomeres in the parental germline may select for genetically or epigenetically altered germline stem cell clones,
our results do not indicate that older parents have shorter telomeres. However we only measured telomere lengths in the caudal fin, and it remains possible that telomere lengths in the adult germline are shorter in older *tert* heterozygous fish. Investigating this hypothesis will require telomere length assessment of gonads in a future study.

The authors discuss that the “accumulation of DNA damage could however be independent of telomere length as it has been shown telomere associated DNA-damage response proteins do not differ in length from those that are negative for DNA damage markers (51).” This paper investigates the chromatin response (de-compaction) to telomere uncapping induced by Trf1 and Trf2 deletion. This paper does not investigate the formation of DNA damage foci in response to telomere shortening – the most relevant process in call aging. There is instead good evidence that the induction of DNA damage responses is indeed telomere shortening dependent and occurs at the critically short telomeres, as the authors discuss in the following paragraph of the discussion. The cited part of the discussion should be corrected as it is misleading.

Response: We apologise for this error and have removed this sentence from the discussion.

**Competing Interests:** No competing interests were disclosed.
phenotypes of a reduction in head and a short tail. The phenotypes may lead to less eating compared to the wild-type and the heterozygotes when they are in the same dish. This condition may cause the male sex bias of the homozygous mutants.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Partly

If applicable, is the statistical analysis and its interpretation appropriate?
I cannot comment. A qualified statistician is required.

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Developmental biology, spermatogenesis

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 12 Feb 2018

**Elisabeth Busch-Nentwich,** Wellcome Trust Sanger Institute, Wellcome Genome Campus, Hinxton, UK

We are grateful for the effort taken to review our manuscript and the helpful comments and questions. Please find our point-by-point response below.

Scahill *et al.* report unique phenotypes of tert mutant zebrafish. The mutant homozygotes die prematurely and the males become prematurely infertile. Interestingly, the age of heterozygous parents influences the adult phenotype of their offspring. The authors estimated average telomere lengths by whole genome sequencing data, and conclude that the phenotypes of offspring from older parents cannot be explained by telomere length. These are findings of importance to researchers in the field.

**Response:** We would like to thank you for concluding that this work is of importance to the field.

I have a concern about condition of fertilized eggs from heterozygous mutant parents. During
maintenance of zebrafish lines we experience that old parents, especially in the female, produce non-healthy eggs, some of which develop to small the adults. The clutch often shows the male sex bias. Is there any possibility that the adult phenotypes of offspring from heterozygous mutants are from defects in oocytes of aged heterozygous mutants because the heterozygous mutants seem to age faster than the wild-type? Alternatively, germ cells of the heterozygous may spoil faster than those of the wild-type. I recommend to add histology of gonads in heterozygous mutants and data about the numbers of spawning eggs, fertilized eggs, and developed embryos to ~3 dpf in each intercross.

Response: The reviewer is concerned that the heterozygous fish might be aging prematurely and produce unhealthy oocytes that affect the adult phenotype. We do not find any overt signs of premature aging of the heterozygous fish such as reduced survival rates (Figure 2D and Supplementary Figure S3) or morphological signs of premature aging. However, following the suggestion of the reviewer, we have assessed the egg production and quality of heterozygous females over three months in comparison to wild-type siblings. We do not find a statistically significant difference between heterozygous and wild-type fish in fertilisation rates and clutch size and no deterioration over time (Supplementary Figure 4A). Heterozygous and wild-type females are both capable of laying in excess of 900 eggs per mating. Phenotypic analysis of multiple clutches derived from heterozygous and wild-type intercrosses reveals no difference (Welch two sample t-test, p-val=0.49) in the number of abnormal embryos up to 5 dpf. This is included in the results section: “Intercrosses from heterozygous fish (965 embryos from 3 separate clutches) showed no difference in the number of phenotypic embryos compared to wild-type sibling intercrosses (614 embryos from 4 separate clutches).” This is in contrast to a mutant we have recently published where egg quality diminishes over time (Scahill et al., 2017, DOI: 10.1371/journal.pgen.1006959).

Gonad histology might identify differences that are not reflected in egg production and quality. For a meaningful result this would require sampling gonads from young wild-type females and then again their aged siblings to establish a baseline catalogue of normal gonad morphology in young and old wild-type fish. The same process would be required for tert heterozygous and homozygous females. We feel that the required time frame and amount of work goes beyond the scope of this publication. Furthermore we are concerned that the lack of an overt oogenesis defect would make it difficult to justify sacrificing such a substantial number of adult fish.

Finally, in our facility we haven’t made the same observations as the reviewer that old fish generally give rise to smaller fish and a male skew, although we agree that productivity tends to decrease with increasing age of females. As we show in Figure 2G, in contrast to offspring from 16 month old tert heterozygous parents, offspring from very old (30 months) wild-type fish do not show reduced body weight although we do see increased variance. To also systematically assess the impact of parental age on sex ratios, we have used sex ratio data routinely collected in our facility. The analysis of sex ratios across 74 different lines in relation to the age of their parents (Supplementary Figure 4B) finds no correlation of skewed sex ratios with either the paternal or maternal age. We would like to thank the reviewer for raising these questions, since we feel that these additional experiments have strengthened our conclusion that tert heterozygosity interacts with age.

In addition, authors should describe how offspring is raised. The homozygous mutants display phenotypes of a reduction in head and a short tail. The phenotypes may lead to less eating compared to the wild-type and the heterozygotes when they are in the same dish. This condition
may cause the male sex bias of the homozygous mutants.

Response: The phenotypic embryo in Figure 1F is derived from tert-/ intercrosses, not tert+/ parents, and is therefore a maternal-zygotic (MZtert) mutant. We have changed the labelling to make this more explicit. Generally, embryos displaying such strong early morphological phenotypes do not survive beyond 3-5 dpf. The clutches raised and analysed for this study were all derived from heterozygous parents. Only morphologically normal larvae with filled swim bladders were raised for this study. We have emphasised this in the methods section.

Competing Interests: No competing interests were disclosed.