Detection of Meningeosis neoplastica by real-time quantitation of telomerase activity

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Abstract

Background: Analysis of cerebrospinal fluid (CSF) to discriminate between benign and malignant conditions is of fundamental importance for the physician and the patient because of the differential therapeutic options and resulting morbidity and mortality. Most human tumours demonstrate increased telomerase activity (TA). Recent technical advances in the detection of TA allow for sensitive and specific detection within 4 h. Thus, the detection of TA is suitable for routine clinical testing.

Methods: This study examines TA in cellular proteins in CSF from 111 patients compared to cytomorphological and laboratory examination.

Results: A positive result for TA in cellular proteins of CSF was correlated significantly with Meningeosis neoplastica, but not with non-malignant conditions.

Telomerase was not detected in CSF supernatant, despite positive results in cellular proteins from identical patients. Furthermore, a 48-h time delay during the pre-analytic processing is not critical for detection of TA detection in native CSF when stored at room temperature.

Conclusions: We conclude that TA is a promising marker for the detection of Meningeosis neoplastica and warrants further study.


Keywords: cerebrospinal fluid; Meningeosis neoplastica; real-time quantitative telomere repeat amplification protocol; telomerase activity.

Introduction

Cytological analysis of cerebrospinal fluid (CSF) is the standard procedure for the diagnosis of meningitis or Meningeosis neoplastica. The incidence of Meningeosis neoplastica diagnosed by cytology in a large non-selected series of 5951 specimens by Prayson and Fischler (1) was 5% in adults and 8% in paediatric patients. Despite the low incidence of Meningeosis neoplastica, discrimination between non-malignant and malignant cytology is of vital importance for the patient due to fundamentally differential therapeutic regimens (anti-infective, immunosuppressive or no therapy vs. anti-neoplastic therapy). Leptomeningeal involvement is present in about 8% of patients diagnosed with acute lymphatic leukaemia (2) and develops in about 5% of patients with solid tumours (3, 4).

In spite of substantial advances in prophylaxis and therapy, the detection of Meningeosis neoplastica usually requires an intensification of anti-neoplastic therapy and predicts an unfavourable outcome in patients resulting from increased disease and therapy related morbidity and mortality (5, 6). Alternative diagnostic technologies including gadolinium imaging (7) or mass spectroscopy (8) are inferior to CSF cytology. For lymphoid neoplasms, multiparameter flow cytometric analysis of CSF (9, 10) may be helpful as an adjunct to cytology, which remains the gold standard for the diagnosis of Meningeosis neoplastica (11).

The enzyme telomerase consists of a catalytic protein subunit human telomerase reverse transcriptase (hTert) and a ribonucleic acid (RNA) template human telomerase RNA (hTr). Telomerase is capable of telomere repeat elongation in vivo involving a unique reverse transcriptase activity, able to regulate telomere length in vivo (12). However, functionally active telomerase is restricted to embryonic stem cells (13), germ line cells (14), rarely in differentiated cells like...
lymphocytes or adult haematopoietic stem cells (15–17) and, notably, in neoplastic cells. Because telomeres progressively shorten with each cellular division as a sign of replicative senescence (18), stabilisation of telomeres by activation of telomerase is presumed to be a fundamental mechanism for indefinite proliferation in neoplasia. Up to 90% of human tumours express telomerase (19), whereas the remaining tumours have developed alternative mechanisms to stabilize their telomeres (20).

Here, we exploited the fact that neoplastic cells are characterised by highly active telomerase in order to facilitate a more rapid diagnoses of Meningeosis neoplastica. With the development of a fast and reliable real-time polymerase chain reaction (PCR) based methodology (real-time quantitative telomere repeat amplification protocol (RQ-Trap) assay as reported by Wege et al. (21)) the detection of telomerase activation may be potentially useful in clinical practice. We have previously shown that telomerase activity (TA) can be measured in lymphoblasts from peripheral blood or CSF using the RQ-Trap protocol (16). The present study was designed to assess the feasibility and diagnostic accuracy of measuring TA on a routine basis for the detection of Meningeosis neoplastica in diagnostic CSF.

Materials and methods

The study was approved by the Local Ethics Committee and all study participants gave written informed consent. CSF was drawn from patients undergoing routine lumbar puncture and was centrifuged at 800 g for 5 min, within 30 min following collection of CSF. The cell pellet and the supernatant were stored at −80°C until further analysis. Samples for TA assays were extracted following standard protocols (22). Cytological examination was performed by an expert cytopathologist who was not aware of other laboratory test results.

Telomerase activity

TA was quantified under real-time PCR conditions utilizing the RQ-Trap procedure (Lightcycler instrument, Roche, Mannheim, Germany) as reported by Wege et al. (21) with minor modifications. In brief, duplicates of protein extracts from 1000 cells or CSF supernatant were mixed with 0.1 μg of Primer TS (template) and 0.05 μg Primer ACX (23) in 20 μL Lightcycler Fast Start SYBR Green PCR Master Mix (Roche, Mannheim, Germany) containing 1.5 mM MgCl₂. The mixture was incubated at 30°C for 30 min to allow template elongation by TA. Immediately following transfer to the Lightcycler instrument, telomerase was heat-inactivated and hotstart DNA polymerase activated by incubation at 95°C for 10 min. Thirty-six cycles of amplification were performed with 20 s denaturation at 95°C, annealing for 30 s at 60°C, and elongation for 50 s at 72°C. TA was displayed as relative TA of 1% (equally to a TA of 10/1000 HEK293 cells). Therefore, the cut-off for a positive test result was defined as a relative TA of 1% (equally to a TA of 10/1000 HEK293 cells). Heat inactivated lysate from 1000 HEK293 cells served as a negative control and telomerase positive CD19⁺ lymphoblasts (16) served as the positive control. Quantitative levels of TA were obtained from cellular proteins of 1000 cells, serial dilutions of cellular proteins or supernatant.

Tissue culture

Cell lines (Jurkat, Daudi, Karpas, K562, L540 and HEK293) were purchased from ATCC (Wesel, Germany) and grown in RPMI 1640 (supplemented with 10% fetal calf serum, glutamine and penicillin and streptomycin, all from Gibco, Karlsruhe, Germany). Lines were maintained in a 37°C, 5% CO₂ atmosphere and divided split two times per week.

Statistical analyses

Graphs and statistical analyses were performed using SigmaPlot (Version 7.0) and SigmaStat (Version 2.03; both from SPSS Inc, Chicago, IL, USA). Repeated measure ANOVA was used for the analysis of TA at different time points during the analysis experiment. Differences in TA from cellular proteins or tissue culture supernatant were analysed using the unpaired Student’s t-test. Correlation between (telomerase-) elongated template and cycle number was determined with linear regression analysis.

Results

TA determined by RQ-Trap assay

Serial dilutions of HEK293 cells were assayed for TA to generate a standard curve for comparison with other cell lines and clinical samples. The RQ-Trap assay showed reproducible linearity from 1000 to 8 HEK293 cells ([Figure 1: R² = 0.99, Figure 2 see online Supplementary data]).
TA in *Meningeosis neoplastica*

To test our hypothesis that the RQ-Trap assay is appropriate for clinical usage in the detection of *Meningeosis neoplastica*, we have studied 111 patients during routine clinical CSF examination. Diagnosis of *Meningeosis neoplastica* was established cytologically in 15 cases by a local neuropathologist (Table 1). All of these specimens had detectable TA. A negative result in the RQ-Trap assay correlated with non-malignant CSF in 96 patients. However, one patient had a positive result without clinical signs of *Meningeosis neoplastica*. This patient was a 6-year-old male patient who was in clinical remission with T-cell non-Hodgkin’s lymphoma (NHL) and undergoing lumbar puncture for prophylactic intrathecal methotrexate chemotherapy. Routinely obtained CSF of this asymptomatic patient showed an increased cell number (23/μL, upper normal value 3/μL) in combination with increased total protein (2.53 g/L, upper normal value 0.45 g/L). There was no clinical correlation to an infection or malignant condition. Interestingly, the patient had received prophylactic intrathecal chemotherapy 2 weeks prior to lumbar puncture. The patient never developed meningeal involvement of his T-cell NHL and remains in continuous complete remission. The sensitivity to detect *Meningeosis neoplastica* with the RQ-Trap assay was 100% (15/15 samples), whereas the specificity was 98.9% (95/96 samples). This resulted in a positive predictive value (positive result for TA) of 0.94 and a negative predictive value of 1.0 for the diagnosis *Meningeosis neoplastica* (Table 2). Furthermore, there was a highly significant association between quantitative values of TA and a diagnosis of *Meningeosis neoplastica* (Figure 3, p < 0.001).

TA in supernatants

In order to potentially ease and more rapidly process CSF specimens, we aimed to quantify TA in CSF supernatant. TA was measured in eight patients with *Meningeosis neoplastica* as well as from tissue culture supernatants (Karpas, Jurkat, Daudi, K562) showing detectable TA, proving that TA can be detected in supernatants from different cell lines (n = 5). None of the patients with clinically diagnosed *Meningeosis neoplastica* had detectable TA in their CSF supernatant, despite traceable TA in corresponding CSF cellular proteins (data not shown). In contrast, 4/5 tissue culture supernatants (Karpas, Jurkat, Daudi, K562) showed detectable TA, proving that TA can be measured directly in the supernatant (Figure 4).

Telomerase stability during the pre-analytic process

Finally, we examined the potential limitations of the handling process of diagnostic CSF samples. Inactivation of telomerase during any time delay in the pre-analytic process may compromise the assay sensitivity resulting in false negative results. We attempted to mimic delays in processing by incubating HEK293 cells or their protein extracts from 0 h to 48 h at room temperature before measuring TA (Figure 5). Stable TA levels were documented when whole cells were incubated at room temperature, whereas protein isolation extraction followed by incubation at room temperature led to rapidly decreasing TA levels (p < 0.001).

Discussion

*Meningeosis neoplastica* is one of the most deleterious complications of neoplastic diseases requiring a prompt and reliable diagnosis. To the best of our knowledge, this is the first report to correlate TA measured by RQ-Trap to the cytopathological diagnosis of *Meningeosis neoplastica* in a routine clinical setting.

The test protocol requires only a small amount of CSF (250 μL for CSF pleocytosis, depending on the cell number) which can be obtained during routine laboratory evaluation. The baseline sensitivity level of 8 malignant cells/1000 cells that we found is comparable to a recent report of Jakupciak et al. (24), who reported a positive result of TA in 10/1000 A594 cells with the RQ-Trap protocol. Notably, the RQ-Trap protocol was about 10 times more sensitive compared to an hTert mRNA quantification in the same cells. Therefore, it will be of interest in future studies to correlate results of TA to markers for minimal residual disease in selected patients. With the rapid and reliable assay presented in this study, 15/15 patients with *Meningeosis neoplastica* were identified correctly. Furthermore, considering the high negative predictive value of 1.0, this assay may serve as a screening test for meningeosis neoplastica. However, the results of this pilot study are preliminary due to the limited patient number and require confirmation using a larger population. Kleinschmidt-DeMasters et al. (25) reported a sensitivity of 60% and specificity of 87% for the detection of *Meningeosis neoplastica* with assessment of TA using conventional TRAP. The increased sensitivity and specificity in our study may be due to improvements in the assay system for the detection of TA. Compared to conventional TA detection methods that rely on end time PCR (e.g., PCR-ELISA or gel detection systems), the RQ-Trap assay provides a greater linear range of analysis that is more sensitive when compared to commercially available PCR-ELISA applications (21). Moreover, with the development of more specific primers to reduce primer-dimers, the RQ-Trap provides a higher specificity. Finally, using RQ-Trap, the total analysis time is reduced to <4 h with hands-on time <20 min. Also, post-PCR carryover sample analysis errors are largely reduced due to the closed tube application.

Our results show a correlation of TA with the presence of malignancy in CSF and are comparable to studies using other body fluids (26–28). This indicates that TA may be an easy to measure marker of malignant cells from a broad spectrum of routine diagnostic samples. However, there are limitations in sensitivity and specificity depending on the sample source. Attempts to detect TA in the CSF supernatant failed for patients’ samples. However, tissue culture
Table 1  Patient characteristics.

<table>
<thead>
<tr>
<th>Meningeosis neoplastica</th>
<th>Tumour patients without Meningeosis neoplastica</th>
<th>Control subjects without tumour</th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients</td>
<td>n = 15</td>
<td>n = 57</td>
<td>n = 39</td>
</tr>
<tr>
<td>Age (years, ± SD)</td>
<td>47.9 (± 15.29)</td>
<td>36.7 (± 26.7)</td>
<td>46.1 (± 23.0)</td>
</tr>
<tr>
<td>Sex (male/female)</td>
<td>12/3</td>
<td>40/17</td>
<td>22/17</td>
</tr>
<tr>
<td>CSF cells (μL, ± SD)</td>
<td>76 (± 146)</td>
<td>3 (± 3)</td>
<td>104 (± 132)</td>
</tr>
<tr>
<td>Underlying diagnosis</td>
<td>DLBCL (n = 13)</td>
<td>Acute lymphatic leukaemia (n = 1)</td>
<td>Intraventricular haemorrhage (n = 9)</td>
</tr>
<tr>
<td></td>
<td>Acute myeloid leukaemia (n = 3)</td>
<td>Peripheral T-NHL (n = 3)</td>
<td>Subarachnoid haemorrhage (n = 6)</td>
</tr>
<tr>
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<td>Follicular lymphoma (n = 3)</td>
<td>Acute myeloid leukaemia (n = 3)</td>
<td>Viral meningitis (n = 5)</td>
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<td>Hydrocephalus occulus (n = 4)</td>
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<tr>
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<td>Mantle cell lymphoma (n = 2)</td>
<td>Shunt infection (n = 3)</td>
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<td>Anaplastic T-NHL (n = 2)</td>
<td>Anaplastic T-NHL (n = 2)</td>
<td>Spinal stenosis (n = 3)</td>
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<td>Seminoma (n = 2)</td>
<td>Idiopathic facial paresis (n = 2)</td>
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<td>Fungal meningitis (n = 1)</td>
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<td>Encephalitis (n = 1)</td>
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<tr>
<td></td>
<td>Breast cancer (n = 1)</td>
<td>Breast cancer (n = 1)</td>
<td>Multiple sclerosis (n = 1)</td>
</tr>
</tbody>
</table>

CSF, cerebrospinal fluid; ns, not significant; DLBCL, diffuse large B-cell lymphoma; NHL, non-Hodgkin’s lymphoma; B-CLL, B-chronic lymphatic leukemia; SD, standard deviation.
Table 2  Contingency table.

<table>
<thead>
<tr>
<th>Meningeosis neoplastica, n</th>
<th>No Meningeosis neoplastica, n</th>
</tr>
</thead>
<tbody>
<tr>
<td>Telomerase activity, n</td>
<td>15</td>
</tr>
<tr>
<td>No telomerase activity, n</td>
<td>0</td>
</tr>
<tr>
<td>Sum, n</td>
<td>15</td>
</tr>
</tbody>
</table>

Figure 3  Relative telomerase activity (TA) in Meningeosis neoplastica and from non-neoplastic CSF.
The level of TA is displayed in % as relative TA compared to that of 1000 HEK293 cells and expressed as box plots, displaying the median as horizontal lines and boxes as 25th and 75th percentile as well as whiskers indicating 10th and 90th percentile. The real-time quantitative telomere repeat amplification protocol (RQ-Trap) assay for relative TA discriminated Meningeosis neoplastica from non-malignant conditions (*p < 0.001).

Figure 4  Telomerase activity (TA) in cellular proteins or tissue culture supernatant.
Relative TA was quantified in cellular proteins (1000 cells) of 5 different malignant cell lines (Jurkat, Daudi, Karpas, K562 and L540) as well as in tissue culture supernatant. TA in tissue culture supernatant was detectable in 4/5 cell lines at lower levels compared to TA from cellular protein (p < 0.03).

supernatants of malignant cell lines showed reproducibly increased levels of TA. This suggests that TA in CSF supernatant may either be diluted by the physiological CSF circulation (29), or our assay system may not be sensitive enough to detect minimal amounts of TA in CSF. Furthermore, extracellular TA may be rapidly inactivated in a time dependent manner if samples are not analysed immediately. This was illustrated by the time-dependent decrease in TA from protein extracts over time. Despite this, TA is readily detectable in tumour cells from CSF for at least 48 h at room temperature.

In conclusion, we demonstrate a fast, reliable and robust method for the detection of TA in a clinical pilot study for the detection of Meningeosis neoplastica. Conceivably, RQ-Trap may function as a “CSF quick test” to identify high-risk patients with leptomeningeal tumour involvement and may, thus, supplement standard cytomorphological examination. The assay is feasible for routine clinical usage and, therefore, warrants further study in order to assess the precise role of TA in CSF, as well as in the context of repetitive testing during anti-tumour therapy.

Supplementary data associated with this article can be found in the online version at: http://www.reference-global.com/doi/suppl/10.1515/CCLM.2009.258.

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