Preclinical analysis of MTOR complex 1/2 inhibition in diffuse intrinsic pontine glioma

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Abstract. Diffuse intrinsic pontine glioma (DIPG) is an incurable childhood brain tumor. The mechanistic target of rapamycin (MTOR), a key oncogene, functions as two distinct signaling complexes, MTORC1 and MTORC2. We set out to determine the preclinical efficacy and mechanism of action of MTOR inhibitors in DIPG. We evaluated the MTORC1 inhibitor everolimus and the MTORC1/2 inhibitor AZD2014 in three patient-derived DIPG cell lines using cell culture models. We created dose-response curves for both compounds. We measured phenotypic effects on cell self-renewal, apoptosis, cell cycle, differentiation, senescence, and autophagy. We assessed the effects of each compound on the AKT pathway. Finally, we measured the efficacy of AZD2014 in combination with radiation therapy (RT) and a panel of FDA-approved chemotherapy drugs. While everolimus showed minimal antitumor efficacy, AZD2014 revealed IC50 levels of 410-552 nM and IC90 levels of 1.30-8.86 µM in the three cell lines. AZD2014 demonstrated increased inhibition of cell self-renewal compared to everolimus. AZD2014 decreased expression of phospho-AKT, while no such effect was noted with everolimus. Direct AKT inhibition showed similar efficacy to AZD2014, and induction of constitutive AKT activity rescued DIPG cells from the effects of AZD2014. AZD2014 exhibited synergistic relationships with both RT and various chemotherapy agents across classes, including the multikinase inhibitor ponatinib. MTORC1/2 inhibition shows antitumor activity in cell culture models of DIPG due to the effect of MTORC2 inhibition on AKT. This strategy should be further assessed for potential incorporation into combinatorial approaches to the treatment of DIPG.

Introduction

Diffuse intrinsic pontine glioma (DIPG) is an incurable brain tumor that mostly affects young children (1). Treatment with radiation therapy (RT) is usually transiently effective, but median progression-free survival is only 7 months, and median overall survival is 11 months (2). Hundreds of clinical trials studying systemically delivered cytotoxic and targeted chemotherapy agents have been conducted without showing any clinical benefit, potentially due to inadequate tumor penetration (3). Investigators are therefore studying novel drug delivery methods for these patients (4,5), and some of these have reached clinical trials (6,7). As these studies advance, better targeted therapeutic options will be needed, especially as part of combination treatments.

The mechanistic target of rapamycin (MTOR) is a serine-threonine kinase that has crucial roles in many cellular pathways dysregulated in cancer, including metabolism, growth, survival, and response to stress. It functions as part of two distinct signaling complexes, MTOR complex 1 (MTORC1), which also contains the regulatory associated protein of MTORC1 (RPTOR), and MTOR complex 2 (MTORC2), containing the RPTOR independent companion of MTORC2 (RICTOR). MTORC2 activates the V-Akt murine thymoma viral oncogene (AKT) through phosphorylation at S473, which mediates many of the cellular effects of MTORC2 (8-10). Preclinical and clinical data suggest a role for MTOR in gliomagenesis (11). First generation MTOR inhibitors (rapamycin analogues or rapalogs), such as everolimus, inhibit only MTORC1, and their clinical use in high-grade glioma has been disappointing (12). Contrastingly, Kahn et al
reported preclinical findings showing that the MTORC1/2 inhibitor AZD2014 enhanced the efficacy of RT in adult glioblastoma stem cells in vitro and in vivo (13), generating hope that combined MTOR inhibition may hold more clinical promise. DIPG has significant biological differences from adult glioblastoma, however (14,15), and prior to undertaking our project, MTOR inhibition of any kind in DIPG had not been addressed in the literature.

In this study, we examined the effects of MTOR inhibition preclinically in DIPG. We hypothesized that the addition of MTORC2 inhibition would increase the antitumor efficacy over the targeting of MTORC1 alone. Our overall goal was to determine how MTOR inhibition could contribute to future clinical treatment approaches to this devastating tumor.

Materials and methods

Aim and design. The aim of this study was to determine the efficacy of MTORC1/2 compared to MTORC1 inhibitors in DIPG. We set out to determine the phenotypic and molecular basis for the difference in efficacy noted between inhibitors. We then studied how MTORC1/2 inhibition could be used in combination with other existing treatments. All assays took place using three patient-derived short-term culture cell lines in cell culture models.

Gene expression in tumor bank. We performed gene expression profiling on patient-derived DIPG (n=16) and normal pons (n=2) samples. All tumor samples were collected from consenting pediatric patients undergoing tumor biopsy using an IRB-approved study protocol (COMIRB 95-500). Samples were snap frozen and stored in liquid nitrogen. RNA was extracted from each sample using an RNaseasy or DNA/RNA AllPrep kit (Qiagen, Valencia, CA, USA) according to the manufacturer's instructions. Four hundred nanograms of RNA was processed using the Ambion MessageAmp™ Premier RNA Amplification kit (Applied Biosystems, Foster City, CA, USA), according to the manufacturer's instructions. RNA quality was verified using the Nano Assay Protocol for the 2100 Bioanalyzer (Agilent, Santa Clara, CA, USA) at two time-points: i) after initial extraction of the RNA from the tumor sample, and ii) after preparation of the RNA for chip hybridization. The prepared RNA was hybridized to HG-U133 Plus 2 GeneChips (Affymetrix, Santa Clara, CA, USA), according to the manufacturer's instructions to measure gene expression. Mean and standard error of fold changes for each sample set were calculated for each measure indicated. We also conducted hallmark gene set enrichment analysis (GSEA) on the sample sets.

Gene expression from public data. We performed gene expression analysis using the R2 genomics analysis and visualization platform (http://r2.amc.nl) using the default settings. We compared expression of MTOR, RPTOR, and RICTOR in a set of normal brain samples (n=172, Berchtold set) versus DIPG samples (n=37, Paugh set) using one way analysis of variance.

Tumor lines and culture conditions. Three primary human pediatric (DIPG 4 and DIPG 6, derived from previously irradiated DIPGs at autopsy, provided by Dr Michelle Monje, Stanford University, and SF7761, derived from a biopsy sample, provided by Dr Nalin Gupta, University of California, San Francisco) were grown in neurosphere (suspension) culture conditions in ultra-low attachment flasks (Corning, Corning, NY, USA, DIPG 4 and DIPG 6) or in tissue culture-treated flasks (SF7761; Falcon/Corning) (16). Characteristics of the cell lines are listed in Table 1 (17). For certain experiments, DIPG 4 cells were also grown adherently in tissue culture-treated plates. The identity of all lines was validated by molecular profiling and compared to known results prior to and during this project. The cell lines were maintained in Neurobasal-A medium mixed 1:1 with Dulbecco's modified Eagle's medium/F-12 supplemented 1:100 by volume with HEPES [4-(2-hydroxyethyl)-1-piperazine ethanesulfonic acid] 1 M, sodium pyruvate 100 mM, MEM non-essential amino acids 10 mM, GlutaMAX-I, and antibiotic-antimycotic (all Gibco/Life Technologies, Waltham, MA, USA); B27-A supplement 50X (1:50; Invitrogen), heparin (2 µg/ml; Stemcell Technologies, Inc., Vancouver, BC, Canada) and human EGF, FGFb, and platelet derived growth factor-AB (all 20 ng/ml; Shenandoah Biotech, Warwick, PA, USA). For SF7761 cells, N2 supplement was also added (1:100; Life Technologies). For all experiments, except as noted, cells were plated at a concentration of 200,000 cells/ml of media in ultra-low attachment plates (Corning). Prior to all end-point measurements, neurospheres were dispersed by trituration using a micropipette.

Dose-response curves. Cell proliferation was determined by the MTS [3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium] assay using CellTiter 96 AQueous One Solution (Promega, Madison, WI, USA). Cells were seeded at 5,000 cells per well into a 96-well plate (Costar) and allowed to attach overnight (DIPG 4) or in a 96-well plate (Corning) in neurosphere culture (DIPG 6 and SF7761), in a total volume of 100 µl of media. Twenty-four hours later, the cells were treated with a range of doses of everolimus and AZD2014 (Selleck, Houston, TX, USA) in triplicate. At the end of the drug treatment period (72 h for DIPG 4 and DIPG 6, 120 h for SF7761), 20 µl of MTS reagent was added to each well to make a final volume of 120 µl and allowed to develop. Absorbance values for plate wells were acquired using a BioTek Synergy 2 plate reader at a wavelength of 490 nm after 3 h of incubation, and background absorbance was subtracted. IC50 values were determined experimentally through Prism, and IC90 values were calculated using the IC50 and hill slope value using the Graphpad EC Anytime online calculator.

### Table I. Cell line characteristics.

| Cell line | H3 mutation | Other genetic features | When collected |
|-----------|-------------|------------------------|---------------|
| DIPG 4    | H3. 1K27M   | TP53 WT, ACVR1 G328V   | Autopsy       |
| DIPG 6    | H3. 3K27M   | TP53 mutant            | Autopsy       |
| SF7761    | H3. 3K27M   | Exogenous hTERT        | Biopsy        |
Neurosphere dilution assay. DIPG 4 and SF7761 cells were plated in 96-well format in suspension, with 10 cells per well and 20 wells per condition. Conditions for each cell line included DMSO control, the calculated IC_{50} for AZD2014, or the same concentration of everolimus. The presence and size (area using two perpendicular axes) of a neurosphere in each well was measured after three weeks.

Apoptosis measurement. SF7761 cells in neurosphere culture were treated in a 96-well plate format with the indicated range of AZD2014 and everolimus concentrations compared to the control, in triplicate. Apoptosis was measured according to the manufacturer's instructions using the Caspase Glo 3/7 assay (Promega) at 48 h of treatment after 2 h of incubation with the Caspase-Glo reagent at room temperature.

Cell cycle analysis. SF7761 cells in neurosphere culture were treated in a 6-well format with the indicated range of AZD2014 and everolimus concentrations compared to the control for 24 h. Cells were then removed from compound solutions and everolimus concentrations compared to the control for 1 h at room temperature or overnight at 4°C. SF7761 cells to be stained were fixed and permeabilized in Foxp3/Transcription Factor Fixation/Permeabilization Concentrate and Diluent (Affymetrix EBioscience, Santa Clara, CA, USA) according to the manufacturer's instructions. Cells were incubated in primary antibody to p21, along with permeabilization Buffer (Affymetrix EBiocene). After rinsing to remove the primary antibody, adherent and suspension cells were incubated in secondary antibody (1:500; Alexa Fluor 488; Life Technologies) for 1 h at room temperature. Suspension cells were affixed to slides using a Cytospin 4 Centrifuge (Thermo Fisher Scientific) at 2,000 rpm for 10 min. Slides were fixed using ProLong antifade reagent with DAPI (Life Technologies). Confocal imaging was performed at a magnification of x400 using 405 (DAPI) and 488 nm (Alexa Fluor 488) lasers on a 3i Marianas imaging system (Intelligent Imaging Innovations). Images were obtained using an Evolve 16-bit EMCCD camera (Photometrics, Tucson, AZ, USA).

Senescence measurement by β-galactosidase staining. Adherently growing DIPG 4 cells were plated at a density of 100,000 cells per well in a 6-well plate and allowed 24 h in which to develop adhesion before subjecting them to experimental conditions. Cells were treated for two days with 50 nM, 500 nM or 5 µM of AZD2014 or everolimus, along with DMSO control. After two days of incubation, the cells were placed into normal growth media for five days. Staining for β-galactosidase was performed using a Senescence β-galactosidase Staining kit (Cell Signaling Technology, Inc.) according to the manufacturer's instructions. Cells were washed with 2 ml of PBS, followed by a 15-min incubation using 1X fixative. Cells were again washed twice with PBS and then incubated with 1 ml of β-galactosidase staining solution. The cells were incubated overnight in a dry 37°C incubator without CO₂. Senescent cells were imaged via brightfield microscopy (Nikon DS-L2).

Autophagy measurement. DIPG 4 and SF7761 cells were plated in neurosphere culture in 6-well plates and exposed to one of the following conditions for 4 h: 0.2% DMSO, 10 µM chloroquine (MP Biomedicals), 1 µM AZD2014, 1 µM everolimus, 1 µM AZD2014 combined with 10 µM chloroquine, or 1 µM everolimus combined with 10 µM chloroquine. Immediately following treatment, the cells were lysed, and western blotting was conducted as described below.

AKT/phospho-AKT measurement. DIPG 4 and SF7761 cells in neurosphere culture were treated in a 6-well format with the indicated range of AZD2014 and everolimus versus the control for 2 h. Western blotting was then conducted as below using antibodies specific to phosphorylated and unphosphorylated AKT.

Western blotting. Cells were lysed in 200 µl radioimmunoprecipitation assay buffer with 100X protease and phosphatase inhibitor (Pierce; Thermo Fisher Scientific) by gentle vortexing for 5 sec. Cells were incubated on ice for 5 min before
centrifugation at 4°C at 14,000 x g for 10 min. Protein concentration was measured using the Pierce BCA assay. Pre-cast 4-20% Mini-PROTEAN TGX gels (Bio-Rad, Hercules, CA, USA) were loaded with ~30 µg of protein per lane and then run for 90 min at 125 V. Transfer was then performed to a polyvinylidene fluoride membrane via a Bio-Rad transfer set at 4°C at 50 V for 90 min. Blocking was carried out in 5% non-fat dry milk, as was all staining. Primary staining was performed as indicated for α-tubulin (#2125 rabbit anti-human mAb at 1:1,000 overnight at 4°C; Cell Signaling Technology, Inc.); pAKT (#4060 rabbit anti-human mAb at 1:1,000 overnight at 4°C; Cell Signaling Technology, Inc.); and total AKT (#9272 rabbit anti-human pAb at 1:1,000 overnight at 4°C; Cell Signaling Technology, Inc.). Secondary staining was carried out with anti-rabbit HRP at 1:3,000 for 1 h at room temperature. Blots were developed using Western Lightning Plus-ECL chemiluminescent (PerkinElmer, Waltham, MA, USA).

**AKT small molecule inhibition.** DIPG 4 and SF7761 cells in neurosphere culture were treated in a 96-well format in triplicate at the indicated concentration range of MK-2206 (Selleck Chemicals, Houston, TX, USA) compared to the control for 120 h. Cell viability was then determined by MTS assay as above.

**Constitutive AKT activation.** Introduction of a mutation from serine to aspartate at amino acid 473 of AKT (S473D) mimics serine phosphorylation at this position and results in constitutive activation of the protein (18). DIPG 4 cells were plated at a density of 200,000 cells per well in 2 ml of medium in a 6-well plate and allowed to adhere overnight. AKT cDNA containing the S473D mutation (gift of Dr. N. Rosen) was transfected into the cells using jetPRIME® transfection reagent (2 µg cDNA per well mixed into 200 µl jetPRIME® buffer, followed by 4 µl of jetPRIME® reagent; 5-min incubation; Thermo Fisher). The medium was changed at 18 h following transfection. At 48 h following transfection, western blotting was conducted as above. Transfected and wild-type cells were then plated in a 96-well format, allowed to adhere, and then treated for 120 h as follows with the indicated range of AZD2014 concentrations compared to the control, in sextuplicate. Cell viability was then measured by MTS assay as above.

**Radiation combination.** SF7761 cells in neurosphere culture were plated in a 96-well format in triplicate. The next day, they were exposed to irradiation at the indicated dose range using a cesium irradiator. They were then treated starting the next day with the indicated dose range of AZD2014 compared to control for 120 h. Cell viability was then measured by MTS assay as above. The combination index at each dose level combination was determined by the Chou Talalay method (19).

**Immunofluorescence staining.** Immunofluorescence staining for γH2AX was performed on cells exposed to AZD2014 followed by radiation to identify double-stranded DNA (dsDNA) damage patterns. SF7761 cells were plated at a density of 20,000 cells per well in BioCoat Poly-D-lysine/laminin-coated chamber slides (#354688; Corning) and allowed ~24 h in which to develop adhesion before subjecting them to experimental conditions. Cells were treated with vehicle or AZD2014 at 50 and 500 nM dose levels for 4 or 48 h and then irradiated with 4 Gy from a Cs137 source in a single dose. At 4 h post-irradiation, the cells were fixed for 20 min in 37% formaldehyde diluted in 10X PBS (Sigma), permeabilized in 0.1% Triton X in PBS for 10 min, and blocked for 45 min in 4% BSA in PBS supplemented with 0.05% Triton X. Cells were incubated in primary γH2AX antibody (#2577; Cell Signaling Technology, Inc.) diluted 1:600 with 4% BSA, dissolved in PBS and 0.05% Triton X, for 1 h at room temperature. After rinsing, cells were incubated in secondary antibody (1:500; Alexa Fluor 488; Life Technologies) for 1 h at room temperature. Slides were fixed using ProLong antifade reagent with DAPI (Life Technologies). Confocal imaging was performed at a magnification of x400 using 405 (DAPI) and 488 nm (Alexa Fluor 488) lasers on a 3i Marianas imaging system (Intelligent Imaging Innovations). Images were obtained using an Evolve 16-bit EMCCD camera (Photometrics).

**Chemotherapy combination panel.** SF7761 cells in neurosphere culture were plated in a 96-well format. One set of cells was treated with the IC50 concentration of AZD2014, while another set of cells was treated with DMSO. All drugs from the Approved Oncology Drugs Set VII (National Cancer Institute, Bethesda, MD, USA) were then added to one well of each set of cells at a concentration of 1 µM. In addition, 6 wells of cells were treated with the IC50 concentration of AZD2014 alone, and with DMSO alone. Cells were exposed to these compounds for 120 h, and then cell viability was determined by MTS assay as above. The relative cell viability for each combination was then compared to DMSO alone. The therapeutic relationship for each combination was then calculated by dividing this value by the product of the relative cell viability for each drug and the relative cell viability for AZD2014 alone.

**Ponatinib combination.** DIPG 4, DIPG 6, and SF7761 cells were treated in a 96-well format in triplicate with the indicated ranges of AZD2014, ponatinib, and the two compounds together, compared to the control, for 120 h. Cell viability was then determined by MTS assay as above. The combination index at each dose level combination was determined by the Chou Talalay method (19).

**Data analysis.** All data analysis and chart creation was performed using GraphPad Prism. All charts show mean ± standard error of the mean (SEM) unless otherwise noted. Comparison of the percentage of neurospheres in the neurosphere dilution assay was carried out via Chi-square comparison of proportions test. Comparison of mean neurosphere area was performed by the unpaired t-test. Statistical significance is represented as follows: p>0.05 is indicated by ns (not significant); p=0.05-0.01 is indicated by *; p=0.01-0.001 is indicated by **; p=0.001-0.0001 is indicated by ***; p<0.0001 is indicated by ****.

**Results**

**Variable MTOR expression in human DIPG samples versus normal brain.** We first performed measurements of gene expression of MTOR and the MTOR components RPTOR (MTORC1) and RICTOR (MTORC2) in a panel of human
DIPG samples versus normal pons samples in our tissue bank. Multiple measures showed no significant difference in expression between the two sample sets (Fig. 1A). On hallmark GSEA, however, the MTORC1 signaling pathway was enriched in the DIPG versus normal samples (NES 1.3, \(p=0.021\), Fig. 1b). We then compared expression of the same three genes using publically-available sets of normal brain (\(n=172\)) and DIPG (\(n=37\)) samples. MTOR (\(p=4.4\times10^{-6}\)) and RPTOR (\(p=4.2\times10^{-29}\)) were both underexpressed in DIPG compared to normal brain, while RICTOR (\(p=3.3\times10^{-9}\)) was overexpressed (Fig. 1C).

MTORC1/2 inhibition decreases cell survival in vitro versus MTORC1 inhibition alone. We measured cell survival after five days of continuous drug exposure by MTS assay for three primary human DIPG cell lines, DIPG 4, DIPG 6, and SF7761 in neurosphere culture. Antitumor efficacy, especially in terms of the IC\(_{90}\), was far greater for AZD2014 compared to everolimus (Fig. 2). For DIPG 4, the IC\(_{50}\) and IC\(_{90}\) were not reached for everolimus and were 0.425 and 1.90 \(\mu\)M for AZD2014,
respectively (Table II). For DIPG 6, the IC$_{50}$ and IC$_{90}$ were also not reached for everolimus and were 0.552 and 1.30 µM for AZD2014. For SF7761, the IC$_{50}$ and IC$_{90}$ were 6.22 and 1,070 µM for everolimus and 0.410 and 8.86 µM for AZD2014.

Cell self-renewal difference but no clear difference in other phenotypic measures. We next examined the general mechanism of action for differences in the effects between the two drugs. When DIPG 4 and SF7761 cells were plated in 96-well format at 10 cells per well and 20 wells per condition to test cell self-renewal capability, those exposed continuously to the IC$_{75}$ dose of AZD2014 were less likely to form spheres and formed smaller spheres than those exposed to the same concentration of everolimus, or control (Fig. 3A). After 48 h of treatment, everolimus and AZD2014 both caused a decrease in apoptosis compared to the control, as measured by caspase 3/7 luminescence assay (Fig. 3B); there was no significant difference between the two drugs. We noted no significant difference in cell cycle distribution with increasing doses of either drug as measured by propidium iodide flow cytometry (Fig. 3C). We also saw no clear trend to demonstrate an increase or decrease in glial or neuronal differentiation as measured by GFAP or TUBB3 immunofluorescence, respectively (Fig. 3D).

When we measured induction of senescence in DIPG 4 and SF7761 cells by immunofluorescence for the senescence marker p21, neither compound showed a consistent increase in senescence compared to the control (Fig. 4A). We also measured by staining for the senescence marker β-galactosidase in DIPG 4 cells; here, both compounds appeared to increase the percentage of senescent cells in a dose-dependent manner, but there was no clear difference between the two (Fig. 4B). AZD2014 and everolimus both induced greater levels of autophagic flux compared to the control, as determined by western blotting for LC3-II, an autophagy biomarker (Fig. 4C). Neither compound induced consistently more autophagy than the other, however.

Inhibition of AKT phosphorylation is necessary and sufficient for the effect of AZD2014 on DIPG. We next set out to determine the specific mechanism of action differentiating AZD2014 from everolimus, focusing on AKT, which is activated when phosphorylated by MTORC2 but is upstream of MTORC1. We found that increasing doses of AZD2014 caused a dose-dependent decrease in pAKT relative to total AKT, as determined by western blotting (Fig. 5A). We saw no such decrease with everolimus. We then used a direct AKT inhibitor, MK-2206, and demonstrated that exposing DIPG 4 and SF7761 cells in neurosphere culture to this drug for five days resulted in IC$_{50}$ levels of 2.60 µM in DIPG 4 and 1.08 µM in SF7761 cells (Fig. 5B). These values were higher but within the range of those obtained with AZD2014. We then transfected DIPG 4 cells with a CDNA for a mutant form of AKT (S473D) intending to confer constitutive activation. Compared to the wild-type, the transfected cells exhibited
increased apparent pAKT (due to the S473D mutation, which binds the phospho-AKT antibody), accompanied by a decrease in unphosphorylated AKT, suggesting that total AKT (measured by the sum of the pAKT and AKT bands) remained
and resistant to RT, we treated radiation-naïve SF7761 cells with increasing doses of AZD2014 and exposed them to various doses of cesium RT in one fraction. We demonstrated a dose-dependent decrease in cell survival with both treatments as measured by MTS assay (Fig. 6A). Maximal cell killing with RT alone was only in the range of 40%, even though SF7761 is our most radiosensitive cell line, because the single fraction delivered is well below the total dose delivered clinically over many fractions. We then determined the therapeutic relationship at each dose combination by the Chou Talalay method (19) to calculate combination indices. The relationship was variable but was generally synergistic (combination index <1) at higher doses of each treatment and antagonistic (combination index >1) at lower doses (Fig. 6B). We found that, while AZD2014 and RT were synergistic in killing cells at high doses, the level of dsDNA damage caused by this combination did not exceed the damage caused by RT alone as assessed by γH2AX staining (Fig. 6C).

AZD2014 has variable therapeutic relationships with FDA-approved chemotherapy agents, including synergy with ponatinib. Finally, we conducted a drug screen of all FDA-approved chemotherapy agents in combination with AZD2014, using SF7761 in a 96-well format (data not shown; goo.gl/3XCCqk). Aside from antagonistic relationships with microtubule inhibitors, there was no clear trend to AZD2014’s therapeutic relationship with any category of drug. AZD2014 did show synergistic relationships with drugs from multiple classes, however, including microtubule stabilizers, topoisomerase inhibitors, and tyrosine kinase inhibitors. We then validated our screening finding of a potentially synergistic relationship between AZD2014 and the multikinase inhibitor ponatinib using multiple dose levels of the AZD2014-ponatinib combination in DIPG 4, DIPG 6, and SF7761 (Fig. 7A). We demonstrated synergy between AZD2014 and ponatinib at most dose levels in all three cell lines, especially dose levels exceeding the combined IC50 of the two agents (Fig. 7B).

**Discussion**

In this study, we demonstrated that MTORC1/2 inhibition shows greatly increased antitumor efficacy in a panel of patient-derived DIPG cell lines compared to MTORC1 inhibition alone, which showed little to no effect. The phenotype behind this advantage appears to be a decrease in cell self-renewal; assays testing other potential general mechanisms of action showed no significant or consistent differences between the two drugs. On a molecular level, the decrease in AKT phosphorylation caused by MTORC2 inhibition appears to be sufficient and at least partially necessary to the augmented tumor inhibition when MTORC2 is targeted. This phenotype and molecular mechanism are consistent with the known role of AKT in cancer cell self-renewal (20-22). Finally, we showed that AZD2014 has the potential to act synergistically with RT, as well as with cytotoxic and targeted chemotherapeutic agents of various classes.

It is unclear whether the efficacy of AZD2014 in this study depends on overexpression of MTORC2 components. Expression levels of MTOR, RPTOR, and RICTOR were similar in the DIPG samples in our tumor bank compared...
to levels in normal pons, although the MTOR pathway was enriched in DIPG on GSEA. MTOR and RPTOR were under-expressed in a large public dataset of DIPG samples, while RICTOR was overexpressed, which could contribute to an explanation for the difference seen between compounds, since RICTOR is unique to MTORC2. It should be noted in this analysis, however, that the available comparison group was tissue samples from throughout the brain, not the pons alone.

In terms of previous studies, a large study of 43 DIPG samples that examined copy number abnormalities and expression profiles did not identify MTOR in any of the abnormalities (23). This study did find AKT as a potential gene of interest in focal recurrent gains, however. Another group working with a genetically engineered mouse model of DIPG found that AKT was overexpressed in cell lines derived from this model, and when they conducted a high-throughput drug screen against these cells, they identified antitumor activity in a multikinase inhibitor that decreases levels of pAKT (24). These findings support our results showing that AKT inhibition through MTORC1 and MTORC2 is necessary and sufficient to the antitumor effect of AZD2014, and that this is the reason for everolimus’ lack of activity.

RT is currently the only effective primary therapy in DIPG, and clinical trial data are also emerging to suggest its utility at recurrence as well (25,26). Therefore, the therapeutic relationship with RT will be relevant to any new targeted drug proposed. Our findings suggest that AZD2014 and RT have a synergistic therapeutic relationship at higher doses of each. These higher RT doses are used in DIPG, and AZD2014 levels at and above the IC_{50} will also be necessary to achieve therapeutic effect. We did not observe greater levels of dsDNA damage from the combination versus RT alone and continue to investigate the underlying mechanism of synergy. The prior study examining this combination in adult glioblastoma found that the reason for synergy is most likely inhibition of DNA repair (13). On our chemotherapy combination screen, AZD2014 also appeared to show synergy with several other DNA damaging agents. Another candidate drug from our chemotherapy screen, ponatinib, primarily targets BCR-ABL but also inhibits VEGFR, PDGFR, and EGFR (27), all of which have relevance to DIPG (28). Ponatinib may also inhibit AKT (29), raising another possible mechanism of synergy with AZD2014. Given that development of resistance to targeted therapies is a major issue with their efficacy, using combinations such as this to target multiple key oncogenic pathways may be crucial to successful DIPG treatment. Our consistent results with ponatinib in each of the lines validates our screening method. The availability of these preclinical models in DIPG is of great translational value. Besides their lack of efficacy, the drugs used in previous DIPG clinical trials have also caused harm through adverse effects, mandating that future treatments undergo rational preclinical testing before they are deemed worthy of the risk they carry for patients.

Our study has several limitations. We did not show that AZD2014 reverts to an everolimus-like dose-response curve with constitutive AKT activation, although the difference in IC_{50} was statistically significant. This may be because AZD2014 was still able to achieve some level of AKT inhibition due to imperfect transfection efficiency in this experiment. Also, AZD2014’s blood-brain barrier (BBB) penetration is unknown, but given the concerns with drug penetration in DIPG in general, it is likely that local delivery methods that bypass the BBB, such as convection-enhanced delivery, will at least partially obviate this consideration. Finally, we do not yet have in vivo validation of our findings. However, while our study was under review, Miyahara et al published their findings on the effect of another MTORC1/2 inhibitor, TAK228, in DIPG, including its efficacy in a patient-derived xenograft model (30). Our study delves further into the molecular mechanism of action and potential combination chemotherapy
approaches, and together, these two studies on different MTORC1/2 inhibitors provide strong preclinical rationale for this strategy in DIPG.

In conclusion, the present study demonstrates that DIPG does not respond preclinically to MTORC1 inhibition alone but does respond well to combined MTORC1/2 inhibition, due to the inhibitory effect of MTORC2 on AKT. AZD2014, an MTORC1/2 inhibitor, shows synergy with RT and with selected chemotherapy agents in DIPG. This strategy should be studied further as a potential component of combinatorial approaches to treatment of this currently incurable tumor.

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