Detection of alternative lengthening of telomeres by telomere quantitative PCR

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ABSTRACT

Alternative lengthening of telomeres (ALT) is one of the two known telomere length maintenance mechanisms that are essential for the unlimited proliferation potential of cancer cells. Existing methods for detecting ALT in tumors require substantial amounts of tumor material and are labor intensive, making it difficult to study prevalence and prognostic significance of ALT in large tumor cohorts. Here, we present a novel strategy utilizing telomere quantitative PCR to diagnose ALT. The protocol is more rapid than conventional methods and scrutinizes two distinct characteristics of ALT cells concurrently: long telomeres and the presence of C-circles (partially double-stranded circles of telomeric C-strand DNA). Requiring only 30 ng of genomic DNA, this protocol will facilitate large-scale studies of ALT in tumors and can be readily adopted by clinical laboratories.

INTRODUCTION

Telomeres, specialized structures of repetitive DNA sequence (5’-TTAGGG-3’) located at the ends of chromosomes, undergo progressive shortening in replicating cells, which prevents the unlimited proliferation of normal somatic cells. Cancer cells evade this barrier and become immortalized by activating one of the two known telomere length (TL) maintenance mechanisms. These are the telomerase enzyme and alternative lengthening of telomeres (ALT), which synthesize new telomeric DNA from a RNA template via reverse transcription and from a DNA template via homologous recombination-mediated DNA replication, respectively (1). Telomerase activity (TA) is most commonly measured by the PCR-based telomere repeat amplification protocol, and a large number of cancers have been screened for TA by this method (2). In the absence of an enzyme activity assay for ALT, ALT has been detected in tumors by observing telomere-related phenotypic characteristics that have previously been documented in immortalized TA-negative cell lines (3). However, detecting ALT in tumors is significantly more challenging than in cell lines as most assays used to identify these phenotypic features in cell lines cannot be applied to tumors. The ability to diagnose ALT efficiently in clinical settings will facilitate the appropriate selection of telomere maintenance mechanism-targeted drugs in the clinic.

There are currently two main methods for detecting ALT in tumors. The first of these is terminal restriction fragment (TRF) Southern-blot analysis of TL profile. In ALT cells, telomeres range from very short to extremely long in individual cells, with mean TL (>20 kb) being about twice of that of TA-positive or normal somatic cells (5–10 kb) (4). ALT cells are therefore described as having long and heterogeneous telomeres as compared to the shorter and more homogeneous telomeres of ALT-negative cells. The second method is detection by a combined promyelocytic leukemia (PML) immunofluorescence/telomere fluorescence in situ hybridization (FISH) analysis of tumor sections for ALT-associated PML bodies (APBs) (5). APBs are PML bodies that contain telomeric DNA and telomere binding proteins (6). Both of these conventional assays are labor-intensive, making screening of large numbers of tumors challenging. Recently, a variation of the APB assay in which telomere FISH was used to detect ultra-bright telomeric signals was used to analyze more than 6000 tumor specimens, mostly in tissue microarray format (7).

An assay, which reliably detects ALT in cell lines by quantitating C-circles (CC) (8), is currently being assessed for its suitability for detecting ALT in tumors.

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CC are extra-chromosomal circles of telomeric DNA that are partially single-stranded, where the C-rich strand is complete and the G-rich strand is gapped. The presence of abundant CC is ALT-specific, and CC levels were shown to have a quantitative relationship to ALT activity (8). The CC assay is an isothermic polymerase reaction, where CC act as self-primering DNA templates in rolling circle amplification, producing long linear telomeric single-stranded DNA (ssDNA) products. These products are then quantitated by dot-blot analysis using a $^{32}$P-labeled telomeric probe (8).

In this study, a strategy utilizing TL measurement by quantitative PCR (qPCR) was developed and evaluated for detecting ALT in cancer cell lines and tumors. While telomere qPCR has been widely applied as a high throughput technique in population studies to measure TL in leukocyte DNA and is capable of detecting small differences (9–11), it has not been used previously to detect ALT. Here, we demonstrated that telomere qPCR could be employed to quantify both TL and CC concurrently with just 30 ng of DNA and in significantly less time than is required for TRF and APB analysis. Furthermore, we showed that examining two ALT characteristics (TL and CC) concurrently in this assay significantly improved ALT detection in tumors, compared with evaluating just one feature, as 50% of the ALT tumors were found to be positive for only one of the two characteristics.

MATERIALS AND METHODS

Samples

Frozen tumor samples were acquired with institution ethics committee approval. The 23 cell lines include CHLA-90, CHP-100, CHP-134, G-292, GM847, IIICF/c, LAN-2, LAN-5, MCF-7, MDA-MB-231, MeT-4A, MG-63, NB69, NM39, NM179, SK-LU-1, SK-N-AS, SK-N-FI, SH-SY5Y, SK-N-BE2, SK-N-DZ, U-2 OS, and W-V. SK-N-AS, SK-N-FI, SH-SY5Y, SK-N-BE2, SK-N-DZ, U-2 OS, and W-V. SK-N-FI, U-2 OS, and G-292 used for calibration measurements were purchased from ECACC at passage 37, + 4 and 24, respectively. The cell lines MeT-4A, MG-63, NB69, NM39, NM179, SK-LU-1, SK-N-AS, SK-N-FI, SH-SY5Y, SK-N-BE2, SK-N-DZ, U-2 OS, and W-V. SK-N-FI, U-2 OS, and G-292 were included.

DNA extraction and quantitation

Genomic DNA from both cell lines and tumors was extracted by lysis at 37°C with 2% sodium dodecyl sulfate (SDS) buffer containing 50 mM Tris, 20 mM ethylenediaminetetraacetic acid (EDTA) and 200 μg/ml Pronase protease (Sigma), followed by precipitation with 5 M sodium chloride and ethanol. DNA was quantitated with a Qubit Fluorometer (Invitrogen).

TRF analysis

TL measurement by TRF Southern-blot analysis was performed as previously described (12). Briefly, genomic DNA was digested with HinFI and RsaI restriction enzymes (4 U/μg DNA) [New England Biolabs (NEB)] and 25 ng/μg RNase (Roche) and resolved on a 1% agarose gel using pulsed-field gel electrophoresis. The dried gel was denatured, neutralized and hybridized to an end-labeled $^{32}$P-(CCCTAA)$_4$ telomeric probe. The phosphor screen was scanned on a Typhoon Trio Variable Mode Imager (GE Healthcare) and signal intensity quantitated using the LAS 4000 Multi Gauge software.

CC assay

Rolling circle amplification of CC was performed as described (8). A 10 μl CC reaction contains 0.2 μg/ml bovine serum albumin, 0.1% Tween, 4 μM dithiothreitol (DTT), 1 mM each dATP, dCTP, dGTP, dTTP, $\phi$29 DNA polymerase ($\phi$29, 3.75 U/16 ng DNA) (NEB), 1× $\phi$29 buffer and 16 ng genomic DNA, and was incubated at 30°C for 8 h then at 65°C for 20 min. For each sample, the assay was done with and without $\phi$29.

To quantitate CC assay products by slot-blot analysis, 32 ng of DNA was used for each CC assay. The 20 μl CC assay product was then diluted with 100 μl of 2× saline-sodium citrate (SSC) and slot-blotted in duplicate onto a Biodyne B nylon membrane (Pall). After UV-cross linking, the membrane was hybridized overnight at 37°C with end-labeled $^{32}$P-(CCCTAA)$_3$, telomeric probe in PerfectHyb Plus buffer (Sigma) in native (non-denatured) condition. The membrane was then washed four times at 37°C in 0.5× SSC/0.1% SDS buffer. The phosphor screen was scanned on a Typhoon Trio Variable Mode Imager (GE Healthcare) scanner and signal intensity was quantitated using the ImageQuant software.

Quantitative PCR

The CC assay was performed with and without $\phi$29. For each sample, 10 μl of the CC assay product containing 16 ng of input genomic DNA was diluted with 30 μl of Tris–EDTA (10 mM Tris, 0.1 mM EDTA, pH 7.6), of which 5 μl containing 2 ng of input genomic DNA was used for each PCR. Each sample required 12 PCRs: triplicate telomere PCR of CC assay/16 ng of input genomic DNA and was incubated at 37°C with end-labeled $^{32}$P-(CCCTAA)$_3$, telomeric probe in PerfectHyb Plus buffer (Sigma) in native (non-denatured) condition. The membrane was then washed four times at 37°C in 0.5× SSC/0.1% SDS buffer. The phosphor screen was scanned on a Typhoon Trio Variable Mode Imager (GE Healthcare) scanner and signal intensity was quantitated using the ImageQuant software.

Green master mix (Qiagen), 10 mM Tris, 0.5 μl dimethyl sulfoxide, 5 μl DNA template and primer sets. The final primer concentrations were: (i) telomere: forward 300 nM and reverse 400 nM, (ii) VAV2: forward 700 nM and reverse 400 nM, (iii) and reverse CCATTCTATCATCAACGGGTACAA (9) PCR of CC assay/16 ng of input genomic DNA, and was incubated for 20 min. For each sample, the assay was done with and without $\phi$29.

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ATGAC and reverse ATCTGCCCTACCTTCTCAA. The 36B4 (RPLP0) gene, located on chromosome 12q24, encodes acidic ribosomal phosphoprotein P0 and VAV2, located on chromosome 9q34, encodes guanine nucleotide exchange factor. 36B4, which has been applied as a SCG for both cancer (13) and non-cancer cells (9,11), was used for melanoma tumors and cell lines, as well as glioblastoma multiforme (GBM) tumors and soft tissue sarcomas (STS). Comparative Genomic Hybridization studies indicate that 36B4 is an appropriate SCG for melanoma (14) and GBM (15). VAV2 was used for neuroblastoma (NB) tumors and all cell lines except for melanoma. VAV2 has previously been validated as a SCG for NB (16), whereas 36B4 is in a region of the genome where a low level of gene amplification has been reported in NB (17).

All PCRs were performed with the Rotor-Gene Q (Qiagen) real-time cycler. PCR conditions were: (i) telomere: 95°C for 15 min, 33 cycles of 95°C for 15 s and 54°C for 2 min, (ii) VAV2: 95°C for 15 min, 40 cycles of 95°C 15 s, 57°C 30 s and 72°C 1 min and (iii) 36B4: 95°C 15 min, 35 cycles of 95°C 15 s and 58°C 1 min. Rotor-Gene Q series software was used to obtain data and generate standard curves.

**Data analysis**

The unweighted mean TRF length was calculated as

\[ \frac{\sum (OD_i \times L_i)}{\sum ODi} \]

where ODi is the signal intensity above background within interval i and Li is the molecular weight (kb) at the mid-point of interval i. A sample was considered TRF+ (ALT+ by TRF) when mean TRF length was ≥16 kb and semi-interquartile range (SIR) was ≥4 kb (5). SIR of the TRF lengths was calculated as (3rd quartile–1st quartile)/2. For CC assay product quantitation by slot-blot analysis, CC assay level was calculated by subtracting the 32P-probe signal of the CC assay without p29 from that of the CC assay with p29. Results were expressed as average of duplicates and as arbitrary units (AU).

qPCR data were analyzed using the two standard curves method as previously described (9). Telomere product was normalized (norm-TEL) with the SCG product. The relative telomeric DNA content (TC) of a sample was the norm-TEL obtained from the CC assay without p29 (norm-TEL/p29–) and was relative to the TC of the ALT-positive U-2 OS cell line with an arbitrary value of 40. The relative CC assay level of a sample was calculated as [(norm-TEL/p29+)−(norm-TEL/p29−)] and was relative to the CC assay level of U-2 OS cell line with an arbitrary value of 170. A sample was only considered positive for CC when the norm-TEL value of each individual p29(+) triplicate was higher than the norm-TEL of all three of the p29(−) replicates and the calculated CC assay level was ≥cut-off value. All PCR results were expressed as mean of triplicate reactions ± SEM.

Means were compared using unpaired two-tailed Student’s t-test and medians compared by the Mann–Whitney test. Linear regression was analyzed using Prism 4 software (GraphPad). Sensitivity was the number of samples that tested positive divided by the number of samples that were classified as positive by the ‘gold standard’. Specificity was the number of samples that were tested negative divided by the number of samples that were classified as negative by the gold standard. TRF analysis was the gold standard when the performance of telomere qPCR was compared with TRF and 32P detection method the gold standard when telomere qPCR was compared with 32P for quantitation of CC assay level. Cut-off values were chosen to provide the highest concordance rate (both tests in agreement) and the best separation between the test-positive and test-negative group.

**RESULTS**

**Distinguishing ALT+ and ALT− samples by TC**

In this study, 66 samples, consisting of 23 cell lines and 43 tumors (18 NB, 15 GBM and 10 melanomas), were analyzed initially. The telomere maintenance mechanism status has previously been published for 16 of the cell lines (Supplementary Table S1). TRF Southern-blot analysis was used to measure TL. As previously described, a sample was classified as ALT+ by TRF when the mean TRF length is ≥16 and SIR ≥4 kb (5). TC was measured by telomere qPCR. We found a very good correlation (\( R^2 = 0.8531; P < 0.0001 \)) between TC and mean TRF (Figure 1a). Importantly, samples that were ALT+ by TRF (\( n = 14 \)) had significantly higher relative TC than TRF− (ALT− by TRF) samples (\( n = 52; \) median 19.4 versus 4.1; \( P < 0.0001 \)). Using a cut-off of 12.0, TC separated the samples distinctly into TRF+ and TRF− groups (Figure 1b), with a sensitivity of 100% (95% CI: 74–100%) and specificity of 100% (95% CI: 92–100%) (Figure 2).

**Detection of CC assay products by radioactive telomeric probe**

The CC assay is the first ALT assay that appears to have a quantitative correlation with ALT activity levels (8). The CC assay utilizes p29 DNA polymerase to produce long G-rich telomeric ssDNA through rolling circle amplification of telomeric CC that are abundant in ALT cells. CC assay products are routinely detected by a 32P-labeled telomeric probe in a dot-blot analysis. We found that TRF+ samples (\( n = 14 \)) had significantly higher CC assay levels than the TRF− samples (\( n = 52; \) median 352 versus 5.1 AU; \( P < 0.0001 \)). The CC assay levels of all samples were distributed into two clusters that corresponded well to the TRF status (Figure 3a). Ninety-two percent (12 of 13) of the samples in the upper cluster were TRF+ and 96% (51 of 53) in the lower cluster were TRF−. The cut-off for ALT detection was set at 35 AU, which represents the mid-point of the interval between the smallest value of the upper cluster and the largest value of the lower cluster (Figure 3a). With a cut-off of 35.0 AU, the CC assay by 32P detection (CC[32P]) had a 95% concordance with the TRF assay (95% CI: 87–99%; Figures 2 and 3a). The three discordant samples were NB tumors. NB9 and NB11 that were TRF+/CC− had CC assay levels in the range of TRF− samples, despite having high relative TC (average 21.7) and a high mean TRF.
This suggests that not all TRF + samples are CC +. The one TRF/C0/CC + sample (NB18) had borderline mean TRF (14.9 kb)/TC (11.7) and CC[32P] assay level of 52.4 AU (Figure 3b).

Correlation between detection of CC assay products by 32P and telomere qPCR

We next examined whether CC assay products could be measured by telomere qPCR (CC[qPCR]). Genomic DNA was subjected to CC assay with and without [32P]. The TC present in the CC assay with no added [32P] reflects the TC of the input genomic DNA since there is no rolling circle amplification. In comparison, the TC derived from the CC assay with [32P] reflects both TC from the input genomic DNA and the G-rich linear telomeric ssDNA synthesized from rolling circle amplification of CC. Therefore, the difference in TC between the paired [32P]+ and [32P]− samples represents CC assay products. As shown by the standard curves in Figure 4a, the presence of [32P] results in the same reduction in C_T value over a 100-fold range of DNA concentrations for the telomere qPCR, but not for the 36B4 SCG qPCR. The robust standard curves demonstrating PCR amplification efficiencies close to 100% with excellent correlation (Figure 4a and b) support the use of telomere qPCR for measuring TC and CC assay products. Furthermore, there was very good correlation between CC[32P] and CC[qPCR] measurements (R^2 = 0.9093; P < 0.0001; Figure 4c).

Telomere qPCR detects CCs in ALT cells

For CC assay level measurements, we compared the performance of CC[qPCR] with CC[32P] in identifying TRF + samples. Samples that were CC[32P]+ (CC assay level ≥35 AU) had significantly higher CC[qPCR] assay levels than CC[32P]− samples (median 146 versus 0; P < 0.0001) (Figure 5a). The only discrepancy between the two methods occurred with a GBM tumor (GBM 6a) that was CC[qPCR]−, but had a CC[32P] assay level of 79
AU and mean TRF of 20.1 kb/SIR 8.0 kb (Figure 3b). The good correlation ($R^2 = 0.9342$) between the two methods was maintained when the analysis was applied to the 12 samples that were CC[qPCR]+ (CC assay level ≥7.5) (Figure 5b). In addition, samples that were TRF+ had significantly higher CC[qPCR] assay levels than TRF− samples (median 137 versus 0; $P < 0.0001$) (Figure 5c). Applying a cut-off of 7.5, CC[qPCR] has a concordance of 94% (95% CI: 85–98%) with the TRF assay for ALT, which is similar to CC[32P] detection [concordance of 95% (95% CI: 87–99%); Figure 2].

**Validation and calibration of the assay**

To validate the assay, we applied the cut-offs for relative TC and CC[qPCR] to an independent cohort of 15 STS. The ALT status of these STS samples has previously been determined in frozen sections using the APB assay (5). There was 100% (95% CI: 76–100%) concordance between TRF and TC, as well as between CC[32P] and CC[qPCR] (Figure 6a). Of these 15 samples, 5 were CC+ with 100% (95% CI: 76–100%) concordance between APB and CC[qPCR] status (Figure 6a). Of note, the sample that was found to have low frequency of APBs also had the lowest CC[qPCR] assay level of 9.4 in this study. Of the five STS that were CC+, three were TRF/TC+ and the remaining two were TRF/TC− (TRF 14.0 kb/TC 10.3/CC 261 and TRF 7.4 kb/TC 4.9/CC 9.4). This suggests not all CC+ samples have very long telomeres.

Furthermore, we have provided calibration parameters derived from three commercially available ALT cell lines (SK-N-FI, U-2 OS and G-292) at specific passages to facilitate this qPCR-based assay being used by other laboratories (Figure 6b).

**DISCUSSION**

ALT is one of the two known telomere maintenance mechanisms, which play an essential role in the immortalization of cancer cells. While the prevalence and prognostic implications of TA have been reported for a large number of human cancers (2), equivalent data are not yet available for ALT. This is primarily because conventional methods for detecting ALT in tumors are labor-intensive. In addition, the quantity of DNA required for TRF analysis may exceed what is available, especially when the tumor of interest is not treated initially by resection and only a small amount of biopsy material is obtained for clinical diagnosis. A less laborious detection method that requires significantly less tumor material is needed for studying ALT in human cancers.

In this study, we developed a new strategy based on the well-established telomere qPCR and the more recently described CC assay to detect two characteristics of ALT in a cohort of 81 samples. It is a two-step procedure whereby genomic DNA is first subjected to rolling circle amplification by $j_29$, followed by telomere qPCR analysis of the paired amplified/non-amplified samples. By this means, both the TC and the CC[qPCR] assay level can be obtained from 30 ng genomic DNA. We demonstrated good correlation between TC and mean TRF length, as has been reported in population studies (9). We use the term TC, instead of TL, in this context because in ALT cells the abundant extra-chromosomal telomeric repeats also serve as templates for the qPCR. TC distinguished TRF+ and TRF− samples with 100% sensitivity and specificity and therefore can substitute for conventional TRF Southern blots. In addition to consuming large amounts of DNA (1–2 μg), TRF analysis can yield a falsely high mean TRF if care is not
taken to obtain adequate restriction enzyme digestion of the genomic DNA.

To obtain sufficient sensitivity, CC assay product (long linear telomeric ssDNA) has previously been detected with 32P-labeled probes (8) and with this technique assaying duplicate samples with and without φ29 amplification requires 120 ng of DNA, compared with 30 ng for the CC[qPCR] detection method. The correlation between the two quantitation methods was very good. There was one discrepant result in the entire cohort: a GBM tumor that was considered to be CC[32P]+ and CC[qPCR]/C0. This tumor had a 9-fold lower CC[32P] assay level than the mean CC[32P] level of TRF+ samples (79 versus 693 AU), while its relative TC of 23.2 was similar to the mean TC (21.6) of TRF+ samples. This discrepancy between the two detection methods may be related to the difficulty in telomere PCR detecting very low levels of CC due to interference from the high levels of double-stranded TC. The low amount of CC assay telomeric product representing only a very small proportion of the high TC may lead to insignificant differences between the paired φ29(+) and φ29(−) samples. It is also relevant to note that this tumor had a somewhat atypical TRF profile with a mean TRF of 20 kb, but with much less heterogeneity (lowest SIR of 8 kb) than the other 13 ALT-positive samples in this study which had a mean SIR of 17.7 kb (range: 11.1–23.4 kb) (Figure 3b). Overall, qPCR is a reliable quantitation method for CC assay levels and has advantages over the original CC assay protocol (8) as this assay requires less DNA, takes less time and does not involve radio-isotope.

To validate the cut-offs for TC and CC[qPCR], we applied them to an independent cohort of 15 STS with known APB status (5). There was 100% concordance between TRF and TC, between CC[32P] and CC[qPCR] and between APB and CC[qPCR]. This verifies these cut-offs were appropriately set. We found discrepancies between TRF/TC and CC assays for ALT in both the original and validation cohort. Such discrepancies have also been reported in cell line studies. Telomerase overexpression in ALT− cells can cause long heterogeneous telomerases (18) while short telomerases have been
observed in an ALT+ cell line (19). In both cases, the CC assay was the more accurate determinant of ALT status (8). This study is the first report of tumors with long and heterogeneous TL but without elevated levels of CC. We found this in two of the four TRF+ NB tumors and both of these TRF+/CC− NB were APB negative. Conversely, we found a STS in the validation cohort that was CC+ without long telomeres (TRF 7.4 kb/TC 4.9/CC 9.4). This STS was previously reported to have low APB frequency, which correlates well with the low CC[qPCR] assay level. There appears to be a good correlation between APB and CC for ALT detection; however, the APB assay is significantly more labor intensive. The CC assay has been studied extensively in cell lines, but validation of the accuracy of the CC assay in a variety of tumor types is needed.

By combining two tests evaluating different ALT characteristics in this protocol, the problem of equivocal results can also be minimized. For example, in this study, two tumors (NB and STS) with equivocal mean TRF/TC have unequivocally positive CC[qPCR] levels of 52 and 261, respectively. Assessing embryonal tumors such as NB for ALT by TRF or TC can be particularly challenging as the separation between the two groups is often not as distinct as in adult tumors, possibly because less telomere attrition occurs during the genesis of embryonal tumors. In NB, we found that ALT− samples had an average TC of 7.3 and TRF of 11.2 kb, compared with GBM and melanoma that had much lower average TC (4.4) and TRF (8.5 kb). Furthermore, of the 58 tumor samples analyzed, six have TC in the mid-range (TC 8–12), of which two were CC-positive and four were CC-negative. The CC assay is therefore required to segregate tumors with mid-TC levels, which can represent 10% of tumor samples.

Evaluating two characteristics simultaneously also increases both the sensitivity and specificity for ALT detection. In this study, we identified 11 tumors that had TC and/or CC[qPCR] levels above the cut-off for ALT (six from initial cohort and five from validation cohort). However, only five samples had both TC and CC[qPCR] levels above the cut-off. The other six samples had only one parameter above the cut-off (three TC+/CC−; three TC−/CC+). Although the significance of only one ALT...
characteristic being positive is not yet clear, the data suggest that examining only one ALT characteristic can potentially miss 25–30% of ALT+ tumors.

The accuracy of qPCR using genomic DNA as the DNA template relies on appropriate choice of SCG. SCG normalizes for input DNA amount and PCR efficiency. 36B4, albumin and β globin are the most widely used SCG in non-cancerous cells (10,20). However, selecting SCG in cancer cells can be challenging as gene copy number variations are common in cancer cells and there are significant differences in affected chromosome regions among different cancers. Fortunately, comparative genomic hybridization array data are now widely available for most tumor types and can direct the selection of SCG. Care must also be taken when evaluating the quantity of SCG product in each sample to ensure the results are consistent with the behavior of a SCG.

Furthermore, qPCR requires normalization to a reference sample. In the setting of this assay, the cut-offs can only be applied when normalization is accurate. We therefore provided relative qPCR measurements obtained from three commercially available ALT cell lines at specified passage. This allows the user of this qPCR-based assay to perform multi-point calibration for both TC and CC measurements prior to testing unknown samples and to ensure that the cut-off values, normalized to U-2 OS, can be accurately applied.

In addition to research applications, this ALT detection protocol can be readily adopted by clinical laboratories. Many clinical laboratories routinely perform qPCR now, but are not equipped for Southern-blot analysis and for techniques involving radiation. The ability to diagnose ALT efficiently in both research and clinical settings will be crucial for the development of ALT inhibitors as anticaner therapies and for choosing the appropriate telomere maintenance mechanism-targeted drugs in the clinic. As more tumor types are screened for ALT, the prognostic implications of ALT will also be better known. Moreover, CC have been detected in the serum of patients with ALT+ tumors (8), indicating that CC circulate as tumor DNA in the blood stream and are therefore a potential biomarker for ALT activity.

In conclusion, this novel protocol is the first to detect ALT by telomere qPCR and is feasible in clinical pathology laboratories. This radio-isotope-free assay has advantages over the original CC assay (8), which requires more DNA for the detection of one characteristic (CC), uses radioactive probe and is more labor-intensive. The concurrent evaluation of two distinct ALT characteristics increases the sensitivity and specificity of ALT detection and the format of the assay together with its requirement for only a small amount of DNA will facilitate medium-throughput screening of large tumor cohorts for ALT.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online: Supplementary Table S1 and Supplementary References [21–25].
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Conflict of interest statement. J.D.H. and R.R.R. have applied for a patent regarding the CC assay.

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