Functional characterization of novel telomerase RNA (TERC) mutations in patients with diverse clinical and pathological presentations

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ABSTRACT

Background and Objectives

Functional characterization of heterozygous TERC (telomerase RNA component) and TERT (telomerase reverse transcriptase) mutations found in autosomal dominant dyskeratosis congenita (DC) and aplastic anemia (AA) shows that telomerase function is defective and that this is associated with short telomeres. This leads to reduced cell longevity with maximal impact on tissues with high proliferative potential. The aim of this study was to establish the role of TERC in the pathophysiology of uncharacterized patients with AA with some features of DC.

Design and Methods

The TERC gene was screened for mutations by denaturing high performance liquid chromatography. To determine the functional significance of TERC mutations telomerase activity was assessed in an in vitro (TRAP) assay and telomere length of patients’ samples was determined using Southern blot analysis.

Results

This study led to the identification of four novel TERC mutations (G178A, C180T, ∆52-86 and G2C) and a recurrent TERC mutation (∆110-113GACT).

Interpretation and Conclusions

Two of the de novo TERC mutations (G178A and C180T) found uniquely produce a clinical phenotype in the first generation, differing from previously published cases in which individuals in the first generation are usually asymptomatic. Curiously these mutations are located near the triple-helix domain of TERC. We also observed that the recurrent ∆110-113GACT can present with AA, myelodysplasia or leukemia. The ∆52-86 is associated with varied phenotypes including pulmonary disease (pulmonary fibrosis) as the first presentation. In summary, this study reports the functional characterization of several novel TERC mutations associated with varied hematologic and extra-hematologic presentations.

Key words: aplastic anemia, de novo, dyskeratosis congenita, haplo-insufficiency, telomerase, TERC.

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Dyskeratosis congenita (DC), a rare bone marrow (BM) failure syndrome, has been linked to mutations in DKK1, TERC (telomerase RNA component) and TERT (telomerase reverse transcriptase). As these molecules associate to form the telomerase complex, it suggests that the various forms of DC are due to defective telomerase function, which results in loss of cell longevity in highly proliferative tissues.

Heterozygous TERC mutations have been identified in a subset of patients with autosomal dominant DC, as well as other related BM failure syndromes, suggesting that disruption of the telomerase complex results in defective hematopoiesis. Previous reports have highlighted that the deletions and small base-pair substitutions in TERC impede telomerase function due to haplo-insufficiency. This occurs from either direct loss of catalytic activity or through dissociation of the telomerase complex itself. Subsequent studies showed that disease anticipation, heterogeneous inheritance patterns and variable penetrance, families with diverse clinical features can be linked to a TERC mutation. While there has been speculation as to the mechanism behind each defective telomerase complex, it has been suggested that disease anticipation occurs due to the gradual erosion of the telomeres in subsequent generations.

In this paper, we report on the functional characterization of several novel TERC mutations, including a recurring 4bp pseudoknot deletion and two de novo substitutions associated with clinical features in the first generation. The latter observation is unique to these families and has not been observed with other TERC mutations published to date. We also observed that one of the mutations was primarily associated with pulmonary disease in some of the affected individuals. This study highlights that in addition to previously documented hematologic pathology, TERC mutations may present with disease features due to pathology in non-hematopoietic tissues.

**Design and Methods**

**Screening of TERC in patients with AA and related syndromes**

Clinical information was collected from many patients who have AA with features overlapping those of DC. These studies have been approved by the Local Research Ethics Committee and informed consent was gained in accordance with the Helsinki Declaration. TERC was screened by denaturing high performance liquid chromatography analysis and abnormal patterns were subjected to direct sequence analysis as previously described. Any mutations identified were confirmed by either sequencing the reverse strand or by re-amplification and restriction enzyme digestion using sites that are created or destroyed by the presence of the TERC mutation in question.

**Table 1. Primers used during TERC plasmid mutagenesis.**

| TERC mutation | Sequence
|---------------|------------------|
| c.180C→T (C180T) | caaacaacatatggcctgccaggccagtaccttgttccttt | cagggggagagtccttggtctttgg
| c.112C→T (C112T) complement to c.178G?A | gtttttcctgggtttcaggggc | gcgcgcacagctgcagagaac
| c.110G→A (G110A) complement to c.180C?T | gtttttcctgtttactcaggggc | gcgcgtgaagcctgcagagaac
| c.2G→C (G2C) | gcagccacagtgtttagggaggg | acccccacacccggggagggg

*Nucleotide number in TERC gene where G of the initial RNA sequence GGG is depicted as number 1; Primer sequence 5’→3’ with the forward primer above the reverse primer. The mutated base in each primer is highlighted in bold.

**Telomere length measurement**

Telomere length was measured as previously described. A linear regression line was calculated for telomere length against age in unaffected siblings and spouses in families in which DKK1 mutations have been characterized. 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.

**Micro-satellite analysis**

Paternity was assessed through the analysis of ten short tandem repeat (STR) loci (FGA, VWA, TH01, D13S1858, D16S539, D2S1338, D8S1179, D21S11, D18S51 and D19S433) provided within the AmpFlSTR SGM Plus PCR amplification kit (Applied Biosystems). The tetranucleotide STR loci were amplified in a single polymerase chain reaction (PCR), separated on a 3130xl Genetic Analyzer and visualised using GeneMapper software (Applied Biosystems).

**TERC plasmid constructs and mutagenesis**

Wild type (WT) TERC and TERT plasmids were constructed as previously described. TERC mutations were produced by a two-stage PCR approach or by using the QuikChange site-directed mutagenesis kit (Stratagene, CA, USA). Complementary overlapping primers were designed (Table 1) for each mutation and were added to 1x reaction buffer, 10 ng of WT TERC, 2% dNTP mix, 6% QuikSolution mix and 1.25 units of FfuTurbo DNA polymerase. Each reaction was denatured at 95°C for 30secs, cycled 14 times at 95°C for 30secs, 55°C for 1 min, and 68°C for 9 min and completed with an extension cycle for 7 min at 68°C. Competent cells were transformed with DpnI-treated DNA following the manufacturer’s instructions. Resulting colonies from each transformation were selected for plasmid DNA extraction (Qiagen, CA, USA) and the TERC sequence of each construct was verified.
Characterization of de novo TERC mutations

Table 2. Hematologic data (at their last clinic visit) from affected cases with TERC mutations.

<table>
<thead>
<tr>
<th>Family and TERC mutation</th>
<th>Age (years)</th>
<th>Hb (g/dL)</th>
<th>MCV (fL)</th>
<th>HbF (%)</th>
<th>PLT (x10^12/L)</th>
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<tr>
<td>A: G178A</td>
<td>5</td>
<td>10.7</td>
<td>97.6</td>
<td>5.7</td>
<td>29</td>
<td>6.6</td>
<td>2.1</td>
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<tr>
<td>B: C180T</td>
<td>35</td>
<td>13.9</td>
<td>117</td>
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<td>37</td>
<td>2.8</td>
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<tr>
<td>C: Δ110-113GACT</td>
<td>17</td>
<td>11.7</td>
<td>104</td>
<td>3.3</td>
<td>90</td>
<td>3.4</td>
<td>1.6</td>
</tr>
<tr>
<td>D: Δ53-87</td>
<td>26</td>
<td>10.8</td>
<td>125</td>
<td>2.9</td>
<td>35</td>
<td>3.6</td>
<td>2.1</td>
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<tr>
<td>E: G2C</td>
<td>27</td>
<td>8.3</td>
<td>117</td>
<td>3.0</td>
<td>35</td>
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Normal ranges

Child 2.6, 12.5±1.5, 81±6 2 years
Adult Males 13.5±1.5, 92±9 <1 years
Females 15.0±2.0, 280±130, 4.5±2.5

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Results

Identification of de novo autosomal dominant TERC mutations

The index case of family A presented at 2 years old with abdominal pain and easy bruising. He also had mild developmental retardation with delayed speech attributed to a hearing impairment secondary to recurring otitis media, for which he underwent grommet insertion. Investigations demonstrated severe BM hypocellularity. He had a normal 46XY karyotype and normal chromosomal breakage scores with DNA cross-linking agents. His parents were asymptomatic so he was diagnosed as having AA (Figure 1A; Table 2). The index case of family B also presented with features of AA. He had avascular necrosis of the hips, patches of hypo-pigmentation on his back and gray hair from the age of 13 years old, but no other signs of DC. He responded poorly to immunosuppressive treatment (anti-lymphocyte globulin (ALG) and cyclosporine) but has had some response to oxymetholone. His parents were clinically normal (Figure 1A; Table 2). TERC screening found two novel heterozygous substitutions c.178G→A (G178A) and c.180C→T (C180T) in the index cases of family A and B, respectively (Figure 1A). Both these mutations are in the P3 pseudoknot stem of TERC. Subsequent analysis of the TERC gene in both sets of parents showed that they were normal with respect to these mutations. Family histories from both cases found no evidence of non-paternity. Analysis of ten STR loci in the two families showed consistent segregation of alleles. This indicated a relative chance of kinship of over 99.999% in both cases, providing virtual proof of the declared paternity. We, therefore, concluded that the TERC mutation arose as a de novo event in both of these cases. G178A was also detected in the buccal smear of the index case indicating that this muta-
De novo autosomal dominant TERC mutations reduce telomerase activity by haplo-insufficiency

Telomere length measurements established that the two index cases had reduced Δtel values of -2.16 and -4.51 for G178A and C180T, respectively (Figure 2). These results placed the index case of both family A and B below the 90% deviation which suggests that these telomere lengths are below the 5th percentile when compared to the normal control panel. The telomere lengths of the parents of both families were located within the 68% percentile or higher which is between the 50th and 68th percentile when compared to the normal control panel (Figure 2).

The functional consequences of these two novel TERC substitutions were investigated by telomerase reconstitution analysis in telomerase-negative cells. The telomerase activity was found to be reduced to approximately 10% and 25% of WT control levels for G178A and C180T, respectively (Figure 1B). Furthermore, no significant evidence of a dominant negative effect was found when either TERC mutation was mixed in equal concentration with WT TERC (Figure 1C). These data, therefore, show that the presence of either novel substitution is capable of reducing telomerase activity via haplo-insufficiency and leads to prematurely reduced telomere lengths.

Additional functional characterization of de novo TERC mutations shows that the primary sequence is as critical as the secondary structure for normal activity

In previous investigations of TERC stem mutations, the reduction of telomerase activity was restored when the stem structure was reconstituted.7A15 When we investigated the two de novo TERC stem mutations, we observed that the two mutations acted in a different manner (Figure 3). G178A had significantly reduced telomerase activity compared to WT controls (Figure 1B, Figure 3), while telomerase activity was close to WT levels when either the complementary stem mutation c.112C→T (C112T) was present alone (Stem in Figure 3) or when both of these mutations were present in cis, restoring the stem structure (Double in Figure 3). For C180T, the presence of the original mutation, the stem mutation (c.110G→A (G110A)) or both mutations in cis (Double in Figure 3) showed reduced telomerase activity.
in comparison to WT controls. This suggests that the primary sequence at positions G178 and C180 is as critical as the intact secondary stem structure at this location.

**Other novel TERC mutations in patients with diverse clinical presentations**

In family C, a girl presented at 6 years old with chronic iron deficiency anemia and dysphagia. Her BM was markedly hypocellular with no evidence of myelodysplasia (MDS) associated with peripheral cytopenia (Table 2). This patient had congenital deafness, learning difficulties, short stature, scoliosis, dental abnormalities and dental caries. She had normal nails but spider naevi on her neck. Her family history is noteworthy: two of her four brothers had congenital deafness and schizophrenia, one of whom had AA and died from carcinoma of the tongue. A third brother died following a BM transplant for transforming MDS. Her fourth brother has normal hearing, a normal blood count and is not schizophrenic. Her mother was also deaf and her father has normal hearing, a normal blood count and is not associated with constitutional AA. Eventually he became refractory to oxymetholone and underwent reduced intensity matched unrelated BM transplant at the age of 25. This was associated with complications (recurrent gastrointestinal bleeding, ascites, infections and pulmonary complications) and he died 2 years after the transplant. Post-mortem examination showed an abnormal liver with focal nodular hyperplasia and oesophageal varices. Pulmonary histology showed cryptogenic organizing pneumonia and obliterator bronchiolitis. The patient’s father had cryptogenic fibrosing alveolitis for which he had a successful heart/lung transplantation. The patient’s paternal grandmother had a history of anaemia and was diagnosed with fibrosing alveolitis which was treated unsuccessfully with steroids and other immuno-suppressive agents. She was described as having *dry skin* and died at

heterozygous for a c.110-113delGACT deletion located within the conserved helix P3 stem domain of the TERC pseudoknot. This TERC mutation has not been identified among healthy individuals including the fourth asymptomatic brother but has been observed in another family with autosomal dominant DC. The family was described to have non-severe AA, elphin appearance and significantly reduced telomerase activity via haplo-insufficiency. The index case of family C has a reduced Δtel value of -5.32, finding that is complementary to the results of the TRAP analysis (Figure 5A). When compared to the normal telomere length range, this index case is located well below the 95% deviation range which suggests that this telomere length is within the 1st and 5th percentile when compared to the normal control panel (Figure 2).

The index case in Family D presented with AA in childhood. His blood count was maintained with oxymetholone for several years and he was diagnosed as having constitutional AA. Eventually he became refractory to oxymetholone and underwent reduced intensity matched unrelated BM transplant at the age of 25. This was associated with complications (recurrent gastrointestinal bleeding, ascites, infections and pulmonary complications) and he died 2 years after the transplant. Post-mortem examination showed an abnormal liver with focal nodular hyperplasia and oesophageal varices. Pulmonary histology showed cryptogenic organizing pneumonia and obliterator bronchiolitis. The patient’s father had cryptogenic fibrosing alveolitis for which he had a successful heart/lung transplantation. The patient’s paternal grandmother had a history of anaemia and was diagnosed with fibrosing alveolitis which was treated unsuccessfully with steroids and other immuno-suppressive agents. She was described as having *dry skin* and died at
the age of 64 from respiratory failure. The index case’s younger sister also developed AA (Figure 4; Table 2). The 35nt index case’s sister and father were found to have a novel heterozygous deletion spanning nucleotides 53 to 87 of TERC, while his mother was normal. This deletion removes the highly conserved alignment region of the RNA template as well as the P2a.1 stem of the pseudoknot. TRAP analysis of this deletion showed that telomerase activity was abolished in comparison to that in normal controls with no evidence of a dominant negative effect on WT TERC (Figure 5). Due to the nature of this deletion, it is not possible to determine whether the loss of telomerase activity is due to the inability of the resulting complex to align itself on the telomere and/or the loss of the P2a.1 stem. Telomere length data suggest that this deletion is capable of reducing telomerase function in vivo as the father has Δtel values of -1.60 and the sister has Δtel values of -2.82 in comparison to the normal population (within 68% and 90% deviation, respectively, which relates to the 5th percentile, in Figure 2).

The index case from family E presented with non-severe AA at the age of 25 years (Table 2). He had no obvious clinical features suggestive of DC but was put forward for a TERC screen in an attempt to rule out this diagnosis since he had failed to respond to immunosuppressive therapy. His blood count responded to oxymetholone treatment. His sister and mother were both asymptomatic with normal blood counts although his sister had some patches of skin hyper-pigmentation and spider naevi. His father died at 51 years old from a myocardial infarct (Figure 4).

TERC screening showed that the index case, mother and sister were heterozygous for a novel c.2G→C (G2C) substitution. Telomere length analysis in family E showed that only the two siblings have short telomeres that are below the 5th percentile when compared to the normal control panel (Figure 2). TRAP analysis revealed that this substitution had apparent WT activity when compared to normal controls, with no evidence of a dominant negative effect (Figure 5). The TRAP data suggest that the G2C mutation is a rare polymorphism although the possibility that it could act as a disease risk factor cannot be completely ruled out. In this respect, it is intriguing that the asymptomatic sister, like the index case, has very short telomeres.

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**Discussion**

We describe the functional characterization of four novel heterozygous mutations in TERC: G178A, C180T, 53-87del, G2C and a recurrent TERC mutation (110-113delGACT). G178A and C180T are the first reported de novo TERC mutations that result in a definitive pathophysiology of AA with indications of additional characteristics of DC (Figure 1A). As with previously reported TERC mutations identified in AA/DC families, telomerase functional analysis showed that both of these substitutions were capable of reducing telomerase activity in reconstituted telomerase assays (Figure 1B). The surprising aspect of these results however was the unique observation that these mutations produced a phenotype in the first generation. This feature has not been previously described in TERC families, in which disease anticipation has been observed. One possible explanation of this new phenomenon was that these particular TERC mutations exerted a dominant-negative effect on telomerase activity. This theory was rejected since further analysis found that the defective telomerase induced by these substitutions appeared to result from haplo-insufficiency in these experiments (Figure 1C). Therefore the severity of clinical phenotype in the first generation in these two case studies is not easily explained. One possible explanation is that although telomerase activity can be reconstituted in vitro, these mutations still have some dominant negative effect on telomere maintenance in vivo. Further studies are required to determine whether this is the case.

The G178A and C180T mutations, along with the previously reported 96-97delCT, GC107-8AG, 110-113delGACT, C116T, and A117C, lie close to each other both in the primary sequence and the secondary structure of TERC. This stem region has over 80% shared conservation with other TERC RNA species, as defined by mutagenesis and phylogenetic analysis, and is stabilized by a triple helix that surrounds the two stems (Figure 6). Although the mutations are contained within the minimal pseudoknot domain, they appear to function in different ways. One report suggests that the P3 helix, and not the intraloop base-pairings, is essential for telomerase activity, where the sequence of the J2b/3 loop rather than its base-pairing ability within the loop region is required for activity. As these two substitutions localize to an area of the pseudoknot domain that involves a proposed triple helix structure, it is possible that the physical location of these novel substitutions disrupts molecular switching in a different way to that of previously reported TERC mutations (Figure 6).

Another hypothesis is that functional differences within the P3 stem region could correlate with the ability of the resulting mutant TERC to either dimerize with other TERC molecules or telomerase-associated proteins. In vitro research shows that human telomerase appears to be highly processive such that a single telomerase enzyme usually continues to extend a single telomere rather than the enzyme being released at the end of each round of repeat extension. It has been hypothesized that the conserved TERC pseudoknot structure is required for template recognition and for defining the template boundaries for telomere synthesis. TERC control over processivity could occur through base pair interactions between the telomeric
The only other recurring mutations had been described in the 53-87del mutations result in a variety of BM failure syndromes? The patients reported here do not present with the classical triad of mucocutaneous features and therefore would not be classified as having DC based on clinical criteria. It is possible that these additional clinical features will present over time, but this is not always the case.

The G2C change identified in family E had no observable effect on telomerase activity in vitro (Figure 5). It is located in the 5' region of TERC, which shows low sequence conservation among vertebrates. Prior to this report, five other TERC mutations had been described that appeared to have no effect on TRAP activity. Two of these (G58A and G228A) are polymorphic, but three others (28-34del, A37G and G450A) have been observed only once in patients with AA, DC (in trans to a deleterious 216-229del mutation) and severe AA. Although it may well be that these TERC mutations are

ciliate, yeast and human TERC templates. In some cases template mutations appear to alter the in vitro and in vivo nature of the telomerase enzyme. The 53-87del TERC mutation identified in family D removes a significant proportion of the template and pseudoknot domain in affected members. Even with such a dramatic loss of TERC structure, affected members of this family initially presented with pulmonary fibrosis (grandmother and father of index case) rather than the classical mucocutaneous features of DC. The finding of pulmonary disease as the primary presentation of disease in two members of this family suggests that screening for TERC mutation should be conducted in other patients presenting with cryptogenic/idiopathic pulmonary fibrosis.

A surprising result from this subset of patients was the discovery of a recurring pseudoknot deletion, 110-113delGACT, in family C. Comparisons between a previously reported family and family C in this study show that this 4bp deletion in TERC can induce a variety of clinical phenotypes, ranging from AA to MDS or AML. These findings suggest that it might be useful to study the TERC gene, not only in patients presenting with AA or MDS, but also in patients presenting with acute leukemia, especially since 10-30% of AA patients develop AML and/or MDS. This is the second report of a conclusively pathogenic recurring TERC mutation. The C116T mutation has also been reported in two apparently unrelated families in which the index cases presented with thrombocytopenia or were diagnosed as having acquired AA. The only other recurring TERC mutations that have been described to date are G58A and G228A, which have been found to be polymorphic, and C-99G, the functional significance of which is not entirely clear. This highlights the diversity of clinical phenotypes resulting from TERC mutations. This also raises the semantic question of whether mutations in TERC result in DC, whose clinical phenotype can vary between cryptic presentations similar to AA through to more severe disease presentations, or whether TERC mutations result in a variety of BM failure syndromes?

The G2C change identified in family E had no observable effect on telomerase activity in vitro could be due to fact that G178 and C180 are localized to the internalized strand of the triple helix, in comparison to nucleotides G107, C108, C116, and A117 which rest on an external strand of the triple helix (Figure 6). Further studies are required to fully elucidate TERC pseudoknot function and the resulting telomerase complex activity in the presence of WT and/or TERC mutations. The identification of template and pseudoknot TERC mutations (53-87del and 110-113delGACT) enhances the observation that disruption of telomerase activity and resulting haplo-insufficiency can cause a variety of clinical presentations. Single-point substitutions, deletions, insertions and complete substitutions have been previously made in
References


