- 1 Disruption of Ataxia telangiectasia mutated kinase enhances radiation therapy
- 2 <u>efficacy in spatially-directed diffuse midline glioma models.</u>
- 3 Author List based on Contribution:
- 4
- 5 Avani Mangoli\*1
- 6 Vennesa Valentine\*<sup>2</sup>
- 7 Spencer Maingi<sup>2</sup>
- 8 Sophie R. Wu<sup>2</sup>
- 9 Harrison Q. Liu<sup>2</sup>
- 10 Michael Aksu<sup>3</sup>
- 11 Vaibhav Jain<sup>3</sup>
- 12 Bronwen E. Foreman<sup>2</sup>
- 13 Joshua A. Regal<sup>2</sup>
- 14 Loren B. Weidenhammer<sup>2</sup>
- 15 Connor E. Stewart<sup>2</sup>
- 16 Maria E. Guerra Garcia<sup>2</sup>
- 17 Emily Hocke<sup>2</sup>
- 18 Karen Abramson<sup>3</sup>
- 19 Tal Michaeli<sup>1</sup>
- 20 Nerissa T. Williams<sup>2</sup>
- 21 Lixia Luo<sup>2</sup>
- 22 Megan Romero<sup>4</sup>
- 23 Katherine Deland<sup>2</sup>

- 24 Samantha Gadd<sup>5</sup>
- 25 Eita Uchida<sup>5</sup>
- 26 Laura Attardi<sup>6</sup>
- 27 Kouki Abe<sup>5</sup>
- 28 Rintaro Hashizume<sup>5</sup>
- 29 David M. Ashley<sup>1</sup>
- 30 Oren J. Becher<sup>4</sup>
- 31 David G. Kirsch<sup>7</sup>
- 32 Simon G. Gregory<sup>3\*\*</sup>
- 33 Zachary J. Reitman<sup>2\*\*\*</sup>
- 34
- 35 \*Co-First Author
- 36 \*\*Co-Corresponding Author
- 37
- 38 \*\*\*Corresponding author
- 39
- 40 **Corresponding Author:**
- 41 Zachary J. Reitman, MD, PhD
- 42 30 Duke Medicine Circle, Box 3085,
- 43 Durham, 27710, NC, USA
- 44 Email: zjr@duke.edu
- 45 Phone: (919) 668-7336
- 46

- 48 **Co-corresponding author:**
- 49 Simon G. Gregory, PhD
- 50 300 N. Duke Street, DUMC 104775
- 51 Durham, 27701, NC, USA
- 52 Email: <u>simon.gregory@duke.edu</u>
- 53 Phone: (919) 684-0726
- 54
- <sup>55</sup> <sup>1</sup>The Preston Robert Tisch Brain Tumor Center, Duke University, Durham, North
- 56 Carolina, United States 27710
- <sup>57</sup> <sup>2</sup>Department of Radiation Oncology, Duke University, Durham, North Carolina, United
- 58 States 27710
- <sup>59</sup> <sup>3</sup>The Preston Robert Tisch Brain Tumor Center Omics Program, Duke University,
- 60 Durham, North Carolina, United States 27710
- <sup>4</sup>Department of Pediatric Hematology Oncology, Mount Sinai Kravis Children's
- 62 Hospital, 1468 Madison Avenue, New York, NY, United States 10029
- <sup>5</sup>Department of Pediatrics, University of Alabama at Birmingham, Birmingham,
- 64 Alabama, United States 35233
- <sup>65</sup> <sup>6</sup>Departments of Radiation Oncology and Genetics, Stanford University School of
- 66 Medicine, 269 Campus Drive, Stanford, California, United States 94305-5152
- <sup>67</sup> <sup>7</sup>Princess Margaret Cancer Centre, University of Toronto, 610 University Avenue,
- 68 Toronto, Ontario, Canada M5G 2M9
- 69

70 Abstract:

72	Diffuse midline gliomas (DMGs) are lethal brain tumors characterized by p53-
73	inactivating mutations and oncohistone H3.3K27M mutations that rewire the cellular
74	response to genotoxic stress. We used RCAS/tv-a retroviruses and Cre recombinase to
75	inactivate p53 and induce native H3.3K27M mutations in a lineage- and spatially-
76	directed manner. We generated primary mouse tumors that recapitulate human DMG.
77	Disrupting ataxia-telangiectasia mutated kinase (ATM) enhanced the efficacy of
78	radiation therapy in murine and patient-derived DMG models which increased survival
79	Microscopy-based in situ sequencing was used to spatially resolve transcriptional
80	profiles in >750,000 single cells with or without ATM disruption and radiation therapy,
81	revealing altered immune-neoplastic and endothelial cell interactions after treatment. An
82	allelic series of primary murine DMG models with different p53 mutations confirmed that
83	transactivation-independent p53 activity is a key mediator of radiosensitivity after ATM
84	disruption. Our findings contribute primary DMG mouse models with deep profiling and
85	reveal the mechanisms of treatment response to an actionable therapeutic strategy.
86	

#### 88 Introduction

89 Diffuse midline gliomas (DMGs) are lethal brain tumors in children and young adults. 90 These tumors are localized in essential midline brain structures, such as the brainstem 91 and thalamus, making them surgically inoperable and unresponsive to conventional 92 chemotherapy. The median overall survival of patients with DMGs is less than two 93 years. Although radiation therapy may improve symptoms and extend life, it remains 94 palliative. Somatic activation of lysine 27 to methionine mutations in histone variant 3.3 95 (H3.3K27M) is a defining feature of DMG (1, 2). Approximately 70% of DMGs harbor 96 inactivating mutations in the tumor suppressor TP53 (1-3), which are associated with 97 radioresistance in patients and preclinical models (4, 5).

98

99 A key limitation of current primary DMG preclinical models is the ability to induce K27M 100 mutations in the native H3f3a locus in a spatial-, lineage-, and temporally controlled 101 manner. Patient-derived xenografts (6), patient-derived cell lines (7), in utero 102 electroporation (7), and syngeneic mouse models (8, 9) have provided key insights into this disease. A conditional H3f3a-loxP-Stop-loxP-K27M-Tag allele (H3f3a<sup>LSL-K27M-Tag</sup>) 103 104 has also been generated that allows the expression of H3.3K27M from the endogenous 105 mouse H3f3a locus in the presence of Cre recombinase (10). However, this model has 106 been limited by cell lineages that can be interrogated with existing Cre driver lines, such 107 as Nestin-Cre (10). To date, the conditional H3.3K27M alleles have not been 108 investigated in an entirely spatially controlled manner. We and others have used the 109 RCAS/tv-a retroviral system for spatially-directed modulation of glioma tumorigenesis in 110 mice (5, 11-15). The RCAS/tv-a platform was used to deliver an exogenous H3.3K27M

(13, 14, 16), but to our knowledge it has not been used to edit the endogenous *H3f3a*allele. A variety of model systems have been used to investigate the mechanisms
associated with the development of DMG and to assess therapeutic strategies.

114

115 Inhibition of ataxia-telangiectasia mutated kinase (ATM) has emerged as a strategy to 116 enhance the efficacy of radiation therapy in DMG (17). ATM is a master orchestrator of 117 the DNA damage response to double strand breaks (17). Patients with hereditary loss-118 of-function ATM variants, and tumors containing ATM variants, are extremely sensitive 119 to radiation therapy (17). Consequently, a brain-penetrant ATM inhibitor has entered 120 clinical trials for adult brain tumors (NCT03423628) (18). A recent study identified ATM 121 inhibition as a potent radiosensitization strategy in various patient-derived pediatric 122 high-grade glioma models (6). We found that functional ATM loss radiosensitized 123 primary mouse models of DMG were driven by p53 loss, but not p53 wildtype (5, 11). 124 ATM loss increases tumor sensitivity to radiotherapy via radiosensitization of neoplastic 125 cells rather than the vasculature (12). However, it remains uncertain whether H3.3K27M 126 affects the ability of Atm loss to radiosensitize primary DMG. This is of particular 127 importance since H3.3K27M regulates the p16 molecular checkpoint that regulates G1-128 to-S cell cycle progression (13) and could thereby influence the radiation response. 129

Here, we examined strategies to exploit the genomically-stressed cell state in
H3.3K27M/TP53-altered DMG. We improved upon previous models that delivered
H3.3K27M from an exogenous RCAS payload (11, 13) by combining the RCAS/tv-a
system with H3f3a<sup>LSL-K27M-Tag</sup> mice to express H3.3K27M from the endogenous *H3f3a*

134 locus. This autochthonous mouse model enabled us to analyze the impact of Atm loss 135 in the context of H3.3K27M/TP53-altered brain tumors to mimic human DMG (10). We 136 found that primary DMGs expressing H3.3K27M driven by p53 loss were 137 radiosensitized by Atm loss. To explore the resistance mechanisms in specific tumor 138 cells, we examined primary mouse DMGs after focal brain irradiation using high-139 resolution single cell in situ sequencing (ISS). The results identified the overexpression 140 of the cell cycle regulator Cdkn1a as a putative resistance factor in Atm-intact DMG. We 141 showed that Cdkn1a, or the transcriptional activity of p53 in general, was dispensable 142 for DMG radiosensitization by Atm loss. Therefore, the non-transactivation functions of 143 p53 may determine the sensitivity of DMGs to combinations of ATM inhibitors and 144 radiation therapy. The high-resolution results describe a genetically faithful and flexible 145 primary mouse model of DMG, identifying the mechanisms of resistance to a 146 therapeutic strategy currently in clinical trials.

148 **Results** 

# Conditional p53 loss and H3.3K27M expression in retrovirus-induced mouse DMGs.

151 To express H3.3K27M from the endogenous H3f3a locus in retrovirus-induced primary 152 mouse gliomas, we used a H3f3a<sup>LSL-K27M-Tag</sup> allele that expresses H3.3K27M in the presence of Cre recombinase (10). To incorporate the H3f3a<sup>LSL-K27M-Tag</sup> allele into the 153 154 RCAS/tv-a retrovirus system, mice were bred with Nestin<sup>TVA</sup> mice to allow RCAS 155 retroviruses to specifically transduce TVA+ Nestin-expressing neural stem cells. To 156 investigate the deletion of p53 specific to tumors, we crossbred a p53 variant in which 157 critical exons were flanked by loxP sites (floxed or FL) allowing for functional deletion of 158 p53 in the presence of Cre recombinase. We first introduced retroviruses into Nestin<sup>TVA</sup>; p53<sup>FL/FL</sup>; H3f3a<sup>LSL-K27M-Tag/+</sup> mice (nPH) and compared them to matched mice lacking the 159 H3f3a<sup>LSL-K27M-Tag</sup> allele (nP, Figure 1A). We induced DMGs by injecting mice with RCAS 160 161 retroviruses expressing Cre recombinase, firefly luciferase, and the oncogene platelet-162 derived growth factor ligand beta (PDGF-B) and monitored for tumor formation via in 163 vivo imaging. Using luciferase-based bioluminescent imaging to detect tumors, we determined that there was no difference in time to tumor formation in H3f3aLSL-K27M-Tag/+ 164 165 mice compared to matched mice lacking the H3f3a<sup>LSL-K27M-Tag</sup> allele (Figure 1B). To investigate the effects of *Atm* deletion in these tumors, we also generated Nestin<sup>TVA</sup>; 166 p53<sup>FL/FL</sup>; H3f3a<sup>LSL-K27M-Tag /+</sup>; Atm<sup>FL/FL</sup> (nPHA<sup>FL/FL</sup>) mice and littermate controls with intact 167 Atm in their tumors of genotype Nestin<sup>TVA</sup>; p53<sup>FL/FL</sup>; H3f3a<sup>LSL-K27M-Tag /+</sup>; Atm<sup>FL/+</sup> 168 (nPHA<sup>FL/+</sup>) (Figure 1C-E, see description of *Atm* loss results below). Tumors exhibiting 169 170 hypercellularity and diffuse infiltration of the nearby normal brain on H&E formed within

171 4-8 weeks with high penetrance (Figure 1F). We detected HA expression indicating the 172 presence of the HA tag on both H3.3K27M and PDGF-B constructs (Figure 1G). As expected, p53 was not detected in p53<sup>FL/FL</sup> tumors by IHC (Figure 1H). Histone 3 lysine 173 27 trimethylation was significantly decreased by IHC in H3f3a<sup>LSL-K27M-Tag /+</sup> tumors 174 175 compared to controls (mean 50.49 % (nP) vs. 5.757 % (nPH) of cells staining positive, 176 p-value <0.001, Figure 1I and Supplemental Figure 1), indicating that H3.3K27M could 177 functionally deplete H3K27me3 as predicted (19). Differentially methylated features 178 between nP versus K27M-bearing tumors showed hyper- and hypo- methylated 179 features within promoters (Supplemental Figure 2A) and enhancers (Supplemental 180 Figure 2B) Additional analysis shows a difference in percentage of methylation within 181 hypomethylated tiles and de-novo tiles in K27M tumors compared to nP and normal 182 murine tissue (Supplemental Figure 2C and D). Differentially methylated genes yielded 183 from the hypomethylated genomic regions were most enriched for processes involving 184 neuronal development and differentiation, suggesting developmental properties for 185 promoter and enhancer tiles (Supplemental Figure 2E). This finding is consistent with 186 other tissues DNA methylation state of these specific tiles and with the role of K27M in 187 regulating oncogenic and developmental processes (20) Sequence motif analysis 188 identified differential methylation of motifs associated with transcription factors Hoxd13 189 and Hoxa11 (Supplemental Figure 2F) which are known to be involved in hindbrain 190 development. Ki67 was elevated in >50% of tumor cells regardless of H3.3K27M status 191 (Figure 1J). Anti-Flag immunohistochemistry confirmed the presence of the Flag tag on 192 the H3.3K27M construct (Figure 1K). Flag IHC demonstrated that H3.3K27M-Tag+ cells 193 diffusely infiltrated from a hypercellular tumor core into the brain parenchyma

194 suggesting the diffuse, infiltrative biology seen in human DMG. These results

demonstrate that RCAS/tv-a and a conditional H3f3a<sup>LSL-K27M-Tag</sup> allele can be combined

196 to target K27M to H3f3a gene in time, lineage, and space to generate primary mouse

197 DMGs that recapitulate human disease.

198

#### 199 *Atm* loss radiosensitizes primary p53-null/H3.3K27M DMGs.

200 Targeting ATM kinase has emerged as a potential strategy to increase the efficacy of 201 standard-of-care radiation therapy for brain tumors (5, 6, 17). We sought to determine 202 whether disruption of ATM could radiosensitize primary mouse DMGs with p53 and 203 H3.3K27M alterations. Previously, we established that H3f3a-wildtype brainstem 204 gliomas lacking Atm in tumor cells were radiosensitized compared to littermate controls 205 with a functional Atm allele in their tumors(5). However, these mice lack H3.3K27M 206 which disrupts the G1-to-S cell cycle checkpoint (13) and may thereby affect the 207 downstream effects of ATM deficiency (17). We hypothesized that Atm inactivation in 208 the presence of the H3.3K27M allele would also radiosensitize tumors. To test this, we examined the tumor-free survival of nPHAFL/FL mice and compared them to controls with 209 intact ATM in their tumors of genotype nPHAFL/+ (Figure 1C). There was no difference in 210 tumor-free survival between nPHA<sup>FL/FL</sup> and nPHA<sup>FL/+</sup> mice in the absence of irradiation 211 212 (Figure 1D). To test whether Atm deletion radiosensitizes p53-null/H3.3K27M DMGs, 213 we delivered three daily fractions of 10 Gy focal brain irradiation to mice using the Small Animal Radiation Research Platform (SARRP). nPHAFL/FL mice had significantly longer 214 median survival than nPHA<sup>FL/+</sup> mice (P-value=0.03 using Mantel Cox (log rank test), 215 216 Figure 1E). Thus, Atm deletion in tumor cells enhances the efficacy of focal brain

irradiation for primary p53-null/H3.3K27M DMGs. H&E confirmed tumor (Figure 1F)
followed by IHC confirming HA expression (Figure 1G), p53 loss (Figure 1H), the
presence of H3.3K27M (Figure 1I), Ki67 (Figure 1J) and anti-FLAG (Figure 1K). These
results show that *Atm* disruption enhances the efficacy of radiation therapy for primary
mouse DMGs that contain p53 loss and the H3.3K27M mutation.

222

# In situ multiplexed microscopy reveals cell cycle and Semaphorin pathway changes after irradiation and Atm disruption.

225 To explore the mechanisms underlying radiation efficacy and resistance, we performed 226 spatially resolved gene expression analyses of primary mouse DMGs. Our previous 227 work identified key differences in the response to irradiation and Atm loss between the 228 neoplastic and vascular compartments within primary mouse tumors(12). To distinguish 229 compartment-specific changes in gene expression, such as vascular and immune cells 230 in specific regions of the tumor and nontumor brain, we needed to profile expression 231 changes at single-cell resolution and in a spatially resolved manner. To achieve such a 232 resolution, we used the 10xGenomics Xenium ISS platform to profile primary p53-233 null/H3.3K27M mouse DMGs. We examined DMG-bearing mice treated with or without 234 focal brain irradiation (10 Gy x 3), with or without tumor Atm loss as depicted in Figure 235 2A. We examined 5 µm mid-sagittal sections of formalin-fixed, paraffin-embedded 236 (FFPE) tumor-bearing brains. We supplemented 10xGenomics' standard mouse brain 237 content with a custom panel containing padlock probes resulting in 298 brain- and 238 DMG-specific mRNA transcript assays (Supplemental Table 1). Individual cells were 239 detected by nuclear DAPI staining and cell boundaries defined by *in silico* segmentation

240 (see Methods). This yielded 790,374 individual cells across the four tumor-bearing 241 brains. Next, we clustered cells based on their transcriptional profiles and compared cell 242 type composition between the samples. Uniform Manifold Approximation and Projection 243 (UMAP)(21) reduction, projection, and harmony integration of differentiated normal and 244 neoplastic brain cells into 20 and 29 clusters per specimen, respectively (Figure 2B and 245 Supplemental Figure 3). Examination of differentially expressed marker genes in each 246 cluster identified neoplastic and normal cells including GABAergic interneurons marked 247 by Gad1 and Gad2; microglia marked by P2ry12, Lyz2 and C1qa; and endothelial cells 248 marked by Cd34, Fn1, and Adgrl4 (Supplemental Figure 4). We used canonical cell-249 type markers and label transfer-based methods to collapse cell clusters into 10 cell 250 archetypes (neoplastic, endothelial, neuron, astrocyte, oligodendrocyte, microglia, T-251 lymphocyte, etc.) that could be directly compared across specimens (Supplemental 252 Table 2 and Supplemental Figure 5). This analysis revealed masslike tumors with 253 infiltrating edges recapitulating diffuse glioma biology (Figure 2C). Notably, an Atm-null 254 post-irradiation tumor was smaller and involuted, which is suggestive of rapid treatment 255 response. The tumor core, periphery, and nontumor areas were contoured using these 256 data to allow comparisons between matching cell types and locations after irradiation or 257 Atm loss (Figure 2C).

258

We used the spatially resolved expression data to identify differentially expressed genes among neoplastic cells within the tumor cores. We first localized the tumor core within the full-brain sagittal sections using canonical DMG neoplastic cell markers, *Olig1*, *Olig2*, and *Pdgfra* (Figure 3A). As expected, we could not detect *p53* in the neoplastic

263 cells within the tumor core in the Tp53<sup>FL/FL</sup> model, whereas low baseline levels could be 264 detected in non-neoplastic cell types (Figure 3B). Similarly, Atm transcripts were nearly 265 undetectable in neoplastic cells from Atm-null tumor (mean fold-change -0.636, 266 P<0.0001 when compared to Atm-intact tumor (Supplemental Figure 6). To identify 267 transcripts that may be differentially expressed after irradiation and/or Atm loss, we 268 interrogated differentially expressed genes in neoplastic cells after focal brain irradiation 269 within Atm intact tumors (Supplemental Table 3) and Atm-null tumors (Supplemental 270 Table 4). Cyclin-dependent kinase 1a (Cdkn1a), which encodes p21 a potent regulator 271 of cell cycle progression at G1, was the most differentially expressed gene after focal 272 brain irradiation among Atm intact tumors (log-fold change 0.8, P-value = 0 by Wilcoxon 273 test, Figure 3C). Cdkn1a was still upregulated, albeit to a lesser degree, after focal brain 274 irradiation among Atm-null tumors (log-fold change 0.6, P-value = 5.46E-08 by Wilcoxon 275 test, Figure 3D). Conversely, transcription factors associated with developmental cell 276 states such as Sox8 and Sox9 were substantially downregulated after irradiation in 277 Atm-intact tumors, while Sox2, Sox4, Pdgfra, and Olig2 associated with early glial 278 differentiation were all substantially downregulated after irradiation in Atm-null tumors. 279 These results identify the differential expression of cell cycle regulators and cell-fate-280 regulating transcription factors after irradiation in a primary DMG mouse model.

281

Irradiation and *Atm* loss were associated with changes in the expression of Semaphorin
genes specifically, *Semaphorin 6A* (*Sema6a*) and *Semaphorin 3D* (*Sema3d*) which
have been implicated in the proliferation and survival of glioma mouse models and
glioblastomas(22, 23). After irradiation in *Atm*-intact tumors, *Sema3d* was significantly

286 increased (log-fold change 1.13, P-value = 0) suggesting that radiation therapy may 287 influence proliferation within the neoplastic core. After irradiation of Atm-null tumors, 288 Sema6a was significantly decreased (log-fold change -0.40, P-value = 6.59E-15). We 289 utilized snRNA sequencing data from additional primary murine models derived via in 290 utero electroporation approaches to validate Semaphorin, p21, and endothelial 291 interactions in orthogonal models (Supplemental Figure 7) (24). However, our single cell 292 spatial transcriptomics provide additional mechanistic insight indicating that specific 293 Semaphorin genes are altered in neoplastic cells after radiotherapy that might play a 294 critical role in glioma biology.

295

# Neighborhood analysis shows altered immune-neoplastic interactions after treatment.

298 Targeting ATM combined with irradiation can bridge innate with adaptive immune 299 processes in extracranial cancers (25, 26). This led us to interrogate the spatial 300 relationship between neoplastic cells and immune microenvironment. We examined 301 whether the proximity between neoplastic cells and normal cells varied across irradiated 302 or Atm-null tumors. Neighborhood analysis guantified the spatial proximity between 303 different cell types and was used to estimate the mean distance between the neoplastic 304 cells and other cell types (Supplemental Figure 8). These data identified increased 305 proximity of neoplastic cells and immune cells, such as antigen presenting cells and 306 microglia, after Atm loss and after treatment with irradiation, which was especially 307 pronounced in the irradiated Atm-null tumor. Co-localization analysis between 308 neoplastic cells and other cell types confirmed that microglia and antigen-presenting

cells were most enriched within 0-500 μm (Figure 3E-H), and these cell types were most
 co-localized in the irradiated, *Atm*-null tumor (Figure 3H).

311

#### 312 Ligand-receptor analysis reveals endothelial cell communications.

313 Next, cell:cell and cell:ligand receptor interactions in primary mouse DMGs established 314 that endothelial cells had the highest frequency of interactions (Supplemental Figure 9 315 and Supplemental Table 5). We evaluated statistically significant ligand-receptor 316 interactions (p-value <0.05) among the tumors and identified the interaction between the 317 endothelium, microglia, and neoplastic cells with the Col1a2:CD93 receptor decreased 318 after Atm loss and irradiation (Supplemental Figure 9). CD93 plays a role in tumor-319 associated vasculature (27) and changes in Col1a2 expression has been observed after 320 radiotherapy in other cancers (28). These results provide insight into the changes in 321 endothelial cell interactions after tumor irradiation. After irradiation of Atm-intact tumors, 322 the cell:ligand interaction of Sema3a:NRP2 between neoplastic cell and microglia 323 decreased. This interaction has been noted to affect glioma cell migration(29) implying 324 a potential alteration in migration with irradiation. The opposite effect was observed in 325 Atm-null tumors after irradiation (Supplemental Figure 9). Thus, ligand-receptor analysis 326 of ISS data suggests that glioma-linked collagen and Semaphorin interactions can be 327 examined in primary DMG mouse models.

328

#### 329 Pharmacologic ATM inhibition deregulates DNA damage responses and improves

**survival.** To validate these findings, we confirmed that pharmacological inhibition of

331 ATM could radiosensitize human patient-derived models of DMG. To do so, we tested

332 whether the brain-penetrant ATM inhibitor AZD1390 (30) combined with focal brain 333 irradiation could similarly improve the survival of a patient-derived xenograft model of 334 H3.3K27M-mutant and p53-mutant diffuse midline glioma, SF8628 (31-34) which lacks 335 a functional ATM mutation (Supplemental Figure 10 and Supplemental Table 6). The 336 combination of AZD1390 and irradiation significantly extended the median survival 337 compared with either treatment alone (Figure 4A). We tested an Atm-intact genetically 338 engineered model with a combination of AZD1930 and irradiation which led to a trend 339 for extended median survival compared to irradiation alone (median 29 days vs. 10 340 days, p-value = 0.1, Log-rank test, Supplemental Figure 11). These results confirmed 341 that pharmacologic or genetic targeting of ATM can radiosensitize multiple types of in 342 vivo DMG models.

343

344 We investigated DNA damage response in the SF8628 line by performing western blots 345 in cells treated with AZD1390 with or without irradiation which showed increased 346 expression of GH2AX up to 24 hours post irradiation (Figure 4B and Supplemental 347 Figure 12). To further interrogate effects of irradiation after treatment with AZD1390, we treated Nestin-Tva Cre p53<sup>FI/FI</sup> PDGF-B H3.3K27M mice (13) with vehicle or drug along 348 349 with 10 Gy irradiation and harvested mouse brain tumors an hour post irradiation. 350 AZD1390 effectively inhibited ATM as indicated by significantly reduced phospho-KAP1 351 expression in the treated group when compared to control (p-value 0.01, Mann-352 Whitney) and increased GH2AX expression (p-value 0.03, Mann-Whitney) in tumor-353 bearing mice (Figure 4C-F). The differential change in GH2AX in tumor bearing mice

treated with AZD1390 when compared to radiation therapy alone indicates a synergisticeffect.

356

#### 357 Atm radiosensitizes Cdkn1a-null primary murine DMGs.

358 Next, we dissected the specific functions of p53 that may affect the radiosensitivity of 359 mouse DMG. Our primary models of DMG indicated that the presence of functional p53 360 is a key determinant of whether tumors are radiosensitized by Atm loss. For instance, 361 primary p53-null/H3.3K27M tumors and primary p53-null/H3f3a-wildtype tumors were 362 radiosensitized by Atm loss (Figure 1E and Deland et al. (5), respectively). Conversely, 363 p53 wildtype primary DMG models driven by Ink4A/ARF or PTEN loss are not 364 radiosensitized by Atm loss (5, 11). However, it is unknown whether the loss of p53 365 transcriptional activation and/or loss of other p53 functions enables radiosensitization by 366 Atm loss. Our ISS data identified increased Cdkn1a expression in neoplastic cells after 367 radiation. Since Cdkn1a (encoding p21) is a major transcriptional target of p53 (35), we 368 hypothesized that the loss of *Cdkn1a* function downstream of p53 may be a key 369 determinant of whether Atm loss can radiosensitize primary DMGs. To test whether 370 Cdkn1a loss allows primary mouse DMGs to be radiosensitized by Atm loss, we 371 examined our model of p53-wildtype DMGs driven by Ink4A/ARF loss, which was not radiosensitized by Atm loss (Nestin<sup>TVA</sup>; Ink4A/ARF<sup>FL/FL</sup>) (5). To test whether p21 loss 372 373 could radiosensitize these mice when Atm was lost, we bred mice with constitutive p21 loss into this genotype (Nestin<sup>TVA</sup>; p21<sup>-/-;</sup> Ink4A/ARF<sup>FL/FL</sup>; Atm<sup>FL/FL</sup> (nlp21A<sup>FL/FL</sup>). We 374 375 tested the effects of tumor-specific Atm loss by comparing these mice with their littermate controls with intact *Atm* with genotype Nestin<sup>TVA</sup>; p21<sup>-/-;</sup>; Ink4A/ARF<sup>FL/FL</sup>; 376

Atm<sup>FL/FL</sup> (nlp21A<sup>FL/+</sup>) (Figure 5A). The time to tumor formation was similar regardless of 377 378 the presence of intact Atm (Figure 5B). Surprisingly, p21-null mice bearing tumors with 379 Atm deletion had shorter survival following fractionated focal brain irradiation compared 380 to littermate controls with intact Atm in the tumors (Figure 5C, P<0.03, log-rank test). 381 We confirmed p21 loss using IHC (Figure 5D-E). p21-null tumors with and without Atm 382 loss had similar proliferation indices as assessed by Ki67 staining (Figure 5F). TUNEL 383 staining of irradiated tumors showed that Atm loss was associated with significantly 384 increased TUNEL staining (P < 0.05, Figure 5G-H), suggesting that tumors lacking both 385 Atm and Cdkn1a were primed for apoptosis. These results show that functional Cdkn1a 386 is not a key mediator of radiosensitization by Atm loss in primary mouse model of DMG. 387

A p53 transactivation domain mutant retains tumor suppressor function in mouse
 DMG.

390 Since Atm loss could not radiosensitize Cdkn1a-null DMGs, we reasoned that regulation 391 of p53 transcriptional targets other than Cdkn1a may cause radioresistance in p53 wild 392 type, Atm-null DMGs. To investigate this possibility, we leveraged the conditional loxP-Stop-loxP-p53<sup>25,26</sup> allele (p53<sup>LSL-25,26</sup>) (36). In the presence of Cre recombinase, this 393 394 allele expresses p53<sup>25,26</sup>, a p53 mutant that is severely compromised for the 395 transactivation of most p53 target genes and cannot induce G1-arrest or apoptosis in 396 response to acute DNA damage (36). Interestingly, p53<sup>25,26</sup> retains tumor suppressor 397 activity in lung tumors (36), however it is unknown whether it retains tumor suppressor 398 activity in brain tumors. We first determined if p53<sup>25,26</sup> retained tumor suppressor activity in DMG. To test this hypothesis, we compared littermate mice with either p53<sup>LSL-25,26/FL</sup> 399

or p53<sup>FL/FL</sup>. All mice harbored Nestin<sup>TVA</sup> and were injected with Cre, luciferase, and 400 401 PDGF-B retrovirus constructs as described above (Figure 6A). We noted a marked delay in tumor presentation in the p53 <sup>LSL-25,26/FL</sup> group compared to that in the p53<sup>FL/FL</sup> 402 403 controls (Figure 6B-C). Immunohistochemical analysis revealed heterogeneous p53 expression in the p53<sup>LSL-25,26/FL</sup> group and apparently absent p53 expression in the 404 p53<sup>FL/FL</sup> group (Figure 6D-E). Thus, a p53 mutant with severely compromised 405 406 transactivation activity retains its tumor suppressor activity in primary mouse brainstem 407 gliomas. These results indicate that p53 transactivation function is dispensable for p53 408 tumor suppression in DMG.

409

#### 410 Atm loss does not radiosensitize mouse DMGs lacking a functional p53

#### 411 transactivation domain.

412 We next sought to determine if Atm loss could radiosensitize DMGs lacking p53 413 transcriptional activity but retaining other non-transcriptional functions of p53. We 414 previously showed that Atm loss did not radiosensitize brainstem gliomas driven by 415 Ink4A/ARF loss, however Atm loss modestly radiosensitized brainstem gliomas with 416 both Ink4A/ARF loss and p53 loss (5). We reasoned that if loss of p53 transactivation 417 domain function is the determinant of radiosensitization by Atm loss, then brainstem 418 gliomas with both Ink4A/ARF loss and expression of a transactivation-deficient p53<sup>25,26</sup> 419 allele would be radiosensitized by Atm loss. To test if mouse DMGs with p53<sup>25,26</sup> and Ink4A/ARF loss were radiosensitized by *Atm* loss, we bred mice of genotype Nestin<sup>TVA</sup>; 420 p53<sup>LSL-25,26/FL</sup>; Ink4A/ARF<sup>FL/FL</sup>; Atm<sup>FL/FL</sup>. To test the effects of *Atm* loss, we compared 421 422 these to littermate controls with the same genotype except an intact Atm allele

(Nestin<sup>TVA</sup>; p53<sup>LSL-25,26/FL</sup>; Ink4A/ARF<sup>FL/FL</sup>; Atm<sup>FL/+</sup>) (Figure 7A). We noted similar time to 423 424 tumor formation in both models (Figure 7D). Atm loss was associated with differential 425 staining of phospho-ATM and phospho-KAP1 after focal brain irradiation (Figure 7B and 426 7C), confirming the loss of ATM functional activity. After subjecting the mice to 427 fractionated focal brain irradiation, no difference in overall survival was appreciated 428 (Figure 7E). These results indicate that the transactivation-independent functions of p53 429 may be the primary determinants of whether mouse DMGs can be radiosensitized by 430 Atm loss. 431

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#### 433 **Discussion**

434 Here, we describe the generation of primary mouse DMGs based on recent advances in 435 murine genetic engineering including the conditional H3.3K27M allele and the RCAS/tv-436 a retrovirus platform (10, 13, 14). We used this model to show that the genetic loss of 437 Atm, an important target for drugs that have entered clinical trials for brain tumor 438 patients(18), radiosensitizes primary DMG models. Our results in p53-null/H3.3K27M 439 mouse DMGs were similar to those reported previously for p53-null mouse brainstem 440 gliomas(5), in which the sole difference is the presence of H3.3K27M expression from 441 the endogenous H3f3a locus in the neoplastic tumor cells in our current model. In 442 addition, we generated several unique genetically engineered mouse models with 443 differential responses based on genotype highlighted in Table 1 which suggests that 444 H3.3K27M is not a primary determinant of the ability to target ATM to enhance the 445 efficacy of radiation therapy in primary mouse DMG models (Supplemental Figure 13).

446

447 Our results from genetic experiments in primary mouse models indicate that p53 is a 448 key determinant of the ability of DMG to be radiosensitized by Atm loss. Almost all p53-449 altered primary mouse models were radioresistant and radiosensitized by Atm loss, 450 including (i) a model driven by p53 loss with wild type H3f3a (5), (ii) a model driven by 451 both p53 loss and loss of Ink4A/ARF(5), and (iii) the H3.3K27M/TP53 mutant model 452 reported here (Figure 1E). In contrast, Atm loss is unable to radiosensitize primary p53-453 wildtype brainstem glioma mouse models, including models driven by Ink4A/ARF 454 loss(5) and models driven by *Pten* loss (11). Notably, a recent study comprehensively

found that pharmacological ATM inhibition radiosensitized both p53-mutant and p53wild type patient-derived models of DMG and pediatric high-grade glioma(6)Also,
H3.3K27M may enhance ATM signaling, increasing radiosensitivity both with and
without ATM inhibition (37). Together, these data suggest that mutational status of p53,
H3.3K27M, and other alterations should be tested in correlative analyses in future
clinical trials of ATM inhibitors in patients with DMG.

461

462 Our ISS data provide the first high-resolution transcriptional analysis at high gene plexy 463 (~300 gene targets) in a mouse tumor model, which is critical to defining the model's 464 tumor vasculature and neoplastic compartments that play distinct roles in therapeutic 465 response (12). Future work will leverage these data to interrogate tumor immune and 466 vascular microenvironment alterations induced by irradiation and *Atm* loss, which may 467 guide the rational design of combinations between radiation therapy, ATM inhibitors, 468 and therapies targeting the immune system or vasculature.

469

470 This work has several limitations. H3.3K27M did not decrease tumor latency in our 471 system as has been observed in other experimental systems (38). This may be due to 472 the highly restricted manner of H3.3K27M induction in our system (ie, from the 473 endogeneous H3f3a locus and only in spatially- and lineage-restricted cells) and/or the 474 use of a relatively strong PDGFB co-driver alteration that could mask more subtle 475 H3.3K27M driver phenotypes in our system. Also, the presence of an HA tag on both 476 HA-PDGFB and H3.3K27M-Flag-HA constructs precludes specific identification of 477 PDGFB in the nPH system. Finally, we observed a trend for improved overall survival

484	
483	effects of the ATM inhibitor.
482	interrogate ATM inhibition effects in Atm-null models to dissect on-target from off-target
481	compared to the pharmacological xenograft experiment. Future studies could
480	significant survival benefit difficult to detect compared to the genetic loss of Atm and
479	heterogeneity in tumor latency and timing of treatment delivery may have made a
478	with ATM inhibition in an Atm-intact genetic model. We hypothesize that the

486 **Conclusion** 

487

488 The current work implicates transactivation-independent mechanisms by which p53 489 mediates radioresistance in Atm-null tumors. Our ISS data showed that irradiation 490 elicited overexpression of Cdkn1a, a key downstream target of p53 that mediates cell 491 senescence and G1-to-S checkpoint arrest, in p53-null tumors suggesting p53-492 independent mechanisms of Cdkn1a expression (39). This finding led us to dissect the 493 contribution of p53 transactivation functions which regulate p21 expression to 494 radiosensitization in Atm-null DMGs. While Atm loss radiosensitizes tumors lacking p53, we found that *Atm* loss could not radiosensitize tumors containing a p53<sup>25,26</sup> allele 495 496 deficient in p53 transactivation function. Similarly, tumors lacking Cdkn1a (p21) could 497 not be radiosensitized by Atm loss. Our findings highlight the importance of carefully 498 considering p21 status in clinical trials involving Atm inhibition given the complex role of 499 p21 in tumor growth and the microenvironment (40). Strikingly, we found that Atm loss 500 made tumors more radioresistant in mice that lacked Cdkn1a, and that this finding was 501 associated with increased apoptosis. These findings implicate the transactivation-502 independent function of p53 as a key determinant of radiosensitivity in Atm-null tumors. 503 Future work will dissect the transactivation-independent functions of p53 such as 504 promoting apoptosis through mitochondrial membrane permeabilization, direct 505 repression of transcription, and/or direct interaction with complexes that detect DNA 506 lesions (41, 42). Our data provides genetic and mechanistic insight that builds upon 507 studies of pharmacological ATM inhibition in patient derived xenograft models (6). In 508 addition, our work identifies ATM inhibition improving response to irradiation leading to

- 509 extended survival. Further studies are needed to determine the transactivation-
- 510 independent mechanisms of p53 and ATM-directed therapies and their impact on
- 511 overcoming resistance to radiation therapy in patients with H3.3K27M-mutant DMG.

#### 513 Methods

514 Detailed workflows for the generation, brain irradiation, and molecular analysis of 515 primary mouse DMG models using RCAS/tv-a and Cre/loxP technologies are found in 516 our recent manuscript(43). Male and female mice were utilized for all murine models. All 517 new reagents, materials, and software are listed in the key resources table (Table 2) . A 518 list of abbreviations utilized is listed in Supplemental table 7.

#### 519 Sex as a biological variant

520 Male and female mice were utilized in all murine experiments in this publication to 521 ensure representation of both sexes. We did not identify any sex specific differences in 522 the data, and all findings were consistent across male and female mice. As such, sex 523 was not considered to be a biological variable in the interpretation of the results. The 524 outcomes of this study are therefore expected to be broadly relevant to both sexes. 525 Mouse strains. 526 Detailed workflows for generation, brain irradiation, and molecular analysis of primary 527 mouse DMG models using RCAS/tv-a and Cre/loxP technologies are found in our 528 recent STAR Protocols manuscript(43). Complex mouse strains were generated by

529 breeding mice with the following alleles: Nes<sup>TVA</sup>, Atm<sup>FL</sup>, and p53<sup>FL</sup> (43), Ink4A/ARF<sup>FL</sup>

530 (43), the  $p53^{LSL(25,26)}$  (36), and the  $p21^{-/-}$  (44). The H3f3a<sup>LSL-K27M-Tag</sup> allele is a gift from

531 Dr. Suzanne Baker(10).

532

533 **DF1 cell culture and retrovirus generation.** 

534	DF1 cells were cultured in Dulbecco's Modified Eagle's medium (DMEM) containing
535	10% fetal bovine serum (FBS) and RCAS/tv-a retroviruses were generated using
536	RCAS-Cre, RCAS-luc, and RCAS-PDGFB plasmids as previously described(43).
537	
538	Mouse brainstem injection.
539	The harvested DF1 cells were injected into the brainstems of mice anesthetized on ice
540	on postnatal day 3-5 as previously described(43). All institutional approvals were
541	obtained prior to the injection. Patient-derived xenografts using the SF8628 model were
542	generated by brainstem injection as described(31-33).
543	
544	Mouse <i>in-vivo</i> imaging
545	Bioluminescence imaging of gliomas within mice was performed by intraperitoneal
546	injection of D-luciferin and IVIS Illumina III as previously described(43).
547	
548	Image Guided focal brain irradiation. Irradiation to gliomas within mice was delivered
549	on a Small Animal Radiation Research Platform (SARRP) using image-guided,
550	opposed-lateral beams as described(43). 10 Gy times three consecutive daily fractions
551	was delivered for primary DMG models. 2 Gy times three days a week to 12 Gy total
552	was delivered for patient-derived xenograft DMG models.
553	
554	ATM inhibitor studies.
555	Patient-derived xenograft model: SF8628 (H3.3K27M DIPG) was obtained from the
556	University of California San Francisco (UCSF) Medical Center, in accordance with an

557 institutionally approved protocol. Establishment of SF8628 cell culture from surgical 558 specimens, and tumor cell modification for expression of firefly luciferase for in vivo 559 bioluminescence imaging, have been described(31-33). SF8628 cells were propagated 560 as monolayers in complete medium consisting of DMEM supplemented with 10% FBS 561 and non-essential amino acids. Short tandem repeats (STR) were obtained to confirm 562 the identity of cell lines. All cells were cultured in an incubator at 37°C in a humidified 563 atmosphere containing 95% O2 and 5% CO2 and were mycoplasma-free at the time of 564 testing. Six-week-old female athymic mice (rnu/rnu genotype, BALB/c background) were 565 purchased from Envigo and housed under aseptic conditions. Pontine injection of tumor 566 cells was performed as previously described (31-33). Each mouse was injected with 1 567  $\mu$ L of the SF8628 cell suspension (100,000 cells/ $\mu$ L) into the pontine tegmentum at a 568 depth of 5 mm from the inner base of the skull. For the efficacy study of AZD1390 and 569 radiation, animals were randomized into four treatment groups: 1) vehicle control (0.5% 570 hydroxymethylcellulose, 0.1% Tween 80, n=6), 2) ADZ1390 monotherapy (oral gavage 571 of 20 mg/kg of AZD1390 for 5 times a week for two consecutive weeks, n=6), 3) 572 radiation monotherapy (2.0 Gy, 3 times a week for two consecutive weeks for a total 573 dose of 12 Gy, n=6), and 4) AZD1390 and radiation combination therapy 574 (n=6). Biweekly bioluminescence imaging was used to monitor tumor growth and 575 response to therapy as previously described(31). Mice were monitored daily and 576 euthanized at the endpoint which included irreversible neurological deficit or a body 577 condition score of less than 2. Genetically Engineered mouse model: Gliomas were 578 generated using the RCAS/Tv-a system as previously described (45, 46). DF1 Cells 579 were transfected with RCAS plasmids (RCAS-PDGF-B, RCAS-Luc, RCAS-Cre and

580 mixed in 1:1:1 ratio prior to intracranial injections. nP mice were intracranially injected 581 with 1.0 µl of RCAS virus producing DF1 cells at postnatal day 3 to 5 (P3-P5) and 582 monitored 3x weekly post-weaning for signs of brain tumor symptoms (enlarged head, 583 ataxia, weight loss up to 20%). Bioluminescence imaging was then used to monitor 584 tumor formation from weeks 4-12 as described previously (39). Following tumor 585 detection via imaging or the onset of neurological symptoms, mice were randomized 1:1 586 into two treatment groups: 1) RT alone or 2) AZD1390 + RT. AZD1390 was obtained 587 from AstraZeneca and resuspended in 0.5% w/v HPMC and 0.1% w/v Tween-80 588 solution. Mice in the AZD1390 + RT group were dosed at 20 mg/kg via oral gavage one 589 hour prior to RT, as suggested by the superior efficacy dose (18). Both treatment 590 groups received three consecutive daily fractions of 10 Gy of focal brain irradiation 591 delivered by the SARRP. Treated mice were monitored for survival until they reached 592 the humane endpoint. (14, 18)

593

#### 594 Immunohistochemistry (IHC)

IHC was performed using methods previously described for Ki67, p-Atm, p-KAP1, total
KAP1, γH2AX, and p53(18, 43). Additional IHC and TUNEL staining were performed by
HistoWiz Inc. (histowiz.com) using a Standard Operating Procedure and a fully
automated workflow for p21 (Cdkn1a), FLAG, and TUNEL on a Bond Rx autostainer
(Leica Biosystems) with enzyme treatment (1:1000) using standard protocols.
Hematoxylin and eosin (H&E) staining was performed using standard protocols.

#### 602 Quantification and Statistical Analysis

603 Statistical significance in volcano plots across ROIs was assessed with the Wilcoxon 604 test on SCTransform normalized count data using the FindMarkers Seurat function. Data plotting and quantification analyses were performed using GraphPad Prism 9. The 605 606 unpaired t-test was utilized to determine significance in quantification by IHC. The log-607 rank test was used to determine the survival rate. The Wilcoxon test was used to 608 determine differences in the time to tumor detection. Individual data points were plotted, 609 and all statistically significant values (p value less than 0.05) are identified with an 610 asterisk (\*).

611

#### 612 Xenium In Situ and Bioinformatics Analysis

613 Tumor-bearing brains subjected to Xenium ISS were detected by *in vivo* imaging 37-48 614 days after birth and collected seven days after tumor detection, either after three daily 615 treatments of 10 Gy initiated within two days of tumor detection, or mock treatment. 616 Initial data generated by the Xenium instrument (47) are processed on board with a built-617 in analysis tool called Xenium Analyzer(47). The Xenium Analyzer is fully automated 618 and includes an imager (imageable area of approximately 12 x 24 mm per slide), 619 sample handling, liquid handling, wide-field epifluorescence imaging, capacity for two 620 slides per run, and an on-instrument analysis pipeline. The analysis pipeline included 621 image pre-processing, puncta detection, transcript decoding and quality score 622 assignment. The pipeline also performed cell segmentation using DAPI images to 623 detect nuclei using a neural network. Each nucleus is then expanded outwards until 624 either a maximum distance of 15 µm is reached or the boundary of another cell is 625 reached. A variety of output files were produced using an on-instrument pipeline. The

essential files for downstream analysis include the feature-cell matrix (HDF5 and MEX
formats identical to those output by single cell RNA tools from 10X
(Cellranger/Spaceranger), the transcripts (listing each mRNA, its 3D coordinates, and a
quality score), and the cell boundaries CSV file.

630

631

632 Xenium output was first imported into R (4.3.1) using the LoadXenium function from

633 Seurat (4.9.9.9050)(48). The four Xenium samples were processed using Seurat(49).

634 Data was loaded and filtered using nFeature\_Xenium>5 and nCount\_Xenium >10 as

635 criteria. Cells without a predicted annotation were then subset out, and the four samples

636 were normalized using SCTransform. PCA was also run for each sample. There four

637 samples were then integrated using the IntegrateLayers function, using

638 HarmonyIntegration as the method(50).

639

640 Specific regions of the tissue were annotated manually using the polygon tool in Xenium 641 Explorer software (development version, 10x Genomics), and the polygon coordinates 642 were exported as csv files. Cells with zero counts were then filtered, and the points 643 within the polygon coordinates were identified using the point in polygon function in the 644 sp (2.0.0) R package. Further plots were generated using Seurat, and deconvolution 645 was performed with spacexr (2.2.1)(51) using a custom annotated single cell reference 646 from a previous experiment. DGE analysis across ROIs was assessed using the 647 Wilcoxon test on SCTransform normalized count data using the FindMarkers Seurat 648 function. Co-occurrence and neighborhood enrichment plots were made using the

649 Python package Squidpy (1.2.3)(52), and trajectory and cell-cell-interaction analyses
650 were performed using the Python package STLearn (0.4.12) (53).

651

All the cells on the entire slide were used to determine the different types of cell clusters. All unlabeled cells were removed for Squidpy and STLearn analyses. Cooccurrence and neighborhood enrichment analyses were conducted on all cells within the entire slide. The tumor core and periphery were utilized to identify cell:cell and cell:ligand interactions. Differentially expressed genes were analyzed in the tumor core.

657

#### 658 Whole Genome Bisulfite Sequencing and Bioinformatics Analysis

659 Bisulfite methylation sequencing and data analysis were performed by Novogene.

Briefly, K27M mutant (nPH) and matched K27M wildtype (nP) tumor-bearing mice (n = 4

biological replicates per group) were generated as described above, and tumor-bearing

brains embedded in FFPE. Tumor regions were identified in the brains on matched

663 H+E slides, and tumor microdissection was performed . Genomic DNA was isolated,

spiked with lambda bacteriophage DNA (to serve as an internal negative control),

665 fragmented to 200-400 base pairs, and bisulfite treatment was performed to convert

666 unmethylated cytosines into uracil via deamination. Notably, this process does not alter

667 methylated cytosines, allowing identification of these sites downstream. After

668 methylation sequencing adapter ligation, double strand DNA synthesis, and library size

669 selection, PCR amplification was performed followed by Illumina sequencing. FastQC

670 was used for quality control on the raw reads. Bismark software (version 0.24.0;

671 Krueger et al., 2011) was used to perform alignments of bisulfite-treated reads to a

672 reference genome (-X 700--dovetail). The reference genome was firstly transformed into 673 bisulfite-converted version (C-to-T and G-to-A converted) and then indexed using 674 bowtie2 (Langmead et al., 2012). Sequence reads were also transformed into fully 675 bisulfite-converted versions (C-to-T and G-to-A converted) before they are aligned to 676 similarly converted versions of the genome in a directional manner. Sequence reads 677 that produce a unique best alignment from the two alignment processes (original top 678 and bottom strand) are then compared to the normal genomic sequence and the 679 methylation state of all cytosine positions in the read is inferred. The same reads that 680 aligned to the same regions of genome were regarded as duplicated ones. The 681 sequencing depth and coverage were summarized using deduplicated reads. The 682 results of methylation extractor (bismark methylation extractor, -- no overlap) were 683 transformed into bigWig format for visualization using IGV browser. The sodium bisulfite 684 non-coversion rate was calculated as the percentage of cytosine sequenced at cytosine 685 reference positions in the lambda genome. Genes were extracted from genome 686 assembly GRCm39 using Ensembl gene set Ver 111. Promoter regions are defined 687 from 1500 bp upstream of TSS to 500 bp downstream. For each region, methylated 688 CpG reads and unmethylated reads were counted, summed and average methylation 689 level was calculated. Similarly, putative enhancers were extracted from Ensembl 690 regulation Ver 111 of GRCm39 and an average methylation level was calculated for 691 each enhancer. Methylation difference was determined for each feature (promoter and 692 enhancer) between samples of K27M group and samples of nPA group. Significance 693 was estimated by applying ANOVA over a linear mode fit(54). For DNA methylation 694 extraction we used methkit. Motif analysis was carried out by HOMER which is available

- 695 online at http://homer.ucsd.edu/homer. For pathway enrichment analysis STRING was
- 696 used. For "other tissues" we analyzed data from GSE42836. methylation extraction
- 697 using the methkit.
- 698

699	Declarations
700	
701	Study approval
702	
703	All animal studies were approved by the institutional animal protocol. All animal
704	experiments were approved by the Institutional Animal Care and Use Committee at
705	Duke and Northwestern University.
706	
707	Consent for Publication
708	
709	Not applicable
710	
711	Availability of data and materials
712	All data were deposited in the GEO database. The accession number for the in situ
713	sequencing is GSE246584 and whole genome bisulfite sequencing is GSE284759. This
714	study does not report original code. Additional information required to reanalyze the
715	data reported in this paper is available from the corresponding author upon request. All
716	raw values for figures are available in Supplemental table 8.
717	
718	Competing Interests
719	
720	DGK is a cofounder and stockholder in XRAD Therapeutics, which is developing
721	radiosensitizers. DGK is a member of the scientific advisory board and owns stock in

722 Lumicell Inc, a company commercializing intraoperative imaging technology. None of 723 these affiliations represent a conflict of interest with respect to the work described in this 724 manuscript. DGK is a coinventor on a patent for a handheld imaging device and is a 725 coinventor on a patent for radiosensitizers. XRAD Therapeutics, Merck, Bristol Myers 726 Squibb, and Varian Medical Systems have provided research support to DGK, but this 727 did not support the research described in this manuscript. ZJR receives royalties for 728 intellectual property related to brain tumor diagnostic tests that is managed by the Duke 729 Office of Licensing and Ventures and has been licensed to Genetron Health, and 730 honoraria for teaching from Oakstone Publishing and Eisai Pharmaceuticals. The other 731 authors have no conflicts of interests to declare.

732

733 Funding

734

735 We thank sources of funding including NCI K08256045 Mentored Clinician Scientist 736 Development Award, Alex's Lemonade Stand Foundation A Award No. 23-27774, 737 ChadTough Defeat DIPG Foundation, the SoSo Strong Foundation, the Pediatric Brain 738 Tumor Foundation, the Emily Beazley's Kures for Kids Fund, the St. Baldrick's 739 Foundation, Lauren Brescia Memorial Fund, and NCI P50CA190991 Duke SPORE in 740 Brain Cancer developmental funds to ZJR. This work was also supported by 741 7R35CA197616 from the NCI to DGK. We thank Hyundai Hope on Wheels for funding 742 to OJB. 743

744 Author contributions

746 AM prepared the manuscript, designed experiments, and analyzed data. VV designed 747 and led the execution of experiments for the revised manuscript including mouse tissue 748 analyses, whole-genome bisulfite sequencing, and pharmacologic in vivo experiments. 749 AM and VV share the first author position. The listed order of the first authors was 750 determined by alphabetical order of the last names. The order was determined to be A-751 Z based on a coin-flip from the corresponding author ZJR. AM, ZJR, SGG, DGK, DMA 752 and OJB designed the study and experiments. SW, HL, BEF, LW, MEG, DG, LL, KD, 753 VV, SM, MR and ZJR performed mouse experiments and tabulated data. NTW 754 performed mouse irradiations. KA and NH performed patient-derived xenograft and 755 ATM inhibitor pharmacologic experiments. LA contributed p53 transactivation mutant 756 mouse strain and experimental design regarding the strain. EH, KA, LW, and ZJR 757 assisted with in situ sequencing experiments. MA, VJ, JAR, SG and ZJR performed 758 bioinformatic analyses. TM performed whole-genome bisulfite sequencing analyses. 759 AM, ZJR, and SGG prepared the manuscript.

760

#### 761 Acknowledgements

762

We thank Dr. Suzanne Baker, St Jude Research Hospital, Memphis Tennessee for the
gift of the H3f3a-loxP-Stop-loxP-K27M-Tag mice. Funds for ISS data generation were
provided by the Duke Brain Tumor Omics Program, Durham, North Carolina.

766

#### 769 **References**

- Wu G, Broniscer A, McEachron TA, Lu C, Paugh BS, Becksfort J, et al. Somatic histone H3 alterations in pediatric diffuse intrinsic pontine gliomas and nonbrainstem glioblastomas. *Nat Genet.* 2012;44(3):251-3.
- Schwartzentruber J, Korshunov A, Liu XY, Jones DT, Pfaff E, Jacob K, et al.
  Driver mutations in histone H3.3 and chromatin remodelling genes in paediatric
  glioblastoma. *Nature.* 2012;482(7384):226-31.
- Zhang L, Chen LH, Wan H, Yang R, Wang Z, Feng J, et al. Exome sequencing
   identifies somatic gain-of-function PPM1D mutations in brainstem gliomas. *Nat Genet.* 2014;46(7):726-30.
- Werbrouck C, Evangelista CCS, Lobon-Iglesias MJ, Barret E, Le Teuff G,
  Merlevede J, et al. TP53 Pathway Alterations Drive Radioresistance in Diffuse
  Intrinsic Pontine Gliomas (DIPG). *Clin Cancer Res.* 2019;25(22):6788-800.
- 5. Deland K, Starr BF, Mercer JS, Byemerwa J, Crabtree DM, Williams NT, et al.
  Tumor genotype dictates radiosensitization after Atm deletion in primary
  brainstem glioma models. *J Clin Invest.* 2021;131(1).
- 786
  6. Xie J, Kuriakose T, Bianski B, Twarog N, Savage E, Xu K, et al. ATM inhibition
  787 enhances the efficacy of radiation across distinct molecular subgroups of
  788 pediatric high-grade glioma. *Neuro Oncol.* 2023.
- 789
   7. Khadka P, Reitman ZJ, Lu S, Buchan G, Gionet G, Dubois F, et al. PPM1D
   790 mutations are oncogenic drivers of de novo diffuse midline glioma formation. *Nat* 791 *Commun.* 2022;13(1):604.
- du Chatinier A, Meel MH, Das AI, Metselaar DS, Waranecki P, Bugiani M, et al.
   Generation of immunocompetent syngeneic allograft mouse models for pediatric
   diffuse midline glioma. *Neurooncol Adv.* 2022;4(1):vdac079.
- McNicholas M, De Cola A, Bashardanesh Z, Foss A, Lloyd CB, Hebert S, et al. A
  Compendium of Syngeneic, Transplantable Pediatric High-Grade Glioma Models
  Reveals Subtype-Specific Therapeutic Vulnerabilities. *Cancer Discov.* 2023.
- Larson JD, Kasper LH, Paugh BS, Jin H, Wu G, Kwon CH, et al. Histone H3.3
  K27M Accelerates Spontaneous Brainstem Glioma and Drives Restricted
  Changes in Bivalent Gene Expression. *Cancer Cell.* 2019;35(1):140-55 e7.
- Stewart CE, Guerra-Garcia ME, Luo L, Williams NT, Ma Y, Regal JA, et al. The
   Effect of Atm Loss on Radiosensitivity of a Primary Mouse Model of Pten-Deleted
   Brainstem Glioma. *Cancers (Basel)*. 2022;14(18).
- Boland K, Mercer JS, Crabtree DM, Guerra Garcia ME, Reinsvold M, Campos
  LDS, et al. Radiosensitizing the Vasculature of Primary Brainstem Gliomas Fails
  to Improve Tumor Response to Radiation Therapy. *Int J Radiat Oncol Biol Phys.*2022;112(3):771-9.
- 808 13. Cordero FJ, Huang Z, Grenier C, He X, Hu G, McLendon RE, et al. Histone
  809 H3.3K27M Represses p16 to Accelerate Gliomagenesis in a Murine Model of
  810 DIPG. *Mol Cancer Res.* 2017;15(9):1243-54.

- 811 14. Hoeman CM, Cordero FJ, Hu G, Misuraca K, Romero MM, Cardona HJ, et al. 812 ACVR1 R206H cooperates with H3.1K27M in promoting diffuse intrinsic pontine 813 glioma pathogenesis. Nat Commun. 2019;10(1):1023. 814 15. Hambardzumyan D, Amankulor NM, Helmy KY, Becher OJ, and Holland EC. Modeling Adult Gliomas Using RCAS/t-va Technology. Transl Oncol. 815 816 2009;2(2):89-95. 817 16. Tomita Y, Shimazu Y, Somasundaram A, Tanaka Y, Takata N, Ishi Y, et al. A 818 novel mouse model of diffuse midline glioma initiated in neonatal oligodendrocyte 819 progenitor cells highlights cell-of-origin dependent effects of H3K27M. Glia. 820 2022;70(9):1681-98. 821 17. Garcia MEG, Kirsch DG, and Reitman ZJ. Targeting the ATM Kinase to Enhance 822 the Efficacy of Radiotherapy and Outcomes for Cancer Patients. Semin Radiat 823 Oncol. 2022;32(1):3-14. 824 18. Durant ST, Zheng L, Wang Y, Chen K, Zhang L, Zhang T, et al. The brain-825 penetrant clinical ATM inhibitor AZD1390 radiosensitizes and improves survival 826 of preclinical brain tumor models. Sci Adv. 2018;4(6):eaat1719. 827 Lewis PW, Muller MM, Koletsky MS, Cordero F, Lin S, Banaszynski LA, et al. 19. 828 Inhibition of PRC2 activity by a gain-of-function H3 mutation found in pediatric 829 glioblastoma. Science. 2013;340(6134):857-61. 830 Filbin MG, Tirosh I, Hovestadt V, Shaw ML, Escalante LE, Mathewson ND, et al. 20. 831 Developmental and oncogenic programs in H3K27M gliomas dissected by single-832 cell RNA-seq. Science. 2018;360(6386):331-5. 833 21. van der Maarten LH, G. Visualizing Data using t-SNE. Journal of Machine 834 Learning Research. 2008;9:2579-605 835 22. Zhao J, Tang H, Zhao H, Che W, Zhang L, and Liang P. SEMA6A is a prognostic 836 biomarker in glioblastoma. Tumour Biol. 2015;36(11):8333-40. Angelucci C, Lama G, and Sica G. Multifaceted Functional Role of Semaphorins 837 23. 838 in Glioblastoma. Int J Mol Sci. 2019;20(9). 839 24. McNicholas M, De Cola A, Bashardanesh Z, Foss A, Lloyd CB, Hébert S, et al. A 840 Compendium of Syngeneic, Transplantable Pediatric High-Grade Glioma Models 841 Reveals Subtype-Specific Therapeutic Vulnerabilities. Cancer Discov. 2023;13(7):1592-615. 842 843 25. Hu M, Zhou M, Bao X, Pan D, Jiao M, Liu X, et al. ATM inhibition enhances 844 cancer immunotherapy by promoting mtDNA leakage and cGAS/STING 845 activation. J Clin Invest. 2021;131(3). 846 26. Zhang Q, Green MD, Lang X, Lazarus J, Parsels JD, Wei S, et al. Inhibition of 847 ATM Increases Interferon Signaling and Sensitizes Pancreatic Cancer to Immune 848 Checkpoint Blockade Therapy. Cancer Res. 2019;79(15):3940-51. 849 Tossetta G, Piani F, Borghi C, and Marzioni D. Role of CD93 in Health and 27. 850 Disease. Cells. 2023;12(13). Yao G, Zhao K, Bao K, and Li J. Radiation increases COL1A1, COL3A1, and 851 28. COL1A2 expression in breast cancer. Open Med (Wars). 2022;17(1):329-40. 852 Nasarre C, Koncina E, Labourdette G, Cremel G, Roussel G, Aunis D, et al. 853 29. 854 Neuropilin-2 acts as a modulator of Sema3A-dependent glioma cell migration. 855 Cell Adh Migr. 2009;3(4):383-9.
  - 856 **!!!** INVALID CITATION **!!!** (18). 30.

- 857 31. Katagi H, Louis N, Unruh D, Sasaki T, He X, Zhang A, et al. Radiosensitization
  858 by Histone H3 Demethylase Inhibition in Diffuse Intrinsic Pontine Glioma. *Clin*859 *Cancer Res.* 2019;25(18):5572-83.
- Bashizume R, Andor N, Ihara Y, Lerner R, Gan H, Chen X, et al. Pharmacologic
  inhibition of histone demethylation as a therapy for pediatric brainstem glioma. *Nat Med.* 2014;20(12):1394-6.
- 33. Aoki Y, Hashizume R, Ozawa T, Banerjee A, Prados M, James CD, et al. An
  experimental xenograft mouse model of diffuse pontine glioma designed for
  therapeutic testing. *J Neurooncol.* 2012;108(1):29-35.
- 866 34. Pal S, Kozono D, Yang X, Fendler W, Fitts W, Ni J, et al. Dual HDAC and PI3K
  867 Inhibition Abrogates NFκB- and FOXM1-Mediated DNA Damage Response to
  868 Radiosensitize Pediatric High-Grade Gliomas. *Cancer Res.* 2018;78(14):4007869 21.
- 870 35. Brugarolas J, Chandrasekaran C, Gordon JI, Beach D, Jacks T, and Hannon GJ.
  871 Radiation-induced cell cycle arrest compromised by p21 deficiency. *Nature*.
  872 1995;377(6549):552-7.
- Brady CA, Jiang D, Mello SS, Johnson TM, Jarvis LA, Kozak MM, et al. Distinct
  p53 transcriptional programs dictate acute DNA-damage responses and tumor
  suppression. *Cell.* 2011;145(4):571-83.
- 876 37. Peterson E, Parsels LA, Parsels JD, Zhao X, Castro MG, Lawrence TS, et al.
  877 Inhibition of H3K27M-enhanced ATM signaling increases radiation efficacy in 878 diffuse midline glioma. *bioRxiv.* 2024:2024.11.01.621526.
- 879 38. !!! INVALID CITATION !!! (10, 16, 38).
- 39. Galanos P, Vougas K, Walter D, Polyzos A, Maya-Mendoza A, Haagensen EJ, et
  al. Chronic p53-independent p21 expression causes genomic instability by
  deregulating replication licensing. *Nat Cell Biol.* 2016;18(7):777-89.
- 40. Hukkelhoven E, Liu Y, Yeh N, Ciznadija D, Blain SW, and Koff A. Tyrosine phosphorylation of the p21 cyclin-dependent kinase inhibitor facilitates the development of proneural glioma. *J Biol Chem.* 2012;287(46):38523-30.
- 886 41. Boutelle AM, and Attardi LD. p53 and Tumor Suppression: It Takes a Network.
   887 Trends Cell Biol. 2021;31(4):298-310.
- 42. Ho T, Tan BX, and Lane D. How the Other Half Lives: What p53 Does When It Is Not Being a Transcription Factor. *Int J Mol Sci.* 2019;21(1).
- Weidenhammer LB, Liu HQ, Luo L, Williams NT, Deland K, Kirsch DG, et al.
  Inducing primary brainstem gliomas in genetically engineered mice using
  RCAS/TVA retroviruses and Cre/loxP recombination. *STAR Protocols.*2023;4(1):102094.
- 44. Deng C, Zhang P, Harper JW, Elledge SJ, and Leder P. Mice lacking
  p21CIP1/WAF1 undergo normal development, but are defective in G1 checkpoint
  control. *Cell.* 1995;82(4):675-84.
- Misuraca KL, Barton KL, Chung A, Diaz AK, Conway SJ, Corcoran DL, et al.
  Pax3 expression enhances PDGF-B-induced brainstem gliomagenesis and characterizes a subset of brainstem glioma. *Acta Neuropathol Commun.* 2014;2:134.

- 46. Misuraca KL, Hu G, Barton KL, Chung A, and Becher OJ. A Novel Mouse Model
  of Diffuse Intrinsic Pontine Glioma Initiated in Pax3-Expressing Cells. *Neoplasia*.
  2016;18(1):60-70.
- 47. Amanda J, Robert S, Andrew G, Florian W, Morgane R, Ghezal B, et al. High
  905 resolution mapping of the breast cancer tumor microenvironment using
  906 integrated single cell, spatial and in situ analysis of FFPE tissue. *bioRxiv*.
  907 2022:2022.10.06.510405.
- 48. Yuhan H, Tim S, Madeline K, Saket C, Paul H, Austin H, et al. Dictionary learning
  for integrative, multimodal, and scalable single-cell analysis. *bioRxiv*.
  2022:2022.02.24.481684.
- 49. Hao Y, Stuart T, Kowalski MH, Choudhary S, Hoffman P, Hartman A, et al.
  912 Dictionary learning for integrative, multimodal and scalable single-cell analysis.
  913 Nature Biotechnology. 2024;42(2):293-304.
- 50. Korsunsky I, Millard N, Fan J, Slowikowski K, Zhang F, Wei K, et al. Fast,
  sensitive and accurate integration of single-cell data with Harmony. *Nature Methods.* 2019;16(12):1289-96.
- 51. Cable DM, Murray E, Shanmugam V, Zhang S, Zou LS, Diao M, et al. Cell typespecific inference of differential expression in spatial transcriptomics. *Nat Methods.* 2022;19(9):1076-87.
- 920 52. Palla G, Spitzer H, Klein M, Fischer D, Schaar AC, Kuemmerle LB, et al.
  921 Squidpy: a scalable framework for spatial omics analysis. *Nat Methods.*922 2022;19(2):171-8.
- 53. Duy P, Xiao T, Jun X, Laura FG, Pui Yeng L, Arti R, et al. stLearn: integrating
  spatial location, tissue morphology and gene expression to find cell types, cellcell interactions and spatial trajectories within undissociated tissues. *bioRxiv*.
  2020:2020.05.31.125658.
- 92754.Falick Michaeli T, Sabag O, Fok R, Azria B, Monin J, Nevo Y, et al. Muscle injury928causes long-term changes in stem-cell DNA methylation. Proceedings of the929National Academy of Sciences. 2022;119(52):e2212306119.
- 930
- 931

933 Figure 1

934





936 Figure 1. Atm loss improves radiosensitivity of primary murine diffuse midline



938 mouse genotypes used to generate primary mouse DMGs with p53 loss (Nestin<sup>TVA</sup>;

939	p53 <sup>FL/FL</sup> , nP) and mouse DMGs with p53 loss and H3.3K27M (Nestin <sup>TVA</sup> ; p53 <sup>FL/FL</sup> ;
940	H3f3a <sup>loxP-Stop-loxP-K27M-Tag/+,</sup> , nPH) with or without conditional H3.3K27M allele. Mice also
941	contained one intact and one floxed allele of Atm (Atm <sup>FL/+</sup> , not shown).
942	(A) Dot plot showing time to tumor formation between nPH and nP mice
943	without any statistical significance. Statistical test utilized Welch's t-test.
944	(B) Schematic showing nPHA <sup><math>FL/+</math></sup> (Atm <sup><math>FL/+</math></sup> ) and nPHA <sup><math>FL/FL (Atm<math>FL/FL</math></math></sup> ) within
945	RCAS/TVA retrovirus and conditional H3K27M allele.
946	(C) Time to tumor formation showing no statistical difference between
947	nPHA <sup>FL/FL</sup> and nPHA <sup>FL/+</sup> mice. Statistical test utilized Welch's t-test.
948	(D)Overall survival of between $nPHA_{FL/FL}$ and $nPHA_{FL/+}$ mice following three
949	daily fractions of 10 Gy image-guided focal brain irradiation shows
950	significantly longer median survival in nPHA <sup>FL/FL</sup> (p-value = 0.03) using
951	Mantel Cox (log rank test).
952	(E) Whole mount and H&E slides showing tumor within nP,nPH, nPHA $^{_{\text{FL}^+}}$ and
953	nPHA <sup>F⊔FL</sup> (top to bottom) mice exhibiting hypercellularity and infiltration of
954	normal brain.
955	(F) IHC for HA expression indicating the presence of the PDGF-B HA tag.
956	(G)IHC for p53.
957	(H)IHC for histone 3 lysine 27 trimethylation (H3K27me3).
958	(I) IHC displaying the Ki67 proliferation of tumors.
959	(J) Anti-Flag IHC confirmed the presence of the Flag-HA tag.
960	Scale bar for all H&E and IHC images (C-H) = 50 $\mu$ m. Scale bar for Whole Mount
961	= 200 uM. Scale bar for FLAG IHC infiltrating biology (I) = 500 $\mu$ m



964	Figure 2. Spatial clustering in primary mouse DMGs treated with focal brain
965	irradiation or tumoral <i>Atm</i> deletion.
966	(A) Schematic of DMG-bearing mice with Atm intact or null within the tumors,
967	with and without focal brain irradiation that underwent in situ spatial
968	transcriptomic sequencing (ISS). All mice were Nestin <sup>TVA</sup> ; p53 <sup>FL/FL</sup> ;
969	H3f3a <sup>loxP-Stop-loxP-K27M-Tag/+</sup> with either <i>Atm</i> intact (Atm <sup>FL/+</sup> ) or <i>Atm</i> null
970	(Atm <sup>FL/FL</sup> ) in the tumor.
971	(B) Harmony integration showing clustering of n=4 tumor bearing mice with
972	H3f3a <sup>loxP-Stop-loxP-K27M-Tag/+</sup> with either <i>Atm</i> intact (Atm <sup>FL/+</sup> ) or <i>Atm</i> null
973	(Atm <sup>FL/FL</sup> ) in the tumor.
974	(C)Spatial clustering of cells into n=10 cell archetypes based on label transfer
975	in n=4 tumor-bearing mouse brains ( <i>left</i> ), H&E of whole brain ( <i>middle</i> ),

976	distribution of cells within normal brain, tumor periphery and tumor core
977	annotated in bar graph ( <i>right</i> ). Top row indicates Atm intact with and
978	without irradiation. Bottom row indicates Atm null with and without
979	irradiation. Color legend to the right corresponds to individual cell type
980	noted on bar graph.
981	(D) Spatial identification of tumors by expression of Pdgfra, Olig1, and Olig2
982	within all conditions: Atm intact, Atm intact with irradiation, Atm null, Atm
983	null with irradiation (top to bottom).
984	(E) Spatial identification of p53 loss in all tumor conditions: Atm intact without
985	and with irradiation (top row, left to right). Atm null without and with
986	irradiation (bottom row, left to right).
987	(F) Key differentially expressed genes in Atm-intact neoplastic tumor cells
988	treated with and without focal brain irradiation. Log $_2$ fold change and p-
989	value for all genes in Table S3.
990	(G)Key differentially expressed genes in Atm-null neoplastic tumor cells
991	treated with and without focal brain irradiation. Log $_2$ fold change and p-
992	value for all genes in Table S4.
993	
994	All cells within entire slide were used for distribution of cells identified in Figure 3b.
995	Tumor core and periphery were utilized to identify the key differentially expressed genes
996	(Figure F-G)

998 **Figure 3** 



999

Figure 3. Differentially expressed genes and neighborhood analysis of primary
 mouse DMGs with tumoral *Atm* loss and/or focal irradiation.

(A) Spatial identification of tumors by expression of *Pdgfra*, *Olig1*, and *Olig2*within all conditions: *Atm* intact, *Atm* intact with irradiation, *Atm* null, *Atm*null with irradiation (*top to bottom*).

1005 (B) Spatial identification of p53 loss in all tumor conditions: *Atm* intact without 1006 and with irradiation (*top row, left to right*). *Atm* null without and with

- 1007 irradiation (*bottom row, left to right*).
- 1008 (C)Key differentially expressed genes in *Atm*-intact neoplastic tumor cells
- 1009 treated with and without focal brain irradiation. Log<sub>2</sub> fold change and p-
- 1010 value for all genes in Table S3.

- 1011(D)Key differentially expressed genes in Atm-null neoplastic tumor cells1012treated with and without focal brain irradiation. Log2 fold change and p-1013value for all genes in Table S4.
- 1014 (E) Co-occurrence plot of Atm-intact (nPHA FL/+) tumor showing number
- 1015 compared to distance of various cell types in relation to neoplastic cells.
- 1016 (F) Co-occurrence plot of *Atm*-intact (nPHA<sup>FL/+</sup>) tumor with irradiation showing
- 1017number compared to distance of various cell types in relation to neoplastic1018cells.
- 1019 (G)Co-occurrence plot of *Atm*-null (nPHA<sup>FL/FL</sup>) tumor showing number
- 1020 compared to distance of various cell types in relation to neoplastic cells.
- 1021 (H)Co-occurrence plot of *Atm*-null (nPHA<sup>FL/FL</sup>) tumor with irradiation showing
- 1022 number compared to distance of various cell types in relation to neoplastic
- 1023 cells. Red arrow indicates increased frequency of immune cells compared1024 to neoplastic cells
- 1025

Color Legend for Figure (E-H) on right side panel. APC – Antigen Presenting Cell. OPC
– Oligodendrocyte precursor cell. TLC- T Lymphocyte cell. Neighborhood enrichment
and co-occurrence analysis were conducted on entire slide. All unlabeled cells were
removed for analysis.

1030



ZD1390			-					+			
R (hour) <sup>γ</sup> H2AX	0	1	3	6	24	0	1	3	6	24	
BRCA1	-	-	-	-	-	-	2014	(Het	196	inia	
Rad51	-		-	-	-	- Contraction	-terris		ang.	(469)	
Rad50		-	-	-	-						
GAPDH	-	-	-	-	-	-	-	-	-	-	

С





1035	Figure 4. Pharmacologic inhibition and DNA damage response signaling in
1036	primary mouse DMGs with tumoral Atm loss and/or focal irradiation.
1037	(A) Overall survival of mice bearing SF8628 diffuse midline glioma patient-
1038	derived xenografts were treated with 20mg/kg of AZD1390 for 2 weeks
1039	(ATMi, 5 days a week x 2 weeks) and/or focal brain irradiation (RT, 2 Gy x
1040	3 days a week for 12 Gy total).
1041	(B) Western blot of SF8628 followed by AZD1390 treatment with and without
1042	radiation (0h, 1h, 3h, 6h, 24h)
1043	(C)Representative IHC of Nestin <sup>TVA</sup> p53 <sup>FL/FL</sup> PDGF-B +H3.3K27M +Cre
1044	(PKC) treated with vehicle (V) or ATM inhibitor drug (D), AZD1390 with
1045	and without irradiation of 10 Gy for the following antibodies (Top to
1046	bottom): H&E, pKAP1, total KAP1, yH2AX
1047	(D)PDGF-B +H3.3K27M + p53 <sup>FL/FL</sup> (n=5 per treatment group) stained with
1048	phospho-KAP1 demonstrate increased phosphor-KAP1 expression in
1049	samples treated with one dose of 10Gy and show significance in vehicle
1050	treated samples (PKC + V +/- irradiation (RT)) with a p-value = 0.0079
1051	(Mann-Whitney). This expression is significantly reduced in drug (D) with
1052	irradiation (RT) when compared to vehicle (V) with irradiation (RT) samples
1053	suggesting ATM inhibitor, AZD 1390 sensitizes the DIPG tumor bearing
1054	mice to irradiation (p-value = 0.0159, Mann-Whitney test).
1055	(E) PDGF-B +H3.3K27M + p53 <sup>FL/FL</sup> (n=5 per treatment group) stained with
1056	total KAP1 show unchanged levels across all treatment groups.

1057	(F) PDGF-B +H3.3K27M + p53 <sup>FL/FL</sup> (n=5 per treatment group) stained with
1058	yH2AX demonstrates increased expression in samples treated with
1059	irradiation with one dose of 10Gy when compared to their respective non-
1060	RT treated samples (p-value = 0.0079, Mann Whitney test for vehicle
1061	treated groups and p-value = 0.0317, Mann Whitney test for drug treated
1062	groups).
1063	Scale bar = 50uM on IHC
1064	
1065	
1066	
1067	
1068	



## 1072 Figure 5. Effect of tumor-specific *Atm* loss in primary DMGs in a *Cdkn1a*-null

## 1073 background (p21-/-)

1074	(A) Overview of p21-/- genotypes analyzed
1075	(B) Tumor free survival of nlp21A mice with and without intact Atm using log-
1076	rank test.
1077	(C)Post-focal brain irradiation survival of nIp21A mice with and without intact
1078	Atm indicating a statistically significant survival benefit in nlp21A FL/+ mice
1079	(P-value < 0.05) using log-rank test.
1080	(D)IHC showing p21 expression in nIp21A mouse brains. Nestin <sup>TVA</sup> ;
1081	Pten <sup>FL/FL</sup> ; Atm <sup>FL/+</sup> (nPtenA) tumor-bearing brain generated with identical
1082	RCAS viruses shown as control.
1083	(E) Plot indicating percentage of tumor cells staining positive for p21
1084	compared to total cell count.
1085	(F) IHC with Ki67 showing proliferation for nlp21A <sup>FL/+</sup> and nlp21A <sup>FL/FL</sup>
1086	(G)TUNEL staining of tumor-bearing brains of nIp21A mice with and without
1087	intact Atm in tumors collected one hour post-focal brain irradiation.
1088	(H)Quantification of TUNEL staining in nlp21A $^{FL/+}$ mice (P-value < 0.05) based
1089	on unpaired t-test.
1090	Scale bar = 50uM on IHC.
1091	
1092	





1095 Figure 6. Tumor formation in mice expressing a p53 transactivation domain 1

**mutant.** 

1097	(A) Schematic for conditional p53 transactivation domain 1 mutant, and mice
1098	genotypes for expression of a p53 transactivation domain 1 mutant.
1099	(B) Tumor free survival in the np53 <sup>(25,26)</sup> compared to the np53 control based
1100	on log-rank test.
1101	(C) Time to tumor presentation in the $p53^{25,26/FL}$ group compared to the p53-
1102	FL/FL controls with Wilcoxon test.
1103	(D)IHC for Anti-HA in p53 and p53 <sup>(25,26)</sup> group. Scale bar = 50 uM
1104	(E) IHC for p53 expression in p53 (Scale bar = 100 uM) and p53 $^{(25,26)}$ (50 uM)
1105	group.
1106	
1107	
1108	



#### 1111 Figure 7. Effect of *Atm* loss on survival after fractionated focal brain irradiation in

#### 1112 mouse DMGs expressing a p53 transactivation domain 1 mutant.

- 1113 (A) Schematic showing p53<sup>LSL(25,26)</sup> allele and genotypes with Nestin<sup>TVA</sup>;
- 1114 p53<sup>LSL(25,26)/FL</sup>; Ink4A/ARF<sup>FL/FL</sup> mice with either Atm<sup>FL/FL</sup> or Atm<sup>FL/+</sup>

1115	(B) IHC showing phosphor-Atm in Atm $^{FL/+}$ and Atm $^{FL/FL}$ tumors. Scale bar =
1116	50uM.
1117	(C) IHC showing phosphor-Kap1 expression in $Atm^{FL/+}$ and $Atm^{FL/FL}$ tumors.
1118	Scale bar = 50 uM.
1119	(D)Time to tumor formation in Nestin <sup>TVA</sup> ; p53 <sup>LSL(25,26)/FL</sup> ; Ink4A/ARF <sup>FL/FL</sup> mice
1120	with either Atm <sup>FL/FL</sup> or Atm <sup>FL/+</sup> (dot plot). ns, no statistical significance by
1121	Wilcoxon test.
1122	(E) Overall survival following fractionated brain irradiation in mouse DMGs
1123	expressing a p53 transactivation domain 1 mutation with or with Atm loss,
1124	with P-value based on log rank test.
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## **Tables**

- 1136 Table 1. Summary of the effect of Atm loss on radiation sensitivity in genetically
- 1137 engineered DMG mouse models.

Genotype	Baseline Radiation	Effect of Atm loss on response to
(Nestin <sup>™A</sup> + Cre, Luc, PDGFB)	Therapy sensitivity	radiation therapy
p53 <sup>FL/FL</sup>	Resistant	More sensitive(5)
Ink4A/ARF <sup>FL/FL</sup>	Sensitive	No effect(5)
Pten <sup>FL/FL</sup>	Sensitive	No effect(11)
p53 <sup>FL/FL</sup> ; Ink4A/ARF <sup>FL/FL</sup>	Very Resistant	More sensitive(5)
H3f3a <sup>LSL-K27m</sup> /+; p53 <sup>FL/FL</sup>	Resistant	More sensitive
p21 <sup>-/-</sup> ; Ink4A/ARF <sup>FL/FL</sup>	Sensitive	More resistant
p53 <sup>LSL-25,26/FL</sup> ; Ink4A/ARF <sup>FL/FL</sup>	Resistant	No effect

# 1140 Table 2 - Key resources table

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit polyclonal HA-probe	Santa Cruz	Cat# sc-805
	Biotechnology	RRID: AB_631618
Mouse monoclonal Ser1981	MilliporeSigm	Cat# 05740
phosphorylated ATM	а	RRID: AB_2062670
Rabbit anti-mouse Ser824	ThermoFisher	Cat# A300-767A
phosphorylated Kap1		RRID:AB_2779445
Rabbit polyclonal IgG p21	Santa Cruz	Cat# sc-471
	Biotechnology	RRID:AB_383248
Mouse TotalKap1	Bethyl	Cat#A300-775A
	Laboratories	
Mouse γH2AX	Millipore	Cat# 05-636
Chemicals, peptides, and		
recombinant proteins		
D-luciferin, potassium salt	Gold	Cat# LUCK-1G
	Biotechnology	
Critical commercial assays		
PicoPure DNA extraction kit	ThermoFisher	Cat# KIT0103
Cell Media and supplements		
Dulbecco's Modified Eagle's medium	ThermoFisher	Cat# 11965092
10% Fetal Bovine Serum	ThermoFisher	Cat# A31604-02

Non-Essential Amino Acids	ThermoFisher	Cat#11140-050
Experimental models: Cell lines		
UMNSAH/DF-1 chicken fibroblast cells	ATCC	CRL-12203™
SF8628 Human Cell line DIPG H3.3-	University of	Developed by Dr. Rintaro
К27М	California San	Hashizume at UCSF
	Francisco	
	(UCSF)	
Experimental models:		
Organisms/strains		
Mouse: <i>Nestin<sup>TVA</sup></i> : Tg(NES-	The Jackson	RRID:IMSR_JAX:003529
TVA)J12Ech/J	Laboratory	
Mouse: H3f3a-LSL-K27M-Tag	Lab of Dr.	N/A
	Suzanne	
	Baker	
Mouse: LSL-p53-25,26	Lab of Dr.	N/A
	Laura Attardi	
Mouse: p21-/-	The Jackson	RRID:IMSR_JAX:016565
	Laboratory	
Mouse: <i>p53<sup>fl</sup></i> : B6.129P2-	The Jackson	RRID:IMSR_JAX:008462
Trp53tm1Brn/J	Laboratory	
Mouse: <i>ATM<sup>fl</sup></i> : 129-Atmtm2.1Fwa/J	The Jackson	RRID:IMSR_JAX:021444
	Laboratory	

Mouse: Ink4a/Arf <sup>fl</sup> : Cdkn2atm1Rdp	Mouse	MGI: 1857942
	Genome	
	Informatics	
Athymic Mice (rnu/rnu genotype,	Envigo	Code: 069
BALB/c background)		
Oligonucleotides		
Primers for <i>p53<sup>fl</sup></i> alleles	Weidenhamm	N/A
	er et al., 2023	
Primers for Ink4a/Arf <sup>fl</sup> alleles, see	Weidenhamm	N/A
table S1	er et al., 2023	
Atm recombined probe, 5'-	Weidenhamm	N/A
ACACATGCATGCAGGCAGAGCATC	er et al., 2023	
CCT-3'		
Atm-floxed probe, 5'-	Weidenhamm	N/A
AGCTGTTACTTTTGCGTTTGGTGTG	er et al., 2023	
GCG-3'		
p53 recombined probe, 5'-	Weidenhamm	N/A
CTTGATATCGAATTCCTGCAGCCC	er et al., 2023	
GGG-3'		
p53 floxed probe, 5'-	Weidenhamm	N/A
ATGCTATACGAAGTTATCTGCAGC	er et al., 2023	
CCGG-3'		

Ink4a/Arf recombined probe, 5'-	Weidenhamm	N/A
CATTATACGAAGTTATGGCGCGCC	er et al., 2023	
C-3'		
Ink4a/Arf floxed probe, 5'-	Weidenhamm	N/A
CTCTGAAAACCTCCAGCGTATTCT	er et al., 2023	
GGTA-3'		
Recombinant DNA		
Plasmid: RCAS-Cre	Barton KL, et	N/A
	al., 2013	
Plasmid: RCAS-Luc	Laboratory of	N/A
	Oren Becher	
Plasmid: RCAS-PDGFB	Barton KL, et	N/A
	al., 2013	
Software and algorithms		
ImageJ	NIH	https://imagej.nih.gov/ij/
Prism 7	GraphPad	https://www.graphpad.com/
	Software Inc.	scientific-software/prism/

QuantaSoft	Bio-rad	https://www.bio-
		rad.com/en-us/life-
		science/digital-pcr/qx200-
		droplet-digital-pcr-
		system/quantasoft-
		software-regulatory-
		edition?ID=1864011
Xenium Analyzer	10xGenomics	https://www.10xgenomics.c
		om/instruments/xenium-
		analyzer
Seurat	Satija Lab	https://satijalab.org/seurat/
StLearn	Stlearn	https://stlearn.readthedocs.i
		<u>o/en/latest/</u>
Squidpy	Squidpy	https://squidpy.readthedocs
		<u>.io/en/stable/</u>
STRING	STRING	https://string-db.org
HOMER Motif Analysis	HOMER	http://homer.ucsd.edu/homer
Trimmomatic	Trimmomatic	Version 0.36
Bismark Software	Bismark	Version 0.24.0
Other		
Biological Safety Cabinet	Thermo Fisher	Cat#13-261-222
	Scientific	

IVIS Lumina III In Vivo Imaging	PerkinElmer	Cat#CLS136334
System		
CO <sub>2</sub> Incubators	Thermo Fisher	Cat#4110
	Scientific	
EVOS M7000 Imaging System	Thermo Fisher	Cat#AMF7000
	Scientific	
Hamilton Syringe	Hamilton	Cat#84851
Isoflurane Vaporizer	Kent Scientific	Cat#VetFlo-1205S
Oxygen Concentrator	Fisher	Cat#04-777-122
	Scientific	
Sure-Seal Large Mouse/Rat Induction	World	Cat#EZ-1785
Chamber	Precision	
	Instruments	
Sterile Sleeves	VWR	Cat#414004-510
TUNEL DeadEnd Colorimetric System	Promega	Cat#G7360
AZD1390	Astrazeneca	N/A
Mycoplasma Detection Kit	InvivoGen	Cat#rep-mys-10
Short Tandem Repeat	Promega	Cat#DC2101
	PowerPlex 16	
	HS System	

## 1141 Materials and equipment

### **D-Luciferin Stock Solution**

Reagent	Final concentration	Amount
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D-Luciferin, Potassium	n/a	1 g
Salt		
Dulbecco's Phosphate	1X	66.6 mL
Buffered Saline without		
Ca <sup>2+</sup> and Mg <sup>2+</sup>		
Total	15mg/mL	66.6 mL

1143 Store at -80°C; expires after 1 year

- 1145 Alternatives: D-Luciferin Sodium Salt and L-Luciferin Potassium Salt can be substitute
- 1146 for D-Luciferin, Potassium Salt