Telomerase reverse-transcriptase homozygous mutations in autosomal recessive dyskeratosis congenita and Hoyeraal-Hreidarsson syndrome

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Dyskeratosis congenita (DC) is a multisystem bone marrow failure syndrome characterized by a triad of mucocutaneous abnormalities and an increased predisposition to malignancy. X-linked DC is due to mutations in DKC1, while heterozygous mutations in TERC (telomerase RNA component) and TERT (telomerase reverse transcriptase) have been found in autosomal dominant DC. Many patients with DC remain uncharacterized, particularly families displaying autosomal recessive (AR) inheritance. We have now identified novel homozygous TERT mutations in 2 unrelated consanguineous families, where the index cases presented with classical DC or the more severe variant, Hoyeraal-Hreidarsson (HH) syndrome. These TERT mutations resulted in reduced telomerase activity and extremely short telomeres. As these mutations are homozygous, these patients are predicted to have significantly reduced telomerase activity in vivo. Interestingly, in contrast to patients with heterozygous TERT mutations or hemizygous DKC1 mutations, these 2 homozygous TERT patients were observed to have higher-than-expected TERC levels compared with controls. Collectively, the findings from this study demonstrate that homozygous TERT mutations, resulting in a pure but severe telomerase deficiency, produce a phenotype of classical AR-DC and its severe variant, the HH syndrome. (Blood. 2007;110:4198-4205)

Introduction

Dyskeratosis congenita (DC) is a genetically and clinically heterogeneous inherited bone marrow (BM) failure syndrome that is classically characterized by a mucocutaneous triad of abnormal skin pigmentation, nail dystrophy, and leukoplakia, as well as a wide range of other cutaneous abnormalities.1 The X-linked recessive form of DC was linked to mutations in DKC1, which encodes the highly conserved multifunctional protein dyskerin.2-7 Dyskerin associates with a subclass of small nucleolar RNAs (snoRNAs) through the RNAs H/ACA domain.8-10 These 2 molecules ultimately form a ribonucleoprotein (RNP) complex with the additional proteins GAR1, NHP2, and NOP10.11-13 This complex has at least 2 significant biologic functions. First, it is responsible for the pseudouridylation of ribosomal RNA at residues specified by the snoRNA contained within the RNP.14 Second, in addition to these pseudouridylation functions, the vertebrate version of this RNP complex associates with a reverse transcriptase (RT) to form the telomerase complex.15-17

Telomeres are unique tandem repeat structures located at the ends of linear eukaryotic chromosomes.18 Their presence serves many functions including prevention of chromosome end-to-end fusion, maintenance of chromosomal stability, and prevention of chromosomal degradation.19 Telomerase is a specialized RNP complex that elongates the G-rich telomeric repeats.20,21 It is proposed that telomerase acts after DNA replication where this specialized enzyme extends the leading DNA strand, reestablishing the 3′ overhang that is lost during normal cell division due to end processing and the “end replication problem.”22,23 The 2 essential components of this telomerase RT are the RNA component (TERC) that contains the RNA template that the RT (TERT) uses to synthesize 6-bp repeats on the 3′ terminal end of telomeric DNA.24 As the telomerase complex also requires dyskerin, the possibility that mutations in other components of the telomerase complex could cause DC was raised.25 Heterozygous TERC mutations have subsequently been characterized in a subset of patients with autosomal dominant (AD) DC, as well as other related BM failure syndromes, suggesting that disruption of the telomerase complex results in defective hematopoiesis.26-35 The deletions and base-pair substitutions that have been identified in TERC have been shown to alter telomerase function via haploinsufficiency of the resulting telomerase complex through either loss of catalytic function or dissociation of the telomerase complex itself. The more recent discovery that heterozygous TERT mutations also result in a disease resembling AD-DC raises the question of whether telomerase catalytic activity is the critical function that is disrupted in these BM failure syndromes.34,36-39

During our investigations into telomerase dysfunction in the DC registry (DCR), we have found that a rare subset of DC patients has homozygous TERT mutations. We note, however, that these mutant alleles are not always fully penetrant and do not always segregate with the disease in an AD fashion.33,37 We have now identified 2 consanguineous families in which novel TERT mutations are segregating and that result in affected homozygous payment. Therefore, and solely to indicate this fact, this article is hereby marked “advertisement” in accordance with 18 USC section 1734.

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children. In this paper, we describe the clinical features of these 2 families and the functional characterization of these homozygous TERT mutations. The findings from these 2 families demonstrate that homozygous TERT mutations resulting in a “pure” telomerase deficiency can produce a phenotype of classical autosomal recessive (AR) DC and its severe variant, the Hoyeraal-Hreidarsson (HH) syndrome.

Patients, materials, and methods
Screening of patients within the DC Registry
Clinical and genetic information has been collected from 277 DC families worldwide. Approval for these studies was obtained from the Research Ethics Committee of the Hammersmith Hospital National Health Service (NHS) Trust and Barts and The London NHS Trust. Informed consent was obtained in accordance with the Declaration of Helsinki. To date, the TERT gene has been screened for mutation in 148 cases of DC in which the genetic basis is unknown, and 23 of these appeared to show an AR pattern of inheritance. The 2 families characterized in this paper have been entered into the DC Registry (DCR) as each index case had clinical features that led to a diagnosis of either DC or HH. Mutation screening of the TERT gene was performed by denaturing high-performance liquid chromatography (dHPLC) as described previously. DNA fragments that displayed an abnormal pattern of elution in this analysis were reamplified and subjected to direct sequence analysis by BigDye chain termination cycle sequencing.

TERT plasmid constructs and mutagenesis
The pcDNA3.1+ wild-type (WT) TERC and pcDNA3.1+ WT TERT plasmids were constructed as previously described. Both TERT cDNA mutations were produced using the QuikChange XL site-directed mutagenesis kit (Stratagene, La Jolla, CA) where 2 complementary overlapping oligonucleotides were designed to contain the mutation to be induced in the original WT plasmid template) following the manufacturer’s instructions. For each TERT variant, 125 ng of the appropriate forward and reverse mutagenesis primers was added to 1× reaction buffer, 10 ng WT p3.1+ TERT, 2% dNTP mix, 6% QuickSolution mix, and 1.25 units PfuTurbo DNA polymerase. Each reaction was denatured at 95°C for 1 minute, cycled 18 times at 95°C for 50 seconds, 60°C for 50 seconds, and 68°C for 11 minutes, and completed with an extension cycle for 7 minutes at 68°C. Competent DH5α cells were then transformed with the resulting DpnI-treated DNA (to remove the original WT plasmid template) following the manufacturer’s instructions. Resulting colonies from each transformation were selected for plasmid DNA extraction (Qiagen, Valencia, CA), and the TERT cDNA coding sequence of each construct was verified.

Telomerase reconstitution in transfected WI-38 VA13 cells
Confluent WI-38 VA13 cells (human “fibroblast-like” cells, which can be grown in the laboratory and do not express telomerase) were plated out at 2 × 10^5/cm² plate in antibiotic-free Dulbecco modified Eagle medium (DMEM) containing 10% fetal calf serum (FCS; Sigma, St Louis, MO). TERT plasmid transfection experiments showed that 50 ng TERT plasmid per transfection was in the middle of the telomere repeat amplification protocol (TRAP) linear range (data not shown). Therefore a total of 50 ng p3.1+ TERT plasmid was mixed with 8 μg WT p3.1+ TERC plasmid and 8 μg pBRL-TK (luciferase internal control; Promega, Madison, WI) and made up to 24 μg total DNA with p3.1+ vector backbone in 1.5 mL serum-free DMEM. In some experiments, 25 ng of each mutated p3.1+ TERT plasmid was mixed with 25 ng WT p3.1+ TERT plasmid. Each plasmid mix was transfected into the telomerase-negative cells using lipofectamine 2000 (Invitrogen, Frederick, MD) and incubated at 37°C and 5% CO₂ overnight. After 24 hours, the medium was changed to DMEM containing 10% FCS and the cells were collected 48 hours after transfection. The cells were split into 1× CHAPS buffer for TRAP analysis, 1× lysis buffer for luciferase analysis, and RTL buffer (Qiagen) for RNA extraction and subsequent cDNA preparation.

TRAP analysis
The TRAP lysates were assayed for protein concentration using a DC protein assay (Bio-Rad, Hercules, CA), normalized to 650 ng/μL, and diluted 1:10, 1:40, and 1:160 using 1× CHAPS buffer prior to TRAP analysis. TRAP lysates were assayed using the TRAPese telomerase detection kit (Intergen, Burlington, MA) as previously described. The resulting TRAP products were separated on 12.5% acrylamide-0.5× TBE gels and exposed to X-ray film. TRAP activities were derived from comparing densitometry reading of a specific TRAP ladder repeat from each serial dilution of mutant samples with those from WT samples of at least 3 separate transfection and subsequent TRAP experiments using Quantity One software (Bio-Rad). The luciferase lysates were analyzed to ensure concordant transfection efficiency within each transfection experiment using the Renilla luciferase assay protocol (Promega) as previously described.

Quantitative real-time polymerase chain reaction on TRAP samples
RNA samples were HhaI (New England Biolabs, Beverly, MA) digested and DNaseI (Promega) treated to remove any contaminating genomic or plasmid DNA. The samples were then reverse transcribed by Moloney murine leukemia virus (M-MLV) RT (Invitrogen) into cDNA with random hexamer primers using standard procedures. TERC and ABL mRNA expression was determined using the ABI PRISM 3700 sequence detection system (Applied Biosystems) using primers and probes described elsewhere. Each quantitative polymerase chain reaction (PCR) reaction was prepared in Taqman Universal PCR master mix (Applied Biosystems) containing 2 μL cDNA with either 300 nM TERC primers and 200 nM TRAP probe or 100 nM ABL primers and 300 nM ABL probe. Baselines and thresholds were automatically set by the software and used after manual inspection. Samples were normalized using the ABL data, and each result was expressed as a relative percentage compared with the WT sample. PCR efficiencies for each probe set were derived from standard curves using a dilution series covering a 7-log range starting from one randomly selected undiluted cDNA sample (R² = 0.9987 and R² = 0.9996 for ABL and TERC, respectively). We selected ABL as a control gene in these experiments because it is the most widely used and validated for quantitative measurements in peripheral blood samples, being the gene of choice in leukemia minimal residual disease (MRD) studies. A multicenter study has shown that, in contrast to other control genes, the ABL gene transcript was similarly expressed in normal and leukemic samples, reflecting considerably different proliferation states. We have also validated the use of the ABL probe to 2 other Taqman probes (GAPDH and HPRT) using our own samples, which gave concordant data to that seen with ABL (data not shown). All reactions were run in duplicate or triplicate.

Telomere length measurement
Telomere length was measured by Southern blot analysis using a subtelomeric probe from chromosome 7 (pTelBam8) as previously described. The size of peak signal intensity was determined using Gel Blot software (UVP, Upland, CA). Unaffected siblings and spouses in families in which DKC1, TERC, or TERT mutations have been characterized were used as age-matched healthy controls. A linear regression line was calculated for telomere length against age in these healthy individuals. This value was then used to determine the age-adjusted telomere length of affected individuals by expressing the difference between the observed length and the predicted telomere length from the linear regression line (Δtel) as previously described.
Telomere lengths have also been measured by quantitative PCR on a 7500 Real Time PCR System (Applied Biosystems) using primer pairs and PCR conditions as previously described. Briefly, 2 measurements were made in triplicate on each DNA sample, one to determine the telomere repeat copy number (T) and another to determine single copy gene copy number (S). A reference DNA sample was serially diluted on each plate so that relative quantities of T and S could be determined from standard curves. T/S ratios were expressed relative to a single DNA, and mean values were used to establish the assay sensitivity. The reproducibility of the assay was assessed by comparing the T/S ratio determined from 2 different dilutions of 15 randomly selected samples. Measurements on these 2 dilutions showed very good correlation (R² = 0.984). All differences between healthy controls and the mutation groups for the T/S ratio were compared using the Mann-Whitney test.

### Results

#### Clinical presentation of 2 consanguineous families with novel homozygous TERT mutations

The index case of family A was a 13-year-old Libyan girl from a consanguineous marriage (II:4 in Figure 1). She was underweight and short for her age. Investigations showed that she had a hypocellular BM and thrombocytopenia. Her low platelet count was treated initially with prednisolone. This treatment was changed to oxymetholone (an anabolic steroid) on referral to the United Kingdom as by this time she had trilineage hematopoietic defect producing pancytopenia (Table 1). The combination of cutaneous abnormalities observed by physicians in Libya and BM failure led to the diagnosis of putative constitutional aplastic anemia (CAA). Further clinical investigation in our department established that the index case had café-au-lait spots on her trunk and reticulated pigmentation on the nape of the neck. She was also found to have “bluish discoloration” on the tongue with areas of leukoplakia, thin nails on the hands, but more significant nail dystrophy on the toes (Figure 2). Blood analysis at this time found raised IgG but normal IgA and IgM levels, as well as a reduction in all blood lineages but no blasts (Table 1). Chromosomal breakage analysis was found to be normal, ruling out Fanconi anemia (FA). The presence of mucocutaneous features (reticulate skin pigmentation, nail dystrophy, and leukoplakia) in the index case (Figure 2) together with the trilineage BM failure (Table 1) led to the diagnosis of DC. Clinical investigations on the rest of the family found that, while clinically asymptomatic with normal blood counts (Table 1), her parents and an older sister had some mild abnormalities (highlighted in Figure 1). The death of an older sister was from unknown causes at 6 years of age.

| Table 1. Features of families with AR inheritance of homozygous TERT mutations |
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| **Family A:** DC clinical phenotype |
| Position | Age, y | Status | Hb level, g/L | Wbc count, ×10⁹/L | Platelet count, ×10⁹/L | MCV level, fl | TERC/ABL ratio† |
| I:1 | Mother | 53 | Het | n/a | n/a | n/a | n/a | n/a | n/a |
| I:2 | Father | 57 | Het | 161 | 6.09 | 149 | 92 | n/a | n/a |
| II:1 | Sister | 28 | Het | 144 | 6.0 | 317 | 86 | n/a | n/a |
| II:4 | Index | 13 | Hom | 841 | 3.8 | 19 | 101 | n/a | 7.5 |
| **Family B:** HH clinical phenotype |
| Position | Age, y | Status | Hb level, g/L | Wbc count, ×10⁹/L | Platelet count, ×10⁹/L | MCV level, fl | TERC/ABL ratio† |
| I:1 | Mother | 31 | Het | 139 | 5.2 | 189 | 93 | –2.2 | 4.1 |
| I:2 | Father | 36 | Het | 154 | 5.8 | 206 | 96 | –2.3 | 3.9 |
| II:1 | Brother | 7 | Normal | 139 | 6.8 | 309 | 88 | +1.2 | 4.7 |
| II:2 | Index | 3 | Hom | 113§ | 3.1 | 21 | 103 | –6.5 | 8.0 |

Hb indicates hemoglobin; Wbc, white blood cell; MCV, mean corpuscular volume; ∆tel, observed-expected telomere length; DC, dyskeratosis congenita; HH, Hoyeraal-Hreidarsson syndrome; Het, heterozygous; Hom, homozygous; and n/a, not available.

†Relative units of quantitative real-time (QRT)-PCR analysis: normal range is 1.1 to 15.5; mean is 4.68; standard deviation is 3.78.


§On oxymetholone.

§After red blood transfusion.
The index case of family B was a 3-year-old girl with an Iranian-Jewish ethnic background who is also from a consanguineous marriage (II:2 in Figure 1). She was found to have cerebellar hypoplasia, early BM failure, and leukoplakia after further investigations to elucidate her failure to thrive. Microcephaly, gastrointestinal abnormalities (dysphagia), learning disabilities, and developmental delay were also noted. FA was ruled out early on and in view of the above combination of abnormalities she was diagnosed to have HH syndrome, the severe variant of DC. Clinical investigation of the rest of the family found that they were clinically asymptomatic with normal blood counts (Table 1). The index case went on to develop progressive trilineage hematopoietic BM failure with a profound thrombocytopenia (Table 1).

Identification of 2 novel homozygous TERT mutations

Since the index cases were female, it was highly improbable that DKC1 would be mutated in these 2 families. Consanguinity in both families suggested an AR-DC inheritance pattern, although AD-DC could not be disregarded, as disease anticipation may support the lack of clinical phenotype in the first generations. AD-DC due to TERC mutations was subsequently ruled out when screening analysis found that both index cases had WT TERC genes (data not shown). As the core telomerase complex contains TERT and TERC, and in light of the recent heterozygous TERT mutations identified in DC families, TERT was an obvious candidate disease-associated gene in these families.

Initial dHPLC screening of the father of family A indicated that he was heterozygous for a mutation in exon 8 of the TERT gene. Subsequent DNA sequence analysis in family A revealed a homozygous c2431C>T nucleotide change in the index case (Figure 3A). This resulted in a pArg811Cys (R811C) substitution in
the nonconserved region of the RT domain (Figure 3B). In family B, dHPLC screening of the index case revealed a subtle abnormality in exon 11 of the TERT gene. The index case of family B had a homozygous 2701C→T nucleotide change (Figure 3A) that resulted in a pArg901Trypt (R901W) substitution in the conserved D motif of the RT domain (Figure 3B). Neither of these homozygous TERT mutations has been identified as either a homozygous or heterozygous mutation in any of the other 158 subjects of mixed ethnic origin who we have investigated or in any of the other reported TERT screens, including one of 282 control subjects, suggesting these single base-pair substitutions are unlikely to be rare polymorphisms. The segregation of each TERT mutation was confirmed by dHPLC and direct DNA sequence analysis that showed that the parents are heterozygous for their specific mutation and that the presence of the homozygous single base change segregated with the disease phenotype (Figure 1). As the rest of the family members appear asymptomatic in family B, it suggests that the novel R901W TERT mutation induces the clinical phenotype through an AR inheritance pattern.

**Novel TERT amino acid substitutions reduce telomerase activity and telomere lengths**

It was not possible to measure telomerase activity in primary cells such as lymphocytes because of lack of appropriate primary material. To determine the consequences of the homozygous amino acid substitutions on telomerase activity, these 2 TERT mutations were transfected into telomerase-negative WI38 cells and the resulting lysates analyzed in reconstituted TRAP assays. The resulting telomerase activity was found to be reduced for R811C (less than 50%) and R901W (less than 25%), respectively, compared with WT controls (Figure 4A). While the index cases were homozygous, their parents were heterozygous for these mutations, yet appeared asymptomatic or mildly suggestive of DC (Figure 1). As the rest of the family members appear asymptomatic in family B, it suggests that the novel R901W TERT mutation induces the clinical phenotype through an AR inheritance pattern.

Investigation of telomere lengths (Figure 5; Table 1) shows that the index case from family B has one of the shortest telomere lengths that we have observed in our cohort of patients (DCR). It lies well below the best-fit-line and outside the 95% deviation area of the normal range (black diamond in Figure 5A). The asymptomatic heterozygous R901W parents lie within the 90% deviation area of the normal range (gray diamonds in Figure 5A), and it is therefore possible to speculate that the presence of significantly short telomere lengths in the index case is a result of inheriting shorter-than-expected telomeres from both parents. If this were the case, the homozygous WT brother would also have shorter-than-expected telomere lengths. Interestingly, the brother from family B has a telomere length that lies above the best-fit-line but within the 68% deviation from the normal range (white diamond in Figure 5A). To confirm this result, relative telomere lengths were also investigated in family B using a quantitative PCR assay in which a ratio between telomere repeat copy number (T) and single copy gene copy number (S) is determined. In a random selection of 26 high-quality DNA samples, we found that T/S ratios obtained from this assay showed significant correlation with the telomere length measured by Southern blot analysis (Figure 5C). We observe that the index case with the R901W mutation has a T/S ratio below the normal range, the heterozygous parents have low-normal T/S ratios, and the unaffected sibling has a relatively high T/S ratio (Figure 5D).

Investigations into TERC levels were also carried out. These results show that the amount of TERC is high in both homozygous index cases (Figure 6; Table 1). This is in contrast to DC patients with hemizygous DKC1 mutations, where TERC levels in peripheral blood are significantly reduced compared with healthy individuals (t test: P < .001 compared with controls; Figure 6). Interestingly, patients with heterozygous P721R TERT mutations have
Figure 5. Telomere length analysis in family B. (A) Healthy subjects are indicated by ○. The best-fit-line through this normal range is shown as a black line that corresponds to the equation \( \text{Y} = 17.821 - 0.0407X \). Deviation from the best-fit-line has been highlighted as a dark gray box for 68%, a lighter gray box for 90%, and the palest gray box for 95%. The healthy brother is highlighted as a ○; while the heterozygous R901W parents are ● and the homozygous R901W index case is a ●. (B) The Δtel values from healthy subjects from panel A (n = 112) are represented on a linear graph and compared with the R901W family (family B). (C) A random selection of subjects (n = 26) was also analyzed using the T/S ratio method25 and plotted to show the relationship to their TRF measurements in panel A. A linear trend line was added for correlation analysis (\( R^2 = 0.6917 \)). (D) Family B was also analyzed using the T/S ratio method and compared with the healthy subjects from panel C.

TERC levels in a similar range to healthy controls (t test: \( P = .17 \) compared with controls; Figure 6). Meanwhile, both sets of heterozygous TERT parents and the homozygous healthy brother have TERC levels near the mean value (Figure 6; Table 1). These results demonstrate that while low TERC levels are a consistent feature of X-linked DC, subjects studied with heterozygous TERT mutations have normal TERC levels and, although numbers are small at present, the 2 homozygous TERT patients have raised TERC levels.

Discussion

These are the first reports of homozygous amino acid substitutions in TERT. They result in a clinical phenotype of either classical DC or HH, which is a severe variant of DC. They also represent the first genetic characterization of any case of AR-DC/HH. Individuals with heterozygous TERT mutations published to date33,35-39 do not have the features of nail dystrophy and skin pigmentation abnormalities that have been observed in the index cases described here. This study therefore shows that a clinical phenotype close to classical DC can be produced in patients with homozygous mutations in TERT and can be associated with a “pure telomerase defect.” This defect is highlighted by the presence of very short telomeres observed in the index case of family B. The reduced telomerase activity observed in the TRAP assays for both novel mutations suggests that the index cases in both of these 2 families would have less than 50% of normal telomerase activity in vivo.

TERT is a member of a large family of nucleic acid–dependent polymerases with conserved primary sequence motifs. While the tertiary structure of the TERT RT domain is unknown, telomerase-specific areas have been located in both the N- and C-termini of the protein.51-53 Key residues in the TERT peptide (Figure 3B), specific areas have been located in both the N- and C-termini of the protein.51-53 Key residues in the TERT peptide (Figure 3B), specific areas have been located in both the N- and C-termini of the protein.51-53 Key residues in the TERT peptide (Figure 3B). These conserved K902 amino acid that was found to be mutated in another DC family,36 although the R901 amino acid itself is not highly conserved between the 5 species investigated. Of the 2 mutations that we have identified here, the R901W amino acid substitution resulted in a lower telomerase activity and is associated with a more severe clinical phenotype (HH). It is possible that homozygous mutations in highly conserved residues would be lethal as these would be predicted to result in negligible telomerase activity in vivo, and therefore would be incapable of maintaining normal growth and development.

DC is a genetically heterogeneous genetic disorder.56 But regardless of the inheritance pattern, progressive loss of cellular renewal associated with short telomeres has been observed. In cells from patients with X-linked recessive DC, lower TERC levels have been observed along with short telomeres.25,57 Haploinsufficiency of TERC leads to the AD inheritance of DC.26,33 This also appears to be the case with homozygous inheritance of mutated TERT.33,35-39

In this paper, we demonstrate that homozygous TERT mutations lead to reduced telomerase activity, shorter-than-expected telomeres, and a suggested increase of TERC levels. It has been
documented that both X-linked DC and AD-DC may arise due to limitation of TERC levels. Our study now shows that classical DC can also arise due to severe deficiency of TERT, but that low TERC levels do not appear to be a uniform feature of DC. Indeed the index cases of both families with homozygous TERT mutations were found to have raised TERC levels reflecting possible feedback mechanisms to compensate for the severe TERT deficiency. Therefore it seems that impaired telomerase function through dyskerin, TERC, or TERT defects is sufficient to induce a DC clinical phenotype.

It is noteworthy that some of the heterozygous individuals in these families had subtle features of DC (Figure 1). While the heterozygous R901W parents in family B had relatively short telomeres but were asymptomatic, the parents in family A had mild nail or skin pigmentation abnormalities. As there is a 20-year age difference between these 2 sets of parents, it is possible that the clinical features in family B simply require more time to develop, in a similar manner to the disease anticipation observed in AD-DC TERC families. We also note that while the presence of the homozygous R901W TERT mutation led to a significant reduction in telomere length, this shortening appeared to be reversed when the R901W alleles were absent, as the unaffected healthy sibling has relatively long telomeres.

The clinical observations in heterozygous individuals of both families are consistent with previous reports of a variable range of presentation in individuals carrying heterozygous TERT mutations. The mild but variable phenotypic expression in part probably reflects the position of the different mutations within the TERT molecule as well as the effect of other environmental and/or genetic factors and therefore the various functional consequences that they might have on TERT activity, stability, or accumulation. The fact that as a group these heterozygotes are usually phenotypically mild compared with the classical presentation of DC, which we see in the homozygous index cases described here, highlights the dosage effect of having 1 versus 2 mutant TERT alleles.

It is unusual, but not unprecedented, for mutations in the same gene to give rise to both dominant and recessive forms of the same disease. The situation we describe here in TERT is partly analogous to β-thalassemia where heterozygous mutations usually produce only mild hematologic abnormalities, but may on occasion give rise to a dominant β-thalassemia. In contrast, those individuals with biallelic mutations in the β-globin gene usually present with a severe hematologic phenotype necessitating regular blood transfusion therapy. The situation with mutations to the telomerase complex is complicated, however, by the phenomenon of disease anticipation in which the dominant disease appears to worsen through successive generations. However, it is clear that the heterogeneous TERC and TERT families published to date never have a clinical phenotype as severe as seen in X-linked DC and AR-DC, particularly with respect to the cutaneous features. This is in contrast to the cases with the homozygous TERT mutations described here where the phenotype closely mimics that observed in X-linked DC. This therefore highlights that a severe telomerase deficiency alone is capable of producing a classical and severe DC phenotype.

In conclusion, we describe the first clinical and functional characterization of families with homozygous TERT mutations. The findings of this study elucidate the first genetic characterization of AR-DC/HH syndrome and show that a “pure” but severe telomerase deficiency due to constitutional homozygous TERT mutations can produce a classical phenotype of DC.

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Authorship

Contribution: A.M. performed plasmid mutagenesis, transfections, TRAP analysis, and Taqman on TRAP samples and was the main author; A.W. performed dHPLC screening and QRT-PCR analysis. H.T. provided the source of one family’s clinical data; Y.M. performed QRT-PCR analysis; M.K. and R.B. performed additional experimental preparation and analysis; T.V. performed sequence analysis and telomere length analysis and was the assisting author; I.D. provided the source of one family’s clinical data, was the laboratory head, and was the assisting author.

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References


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