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Department of Pathology, University Medical Center Utrecht, Utrecht, The Netherlands; Laboratory of Translational Immunology, University Medical Center Utrecht, Utrecht, The Netherlands; Institute of Clinical Pathology, University Medical Center Freiburg, Freiburg, Germany; Division of Pediatric Neurooncology, German Cancer Research Center (DKFZ) Heidelberg, Heidelberg, Germany; German Cancer Consortium (DKTK), Heidelberg, Germany; Hopp Children’s Cancer Center at NCT Heidelberg (KiTZ), Heidelberg, Germany

ABSTRACT

Pediatric medulloblastomas are the most frequently diagnosed embryonal tumors of the central nervous system. Current therapies cause severe neurological and cognitive side effects including secondary malignancies. Cellular immunotherapy might be key to improve survival and to avoid morbidity. Efficient killing of tumor cells using immunotherapy requires to overcome cancer-associated strategies to evade cytotoxic immune responses. Here, we examined the immune response and immune evasion strategies in pediatric medulloblastomas. Cytotoxic T-cells, infiltrating medulloblastomas with variable activation status, showed no correlation with overall survival of the patients. We found limited numbers of PD1 + T-cells and complete absence of PD-L1 on medulloblastomas. Medulloblastomas downregulated immune recognition molecules MHC-I and CD1d. Intriguingly, expression of granzyme inhibitors SERPINB1 and SERPINB4 was acquired in 23% and 50% of the tumors, respectively. Concluding, pediatric medulloblastomas exploit multiple immune evasion strategies to overcome immune surveillance. Absence of PD-L1 expression in medulloblastoma suggest limited or no added value for immunotherapy with PD1/PD-L1 blockers.

Introduction

Brain tumors are the second most frequent cancers after hematological malignancies in children. The embryonal tumors of the central nervous system (CNS), i.e. medulloblastoma, atypical teratoid rhabdoid tumor, CNS primitive neuroectodermal tumor (CNS-PNET), are the most frequently diagnosed brain tumors in children in the Netherlands, accounting for 40% of pediatric brain cancers in 2015. Embryonal tumors have an incidence of 4.0 per 1,000,000 children in the Netherlands. The survival rate of medulloblastoma patients largely depends on the clinical and molecular features of the tumor varying from >90% 5-years overall survival for patients with a WNT-driven medulloblastoma till <50% for patients with a metastatic Group 3 or SHH-driven tumor with a TP53 mutation.

Despite molecular phenotyping, therapeutic possibilities are limited to radiation therapy, chemotherapy and surgery, causing severe late onset neurological and cognitive side effects including secondary malignancies. In analogy with other malignancies, immunotherapy might be key to improve survival and to avoid these side effects. Preclinical data from mouse models of SHH and Group 3 medulloblastomas revealed that SHH tumors have higher percentages of dendritic cells, T-cells and myeloid cells than Group 3 tumors in mice. Little is known on immune infiltration of medulloblastomas in humans, although a study with six patients demonstrated that infiltrative myeloid cells are more immunosuppressed and T-cell lineages are less frequent than in other pediatric brain tumors.

Next to immune cell infiltration, efficient killing of medulloblastomas during immunotherapeutic protocols can only be achieved when mechanisms to evade recognition or killing are overcome. It has been well established that cancers employ multiple mechanisms to evade our immune system, making them less susceptible for immunotherapy. Studies in gliomas, medulloblastomas, and CNS-PNETs have shown that certain tumors downregulate MHC-I or lack CD1d expression in order to evade T cell-mediated immunity and NKT cell recognition, respectively. Another potential mechanism to evade immune recognition and subsequent cytotoxic killing is expression of intracellular apoptosis inhibitors (e.g. caspase inhibitors) to escape from death receptor-induced apoptosis and granzyme-mediated killing pathways. Granymes are the major tumor killing molecules secreted by cytotoxic cells. In humans, five granymes (i.e. GrA, GrB, GrH, GrK, and GrM) exist with distinct substrate specificity and only partially overlapping routes of apoptosis induction. Certain tumors can

KEYWORDS

medulloblastoma; brain cancer; pediatric; Serpin; PD-1; PD-L1; immune evasion; tumor-infiltrating lymphocytes

CONTACT

Dr. Niels Bovenschen
n.bovenschen@umcutrecht.nl
Department of Pathology, University Medical Center Utrecht, Heidelberglaan 100 3584 CX, Utrecht, The Netherlands.

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express serine protease inhibitors (Serpins) to directly block granzyme activity.\textsuperscript{19–21} Recently, we have demonstrated that CNS-PNETs can induce expression of SERPINB1 (GrH inhibitor), SERPINB4 (GrM inhibitor), and SERPINB9 (GrB inhibitor).\textsuperscript{17}

Interference with the programmed death 1 (PD-1)/programmed death ligand 1 (PD-L1) immune checkpoint is a pivotal pathway for immune escape by tumors.\textsuperscript{22,23} Expression of PD-L1 on tumor cells, diminishes T-cell activity towards the tumor. Therefore, blocking this axis with antibodies directed to PD-1 (nivolumab and pembrolizumab) resulted in improved therapy response in non-small cell lung cancer (NSCLC), melanoma, and their corresponding brain metastases.\textsuperscript{24–26} Recently, Berghoff \textit{et al.} showed that 88% of adult glioblastomas had PD-L1 expression. Given this high frequency of PD-L1 expression, clinical trials are needed to evaluate the clinical benefit of PD1/PD-L1 blockade in glioblastoma.\textsuperscript{27} However Nduom \textit{et al.}, found that 2.77% of glioblastoma cells had PD-L1 expression, meaning that 61% of patients had >1% and only 38% of patients had >5% PD-L1 positive tumor cells. Expression of PD-L1 was a significant prognosticator for poor survival.\textsuperscript{28} Surveying a large set of adult brain cancers revealed that expression of PD-L1 was restricted to grade IV tumors i.e. glioblastoma multiforme (7.8%) and gliosarcoma (25%). Other grade I-III tumors e.g. (anaplastic) astrocytoma, (anaplastic) oligodendroglioma, ependymoma did not express PD-L1, arguing that either brain tumors are not immunogenic or other immune evasion strategies are exploited.\textsuperscript{29}

Expression of CDK5 in medulloblastoma has been shown in mouse models to induce persistent PD-L1 expression, resulting in resistance towards CD4-dependent cytotoxic T-cell activity.\textsuperscript{30} Therefore, CDK5 might be the pivotal factor in PD1/PD-L1 immune response.\textsuperscript{30} A recent paper on PD-L1 expression in medulloblastomas showed that PD-L1 positive tumors had less influx of T-cells and those patients had a worse survival. These data suggest that also medulloblastoma patients may benefit from PD1/PD-L1 axis blockade.\textsuperscript{31} In contrast the study of Aoki \textit{et al} showed that all four analyzed medulloblastomas had no PD-L1 expression.\textsuperscript{32}

The aim of this study is to survey human pediatric medulloblastomas for tumor-infiltrating lymphocytes (TILs), immune checkpoints, and expression of immune evasion molecules, allowing to facilitate prediction of the tumor response to immunotherapy. However, in contrast to previous reports, we found no activated T-cells or PD-L1 expression in pediatric medulloblastomas, suggesting that the added value of immunotherapy with PD1/PD-L1 blockers in this cancer type is limited.

\section*{Materials and methods}

\subsection*{Patients}

We examined by immunohistochemistry the immune response and immune checkpoints in 26 primary pediatric medulloblastomas operated between 1990–2015 at the University Medical Center Utrecht (Utrecht, The Netherlands). Patient characteristics are shown in Table 1. The study material was derived from the archive of the Department of Pathology of the University Medical Center Utrecht, Utrecht, The Netherlands and distributed by the Biobank of the Department of Pathology. The biobank is overseen by the institutional medical ethical review board.

Since we are using archival pathology material which does not interfere with patient care and does not involve physical involvement of the patient, no ethical approval is required according to Dutch legislation.\textsuperscript{33} Use and storage of anonymous or coded left over material for scientific purposes is part of the standard treatment contract with patients and therefore informed consent procedure was not required according to our institutional medical ethical review board.\textsuperscript{34}

Overall survival data were obtained from the Comprehensive Cancer Center of The Netherlands (Integraal Kankercentrum Nederland).

\subsection*{Immunohistochemistry}

Immunohistochemistry was carried out on 4 \textmu m thick formalin fixed paraffin embedded consecutive sections as described before by Vermeulen \textit{et al.}\textsuperscript{35} Immunohistochemistry for PD-1 (Abcam clone NAT105, ab52587) 1:50 and PD-L1 (Abcam clone 28-8, ab205921) 1:50 was performed using an automated immunostainer (Benchmark Ultra, Ventana, Roche). Immunohistochemistry for myeloperoxidase (MPO) (Leica Biosystems Novocastra clone 59A5) was performed manually. Appropriate positive and negative controls were included in all stainings. For classification, all tumors were restained, reevaluated, and molecularly classified using Illumina DNA methylation arrays as described,\textsuperscript{35} according to the 2016 edition of the WHO classification of tumors of the central nervous system.

\subsection*{Scoring of immunohistochemistry}

Reclassification of cases was performed independently by two experienced neuropathologists (WGMS and WVH). All scoring was done blinded to patient characteristics and results of other staining by three independent observers (JFV, WGMS, WVH). Analysis of the immune influx was performed on whole slides at 20x magnification. Immune influx was corrected for the size of tissue on the slide and the tumor percentage as calculated with the manufacturers algorithm based on digitalized immunohistochemical slides using a digital slide scanner (Aperio Technologies Inc.). SerpinB1, SerpinB4, SerpinB9, CD1 d, HLA-A

\begin{table}[h]
\centering
\begin{tabular}{|l|l|l|}
\hline
\textbf{Table 1. Patient characteristics.} & \textbf{N or value} & \textbf{\%} \\
\hline
\textbf{Gender} & Male & 17 & 65.4 \\
 & Female & 9 & 34.6 \\
\hline
\textbf{Age (years)} & Mean ± SD & 8.5 ± 5.3 & \textbf{Range} & 0.4 – 17.8 \\
\hline
\textbf{Histological type} & Classic & 16 & 61.5 \\
 & Desmoplastic nodular & 6 & 23.1 \\
 & Extensive nodular & 3 & 11.5 \\
 & Anaplastic & 1 & 3.8 \\
 & WNT & 1 & 3.8 \\
 & SHH & 8 & 30.8 \\
 & Group3 & 5 & 19.2 \\
 & Group4 & 9 & 34.6 \\
 & Undetermined & 3 & 11.6 \\
\hline
\end{tabular}
\end{table}
and HLA-B expression was scored as present (+) or absent (−), using a cut-off of 5% positivity.

**Statistics**

Statistical analysis was performed using IBM SPSS version 23 (SPSS Inc.). Data are depicted as median and inter quartile range (IQR) or as mean and 95% confidence interval (95% CI) when stated, taking all patients into account. Differences in number of TILs per histological subtype were analyzed using a Kruskal-Wallis test and correlations using Pearson correlation. Overall survival (OS) was used as outcome endpoint and defined as the time between date of surgery and death. Censored patients were confirmed alive at time of censoring. Thus, there was no loss to follow-up. Survival rates were plotted according to the Kaplan-Meier method and associations were analyzed using the log-rank test. High and low number of TILs were stratified after median split. Statistical significance was set at p < 0.05.

**Results**

**Distribution and characterization of tumor infiltrating lymphocytes in medulloblastoma**

Infiltration of CD3+ T-lymphocytes in human pediatric medulloblastoma occurred in different patterns, most frequently intratumoral infiltration coinciding with perivascular infiltration (Fig. 1). Peritumoral infiltration as observed in brain metastases was not seen in our samples. The number of tumor-infiltrating lymphocytes (TILs) in pediatric medulloblastoma was limited and highly variable between tumors i.e. 23.5 (26.2) [median (IQR)] CD3+ T-cells and 1.56 (2.8) CD20+ B-cells per 2mm² tumor tissue, but the number of infiltrating T-cells and B-cell was highly correlated (r = 0.748, p < 0.001). These data show that there are 10 times more T-cells than B-cells in the tumor tissue, suggesting selective infiltration of medulloblastomas by TILs. The number of TILs did not differ between histological subtypes (CD3+ T-cells p = 0.247; CD20+ B-cells p = 0.668), nor between molecular subtypes (CD3+ T-cells p = 1.0; CD20+ B-cells p = 1.0) of medulloblastoma.

To further characterize the composition of TILs, we analyzed T-cell subtypes. CD8+ T-cells were the main contributing T-cell population with 52% (47-58%) [mean (95% CI)] followed by CD4+ T-cells 35% (28-43%) and FOXP3+ regulatory T-cells (Treg) 2.5% (1.3-3.8%) to the total number of TILs. Furthermore we found 1.70 (1.45) [median (IQR)] times more CD8+ than CD4+ T-cells in the tumors. The influx of these T-cell subsets was correlated to the number of CD3+ T-cells (CD8+ ρ = 0.954, p < 0.001; CD4+ ρ = 0.841, p < 0.001; Treg ρ = 0.422, p = 0.032) as expected. Next, we studied the influx of γδ-T-cells, an additional cytotoxic lymphocyte subset.36,37 We detected low numbers of γδ-T-cells in our cohort of pediatric medulloblastoma [0.05 (0.11)] per 2mm² tumor tissue. Recruitment of NK cells was limited to single NK cells, which is in line with results in other CNS tumors.38,39 Furthermore, the influx of neutrophilic granulocytes was highly variable between tumors, i.e. 17.5 (19.5) per 2mm² tumor tissue and was not correlated to infiltrating T-cells.

**Immune evasion strategies and PD-1/PD-L1 axis in medulloblastoma**

The activity of cytotoxic T-lymphocytes (CTLs) is one of the prerequisites for cytotoxic killing of tumor cells. Therefore, we examined the activation status of CTLs by Granzyme B (GrB) positivity. The GrB-positive CTL fraction was low 3.9% (6.8%), while some cases had up to 35% GrB-positive CTLs. Given the low CTL activation, we questioned whether medulloblastomas avoid immune recognition by downregulation of MHC-I (HLA-A and HLA-B) and/or CD1 d in order to escape CTL, NKT, and/or (CD1 d-restricted) γδ-T-cell recognition.40 Whereas blood vessels, and TILs readily express these HLA molecules and CD1 d as expected, we found that HLA-A, HLA-B, and CD1 d expression was negative in all our pediatric medulloblastomas (Fig. 2A).

Expression of intracellular apoptosis inhibitors (e.g. caspase inhibitors) to escape from death receptor-induced apoptosis and granzyme-mediated killing pathways are alternative immune evasion pathways employed in multiple cancers.13 Granzymes are the major tumor killing molecules secreted by cytotoxic cells with partially overlapping routes of apoptosis induction.18 Tumors can express Serpins to directly avoid immune recognition by downregulation of MHC-I expression of SerpinB1 (23%) and 13 out of 26 tumors, this results in functionally impaired TILs.41,42 SerpinB1 (r = 0.225, p = 0.042) but not to SerpinB4 expression (r = 0.418, p = 0.042) nor to SerpinB1 (r = 0.225, p = 0.042).

In addition to immune escape by anti-apoptotic pathways, we wondered whether these CTLs are functionally impaired by e.g. the PD-1/PD-L1 immune checkpoint. CD8+ T-cells are known to express PD-1 and upon PD-L1 expression by tumor cells, this results in functionally impaired TILs.41,42 First, we examined PD-1 expression and found that PD-1 positive TILs had predominantly a perivascular localization pattern (Fig. 2C). The fraction of PD-1+ TILs [4.6% (6.4%)]
was low and correlated to GrB-positive CTLs ($r = 0.646$, $p < 0.001$), although it was not correlated to total number of CTLs ($r = -0.209$, $p = 0.305$). The influx of PD1$^+$ TILs was not correlated to the (molecular) subtype of medulloblastoma. Furthermore, we found that none of the analyzed pediatric medulloblastomas expressed PD-L1 (Fig. 2C), suggesting that the PD-1/PD-L1 axis likely is not of great importance in pediatric medulloblastoma and thereby limiting the therapeutic potential of PD-1 blockers.

**TILs are not associated with patient survival**

Depending on the molecular characteristics patients have a prognostic beneficial or worse phenotype of pediatric medulloblastomas. Potentially immunotherapeutic strategies can contribute to better survival. For that reason we analyzed our patient cohort regarding the association between overall survival and TILs. We stratified for low and high expression of each factor. In our cohort we did not find an association between TILs and patients survival (Fig. 3), although there is a trend towards beneficial outcome for patients with low TILs. However, we found that patients with high numbers of GrB$^+$ CTLs have worse survival than patients with low numbers of GrB$^+$ CTLs ($p = 0.015$). High expression of anti-apoptotic molecules *i.e.* Bcl-2 and Survivin did not contribute to survival (data not shown). Overall survival of patients was not dependent on the expression of SerpinB4 ($p = 0.974$). In contrast to oropharynx squamous cell carcinomas, high expression of SerpinB1 in pediatric medulloblastoma was associated with better survival ($p = 0.050$).

**Discussion**

Pediatric medulloblastomas are the most frequently diagnosed embryonal tumors of the CNS in children. The prognosis is mainly dependent on the molecular subtype of the tumor, although therapeutic strategies are limited to conventional radiation therapy, chemotherapy, and surgery. Because of severe neurological side effects, additional strategies such as immunotherapy are being investigated. This study investigates the composition and prognostic value of TILs in pediatric medulloblastoma and the immune evasion strategies exploited by these tumors. We show that the main TIL subsets are CD3$^+$, CD8$^+$ T-cells which have predominantly a perivascular and intratumoral infiltration pattern. The CTLs are barely activated given the low percentage of GrB and PD1 positive cells. It has been hypothesized that pediatric and in particular embryonic tumors are not immunogenic, and therefore immunotherapeutic interventions have a limited success rate compared to *e.g.* metastases of NSCLC or melanoma. A recent study compared several pediatric tumors, which revealed that glioblastoma, neuroblastoma as well as the embryonic Atypical teratoid/rhabdoid tumor have increased expression of PD-L1 and increased amounts of TILs. Together this might give a direction why the influx of TILs does not influence the overall survival of medulloblastoma patients in contrast to other tumor types.

One of the mechanisms for immune modulation is the lack of expression of MHC-I and CD1d which is the case in all tumors. Despite the lack of MHC-I in neuroblastoma, still expression of MHC-I results in activation of NK cells, therapeutic strategies exploiting NK cells might be of clinical significance. So far no clinical data is available on the effectiveness of NK cell based therapies in medulloblastoma. In order to engage NK cells to kill medulloblastoma cells, expression of ligand for the activation receptors might be required. Expression of ligands for NKG2D (NK cell activation receptor) are elevated in 50% of medulloblastomas. In line with this, several patients have been intrathecally treated with LAK cells, showing a remission of the medulloblastoma. Recently Castriconi et al., showed for the first time that NK cells can kill medulloblastoma cells *in vitro*, which opens the way to study the potential of NK cell based immunotherapy in medulloblastoma. We showed that
half of the tumors expressed SerpinB4 and 23% SerpinB1 to potentially resist granzyme induced apoptosis. Expression of SerpinB1 but not SerpinB4 had a beneficial effect on survival. High expression of SerpinB1 has been linked to increased sensitivity towards cisplatin-based chemotherapy and increased survival in melanoma.\(^53\) Since cisplatin is a standard chemotherapeutic agent in treatment protocol of pediatric medulloblastoma, the improved survival might be related to increased cisplatin sensitivity, rather than to a SerpinB1 dependent immunomodulatory effect.

The PD-1/PD-L1 immune checkpoint is a pivotal pathway for immune escape by tumors, since expression of PD-L1 on tumor cells, diminishes T-cell activity towards the tumor.\(^54,55\) Therefore, interference in this checkpoint constitutes a promising form of immunotherapy in many tumor types.\(^56–58\) For instance, clinical studies with GBM patients revealed that the majority of patients had tumor cells expressing some PD-L1 of which 38% had >5% PD-L1 positive tumor cells, and activation of the PD1/PD-L1 axis was associated with poor prognosis.\(^28\) In preclinical studies, CD8^+^ T-cells are enriched in murine medulloblastoma models and those were more frequently PD1 positive, and administration of PD1 blocking antibodies indeed have a beneficial survival effect in mice.\(^11\)

In contrast to the very recent study of Murata et al.,\(^31\) that describes worst survival of the patients with high PD-L1 expression, we could not detect PD-L1 expression in any of the

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**Figure 3.** TILs and survival of pediatric medulloblastoma patients. Survival analysis of pediatric medulloblastoma patients stratified for amount of TILs. High and low number of TILs was determined by median split.
human pediatric medulloblastoma tumor samples. Our finding is in line with recent studies, which demonstrated that pediatric tumors in general are not likely to exploit the PD1/PD-L1 axis. The discrepancy between our study and Murata et al might be due to differences in staining procedure and used antibodies. The Murata study used a polyclonal antibody for IHC staining and scored intensity of the staining regardless of antibodies. The discrepancy between our study and Murata et al might be due to differences in staining procedure and used antibodies. Therefore to our opinion this study strongly suggests that the therapeutic potential of immunotherapy with PD1/PD-L1 axis blockers seems limited in pediatric medulloblastomas. Further standardized studies are required to examine the PD-L1 expression in this type of cancer.

Competing interests
The authors have declared that no competing interests exist.

Author contribution
JFV, EJMA, MKJ, RGB, WvH, WGMS, JVH, MK, and NB performed the experiments. JFV, WvH, WGMS, JVH, PF, MK, and NB provided the study material, analyzed and interpreted tumor pathology. JFV and NB designed the experiments and wrote the manuscript. All authors critically reviewed the report and approved the final version of the report for submission. The corresponding author (NB) had access to the primary data, mission. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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ORCID
Rianne G. Bouma [http://orcid.org/0000-0002-6652-1891

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