Research Program, University of Colorado School of Medicine/Children's Hospital Colorado, Aurora, CO, USA

Diffuse intrinsic pontine glioma (DIPG) is a fatal pediatric brain tumor with no curative treatments. Approximately 80% of DIPGs contain an H3K27M mutation. The implications of the mutation and how they may be targeted are not fully understood. We established an H3K27M effectisolating model by transducing H3K27-wildtype lines (HSJD-GBM-001, normal human astrocytes) with lentiviral-packaged H3K27M. We characterized H3K27M-related changes through western blot, phenotypic assays, and RNA-seq. Drug screening of H3K27-wildtype and matched H3K27Mtransduced lines was used to identify targets more effective with H3K27M present. Patient-derived pediatric glioblastoma and DIPG lines (BT-245, SU-DIPG-IV, HSJD-DIPG-007, SU-DIPG-XIII*, SF7761) were used for validation. We observed increased H3K27ac and decreased H3K27me3, as well as increased proliferative and migratory abilities, with the addition of H3K27M to H3K27-wildtype lines. RNA-seq showed downregulation of cell cycle regulation and upregulation of epithelial-mesenchymal transition. GSK1070916, an Aurora kinase B/C inhibitor, was isolated from a synthetic lethality screen with H3K27M. GSK1070916 showed strong efficacy in native H3K27M lines (IC $_{50}$ s=60nM-1250nM), superior to the Aurora kinase A inhibitor alisertib, to which all cell lines showed substantial resistance. Combination of both drugs was not synergistic. GSK1070916 treatment caused increased H3K27me3 and decreased H3S10ph and H3S28ph. GSK1070916 induced apoptosis and S-phase stall. The H3K27M mutation induces epigenetic, phenotypic, and cell cycle regulation changes resulting in relaxation of transcriptional controls and more aggressive growth. Aurora kinase B/C inhibition is a novel therapeutic modality for DIPG that appears capable of reversing some H3K27M-related epigenetic changes, inducing apoptosis, and repressing uncontrolled cellular division.

DIPG-80. CLINICAL AND RADIOGRAPHIC RESPONSE TO ONC201 IN A PEDIATRIC PATIENT WITH A THALAMIC H3K27M AND BRAFV600E MUTANT DIFFUSE MIDLINE HIGH GRADE GLIOMA Elizabeth Duke¹, Jonathan Murnick¹, Rohinton Tarapore², Joshua Allen², and <u>Lindsay Kilburn¹</u>, ¹Children's National Hospital, Washington, DC, USA, ²Oncoceutics, Inc, Philadelphia, PA, USA

Recent improved understanding of the molecular markers of high grade glioma has shifted the approach to these aggressive CNS tumors to increasingly use molecularly guided targeted therapies. Treatment of patients with BRAFV600E mutant high grade gliomas with BRAF inhibitors has shown efficacy, however the impact of concomitant H3K27M mutation is unknown. ONC201 targets dopamine receptor D2 (DRD2), which is shown to be broadly overexpressed in the thalamus as well as multiple tumor types; its antagonism has demonstrated anti-tumor efficacy and immunomodulatory properties in preclinical studies. ONC201 has also demonstrated clinical efficacy in patients with H3K27M mutant gliomas. We present the case of a 9-year-old male with a right thalamic H3.3K27M mutant diffuse midline glioma with a concomitant BRAFV600E mutation with an ongoing partial response to ONC201 treatment. The patient was diagnosed in May 2018. He underwent biopsy, followed by standard focal proton radiation therapy (54Gy) and subsequent treatment with dasatinib, bevacizumab and everolimus over the course of five months. After continued radiographic progression on serial imaging, in April 2019 he started ONC201 375mg orally once per week through an expanded access trial. He has tolerated the medication well with grade 1 nausea and fatigue. Over the next nine months, he demonstrated clinical and radiographic improvement with modest increased use of his left side and MRIs showing progressive decrease in size of the thalamic lesion with a 70 % decrease in the target lesion (measuring 53x62mm prior to treatment, decreased to 38x26mm in January 2020).

DIPG-82. CLINICAL EXPERIENCE OF CONVECTION ENHANCED DELIVERY (CED) OF CARBOPLATIN AND SODIUM VALPROATE INTO THE PONS FOR THE TREATMENT OF DIFFUSE INTRINSIC PONTINE GLIOMA (DIPG) IN CHILDREN AND YOUNG ADULTS AFTER RADIOTHERAPY

Elwira Szychot^{1,2}, David Walker^{3,2}, Peter Collins², Harpreet Hyare^{4,2}, Ananth Shankar^{4,2}, Alison Bienemann⁵, Milo Hollingworth⁵, and Steven Gill⁵; ¹The Royal Marsden Hospital, Sutton, United Kingdom, ²Harley Street Children's Hospital, London, United Kingdom, ³University of Nottingham, Nottingham, United Kingdom, ⁴University College London Hospitals NHS Foundation Trust, London, United Kingdom, ⁵University of Bristol, Bristol, United Kingdom

PURPOSE: Effective treatment of diffuse intrinsic pontine glioma (DIPG) remains a formidable challenge due to inadequate penetration of the bloodbrain barrier (BBB) by systemically administered chemotherapies. The BBB can be overcome by directly infusing drugs into pons using method of convection-enhanced delivery (CED). We describe our clinical experience and what we have learned about the safety and feasibility of treating DIPG with intermittent CED of carboplatin and sodium valproate to the pons through the Renishaw Drug Delivery System (RDDS). METHODS: Retrospective review (2017-2020) of children with DIPG, who following radiotherapy, received compassionate treatment commencing 3,3-10 months post diagnosis (median 4.9 months). They received up to 7 cycles of 3-6 weekly (14.4-28.8mg/ml). RESULTS: 13 children 3–19 years (mean 6.9 years) were treated. There were no surgical complications. With the exception of infusion channels blocking in one device there were no adverse device effects. Two patients developed persistent 6th nerve palsies, which led to drug concentration reduction in the combination therapy. Subsequently infusion/ drug related toxicities were transient. Tumour was controlled in pons in 11/13 patients. Median progression free survival (PFS) was 13.0 months, while median overall survival (OS) was 15.3 months. CONCLUSIONS: Use of the RDDS was safe and well tolerated in all 13 patients. Treatment improved control of pontine disease resulting in longer PFS and OS than reported for conventional therapy and merits further evaluation in a clinical trial.

DIPG-83. USING COPPER CHELATING AGENTS TO TARGET RECEPTOR TYROSINE KINASE SIGNALLING IN DIFFUSE INTRINSIC PONTINE GLIOMA (DIPG)

<u>Filip Michniewicz</u>^{1,2}, Federica Saletta¹, Jourdin Rouen^{1,2}, David Ziegler^{1,3}, and Orazio Vittorio^{1,2}; ¹Children's Cancer Institute, Lowy Cancer Research Centre, Sydney, NSW, Australia, ²UNSW School of Women's and Children's Health, Sydney, NSW, Australia, ³Sydney Children's Hospital, Sydney, NSW, Australia

DIPG is a universally fatal pediatric brain cancer. Receptor tyrosine kinase (RTK) pathway alterations are among the defining characteristics in many patients. Copper is a transition metal essential for cellular signaling, known to impact P13K/AKT and MAPK/ERK pathways. Copper chelating agents are clinically approved for use in children with Wilson's Disease, documented to reduce brain copper levels and are cited as potential cancer therapeutics. Due to copper's wide cellular integration, we propose that targeting copper in DIPG through use of copper chelators is a viable therapeutic strategy and are strong candidates for combination therapy. Cytotoxicity assays performed in a panel of DIPG cell lines using copper chelator tetraethylenepentamine (TEPA) demonstrated a millimolar range of efficacy. To identify copper integrated pathways, western blots were performed on DIPG cell lines dosed with sub-lethal copper concentrations, which increased phosphorylated expression of AKT, ERK1/2, ERK5 and STAT3. Conversely, western blots performed after TEPA treatment demonstrated reduced phosphorylated expression of all these proteins compared to controls. Western blots investigating TEPA in combination with Everolimus and Trametinib demonstrated synergistic targeting of these proteins. Our results indicate that adding copper in the culture media initiated two RTK-mediated downstream signal transductions, including AKT and ERK and additionally STAT signaling. The use of copper chelator TEPA affected copper homeostasis and reduced DIPG cell proliferation. Our study proposes copper plays an important role in RTK-mediated signaling promoting DIPG proliferation. This implies that reducing copper with clinically available chelation agents can represent a potential anti-cancer treatment for DIPG.

DIPG-84. COMPLEMENTARY AND ALTERNATIVE MEDICINE IN DIFFUSE INTRINSIC PONTINE GLIOMA

Fatma El-Khouly^{1,2}, Syed Adil¹, Harry Hendrikse¹, Gertjan Kaspers^{1,2}, Christof Kramm³, Sophie Veldhuijzen van Zanten^{1,2}, and Dannis van Vuurden²; ¹Amsterdam UMC - location VUmc, Amsterdam, Netherlands, ²Princess Maxima Center for Pediatric Oncology, Utrecht, Netherlands, ³University medical center Goettingen, Goettingen, Germany

INTRODUCTION: Diffuse intrinsic pontine glioma (DIPG) is a rare and aggressive childhood brainstem malignancy with a two-year survival rate of ≤10%. In this international survey study we aim to evaluate the use of complementary and alternative medicine (CAM) in this patient population. METHODS: Parents of-, and physicians treating DIPG patients were asked to participate in a retrospective online survey with questions regarding CAM use during time of illness. RESULTS: 120 parents and 75 physicians contributed to the online survey between January and May 2020. Physicians estimated that <50% of their patients used CAM, whereas 69% of the parents reported to have used CAM to treat their child during time of illness. Cannabis was the most widely used form of CAM, followed by vitamins and minerals, melatonin, curcumin and boswellic acid, CAM was mainly used to actively treat the tumor. Other motivations were to treat side effects of chemotherapy, or to comfort the child. Children diagnosed \geq 2016 were more likely to use CAM (χ 2=6.08, p=0.014). No significant difference was found between CAM users and non-users based on ethnicity ($\chi 2=4.18$, p=0.382) and country of residence ($\chi 2=9.37$, p=0.154). Almost 50% of the physicians do not frequently ask their patients about possible CAM use. CONCLUSION: This survey demonstrates that worldwide a considerable number of DIPG patients use CAM. Physicians should be more aware