Optimization of molecular detection of GD2 synthase mRNA in retinoblastoma

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Abstract. Extraocular dissemination is the main cause of death in patients with retinoblastoma (RB) in developing countries, and there are few molecular markers that are useful for the evaluation of minimal disseminated disease. The GD2 ganglioside is known to be expressed by RB cells that metastasize in bone marrow, and the activity of the enzyme responsible for its synthesis, GD2 synthase, can be detected in neuroblasto- toma, which shares many phenotypic features with RB. The purpose of the present study was to optimize the detection of GD2 synthase expression by reverse transcription-polymerase chain reaction (RT-PCR) followed by nested-PCR in human RB cell lines and patient samples. The optimization strategy was carried out using the RB cell lines Y79 and WERI-Rb1 and specific primers designed for the human sequence of GD2 synthase mRNA. We detected GD2 synthase expression with at least 200 and 40 pg of total RNA extracted from cultured RB cells using a first round of RT-PCR amplification or a second round of nested-PCR, respectively. We also confirmed the expression of GD2 synthase by RT-PCR and immunohistochemical detection of the ganglioside in human RB tumors xenotransplanted in nude mice. Using tumor bank specimens from eight RB patients, we were able to demonstrate the presence of GD2 synthase mRNA in blood and cerebrospinal fluid samples in cases of extraocular dissemination of the tumor. The sequence was not detected in samples derived from children with low-risk disease or healthy adult volunteers. Hence, GD2 synthase mRNA detection through an optimized nested RT-PCR assay is a promising tool for the assessment of minimal disseminated disease in enucleated patients.

Introduction

Retinoblastoma (RB) is the most common eye cancer in children, affecting 1 in 15,000-20,000 births (1) and accounting for 4% of all pediatric malignancies (2). It is caused by the accumulation of mutations in both alleles of the RB1 gene, the first cancer gene identified and cloned in 1986 (3). When it disseminates outside the eye, it is associated with a dismal patient prognosis. In developing countries with poor education, socioeconomic conditions and health care systems, delayed diagnosis and suboptimal care constitute the common scenario (4,5). Therefore, while salvage has become a priority in developed countries, this is currently not the case in less developed ones, where death from RB remains a major issue (6-9). In Argentina and other countries with a similar degree of socioeconomic development, there is a greater than 80% chance of survival, but microscopical invasion of the ocular coats, requiring more intensive treatment, is still common. Some of these patients die due to extraocular disease after undergoing enucleation when adequate post-operative treatment is not provided (5). The risk of extraocular relapse is currently estimated by assessing the invasion of ocular coats such as the optic nerve, choroid or sclera, with adjuvant therapy administered to patients based upon this information. However, there is no consensus regarding the value of each of these prognostic factors (10,11).

This estimation, based upon histopathological risk factors, has a limited sensitivity for identifying patients at risk due to minimal disseminated disease (MDD). Thus, some patients needing adjuvant therapy may be overlooked while, conversely, some identified as being at a higher risk by histopathology may not need adjuvant therapy, and are therefore unnecessarily exposed to the adverse effects of high-dose chemotherapy (12-14). Adjuvant therapy usually requires the use of high doses of chemotherapy, including drugs with good penetration into the central nervous system (15,16), where recurrences are more likely. Since Smith et al first reported the use of the reverse transcription-polymerase chain reaction (RT-PCR) methodology for the amplification of tyrosinase transcripts to detect circulating melanoma cells in peripheral blood samples in 1991 (17), molecular detection of MDD utilizing specific target sequences has been proposed for several types of cancer (18) due to its high
specificity and sensitivity. However, in RB a major obstacle for MDD detection is the lack of specific tumor markers. Certain gangliosides are the most frequently used molecular markers for such a purpose in other pediatric malignancies that share a similar antigenic profile with RB, such as neuroblastoma (19). Gangliosides are involved in normal biological functions including cell adhesion, cell–cell interactions and proliferation (20). Specific alterations in the expression of gangliosides after neoplastic transformation, which are likely involved in the metastatic phenotype of malignant cells (21,22), have been found, particularly in tumors of neural crest-derived tissues (23).

One of the most extensively studied molecular markers for the detection of MDD in neuroblastoma is the transcript of the β1,4-N-acetylgalactosaminyltransferase (GD2 synthase) gene (19,24). GD2 synthase is the key enzyme required for the synthesis of the GD2 ganglioside (25), and is commonly expressed in normal tissues such as the brain and in several tumor types, as previously mentioned (23).

To date, molecular markers for RB that can be used for the determination of MDD have received little attention in the scientific literature, probably since extracolonic dissemination is uncommon in more developed countries. The finding of such a molecular marker for MDD detection would not only allow for more appropriate therapeutic decision making before metastasis or a potential recurrence takes place, but would also allow for the monitoring of the outcome of patients during treatment. The possibility of identifying a high-risk subgroup of patients through the detection of this type of marker may result in the prevention of metastasis by the selection of a more personalized adjuvant therapy. As GD2 synthase has proven to be a useful molecular marker in neuroblastoma, and given the fact that the GD2 ganglioside was found to be shed from human RB tumors (26) and was found in the bone marrow of RB patients (27), we suggest that it may be a good candidate for MDD detection in RB.

Based on these findings, our aim was to study the expression of GD2 at the immunohistochemical level and to detect the GD2 synthase transcript in human RB cell lines and patient samples by means of an optimized nested RT-PCR technique. We also pursued the design of novel high-affinity primers for this sequence.

Materials and methods

Cell lines. The human RB cell lines Y79 and WERI-Rbl were cultured in suspension in RPMI-1640 medium (Gibco BRL) with 20 or 10% heat-inactivated fetal bovine serum (PAA Laboratories GmbH), respectively. Cultures were maintained at 37°C in 5% carbon dioxide and at the appropriate conditions of humidity, with change of the culture medium once or twice weekly. The culture medium was supplemented (per liter) with 2.5 g glucose, 2.38 g HEPES, 1 mM sodium pyruvate, 1.5 g sodium bicarbonate and 2 ml gentamycin (80 mg/ml) as specified by the supplier (ATCC). RNA from other human tumor cell lines (MCF7 breast, HeLa cervical and H125 lung human carcinomas) had previously been extracted and stored at our laboratory.

Tumor growth in nude mice. Specific pathogen-free athymic female nude (nu/nu) Balb/c mice were purchased from the Animal Facility of UNLP (Argentina) and maintained in sterile isolated conditions. Water and food was given ad libitum. All animal protocols were supervised and managed by qualified trained personnel according to the international guidelines for animal care.

Y79 cells were harvested by centrifugation, washed twice and resuspended in RPMI-1640 medium or in 10 mg/ml cold liquid Matrigel (BD Biosciences) in a final volume of 250 µl at a concentration of 1x10⁶ cells/mouse. Tumor cells were immediately inoculated in the subcutis of the flank of nude mice with either culture media or Matrigel (28). Animals were monitored twice a week for 4 weeks until tumors reached a mean diameter of ~10 mm. Tumor tissue from growing tumors was either fixed in 10% buffered formalin for immunohistochemical studies or rapidly frozen at -70°C for mRNA isolation.

Immunohistochemical detection of ganglioside GD2. Formalin-fixed paraffin-embedded tissue blocks of RB tumors generated in nude mice were sectioned (3 µm), mounted on siliconized slides, deparaffinized in xylene, blocked in 3% H₂O₂ and processed for antigen retrieval in pre-heated sodium-citrate buffer (pH 6.0). The murine anti-GD2 monoclonal antibody 3F8 was provided by Dr Nai-Kong V. Cheung (Memorial Sloan-Kettering, NY, USA). Samples were incubated overnight with the antibody (1:50) at 4°C in a moist chamber. After washing with Tris-buffer (pH 7.3), a peroxidase-labeled polymer conjugated to secondary anti-mouse antibodies (Dako EnVision Detection System) was applied, and samples were further incubated for 30 min at room temperature and developed with diaminobenzidine as the chromogen. Sections were counterstained with Mayer's hematoxylin, then dehydrated, cleared and mounted.

Sample collection and processing. Peripheral blood (PB), cerebrospinal fluid (CSF) and tumor samples from patients <5 years of age who had undergone enucleation were obtained from the Tumor Bank of the Pediatric Hospital Professor Dr Juan P. Garrahan (Buenos Aires, Argentina) for use in this study. Written informed consent was obtained from all patient guardians, and ethical approval was obtained from the Institutional Review Board of the hospital. The patient samples and PB samples from healthy adult volunteer donors were processed for GD2 synthase mRNA detection. Each sample was stored in a 1:1.5 ratio of sample:guanidinium thiocyanate (GTC) buffer. The buffer contained 6 M GTC (Promega), 0.0375 M sodium citrate and 0.75% N-lauroylsarcosine. After mixing, the sample/GTC mixtures were stored immediately at -70°C until use. In order to achieve a final GTC concentration of 4 M (29), 3 ml of PB and CSF extracted from the patients was collected directly in 4.5 ml of 6 M GTC buffer. Due to the small size, tumor samples were collected in 4 M GTC buffer, on the assumption that the amount of samples would not affect the buffer concentration.

RNA extraction. RNA extraction from cell lines and patient samples was conducted at 4°C based on TRIzol extraction methodology (TRIzol LS Reagent; Invitrogen) according to the manufacturer's instructions. RNA pellets were resuspended in a 30-µl final volume of DEPC-treated water and stored at
-70˚C until use. In the case of small samples such as tumor fragments or CSF, we used ultrapure glycogen (Invitrogen) as a carrier to optimize the extraction. RNA was first quantified spectrophotometrically, and its integrity was assessed electrophoretically after using the obtained RNA as a template for glyceraldehyde 3-phosphate dehydrogenase (GAPDH) mRNA amplification.

**Primer design.** The design of highly specific primers for the human GD2 synthase mRNA sequence (GenBank accession no. NM_001478) was carried out with PrimerSelect™ 5.05 software (DNastar Inc.), with default settings applied. The best score, the lower difference in melting temperature between each pair of primers (<1˚C), and the best sequential and structural features to prevent intramolecular or intermolecular secondary structures were considered. In addition, an inter-exonic design was employed, thereby allowing for the distinction between the amplification of contaminating genomic DNA present in the sample from those amplified from the transcripts. Primer details are presented in Table I.

**RT-PCR optimization.** RNA samples were analyzed by a two-step RT-PCR assay using specific primers for a 347-bp GD2 synthase mRNA fragment, as well as specific primers for a 457-bp fragment from the housekeeping gene GAPDH as a loading and integrity control. The first step of cDNA synthesis was carried out in a final volume of 50 µl using the Illustra™ Ready-To-Go™ RT-PCR Bead kit (GE Healthcare) and 1 µl of random hexamers pd(n) (12 pmol/µl) at 43˚C. In the second step, 1 µl of the forward and reverse primers was added. To optimize the RT-PCR, we evaluated several parameters that are known to influence the outcome of this type of assay. After optimization, a typical 50-µl reaction included 2 µl RNase Out (80 U; Invitrogen), 2 µl Mg²⁺ (25 mM; Promega) and 1 µl DTT (1 mM; Invitrogen). Amplification was carried out for 60 cycles with a 63.7˚C annealing temperature. We also tested different types of first-strand primers during the reverse transcription stage, such as pd(T)_{12,18} (0.5 µg/µl), the reverse PCR primer, and random hexamers pd(n)_n, the latter being the best option. The optimal final concentration of primers used was 0.24 pmol/µl. A final volume of 50 µl was completed with DEPC-treated water. We used 5 ng RNA from the Y79 human RB cell line as a template for each optimization round. PCR products were analyzed by electrophoresis on 2% agarose gels followed by ethidium bromide staining.

**RT-PCR and nested-PCR assay.** For each RT-PCR reaction, both a loading and integrity control (GAPDH) and a negative control (without template) were used, as well as a positive GD2 amplification control (5 ng of total RNA from Y79). For patient samples, 25 µl total RNA was employed as a template for the GD2 analysis (regardless of the RNA concentration obtained), and 4 µl total RNA was used for GAPDH amplification. Second round (nested-PCR) reactions consisted of a 1-µl aliquot of the first round reaction mix as a template, 1 µl (12 pmol/µl) of each specific nested primer and 27 µl Platinum® PCR Supermix (Invitrogen) in a 30-µl final reaction volume. Finally, 45 µl of the first round reaction volume and 30 µl of the second were subjected to electrophoresis in the agarose gels.

The cycling profile for the first round of amplification was carried out as follows: reverse transcription, 1 h at 43˚C; initial denaturation, 5 min at 95˚C; amplification (60 cycles), 1 min at 95˚C, 1 min at 63.7˚C and 30 sec at 72˚C; final extension, 10 min at 72˚C. For the second round, cycling consisted of: initial denaturation, 1 min at 95˚C; amplification (30 cycles), 30 sec at 95˚C, 30 sec at 60.2˚C and 30 sec at 72˚C; final extension, 5 min at 72˚C.

**Results**

In order to determine whether the putative marker, GD2 synthase, was present in an in vivo context, we first tested the expression of the GD2 synthase product, the GD2 ganglioside, using a xenograft model by means of an immunohistochemical method as indirect evidence of enzyme expression in tumors. For this purpose, Y79 cells were injected into the flank of nude mice with culture medium or Matrigel to allow for tumor development. Animals developed subcutaneous RB tumors with a latency of 12 or 4 days, and a volume of ~250 or 1,000 mm³ at 25 days after inoculation, for medium or Matrigel injection, respectively. Evaluation with the specific anti-GD2 antibody 3F8 confirmed the abundant expression of the ganglioside in the cell membranes and cytoplasm of the Y79 tumor cells (Fig. 1A). Other portions of the same tumors were subjected to RNA extraction and subsequently analyzed by an appropriately optimized RT-PCR assay. The first round of amplification detected an almost identical GD2 synthase mRNA level as in the positive control in the xenograft tumors generated either with or without Matrigel (Fig. 1B).

After achieving an easily visible band of the expected size for the target amplicon in 2% agarose gels, we accomplished the optimization process of the RT-PCR assay for the GD2 synthase mRNA (Materials and methods). The sensitivity and specificity of the optimized assay was determined by analyzing serially reduced amounts of template, starting from

### Table I. Primers for amplification of GD2 synthase and GAPDH mRNA sequences.

<table>
<thead>
<tr>
<th>mRNA</th>
<th>Forward primer</th>
<th>Reverse primer</th>
<th>Product size (bp)</th>
</tr>
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<tbody>
<tr>
<td>GD2 RT-PCR</td>
<td>5'‑TCGGCTACGGCTCTCATCACAG‑3'</td>
<td>5'‑CTGAGCGTGAGCCCCGGG‑3'</td>
<td>347</td>
</tr>
<tr>
<td>Nested-PCR</td>
<td>5'‑GAACCTGGCGCTGTCAAGTA‑3'</td>
<td>5'‑CACCACCTTATCGGCAAGCT‑3'</td>
<td>180</td>
</tr>
<tr>
<td>GAPDH RT-PCR</td>
<td>5'‑GGGGAGCCAAAAGGGTCATCATCT‑3'</td>
<td>5'‑GACGCTTGCTTCACCACCTTTG‑3'</td>
<td>457</td>
</tr>
</tbody>
</table>
5 ng of total RNA obtained from human RB cells as well as from other tumor cell lines. As shown in Fig. 2A, we detected the GD2 synthase transcript when using at least 200 pg of total RNA from the Y79 cells in the first amplification reaction. This sensitivity was further improved by nested amplification, in which the GD2 synthase transcript was detected when using 40 pg total RNA. In both rounds, the products obtained showed their predicted sizes (347 bp in the first and 180 bp in the second round of amplification). Similar results were obtained with the WERI-Rb1 cell line. Using the same amount of total RNA (5 ng), we also observed GD2 synthase mRNA amplification in human carcinoma cell lines (Fig. 2B). In all cases, RNA integrity was confirmed by RT-PCR using specific primers for GAPDH mRNA.

We subsequently tested eight tumor bank samples from different RB patients in order to preliminarily examine the potential clinical value of the present RT-PCR assay in detecting disseminated tumor cells. Six of these patients had no histopathology risk factors on examination, one patient had stage 2 disease due to tumor invasion of the optic nerve at the resection margin as well as microscopical choroid invasion, while the remaining patient had stage 4a disease due to bone marrow metastasis (30). In this patient, the optic nerve stump was also involved. The patient with stage 2 disease, who was hence at high risk of relapse, presented no target amplification in PB in the first round of RT-PCR. However, when nested-PCR was performed, a clear GD2 synthase band was detected (Fig. 3A, lane b2). He was treated with adjuvant systemic chemotherapy and orbital irradiation according to the institutional protocol (31), and has been in complete remission for 25 months. The patient with stage 4a disease had GD2-positive cells by immunocytology in the bone marrow (data not shown) and showed a faint band corresponding to GD2 synthase mRNA visible in the first round of RT-PCR amplification in the CSF sample at diagnosis and a clear band in the nested-PCR (Fig. 3C, lane csf8). Neither cytological examination of the CSF centrifugate nor immunocytology for GD2 detected tumor cells. No PB sample was available at the tumor bank for this patient. The patient was treated with intensive chemotherapy including autologous stem cell rescue, but relapsed in the CSF 11 months post-transplantation and succumbed to resistant disease. By contrast, none of the other PB samples, which were derived from patients who did not present histological high-risk features, were positive for GD2 synthase after RT-PCR or nested-PCR (Fig. 3A).

We also analyzed the expression of GD2 synthase in human tumor samples from another three patients from the tumor bank in order to verify whether GD2 synthase was expressed as in the cell lines. As shown in Fig. 3B, only one sample was
positive in the first round of amplification (lane t7). Using a second round of amplification, all tumor samples were positive for GD2 synthase, as expected (Fig. 3B).

In addition, we checked target mRNA expression in PB samples obtained from eight healthy adult donors. All samples were negative for GD2 synthase mRNA expression after either the first or second round of amplification (Fig. 4).

**Discussion**

In the present study, we detected for the first time the expression of GD2 synthase mRNA in RB cell lines and tumor samples from RB patients. The presence of GD2 synthase and the GD2 ganglioside in RB tumors was also confirmed by RT-PCR and immunohistochemical assays using a xenograft model. Despite the fact that one of the three tumor samples was positive in the first round of amplification, the other two required a second round for the detection of GD2 synthase mRNA. This may have been due to a poor harvest of viable cells during tumor sampling. In this regard, the relationship between the mRNA level of glycosyltransferases and the presence of their products was established elsewhere (21). Therefore, as it is now routinely used for neuroblastoma, GD2 synthase may be a suitable molecular marker for the detection of MDD in RB.

To the best of our knowledge, there have been few previous reports concerning the use of molecular markers for minimal residual disease (MRD) assessment by RT-PCR in RB. Instead, other traditional techniques such as cytometric methods, standard light microscopy (32), immunocytochemistry/cytology and fluorescence-activated flow cytometry (27) have been used for MRD analysis in RB.

Yamane et al (33) serially assessed, using RT-PCR, the expression of the neuroendocrine protein gene product PGP9.5 in PB mononuclear cells, bone marrow and mobilized PB stem cells from a single RB patient with metastatic disease. The aim was to use PGP9.5 expression as a predictable marker for detecting MRD after intensive therapy and for evaluating therapeutic effects. However, the specificity of PGP9.5 gene transcripts in the detection of micrometastasis in blood and bone marrow in neuroblastoma was not convincing, since the marker was also found in control samples in another investigation (34).

More recently, Yamashita et al (35) presented a case report of a patient with disseminated RB and evaluated three different gene sequences: cone α-subunit of cGMP phosphodiesterase, rod β-subunit of cGMP phosphodiesterase (rod β-PDE) and interphotoreceptor retinoid-binding protein (IRBP). The authors concluded that molecular detection of IRBP is useful in cases of disseminated RB to ascertain the safety of stem cell transplantation, and that both IRBP and rod β-PDE are more suitable for monitoring the effectiveness of chemotherapy. However, only case studies have been reported, and a large number of RB cases are needed to define the clinical utility of these and other molecular markers. Both reports dealt with the study of MRD in patients with metastatic disease who underwent intensive treatment. However, this condition is very uncommon in our setting. Molecular markers of disseminated disease may be useful for detecting MDD at diagnosis in patients with histopathological risk factors for extraocular relapse, and could aid the clinician in prescribing adjuvant therapy in these cases, which are common in developing countries such as Argentina.

After optimizing our nested RT-PCR assay, we achieved a final sensitivity for GD2 synthase mRNA detection of at least 200 pg of total RNA in the first round of amplification, and 40 pg in the second. In comparison with the aforementioned study on PGP9.5, IRBP and rod β-PDE RT-PCR detection, we obtained a much better sensitivity starting from an amount of RNA template three orders scarcer. Thus, our optimization is a very good option for MDD detection in RB, and should encourage others to continue this type of practice in RB or other malignancies.

The sensitivity achieved allowed us to detect malignant RB cells in a particular PB sample through GD2 synthase mRNA amplification, a task that is often difficult as malignant cells circulate at low concentrations (36). The sample corresponded to a patient who had a tumor at the resection margin of the optic nerve after enucleation. No prognostic implications were drawn from this case since only samples from our tumor bank were used and there was no prospective follow-up. Furthermore, all blood samples from healthy volunteers were negative for GD2 synthase, indicating that this mRNA may be a valuable molecular marker for MDD.

Molecular analysis is difficult in biological compartments such as CSF due to its low cellularity, but is potentially important since CSF is the most common site of the recurrence of metastatic retinoblastoma. Tumor-specific RT-PCR assays have been widely used for the detection of MDD in several biological compartments, but this is the first time that RT-PCR and GD2 synthase were successfully used to identify the presence of tumor cells in CSF in a patient with metastatic RB in bone marrow. While the use of GD2 synthase as a marker for meningeal infiltration has been previously reported in neuroblastoma (37), to our knowledge it has never been described in RB.

The number of patients studied was an intrinsic limitation of our study and others. Thus, though suggestive, our data are insufficient to reliably demonstrate that the amplification of GD2 synthase mRNA is a valuable molecular marker for MDD in RB. This marker is currently being prospectively evaluated in a larger cohort to confirm its usefulness. Moreover, the sensitivity and the prognostic value of PCR-based assays for the detection of occult tumor cells can be improved by serial sampling and the use of multiple markers.

The ideal MDD marker is tumor specific; it should not be expressed in normal compartments such as bone marrow or PB. However, the potential molecular marker that we
investigated in this report has been found to be expressed in normal tissues. Indeed, recent publications confirm this fact and propose the utilization of other molecular markers instead (38,39). Quantitative RT-PCR (qRT-PCR) is an indispensable tool for the determination of the threshold between normal and abnormal expression of any type of marker. Certainly, the development of real-time qRT-PCR is valuable for monitoring cancer progression (40), and is currently being carried out at our laboratory for this particular molecular marker. Despite the ubiquitous character of the enzyme, its expression level is low and, to be detected, it requires a system with very high sensitivity. Again, qRT-PCR will prove useful in determining a threshold at which levels of GD2 synthase expression become clearly abnormal.

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