

Immunotherapy for Medulloblastoma: Current Perspectives

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Background: Immune-mediated therapies have transformed the treatment of metastatic melanoma and renal, bladder, and both small and non-small cell lung carcinomas. However, immunotherapy is yet to demonstrate dramatic results in brain tumors like medulloblastoma for a variety of reasons. Recent pre-clinical and early phase human trials provide encouraging results that may overcome the challenges of central nervous system (CNS) tumors, which include the intrinsic immunosuppressive properties of these cancers, a lack of antigen targets, antigenic variability, and the immune-restrictive site of the CNS. These studies highlight the growing potential of immunotherapy to treat patients with medulloblastoma, a disease that is a frequent cause of morbidity and mortality to children and young adults.

Methods: We conducted an inclusive review of the PubMed-indexed literature and studies listed in clinicaltrials.gov using combinations of the keywords medulloblastoma, immunotherapy, CNS tumors, brain tumors, vaccines, oncolytic virus, natural killer, and CAR T to identify trials evaluating immunotherapy in preclinical experiments or in patients with medulloblastoma. Given a limited number of investigations using immunotherapy to treat patients with medulloblastoma, 24 studies were selected for final analysis and manuscript citation.

Results: This review presents results from pre-clinical studies in medulloblastoma cell lines, animal models, and the limited trials involving human patients.

Conclusion: From our review, we suggest that cancer vaccines, oncolytic viral therapy, natural killer cells, and CAR T therapy hold promise against the innate immunosuppressive properties of medulloblastoma in order to prolong survival. There is an unmet need for immunotherapy regimens that target overexpressed antigens in medulloblastoma tumors. We advocate for more combination treatment clinical trials using conventional surgical and radiochemotherapy approaches in the near-term clinical development.

Keywords: medulloblastoma, immunotherapy, vaccines, oncolytic virus, natural killer, CAR T, review

Introduction

Primary brain and CNS malignancies are among the most common solid tumors in the pediatric population, and medulloblastoma is the most prevalent brain tumor in children.¹ Medulloblastomas originate from the cerebellar vermis and usually in proximity to the fourth ventricle, commonly metastasizing through cerebrospinal fluid pathways.^{2,3} Medulloblastoma accounts for 8–10% of pediatric brain tumors and the 5-year survival rate in children is 75–85% with conventional treatments.^{4–6}

However, the current standard treatment, which includes surgery with subsequent

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chemotherapy and radiation, often results in severe neurological and endocrine deficits.^{2,7-9} New therapies are vital to improve treatment outcomes but require penetration of the blood–brain barrier. Although the blood–brain barrier remains a significant challenge, activated T cells and other elements of the immune system can traverse the capillary tight junctions formed in the blood-brain barrier, unlike many chemotherapy agents.¹⁰ Immunotherapy is an attractive targeted approach to eliminate cancer cells while simultaneously sparing adjacent brain tissue.² Tumor targeting T cells can be activated *in vivo* via cancer vaccines and oncolytic viruses while other *ex vivo* engineered therapies can be transfused into patients to stimulate the host immune system. Immunotherapy has shown clinical benefit in a variety of cancers like melanoma, lung cancer, and leukemias. Yet several challenges exist in targeting central nervous system (CNS) tumors such as medulloblastoma including the lack of known immunogenic antigens.¹⁰ Encouraging results were demonstrated with immunotherapy for brain tumors including glioblastoma, and recent studies documented the overexpression of certain antigens on medulloblastoma that can potentially serve as targets for vaccines, CAR T, and other forms of immunotherapy.

We conducted a review of PubMed-indexed literature and clinicaltrials.gov using combinations of the keywords medulloblastoma, immunotherapy, CNS tumors, brain tumors, vaccines, oncolytic virus, natural killer, and CAR T to find as many completed and ongoing trials as possible that evaluated immunotherapy as treatment for patients with medulloblastoma. This paper presents a review of these findings with a discussion of the investigations categorized by therapeutic modalities: cancer vaccines, oncolytic viruses, checkpoint inhibitors, natural killer cells, radiotherapy, and CAR-T cell therapy. Current FDA-approved studies evaluating immunotherapy in medulloblastoma will also be discussed and outlined in [Table 1](#).

Background

The World Health Organization classifies medulloblastomas into four histological groups: large cell, anaplastic, nodular desmoplastic, and extensive nodularity.¹¹ In addition, a newer, revised classification system is based on the molecular profile of medulloblastoma and divided into four subgroups: sonic hedgehog activated medulloblastoma (SHH, further subdivided into TP53-mutant and Tp53-wildtype), WNT-activated, group 3, and group 4.¹¹ The first two subgroups are caused by alterations in the SHH and WNT pathways, respectively, while less is

known about the etiology of groups 3 and 4. Furthermore, advancements in gene sequencing have shown that the subgroups can also be divided into 12 subtypes with distinct activated pathways, somatic copy-number aberrations, and clinical outcomes: SHH α , SHH β , SHH γ , SHH δ , WNT α , WNT β , group 3 α , group 3 β , group 3 γ , group 4 α , group 4 β , and group 4 γ .¹² For example, SHH α tumors demonstrate the most MYCN and GLI2 gene amplification and have worse prognosis compared to SHH β tumors, which exhibit increased PTEN gene deletion.¹²

Embryologically, SHH medulloblastomas originate from cerebellar granule neuron precursor cells (GNPCs), while WNT tumors originate from precursor cells within the dorsal brain stem.^{13,14} Group 3 medulloblastomas also appear to originate from GNPCs but through a different pathway than SHH; interestingly, these tumors express photoreceptor-encoding genes resembling rod precursor cells at week 15 of retinal development.¹⁵ Group 4 cells arise from GNPCs as well, and their gene expression is similar to cerebellar glutamatergic granule neurons at late fetal stages of development.¹⁵ WNT medulloblastomas have the best prognosis of 95% 5-year survival but are the least common subgroup.^{16,17} Group 4 medulloblastoma is the most common subgroup.

Most of our current knowledge of CNS immunology is based on glioma research, which showed a deficient immune response in patients and animal models with gliomas.^{2,18-20} Similarly, immunosuppression was first reported over 30 years ago in medulloblastoma patients who had reduced peripheral T cell activity, shown by skin hypersensitivity reactions.^{21,22} However, given the less deleterious side effects of immunotherapy compared to conventional therapies, researchers nonetheless strive to circumvent the immunosuppressive properties of these tumors and elicit a host response. Furthermore, immunotherapy can be used in conjunction with chemotherapy, as various preclinical and clinical studies have shown that vaccination concurrent with chemotherapy improves antigen-specific T cell activity.^{10,23-25} For an immune response to be elicited against brain tumors, appropriate tumor-specific antigens must be targeted. Antigens not expressed on normal tissues are the ideal target; however, these antigens are likely personalized to the tumor in each individual patient. CNS tumors, including medulloblastomas, exhibit a low mutational burden.²⁶ Alternative targets may include normal tissue antigens that are overexpressed on tumor cells.²⁷ As brain tumors, including medulloblastomas, have limited gene mutations, there are a number of epigenetic

Table 1 Current Immunotherapy Clinical Trials in Patients with Medulloblastoma

Title	Treatment	Diseases	Type ^a	N ^b	Status	Trial ID
Vaccine Immunotherapy For Recurrent Medulloblastoma And Primitive Neuroectodermal Tumor (Re-MATCH)	-Total tumor RNA-loaded dendritic cells; -Total tumor RNA-loaded autologous lymphocyte transfer	-MB; -PNET	I/2	Phase 1 : 9; Phase 2: 35	Active, Not Recruiting	NCT 01326104
Vaccination With Dendritic Cells Loaded With Brain Tumor Stem Cells For Progressive Malignant Brain Tumor	-Dendritic cells; -Imiquimod	-MB; -GBM; -Epend; -Anaplastic astro; -Brain tumors	I	8	Completed	NCT 01171469
Combining Decitabine And Vaccine Therapy For Patients With Relapsed Or Refractory Pediatric High Grade Gliomas, Medulloblastomas, And Central Nervous System Primitive Neuroectodermal Tumors (CNS PNETs)	-Autologous dendritic cells; -Decitabine; -Hiltonol	-MB; -Gliomas; -PNET	I/2	I	Terminated	NCT 02332889
High Dose Cyclophosphamide, Cisplatin And Carmustine With Stem Cell Reconstitution Followed By Specific Cellular Therapy In Patients With Recurrent Or Refractory Brain Tumors	-Autologous tumor cell vaccine; -Aldesleukin; -Flgrastim; -Sargramostim; -Autologous lymphocytes; -Carmustine; -Cisplatin; -Cyclophos -Paclitaxel; -Autologous bone marrow transplantation; -Conventional surgery; -Peripheral blood stem cell transplantation	-CNS tumors	2	30	Completed	NCT 00014573
CMV RNA-Pulsed Dendritic Cells With Tetanus-Diphtheria Toxoid Vaccine In Pediatric Patients And Young Adults With WHO Grade IV Glioma, Recurrent Malignant Glioma, Or Recurrent Medulloblastoma	-CMV; -Dendritic cells; -GM-CSF; -Td	-MB; -GBM; -Malignant glioma; -Recurrent pediatric GMB; -Pediatric brain tumor; primary and recurrent	I	I I	Active, Not Recruiting	NCT 03615404
PEP-CMV In Recurrent Medulloblastoma/ Malignant Glioma	PEP-CMV	-Recurrent MB; -Recurrent pediatric brain tumor; -Malignant glioma	I	30	Recruiting	NCT 03299309
Phase Ib Study PVSRIPO For Recurrent Malignant Glioma In Children	Polio/ rhinovirus recombinant (PVSRIPO)	-MB; -Malignant glioma; -Anaplastic astro; -Anaplastic oligoastrocytoma; -Anaplastic oligodendroglioma; -GBM; -Gliosarcoma; -ATRT of brain; -Epend; -Pleomorphic xanthoastrocytoma of brain; -Embryonal tumor of brain	I	I 2	Recruiting	NCT 03043391
HSV G207 In Children With Recurrent Or Refractory Cerebellar Brain Tumors	-G207 (HSV viral therapy); -Radiation	-MB; -CNS neoplasms	I	I 5	Recruiting	NCT 03911388

(Continued)

Table 1 (Continued).

Title	Treatment	Diseases	Type ^a	N ^b	Status	Trial ID
Modified Measles Virus (MV-NIS) For The Treatment Of Children And Young Adults With Recurrent Medulloblastoma Or Recurrent Atypical Teratoid Rhabdoid Tumors	-Modified measles vaccine	-MB; -Recurrent ATRT	I	46	Recruiting	NCT 02962167
HSV G207 Alone Or With A Single Radiation Dose In Children With Recurrent Supratentorial Brain Tumors	-G207 (HSV viral therapy); -Radiation	-Supratentorial neoplasms; -Malignant glioma; -GBM; -Anaplastic astro; -PNET; -Embryonal tumor	I	18	Recruiting	NCT 02457845
Replication Competent Reovirus (Reolysin [®]) In Combination With GM-CSF In Pediatric Patients With Relapsed Or Refractory Brain Tumors	-Wild-type reovirus; -Sargramostim	-Recurrent childhood MB; -Childhood astro; -Childhood ATRT; -DIPG; -Glioma recurrent childhood; -Anaplastic oligodendroglioma; -Recurrent childhood brain neoplasm; -Recurrent childhood GBM; -Recurrent primitive PNET; -Refractory brain neoplasm	I	6	Active, Not Recruiting	NCT 02444546
Immune Checkpoint Inhibitor Nivolumab In People With Select Rare CNS Cancers	-Nivolumab	-MB; -Epend; -Pineal tumors; -Choroid plexus tumors; -Malignant meningioma; -Histone mutated glioma; -Gliomatosis cerebri; -ATRT; -Gliosarcoma; -Sarcoma	2	80	Recruiting	NCT 03173950
MK-3475 (Pembrolizumab; Anti-PD-1) In Children With Recurrent, Progressive Or Refractory Diffuse Intrinsic Pontine Glioma (DIPG), Non-Brainstem High-Grade Gliomas (NB-HGG), Ependymoma, Medulloblastoma Or Hypermutated Brain Tumors	-Pembrolizumab	-Recurrent/refractory MB; -Constitutional mismatch repair deficiency syndrome; -Lynch syndrome; -Malignant glioma; -Recurrent/refractory brain neoplasm; -Recurrent childhood epend; -Recurrent/refractory DIPG; -Refractory epend	I/IO	I	Recruiting	NCT 02359565

Study To Assess The Safety, Tolerability, And Pharmacokinetics Of Durvalumab In Pediatric Patients With Relapsed Or Refractory Solid Tumors, Lymphoma, And Central Nervous System Tumors	-Durvalumab	-CNS tumors; -Solid tumors; -Lymphomas	1	36	Recruiting	NCT 02793466
Nivolumab Monotherapy And Nivolumab In Combination With Ipilimumab In Pediatric Subjects With High Grade Primary CNS Malignancies	-Nivolumab; -Ipilimumab	-MB; -DIPG; -High-grade glioma; -Epend	2	170	Active, Not Recruiting	NCT 03130959
Study Of The IDO Pathway Inhibitor, Indoximod, And Temozolomide For Pediatric Patients With Progressive Primary Malignant Brain Tumors	-Indoximod; -Temozolomide; -Conformal radiation; -Cyclophos; -Etoposide	-MB; -GBM; -Glioma; -Gliosarcoma; -Malignant brain tumor; -Epend; -DIPG; -Primary CNS tumors	1	115	Recruiting	NCT 02502708
Indoximod With Chemotherapy And Radiation For Children With Progressive Brain Tumors Or Newly Diagnosed DIPG	-Indoximod; -Full-dose radiation; -Temozolomide; -Cyclophos; -Etoposide; -Lomustine	-MB; -GBM; -Epend; -DIPG	2	140	Recruiting	NCT 04049669
Intraventricular Infusions Of Autologous Ex Vivo-Expanded NK Cells In Children With Recurrent/Refractory Malignant Posterior Fossa Tumors Of The Central Nervous System. NOAH's (New Opportunity, Advancing Hope) Protocol	-Natural killer cell therapy	-Recurrent childhood MB; -Recurrent epend	1	12	Active, Not Recruiting	NCT 02271711
Solid Tumor Immunotherapy Trial Using HLA-Haploidentical Transplant And Donor Natural Killer Cells: The STIR Trial	-Allogeneic HCT; -Donor natural killer cell infusion	-MB; -Ewing sarcoma; -Neuroblastoma; -RMS; -Osteosarcoma; -CNS tumors	2	20	Recruiting	NCT 02100891

(Continued)

Table 1 (Continued).

Title	Treatment	Diseases	Type ^a	N ^b	Status	Trial ID
EGFR806-Specific CAR T Cell Locoregional Immunotherapy For EGFR-Positive Recurrent Or Refractory Pediatric CNS Tumors	-EGFR806-specific chimeric antigen receptor (CAR) T cell	-MB; -Glioma; -Epend; -Pediatric CNS tumors; -Germ cell tumor; -ATRT; -PNET; -Choroid plexus carcinoma; -Pineoblastoma	I	36	Recruiting	NCT 03638167
HER2-Specific CAR T Cell Locoregional Immunotherapy for HER2-Positive Recurrent/Refractory Pediatric CNS Tumors	HER2-specific chimeric antigen receptor (CAR) T cell	-MB; -Pediatric CNS tumors; -Glioma; -Epend; -Germ cell tumor; -ATRT; -PNET; -Choroid plexus carcinoma; -Pineoblastoma	I	36	Recruiting	NCT 03500991
Radioimmunotherapy, Reduced-Dose External Beam Craniospinal Radiation Therapy With IMRT Boost, And Chemotherapy For Patients With Standard-Risk Medulloblastoma	-Cisplatin; -Lomustine; -Vincristine sulfate; -Iodine ¹³¹ I monoclonal antibody 3F8, radiation	CNS Tumors	2	6	Completed	NCT 00058370
Dosimetry-Guided ⁹⁰ Y-DOTA-tyr3-Octreotide (90Y-DOTATOC) Peptide Receptor Radiotherapy (PRRT) In Children & Adults With Neuroendocrine And Other Somatostatin Receptor Expressing Tumors Determined By ⁶⁸ Ga-DOTA-tyr3-Octreotide (⁶⁸ Ga-DOTATOC) PET	- ⁹⁰ Y-DOTA-tyr3-Octreotide; -Dosimetry-guided peptide receptor radiotherapy; -Aminofyllin	-MB; -NET; -Carcinoid tumors; -Neuroblastoma	I	25	Recruiting	NCT 02441088
A Two-Part Study Of TB-403 In Pediatric Subjects With Relapsed Or Refractory Medulloblastoma	-TB-403 (humanized monoclonal antibody against placental growth factor)	-Refractory MB; -Neuroblastoma; -Ewing sarcoma; -Alveolar RMS	I/2	36	Recruiting	NCT 02748135

Notes: ^aType=Level of Clinical Trial; ^bN=Enrollment.

Abbreviations: Astro, Astrocytoma; ATRT, Atypical Teratoid Rhabdoid Tumor; DIPG, Diffuse Intrinsic Pontine Glioma; Epend, Ependymoma; GBM, Glioblastoma; MB, Medulloblastoma; NET, Neuroendocrine Tumor; PEP-CMV, Peptide Vaccine Derived From Cytomegalovirus; Cyclophos, Cyclophosphamide; PNET, Neuroectodermal Tumor/Primary Neuroectodermal Tumor; RMS, Rhabdomyosarcoma; Td, Tetanus Toxoid.

deregulatory pathways that may increase the re-expression of fetal antigens in medulloblastomas.^{28–30} These fetal, developmental antigens may be tumor-specific and targeted as they are not expressed in normal tissue.¹⁰

Until recently, accepted dogma was that the CNS was resistant to immunotherapy based on the limited access to antibodies.^{26,31} However, the occurrence of autoimmune syndromes including multiple sclerosis that is regulated by the immune system suggests that T cell immunosurveillance does occur.²⁶ Furthermore, the incidence of John Cunningham virus progressive multifocal leukoencephalopathy in patients treated with rituximab shows that immunosuppression can lead to a re-activation of viruses in the CNS.²⁶ While it was previously thought that the brain did not contain lymphatics, studies have shown that lymphatics do indeed exist in the dura and meninges and that lymphocytes exit the CNS via the deep cervical lymph nodes.³² T cells do not enter through the blood–brain barrier but rather access the parenchyma through the choroid plexus or pia mater.³³ Thus, theoretically, antigens created through tumor mutations should be detected by the immune system within the deep cervical lymph nodes, and T cells administered through systemic infusion should be able to target tumors via these routes.²⁶ One recent study showed that the disruption of dorsal meningeal lymphatic vessels in mice impaired tumor fluid drainage and dissemination of tumor cells to deep cervical lymph nodes; however, dendritic cell trafficking to these lymph nodes was also diminished.³⁴ Consequently, the disrupted dorsal meningeal lymphatics reduced the efficacy of combined anti-CTLA-4/PD-1 checkpoint immunotherapy in striatal tumor models.

However, many challenges remain in treating brain tumors with immunotherapy. The highly specialized blood–brain barrier, which protects the brain from circulating toxins and viruses, can prevent chemotherapy and large-sized molecules from entering the CNS. The brain also appears to have a different set of immune cells than the periphery and does not generate an immune response in the same manner. Furthermore, components of the immune system, including tumor-infiltrating T lymphocytes, appear depleted surrounding certain tumors including glioblastoma.³⁵ As glioblastoma has been the most widely investigated brain tumor using immunotherapy, much of our knowledge on immunotherapeutic mechanism of action and toxicity within the CNS is based on such research. It is encouraging that clinical trials for glioblastoma have progressed to Phase III; however,

results have so far been less than promising. The first ever Phase III clinical trial evaluating the PD-1 checkpoint inhibitor nivolumab in glioblastoma (Checkmate 143) failed to increase survival compared to bevacizumab. Another phase III clinical trial evaluating a vaccine known as rindopepimut that targets EGFRvIII, overexpressed in many glioblastomas, also failed to improve survival.³⁶

Current treatment regimens for medulloblastoma have been relatively unchanged for the past two decades. Medulloblastomas have typically been more radiosensitive than other pediatric brain tumors, including glioblastomas.⁴ For children older than 3 years, external beam radiation to the spine and brain combined with multidrug chemotherapy and surgery has been standard.^{37,38} Treatment is stratified based on risk: average risk and high risk.³⁹ Average risk includes residual tumor post-surgery less than 1.5 cm² with no metastasis, while high risk includes metastases and greater than 1.5 cm² residual disease.⁴⁰ While both are treated with radiation, higher-risk patients are given larger boosts of radiation. Both groups typically receive four 28-day cycles of cisplatin, vincristine, and cyclophosphamide.^{39,41,42} Children younger than 3 years are given high dose chemotherapy but no radiation due to the adverse effects on the developing brain. However, surgery can result in multiple complications including cerebellar mutism syndrome.⁴³ Radiation in older children has been linked to reduced IQ and induction of secondary cancers, vasculopathy, hearing loss, and future strokes.^{44–48}

The main cause of death from medulloblastoma is due to metastasis. While metastatic tumors have the same molecular subtype as the primary medulloblastoma, the genomics of the secondary tumor often differ.^{49,50} Additionally, pediatric brain tumors differ from adult brain tumors in clinical presentation in addition to gene mutations, embryological origins, and microenvironment of the tumor.⁵¹ Even more problematic, major subpopulations of medulloblastoma patients do not respond to current immunotherapy due to lack of antigenic mutations and/or immuno-resistant properties of medulloblastoma cells.⁵²

However, there have been many advances in our understanding of the mechanisms of immunosuppression in patients with medulloblastoma. Such knowledge will help researchers exploit these pathways to develop immunotherapies that can circumvent the tumor's immunosuppressive properties. Already documented is the finding that immune response correlates with prognosis in patients with medulloblastoma. For example, there is evidence that inflammation and immunological markers correlate

with survival of patients with medulloblastoma. In one study, 144 patients with medulloblastoma were divided into subgroups (SHH, WNT, group 3, group 4) and evaluated for preoperative hematological markers and overall survival.⁵³ The study found that an increased preoperative neutrophil-to-lymphocyte ratio (NLR) as well as platelet-to-lymphocyte ratio (PLR) predicted worse prognosis in childhood medulloblastoma patients but not in adult patients. Preoperative monocyte-to-lymphocyte ratio, mean platelet volume, platelet distribution width, and albumin-to-globin ratio did not correlate with prognosis in either age group. Furthermore, the levels of NLR and PLR in group 3 were higher than in WNT.

The remainder of this paper will discuss the most recent findings on various forms of immunotherapy for the treatment of medulloblastoma.

Vaccine Therapy in Medulloblastoma

Cancer vaccines are a form of immunotherapy that can be categorized into several groups: whole tumor cell, peptide, DNA, RNA, etc. Cancer vaccines must activate an immune system that has become tolerant of the host cancer.⁵⁴ Whole tumor cell and peptide cancer vaccines deliver tumor antigens directly to antigen-presenting cells (APCs) for processing. These APCs then present the antigens as epitopes on the surface of their own MHC class I and II molecules, which are then recognized by T cells.^{10,55} One example is the FDA-approved sipuleucel-T vaccine for prostate cancer, which consists of a fusion protein antigen containing a tumor-specific antigen, prostate acid phosphatase, and expanded with granulocyte-macrophage colony stimulating factor (GM-CSF). The fusion protein is incubated for many days with antigen-presenting cells taken from the patient, then re-infused into the patient.⁵⁶ Sipuleucel-T has increased overall survival by 4 months in hormone-resistant prostate cancer.⁵⁷

In contrast, DNA- and RNA-based cancer vaccines bypass the human leukocyte antigen (HLA) haplotype restriction; this system allows a person's own cellular processes to generate proteins via transcription/translation for more personalized processing.^{10,58–61} One research group generated RNA-loaded autologous dendritic cells from patients with medulloblastoma; these cells were matured *ex vivo* and activated with inflammatory cytokines.⁶² The activated dendritic cells were then incubated with a personalized group of tumor messenger RNA amplified from that individual's cDNA library including tumor-specific transcriptome. The dendritic cells were

cultured with T cells and then re-administered with the activated T cells in the patient. This research is currently in Phase I and II studies involving patients with medulloblastoma (NCT01326104). The challenge of using dendritic cells is that it requires tumor collection from each patient, which results in extensive processing.⁶³ In another study, autologous monocyte-derived dendritic cells were loaded with tumor lysate, which served as a source of tumor antigens.⁶⁴ Patients with high-grade glioma and atypical teratoid-rhabdoid tumors responded more favorably to the vaccine treatment than the medulloblastoma patients.

Pre-clinical trials evaluating vaccines showed promise in medulloblastoma but have yet to be translated into human trials. More favorable responses have been observed in other brain tumors such as glioma thus far. Two of the most successful vaccines used in gliomas include epidermal growth factor receptor variant III (EGFRvIII) junctional epitope combined with GM-CSF and whole tumor cell antigen pools presented by autologous dendritic cells.^{2,65–68} Rindopepimut, the vaccine consisting of EGFRvIII-specific peptide, stimulates the immune system to target the EGFRvIII protein on intracerebral melanoma tumors. In one study, C3H mice with intracerebral melanoma expressing EGFRvIII and treated with the vaccine had a 600% increase in median survival compared to controls.^{56,69} However, a phase III trial of 165 glioblastoma patients with EGFRvIII was stopped due to lack of efficacy.³⁶ EGFRvIII, while specific in a variety of tumors, can downregulate and escape immune system surveillance.^{70,71}

Unfortunately, recent clinical trials to evaluate cancer vaccines in medulloblastoma patients have been relatively unsuccessful. A phase I/II trial assessed a cancer vaccine consisting of autologous dendritic cells targeting tumor proteins NY-ESO-1 melanoma antigen gene-A1 (MAGE-A1) and melanoma antigen gene-A3 (MAGE-A). The vaccine, administered in combination with decitabine and hiltonol to patients with refractory medulloblastoma was terminated after enrolling one patient due to a serious adverse event and/or disease progression (NCT02332889). Another phase I study evaluated the safety and tolerability of a cancer vaccine consisting of autologous dendritic cells loaded with allogeneic brain tumor stem cells and imiquimod (NCT01171469). Eight patients enrolled in this study, and the trial was terminated with no results posted. Similarly, an earlier Phase II trial that evaluated an autologous tumor cell vaccine concurrent with standard chemotherapy (cyclophosphamide, cisplatin, carmustine), stem cell transplantation,

and interleukin-2 was terminated without any results published (NCT00014573). To some degree, the failure of these trials may be attributed to limited viable tumor tissue available for processing and inconsistent expression of tumor antigens.

Yet a number of cancer vaccines with more sophisticated processing combined with adjuvant therapy to enhance the immune response are currently in clinical trial for the treatment of medulloblastoma. In the previously mentioned phase I/II trial using total tumor RNA-loaded dendritic cells, researchers showed that sufficient RNA can be amplified with as few as 500 tumor cells, which allows vaccine preparation from surgical biopsies (NCT01326104). Furthermore, the immunotherapy is infused during recovery from chemotherapy, an advantage as lymphodepletive conditioning is the most effective therapeutic strategy for the treatment of metastatic melanoma. Finally, ex-vivo expanded RNA-loaded autologous lymphocytes will also be infused into the patient to mount an increased immune response, a novel approach distinct from earlier studies. Results are eagerly awaited.

Oncolytic Viral Therapy in Medulloblastoma

Similar to vaccines, oncolytic viral therapy attempts to stimulate the host's innate immune response. Oncolytic viruses are intended to propagate within tumor cells until they are recognized by the immune system and targeted for rejection.^{10,72-77} When a virus-infected tumor cell lyses, a systemic immune response is stimulated that targets tumor antigens that have shed; this process is known as epitope spreading. Viral therapy has been studied in a variety of cancers with encouraging results, and one study showed that oncolytic herpes simplex virus expressing murine IL-12 (interleukin-12) cured most mice in two different glioma models.⁷⁸ In a phase I trial consisting of human patients with recurrent glioma, the oncolytic adenovirus DNX-2401 was injected directly into the primary tumor and reportedly shrunk the tumor by at least 95% in 3/25 of the patients.⁷⁹ Five of the treated patients lived for at least 3 years post-treatment. The researchers theorized that the dramatic tumor response was a direct effect of the virus eliciting a long-lasting immune-mediated response.

In medulloblastoma, oncolytic viral therapy using poliovirus has demonstrated efficacy in vitro. One study evaluated the expression of the poliovirus receptor CD155 in medulloblastoma and the ability of that virus to infect as

well as inhibit tumor proliferation.⁸⁰ A rhinovirus recombinant form of polio (PVSRIPO) was used, in which part of the virus in the live polio vaccine was removed and replaced with a reciprocal part of a common cold virus. The viral replacement with the reciprocal part of the common cold virus ensures that the modified virus does not cause polio in the host. Poliovirus receptor mRNA expression was studied in 763 medulloblastoma samples from two group 3 cell lines: D283 and D341. CD155 was expressed in the medulloblastoma cell lines, and PVSRIPO infection resulted in decreased tumor proliferation in the D341 cell line at 48 hours resulting in cellular death. The study noted that poliovirus receptor is highest in medulloblastoma subtypes WNT α , WNT β , and group 3 γ . This study concluded that PVSRIPO is capable of infecting and killing group 3 medulloblastoma.

Though controversial, there is evidence that human cytomegalovirus (CMV) infection is linked to medulloblastoma, possibly by promoting an inflammatory environment.⁸¹⁻⁸³ This pro-inflammatory state occurs, in part, by an up-regulation of cyclooxygenase-2 (COX-2), and up-regulation of COX-2 has been demonstrated in medulloblastomas. Even more, COX-2 induces synthesis of inflammatory stimulants including prostaglandin E2 (PGE2), which stimulates the proliferation of medulloblastomas.⁸⁴ Human CMV has been found in glioblastomas and up to 99% of brain metastases of colorectal and breast cancers, but it is still controversial whether CMV infection is linked to the development of medulloblastomas.⁸⁵ Several research groups reported CMV infection in human medulloblastomas including cell lines and xenografts.⁸⁶ One study showed that the antiviral drug ganciclovir results in reduced clonogenic capability of medulloblastomas in vitro.^{84,86} The COX-2 inhibitor celecoxib also diminishes clonogenic ability by decreasing PGE2 levels, which reduced stem cell-like ability and increased sensitivity to radiation in medulloblastoma transplanted into mice.⁸⁶⁻⁸⁸ Synergistically, valganciclovir and celecoxib reduce tumor growth of CMV-infected tumor cells both in vitro and in mice engrafted with human medulloblastoma cells.⁸⁶

Currently, a host of modified viruses are under evaluation for the treatment of medulloblastoma. A combination of CMV RNA-pulsed dendritic cells, tetanus-diphtheria toxoid vaccine, and GM-CSF is currently in a phase I trial (NCT03615404). PEP-CMV, a vaccine comprised of a synthetic long peptide from human CMV matrix protein pp65, is another investigational viral agent in phase I trial and given after the patient receives a pre-conditioning tetanus-diphtheria toxoid vaccine

(NCT03299309). The previously mentioned poliovirus with polio/rhinovirus recombinant (PVSRIPO) is also under phase I testing in patients with a variety of brain tumors including medulloblastoma (NCT03043391). In this ongoing study, the recombinant virus is injected directly into the primary tumor in the brain. Even a modified measles vaccine is under investigation in a phase I trial. An altered virus is injected directly into the primary tumor or via lumbar puncture if the disease is disseminated (NCT02962167). Similarly, a re-engineered reovirus known as reolysin, in combination with GM-CSF, is under investigation in patients with high-grade brain tumors (NCT02444546). Finally, a phase I trial is currently investigating an oncolytic herpes simplex virus-1 engineered to introduce viral mutations that enable it to selectively replicate within tumor cells. Lysed tumor cells then disseminate to subsequently target other tumor cells while also mounting a host immune response by exposing shed cancer antigens (NCT03911388). In addition, low dose radiation is administered to enhance viral replication and promote tumor lysis. The safety of this regimen, in addition to radiographic and pathologic evidence of anti-tumor response, has been successfully demonstrated in three phase I trials with adult glioma patients and an ongoing trial in children with recurrent supratentorial brain tumors (NCT02457845).

Checkpoint Inhibitors in Medulloblastoma

Checkpoint inhibitors have shown promise in a variety of cancers including melanoma, small cell lung cancer, and renal cell carcinoma. Activity of checkpoint inhibitors such as nivolumab depends on expression of PD-L1 on tumor cells. However, there are limited data on PD-L1 expression in brain tumors including medulloblastomas. One study with 89 patients in Korea showed that 0% of medulloblastoma patients demonstrated expression of PD-L1 (0 out of 28 patients).⁸⁹ This was in direct contrast to other pediatric brain tumors including 40% of atypical teratoid/rhabdoid tumor (8 out of 20), 20% of ependymoma (4 out of 20), and 19% of high-grade glioma (4 out of 21) that demonstrate expression of PD-L1. This study concluded that medulloblastoma may not be a suitable candidate for PD-1 checkpoint blockade. These results are consistent with another study, which found a complete absence of PD-L1 on medulloblastomas, further suggesting limited value for PD-1 blockade to treat medulloblastoma.⁹⁰

However, PD-1 blockade may depend on the subtype of medulloblastoma. Given that human studies evaluating PD-1 blockade are limited in human medulloblastoma patients, one study using immunocompetent murine models of human SHH-driven and group 3 medulloblastoma showed that PD-1 blockade demonstrated greater anti-tumor efficacy in group 3 medulloblastoma tumors compared to SHH tumors.³¹ This study showed that response to immune checkpoint blockade differs across medulloblastoma subtypes, at least in murine models. Of note, anti-PD-1 antibody binding was evident in peripheral T cells and not tumor-infiltrating lymphocytes within the brain's tumor microenvironment. But this peripheral PD-1 blockade did result in increased CD3 T cells within the tumor microenvironment. This study concluded that effective PD-1 blockade does not require peripheral antibodies to penetrate the tumor microenvironment within the brain.

While certain studies documented limited levels of PD-L1 in medulloblastoma, one study showed that the SHH variant has higher PD-L1 expression than the other groups (and lowest MYC expression).⁹¹ Groups 3 and 4 had the highest levels of MYC expression (and lowest PD-L1 expression). This study subsequently showed in vitro IFN- γ -induced expression of PD-L1 in all cell lines while radiation stimulated variable expression. Altering the expression of MYC did not change the expression of PD-L1. The study concluded that TH1 cytokine induction is the most potent stimulator of PD-L1 expression in vitro, suggesting that an inflamed microenvironment is required for PD-1 activation in medulloblastoma.

Another study showed that in brain tumors resistant to PD-1 blockage, lineage negative hematopoietic stem and progenitor cells (HSCs), derived from bone marrow that expresses C-C chemokine receptor type 2 (CCR2+), reversed resistance by sensitizing mice to PD-1 blockage.⁹² PD-1 blockage with HSC transfer increased T cell activation within preclinical models of medulloblastoma. The HSC transfer also diminished resistance to adoptive cellular therapy against medulloblastoma.

In human trials so far, immune checkpoint inhibitors have failed as monotherapy in recurrent glioblastoma.^{63,93} Lack of efficacy is believed due to minimal T cell infiltration into glioblastoma tumors, with similar findings reported in medulloblastomas.^{63,90} One study showed that cytotoxic T cells infiltrate medulloblastomas with variable activation and do not correlate with increased survival.⁹⁰ Medulloblastoma cells have decreased antigen presentation via downregulated MHC-1 expression of PD-L1. Thus, it is

possible that PD-1/PD-L1 contributes to immune escape in medulloblastoma.⁶³

Nevertheless, the checkpoint inhibitors against PD-1, including nivolumab, pembrolizumab, and durvalumab, are each under investigation in clinical trials recruiting medulloblastoma patients and other CNS tumors (NCT03173950, NCT02359565, NCT02793466). Furthermore, nivolumab with and without ipilimumab, another checkpoint inhibitor that targets a different pathway known as CTLA-4, is also under investigation in a phase II trial of patients with high-grade CNS malignancies including medulloblastoma (NCT03130959). In addition, an inhibitor of the immune checkpoint pathway indoleamine 2,3-dioxygenase (IDO), in combination with radiochemotherapy, is currently in a phase I trial involving pediatric brain tumors. This study was recently expanded into a new phase II trial in August 2019. Although results from the first trial have not been released, the acceleration into phase II investigation is encouraging.

Natural Killer Cells in Medulloblastoma

Adoptive cell therapy, also known as cellular immunotherapy, involves isolation of immune cells from the host for modification and subsequent transplantation back into the host for direct targeting of tumor cells. Natural killer (NK) cells, a subset of the innate immune system often involved in controlling viral infections, have been studied to potentially serve as a type of adoptive cell therapy. Natural killer cells target medulloblastoma cells, primarily through activation of receptors such as natural killer group 2 member D activator receptor (NKG2D) and others (DNAM-1, NKp30, and NKp46), via ligands on medulloblastoma cells.⁹⁴ Resistance to natural killer cell activity may be one mechanism by which medulloblastoma tumors can evade the natural immune response. One study analyzed tumor samples from 54 medulloblastoma patients for ligands of NKG2D, which included UL16 binding protein (ULBP-2) and major histocompatibility complex class I-related chains A (MICA). The study found that a certain medulloblastoma cell line, HTB-186, was relatively resistant to *in vitro* NK cell toxicity.⁹⁵ By blocking the NKG2D receptor on the natural killer cells and blocking MICA/ULBP-2 on the HTB-186 tumor cells, the resistance to natural killer cell lysis increased. On the other hand, blocking the HLA class I on these cells and incubation with NK cells treated with IL-15 resulted in

increased killing of tumor cells *in vitro*. The study found that at least *in vitro*, interactions of NKG2D/MICA-ULBP-2 play a role in natural killer cytotoxicity of medulloblastoma, and increased expression of HLA class I may increase medulloblastoma resistance to natural killer cell toxicity.

Cord blood natural killer cells may potentially be an effective immunotherapy due to their recognition of tumor cells and ability to be expanded exponentially in the laboratory.⁹⁶ Natural killer cells lyse tumor cells without immunization and secrete granules containing granzymes and perforin.^{63,97} However, their efficacy is limited by immunosuppressive cytokines released in the medulloblastoma microenvironment such as transforming growth factor B (TGF-B). One study modified cord blood natural killer cells to express a dominant negative TGF-B receptor.⁹⁶ When unmodified natural killer cells were added to TGF-B rich medulloblastoma media, their cytotoxic ability was reduced. However, the cytotoxic effect with modified natural killer cells was stronger and unaffected by the environment. The study concluded that medulloblastoma may be rendered susceptible to natural killer cytotoxicity with neutralization of TGF-B.

Other studies showed that CD1d, an antigen-presenting molecule for natural killer T cells, is highly expressed on medulloblastoma cells. This molecule is particularly overexpressed in the SHH subtype.⁹⁸ One study showed that 9 of 20 primary medulloblastoma tumors express CD1d, which means it may be a target for immunotherapy. Another study showed that 13 of 38 medulloblastomas (34%) expressed CD1d. The majority (82%) of groups 3 and 4 medulloblastomas in that study were CD1d-negative.

One study evaluated lymphokine-activated killer cells (LAK) gathered by harvesting non-antigen primed monocytes in the peripheral blood of HLA-matched donors or the patients themselves, then expanding the monocytes by stimulation with anti-CD3 antibodies.^{99,100} *In vitro* studies showed that LAK cells target medulloblastomas but spare healthy brain tissue.^{99,101} This therapy was expanded clinically and published in a case series involving 8 patients with medulloblastomas that involved CSF dissemination.² Intrathecal injection of the LAK cells provided a complete response within 3 months and a durable response up to 20 months in three patients.^{100,102} However, only two other reports of successful treatment with LAK have been published in the past 20 years.^{103,104}

Currently, an ongoing phase I clinical trial is evaluating the safety and efficacy of ventricular infusion of autologous ex-vivo expanded NK cells in patients with recurrent CNS tumors of the posterior fossa (NCT02271711). In another phase II trial, NK cells from a donor are administered to patients after allogeneic hematopoietic cell transplantation and reduced intensity radiochemotherapy in order to stimulate graft versus tumor response (NCT02100891).

CAR T Cell Therapy in Medulloblastoma

While oncolytic viral therapy and cancer vaccines aim to stimulate a specific T cell response, a number of steps must be orchestrated including APC activation and presentation to the T cells. The T cells must then penetrate the microenvironment of the tumor. However, tumor cells can downregulate MHC class I, which disables T cell recognition. Engineered chimeric antigen receptor T cells (CAR T) can overcome MHC downregulation by cancer cells and directly target surface antigens on the malignant cells.^{10,105–110} As another form of adoptive cell therapy, CAR T therapy involves use of a patient's own re-engineered T cells and was first reported in the 1980s.^{56,111} The technology has been translated clinically only recently. CAR T therapy consists of hybrid receptors, which fuse T lymphocytes and other cells with an antibody-binding domain. The T lymphocyte can then theoretically target an antigen of choice.¹¹² These synthesized receptors are retrovirally integrated into the patient's own T cells and then reinfused back into the patient. But design of CAR T is complex, requiring a short-chain variable fragment (ScFv) to bind to an antigen of choice, a transmembrane domain, costimulatory domain, hinge-linker region, and intracellular CD3 ζ tail.^{56,112} While first-generation CAR T therapy lacked a costimulatory domain resulting in limited expansion, the incorporation of CD28 and CD137 co-stimulating domains produced second-generation CARs effective against many blood cancers including relapsed B cell acute lymphocytic leukemia (ALL), chronic lymphocytic leukemia, and non-Hodgkin lymphoma.^{112,113} CAR T therapy does not require a systemic immune response and is theoretically useful for tumors that lack a high tumor mutational burden.⁵⁶ One study showed that intrathecal and intraventricular administration of CAR T targeting IL-13 receptor α 2, an antigen associated with glioma, led to tumor regression in

an adult patient with relapsed glioblastoma.¹¹⁴ Of note, the patient relapsed 6 months later when a tumor did not express the antigen. But the therapy was well tolerated without cytokine release syndrome or severe neurotoxicity.

Multiple trials using CAR T targeting the CD19 antigen have proven that CAR T can cross the blood–brain barrier, as these cells were detected in the CSF via immunofluorescence post-treatment.¹¹⁵ Other studies involving glioblastoma and B cell ALL patients also reported CAR T cells in brain tissue following infusion as demonstrated by flow cytometry.¹¹⁶ However, severe grade 3 neurotoxicity was reported in several of the B cell ALL patients following infusion revealing new challenges that must be investigated. CAR T cells essentially contain the cytotoxicity of T cells with specificity of antibodies; however, side effects have been noted in the brain including neurotoxicity, inflammation, and increased intracranial pressure. The development of CAR T cells that are activated only when stimulated by multiple tumor antigens simultaneously may decrease the incidence of side effects in the future.

CAR T cells showed anti-tumor activity against glioblastoma by targeting human epidermal growth factor receptor 2 (HER2).^{114,117} Since HER2 is overexpressed in certain medulloblastomas and anti-HER2 CAR T therapy has been successfully used against other cancers, it is plausible that such therapy can be used against medulloblastoma. One study successfully evaluated HER2-BBz-CAR T cells in mice and non-human primates.¹¹⁸ The CAR T cells cleared the medulloblastoma implanted in the posterior fossa of the mice without significant toxicity. The therapy showed strong efficacy both in vitro and in the murine medulloblastoma models, leading to possible future testing in humans. Another study showed that the antigen B7-H3 (CD276) could successfully be used as a target for CAR T cell therapy in a variety of pediatric cancers.¹¹⁹ The B7-H3 CAR T cells induced tumor regression in xenograft models of medulloblastoma, osteosarcoma, and Ewing sarcoma.

Challenges in developing the next generation of CAR T cell therapy for medulloblastoma include accessing the specialized tumor microenvironment within the CNS and variability of target antigens in tumor cells.^{56,115} So far, current CAR T cell proliferation has been poor with inadequate cytokine secretion. Furthermore, there is the challenge of tumor variants losing antigen, thus theoretically rendering CAR T ineffective without a target to hone in on. A new generation of CAR T cells will need to be engineered to

circumvent these issues. Gene editing technologies are currently in development to increase cytokine overexpression, target a variety of antigens simultaneously, and provide more intricate control of CAR T signaling.

Antigenic escape must also be resolved for CAR T to become an established treatment against medulloblastoma. Antigenic escape has been one of the primary causes of tumor resistance and relapse in the leukemia patients treated with CD19-targeted CAR T therapy.^{56,109,120} This issue will be especially challenging in CNS tumors including medulloblastoma with known antigen heterogeneity. Decreased expression of tumor antigens below the level that can be detected by CAR T cells has been reported in multiple trials including the groundbreaking CAR T trial in brain tumors where a glioblastoma patient relapsed with an IL13-Ra2-negative tumor that lost antigen expression.¹¹⁴ CAR T cells must be designed to target multiple tumor antigens to prevent antigenic escape and relapse. For example, trivalent CAR T cells were able to overcome antigenic variability in patients with glioblastoma.¹²¹

Toxicity of treatment must be seriously considered, particularly in children with developing nervous systems. It is still unclear what the long-term effects of immunotherapy-induced inflammation in the CNS are. Vaccine and antibody therapy have yet to report serious adverse events in the CNS. However, neurotoxicity was reported in multiple CAR T cell trials targeting CD19 and included aphasia, encephalopathy, seizures, and memory loss in leukemia patients.^{109,122–124} The increased neurological disease burden in patients with CNS tumors may demonstrate even greater neurotoxicity risk.¹¹⁵

Currently, HER2-specific CAR T therapy is under investigation in HER2-positive recurrent CNS tumors including medulloblastoma using autologous CD4+ and CD8+ T cells lentivirally transduced to express a HER2 chimeric antigen receptor and EGFRt (NCT03500991). Similarly, autologous CD4+ and CD8+ lentivirally transduced to express an EGFR806 chimeric antigen receptor and EGFRt is currently in a phase I trial recruiting patients with EGFR-positive CNS tumors including medulloblastoma (NCT03638167).

Radiation with Immunotherapy in Medulloblastoma

The future of medulloblastoma therapy may incorporate radiotherapy that synergistically enhances the efficacy of immunotherapy. Studies have demonstrated that ionizing radiation can enhance the surface expression of MHC class

I molecules and alter cellular proteins, enhancing the visibility of treated tumor cells to the immune system.^{125,126} One study evaluated the effect of low dose radiation on the functional immunological responses of medulloblastoma cell lines (DAOY, D283, D341).⁵² The study found that low dose radiation upregulated the expression of HLA class I and HLA class II molecules in medulloblastoma cells by more than 20% in vitro, including subtypes MAGE C1, CD137, and ICAM-1. Even more, an increase in reactive oxygen species resulted in altered cell surface expression of monoclonal antibody target molecules HER2 and vascular endothelial growth factor (VEGF), suggesting that low dose radiation and monoclonal antibodies can create a synergistic effect. While no studies have demonstrated increased survival using combination radiation and immunotherapy in patients with medulloblastoma, one study did show improved response of immunotherapy in patients with melanoma brain metastasis when given within 1 month of radiation.¹²⁷ While various other trials have similarly shown greater efficacy when combining radiation with immunotherapy in lung malignancies, this study shows that the synergistic effect can be demonstrated in malignancies within the CNS.^{128,129} The induction of immune-mediated medulloblastoma cell death coupled with radiation-induced tumor necrosis may 1 day be part of the future paradigm in the treatment of medulloblastoma.

One recent phase II trial completed in June 2019 evaluated the use of radiolabeled monoclonal antibodies to target tumors and deliver radioactive substances to the cells in combination with radiation and chemotherapy in patients who have undergone surgery for medulloblastoma (NCT00058370). Patients first received iodine ¹³¹I monoclonal antibody 3F8 followed by 6 weeks of radiation, vincristine, lomustine, and cisplatin. Six patients enrolled in the trial, and results are pending. In another trial, a radiopharmaceutical, ⁹⁰Y-DOTA-tyr3-octreotide, and its renal protectant, aminosyn II, were infused in patients with somatostatin receptor-positive tumors including medulloblastoma, with the intent to deliver radiation directly into tumor cells (NCT02441088).

Novel Antigenic Targets in Medulloblastoma

One of the major challenges in treating medulloblastoma with immunotherapy is the low immunogenicity and mutational load. Despite the low mutation burden, one study did find immunogenic peptides specific to the medulloblastoma tumors in human patients.¹³⁰ Induction of a CD8 T cell

response was successfully initiated for the neoepitopes derived from neuraminidase 2 (NEU2), programmed cell death 10 (PDCD10), supervillin (SVIL), histidine ammonia-lyase (HAL), tRNA splicing endonuclease subunit 54 (TSEN54), and proprotein convertase subtilisin (PCSK9) variants. Tumor-derived neoantigens were specific to each patient and confirm that T-cell therapy can be designed for the individual patient in the future.

Another recent study in 2018 analyzed 36 medulloblastomas (18 adults and 18 pediatric samples) via next-generation sequencing to identify overexpressed proteins. Although PD-L1 expression was uncommon, there were various other proteins overexpressed that may be used as therapeutic targets.¹³¹ Testing on samples was performed at the discretion of the physician, so not every sample was evaluated for all mutations. Multidrug resistance-associated protein 1 (MRP1) was found in 89% of the medulloblastomas (8 out of 9), tubulin beta 3 class 3 (TUBB3) was found in 86% of samples (18 out of 21), and thymidylate synthase (TS) was found in 80% of the samples (24 out of 31). Phosphatase and tensin homolog (PTEN) was evident in 85% of tumors (28 out of 33), topoisomerase 2A (TOP2A) was overexpressed in 84% of tumors (26 out of 31), and ribonucleotide reductase M1 (RRM1) was found in 71% of tumors (15 out of 21). Even more, topoisomerase 1 (TOP1) was found to be overexpressed in 90% of metastatic tumors (9 out of 10) relative to posterior fossa medulloblastomas that did not metastasize (50%, 10 out of 20). Additionally, PGP expression was found solely in the pediatric medulloblastomas.

Another study found that the tumor-associated antigen PRAME was detectable in 82% of medulloblastomas, independent of subgroup.¹³² Since PRAME has limited expression in normal tissues, this antigen can serve as a potent target for immunotherapy. This study also evaluated PRAME as a viable target in immunotherapy using genetically altered T cells with PRAME-specific T-cell receptors. The T cells were found to control tumor growth in a murine model of HLA-A*02 medulloblastoma, concluding that PRAME-specific T-cell receptor therapy may show promise in the treatment of HLA-A*02 medulloblastoma. Another study analyzed 37 tumor samples and showed that PRAME was overexpressed in 84% of the tumor samples.¹³³ Of note, the study did not find any association between PRAME overexpression and patient survival.

Cancer testis antigens (CTAs) are among the best-described antigens in medulloblastomas, specifically the

subtypes MAGE and GAGE proteins.^{2,134,135} These proteins were first described in melanoma and shown to stimulate T cell anti-tumor responses. These antigens can also be found in lung, breast, esophageal, and hepatocellular cancers.¹³⁴ Even more, expression of these antigens has been found to correlate with aggressiveness of tumor and resistance to chemotherapy.¹³⁴ One study explored expression on eight medulloblastomas.² MAGE-4 expression was evident in 50% of the medulloblastomas, SCP-1 was found in 50% of the medulloblastomas, and the other CTAs were not found. Another study that used Western blot analysis found GAGE, MAGE-A family, and MAGE-A1 in 84%, 62%, and 46% of medulloblastoma samples, respectively. This same study showed that the downregulation of GAGE and MAGE led to an increased response to cisplatin. MAGE has been successfully targeted by vaccine therapy in various other cancers.^{136–139}

CD47, a protein on the surface of a variety of solid tumors including medulloblastoma, is involved in evading the host immune response and may be a viable antigenic target. Preclinical trials have shown that antagonizing CD47 with a humanized monoclonal antibody can activate macrophages to phagocytize cancerous cells in several CNS tumor models.¹⁴⁰ Finally, a subset of the VEGF group, known as placental group factor (PIGF), was secreted by various pediatric brain tumors including medulloblastoma.¹⁴¹ Inhibition of PIGF via the receptor neuropilin 1 pathway delayed tumor growth in murine models of medulloblastoma.

A current, ongoing phase I/II trial is testing various doses of a humanized monoclonal antibody known as TB-403 against PIGF in patients with refractory medulloblastoma (NCT02748135). Patients with other cancers that exhibited PIGF secretion in past studies, including Ewing sarcoma and alveolar rhabdomyosarcoma, are also eligible for this trial. These studies demonstrate the vast potential of immunotherapy when tumor-specific antigens can be identified.

Challenges in Treating Medulloblastoma with Immunotherapy

Various cell lines derived from medulloblastoma have existed since the 1980s.^{51,142} However, tumor implantation models do not reflect the normal events of tumor initiation and growth. Furthermore, experiments are often conducted in immunodeficient mouse models to allow tumors to be transplanted without eradication by the host immune

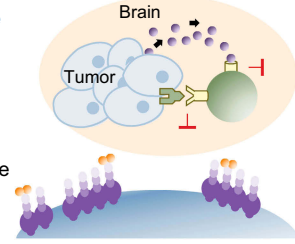
response. Genetically engineered animal models must be developed to introduce genetic mutations that drive the development of tumors and provide more accurate insight

into tumor development and response to treatment. Many genetically engineered models of medulloblastoma have been developed.^{143,144}

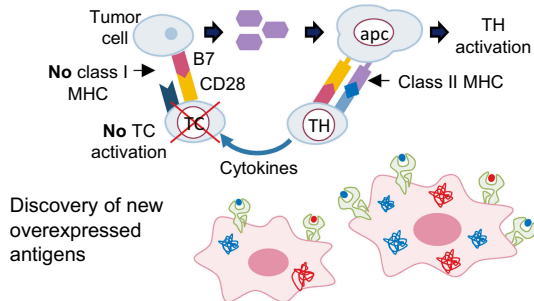
Challenges & Solutions

1-Immunosuppressive properties of tumors

Cord blood natural killer cells expressing negative TGF-β receptor

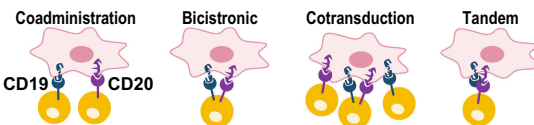


3-Lack of antigen targets

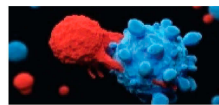


5-Antigenic variation Antigenic escape (down regulation)

New generation CAR T therapy with multiple antigen targets, cytokine over-expression, better control of signaling

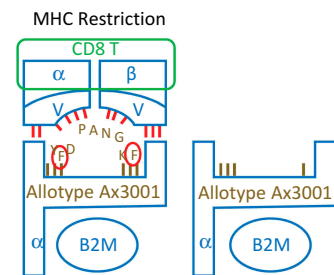


7-Neurotoxicity

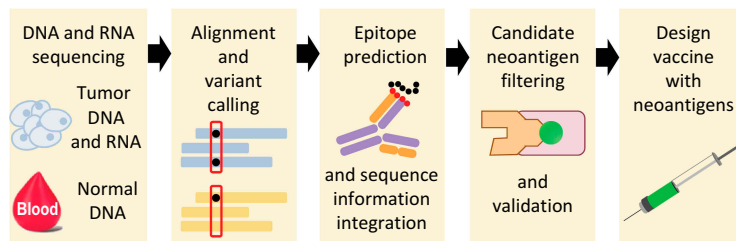


CAR T therapy with simultaneous targets with less specificity to brain, minimize inflammation

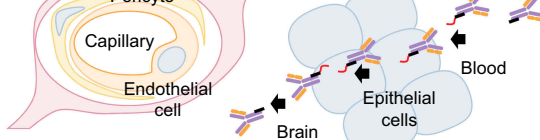
9-HLA haplotype restriction



Using DNA and RNA cancer vaccines

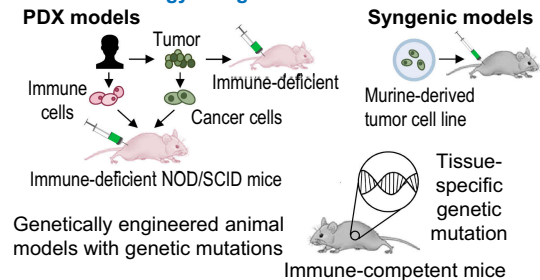


2-BBB

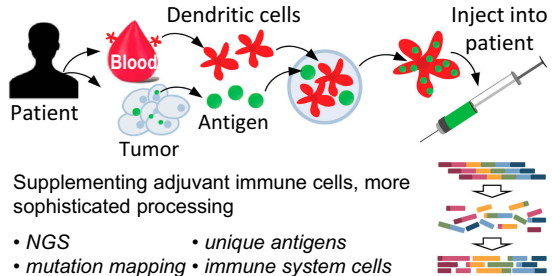


Transcytosis of small peptides across BBB or Radiation to break the BBB

4-Tumor implantation in animals is not applicable to real tumor biology and growth



6-Failure of past cancer vaccines



8-Difficult to gather sufficient tumor

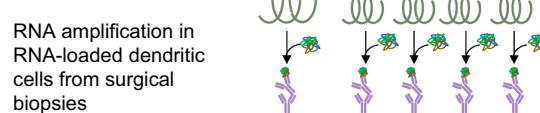


Figure 1 Immunotherapy in medulloblastoma – Challenges and Solutions.

The blood–brain barrier also continues to be a major challenge in targeting brain tumors. IgG antibodies penetrate the blood–brain barrier poorly, due to their large size. Tight junctions of epithelial cells in the blood–brain barrier prevent passage of molecules larger than 500 Da.¹⁴⁵ This may be one of the reasons that checkpoint inhibitors have failed to improve survival in recurrent GBM in large multicenter trials such as the Phase III trial Checkmate-143.¹⁴⁶ On the other hand, more encouraging results were found in clinical trials evaluating checkpoint inhibitors in melanoma and NSCLC that had metastasized to the brain.^{147,148} While it is likely that the difference in trial results is due to differences in the biology of the tumor, it is possible that patients with metastatic brain tumors have a disrupted blood–brain barrier since tumor cells were able to initially cross the barrier.¹⁴⁹ Current trials are evaluating checkpoint inhibitors combined with radiotherapy against glioblastoma multiforme with the hope of disrupting the blood–brain barrier (NCT02617589, NCT03576612).

Smaller peptides less than 50 residues (which can cross cell membranes) have been shown to circumvent the blood–brain barrier via receptor-mediated transcytosis.¹⁵⁰ Anti-human epidermal growth factor (HER2) antibodies conjugated with cell-penetrating peptides were able to cross the blood–brain barrier and prolong overall survival in BT-474 mice with brain tumors.¹⁵¹ It has also been demonstrated that CAR T cells can bypass the blood–brain barrier, particularly when conjugated to cell-penetrating peptides less than 40 residues that can undergo transcytosis. Coupling of vaccines or antibody therapies with cell-penetrating peptides may be necessary to bypass the blood–brain barrier.

Conclusion

Future forms of immunotherapy in medulloblastoma must overcome tumor antigen heterogeneity, the blood–brain barrier, and mechanisms of tumor immunosuppression (Figure 1). Currently, vaccine and CAR T therapy are limited by lack of documented tumor antigens.¹¹⁵ Preclinical research must be continued to identify new antigen targets and engineer therapies that can target multiple antigens simultaneously. By counteracting tumor immunosuppression and activating immunity via vaccines, viruses, and/or adoptive T cell therapy, combination therapy should be the hallmark of future treatment of medulloblastoma to improve survival outcomes.

Disclosure

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