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Title: Efficacy and Safety of Dabrafenib in Pediatric Patients with *BRAF* V600 Mutation–
Positive Relapsed or Refractory Low-Grade Glioma: Results from a Phase 1/2a Study
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56 Low-grade gliomas (LGGs) are the most frequently occurring brain tumors in pediatric 57 patients. This study represents the largest clinical trial demonstrating the activity and safety of a BRAF inhibitor (dabrafenib) in pediatric patients with tumors harboring a 58 BRAF V600 driver mutation. Meaningful clinical benefit with dabrafenib was 59 60 demonstrated in pediatric patients with relapsed or refractory BRAF V600-mutant LGG, with an objective response rate of 44% and a 1-year estimated progression-free survival 61 rate of 85% by independent review. The safety profile was favorable and consistent with 62 adverse events observed in adult patients. These safety and preliminary efficacy data 63 demonstrate the potential of dabrafenib as a novel therapy in a pediatric patient 64 population who have few effective treatment options, providing support for further 65 investigation in patients with BRAF V600 mutation-positive tumors, including LGG. 66

67 **Abstract**

68 **Purpose:** Pediatric low-grade glioma (pLGG) is the most prevalent childhood brain

tumor. Patients with *BRAF* V600 mutation–positive pLGG may benefit from treatment

with dabrafenib. Part 2 of a phase 1/2a study, open-label study (NCT01677741)

explores the activity and safety of dabrafenib treatment in these patients.

Experimental Design: Patients aged 1 to <18 years who had *BRAF* V600–mutant solid tumors (\geq 1 evaluable lesion) with recurrent, refractory, or progressive disease after \geq 1 standard therapy were treated with oral dabrafenib 3.0–5.25 mg/kg/day (part 1) or at the recommended phase 2 dose (RP2D; part 2). Primary objectives were to determine the RP2D (part 1, results presented in a companion paper) and assess clinical activity (part 2). Here, we report the clinical activity, including objective response rates (ORRs) using RANO criteria and safety across parts 1 and 2.

79 **Results:** Overall, 32 patients with pLGG were enrolled (part 1, n = 15; part 2, n = 17).

80 Minimum follow-up was 26.2 months. Among all patients, the ORR was 44% (95%

confidence interval [CI] 26–62) by independent review. The 1-year progression-free

survival rate was 85% (95% CI 64–94). Treatment-related adverse events (AEs) were

reported in 29 patients (91%); the most common was fatigue (34%). Grade 3/4

treatment-related AEs were reported in nine patients (28%).

Conclusions: Dabrafenib demonstrated meaningful clinical activity and acceptable
 tolerability in patients with *BRAF* V600–mutant pLGG.

87 Introduction

Glioma accounts for nearly two-third of all pediatric malignant central nervous system 88 tumors and comprises a diverse spectrum of low-grade (eg, pilocytic astrocytoma, 89 diffuse (fibrillary) astrocytoma, and ganglioglioma) and high-grade (eg. anaplastic 90 astrocytoma and glioblastoma) malignancies (1-3). Patients with pediatric low-grade 91 glioma (pLGG) can be cured with complete surgical resection; however, most patients 92 with incompletely resected tumor will require additional treatment (4). Current options 93 for patients whose tumors are not amenable to definitive surgery or whose tumors have 94 recurred or progressed include radiotherapy and chemotherapy, which may provide 3-95 year progression-free survival rates up to approximately 70% but are associated with 96 significant morbidities (eq, cognitive/neurological dysfunction, secondary malignancies, 97 and infertility) (4-7). 98

A greater understanding of the molecular mechanisms underlying pLGG has led to the 99 100 identification of potential targets that can be evaluated for clinical intervention (8). Genetic alterations that result in constitutive activation of the BRAF kinase, including a 101 nucleotide transversion resulting in the substitution of valine (V; most commonly with 102 glutamic acid [E]) at position 600 [ie, V600E point mutation] or a tandem duplication 103 resulting in the fusion of KIAA1549 and BRAF [ie, BRAF:KIAA1549]), have been 104 implicated in the development of pLGG (9-11). In one large series, BRAF V600E 105 mutations were detected in 19% of pLGGs across a broad range of histologies (12). 106 Pleomorphic xanthoastrocytomas and gangliogliomas have been reported to have the 107 108 highest incidence of BRAF V600E mutations among pLGGs, while pilocytic astrocytoma has the highest incidence of BRAF:KIAA1549 gene fusions (13,14). BRAF V600 109

mutation–positive LGG in pediatric patients has been associated with poor responses to
chemotherapy and radiation as well as shorter duration of response and worse
long-term outcomes vs non-*BRAF* V600 LGG (12); thus, these patients represent an
important subpopulation in need of improved treatment options.

Dabrafenib, a potent and selective BRAF V600 inhibitor, has demonstrated clinical 114 benefit in adult patients across a spectrum of BRAF V600-positive solid tumors, and is 115 currently approved as a single agent and in combination with the MEK inhibitor 116 trametinib in patients with unresectable or metastatic BRAF V600E/K-mutant 117 melanoma. Dabrafenib as monotherapy or in combination with trametinib has shown 118 119 activity against melanoma brain metastases in these patients (15). Additionally, dabrafenib plus trametinib is approved in patients with BRAF V600 mutation-positive 120 non-small cell lung cancer (NSCLC) or anaplastic thyroid cancer and as an adjuvant 121 122 therapy in patients with BRAF V600 mutation-positive resectable melanoma. The efficacy of dabrafenib in these adult populations suggests the potential for clinical 123 benefit in pediatric patients with other tumor types driven by the BRAF V600 mutation. 124 including pLGG. 125

Based on the mechanistic rationale, the ability to screen for the relevant driver mutations, and the availability of an age-appropriate formulation, we conducted a twopart, phase 1/2a, single-arm, open-label trial evaluating the safety, tolerability, and clinical activity of dabrafenib in pediatric patients (>12 months) with advanced *BRAF* V600 mutation–positive solid tumors (16). Part 1 was a dose-escalation study to determine the recommended phase 2 dose (RP2D) of dabrafenib in pediatric patients with advanced *BRAF* V600 mutation–positive solid tumors (including LGG) for

subsequent evaluation in part 2 of the study, and is reported in the companion paper to 133 this manuscript (16). Age-dependent dose escalation of dabrafenib in part 1 established 134 the RP2D at 5.25 mg/kg/day in patients <12 years of age and 4.5 mg/kg/day in patients 135 ≥12 years of age, with no dose-limiting toxicities (DLTs) observed (16). Part 2 included 136 four tumor-specific expansion cohorts of patients with BRAF V600 mutation-positive 137 138 tumors (LGG, high-grade glioma [HGG], Langerhans cell histiocytosis, and other tumors such as melanoma and papillary thyroid carcinoma). Here, we report the activity and 139 safety of dabrafenib treatment in pediatric patients with BRAF V600-mutant relapsed or 140 141 refractory LGG.

142 **Patients and Methods**

143 Study design and participants

144 We performed a phase 1/2a multicenter, open-label study in pediatric patients with advanced BRAF V600 mutation-positive solid tumors (NCT01677741). The completed 145 part 1 is detailed in a separate report (16). The dose-escalation decisions were made 146 based on the DLTs observed during the first 28 days, overall toxicity profile, and 147 pharmacokinetics data. Part 2 was an expansion study conducted in four BRAF V600 148 mutation-positive tumor-specific cohorts at 18 sites in eight countries (Australia, 149 Canada, Denmark, France, Germany, Spain, UK, and USA). Patients enrolled in part 2 150 were treated with the established RP2D from part 1. Patients only participated in part 1 151 152 or part 2 of the study. The study will be completed after the last patient has been treated for ≥ 6 months in the last accruing stratum. 153

Eligible patients with LGG were aged 1 to <18 years and had at least one evaluable 154 BRAF V600 mutation-positive tumor according to RANO criteria, determined locally by 155 a Clinical Laboratory Improvement Amendments (CLIA)-approved laboratory (or 156 equivalent), adequate organ function, and a Karnofsky (for ≥ 16 years of age) or Lansky 157 (for <16 years of age) performance score \geq 50. Baseline evaluable (but not measurable) 158 159 disease was required. Patients were required to have recurrent, refractory, or progressive disease following receipt of ≥1 prior standard therapy. Patients could not 160 have received chemotherapy or radiotherapy within 3 weeks (or 6 weeks for 161 162 nitrosoureas or mitomycin C) or an investigational agent within 28 days (or five half-lives or twice the duration of the biological effect) prior to the first dose of dabrafenib. Only in 163 part 2, patients were excluded if they had received previous treatment with a RAF 164 inhibitor (including dabrafenib) or a MEK inhibitor; previous treatment with sorafenib was 165 permitted. Treatment with dabrafenib was continued until disease progression, lack of 166 clinical benefit from continued treatment, unacceptable toxicity, initiation of a new 167 therapy, or consent withdrawal. 168

The study was conducted in accordance with the provisions of the Declaration of
Helsinki, Good Clinical Practice guidelines, and all applicable regulatory requirements.
The protocol was approved by the institutional review board or human research ethics
committee at each study center. Written informed consent (or assent, for ageappropriate patients according to institutional guidelines) was obtained from each
patient, patient's parent, or legal guardian prior to the performance of any study-specific
procedures.

177 Procedures

For part 1 (see companion paper) (16), the initial cohort received a starting dose of 3.0 178 mg/kg/day, as two divided daily doses. Dabrafenib dose levels evaluated in part 1 were 179 3.0, 3.75 (corresponds to the approved adult dose of 150 mg twice daily), 4.5, and 5.25 180 mg/kg. The total daily dose was split evenly into morning and evening doses to follow 181 the twice-daily regimen as administered in adults. Standard dabrafenib capsule 182 strengths (50 mg and 75 mg) were administered to children who were able to swallow 183 capsules. Lower strength capsules (10 mg and 25 mg) and an oral suspension 184 formulation were used for patients who could not swallow the larger capsules. Follow-up 185 186 dermatologic skin assessments were performed every 2–3 months for 6 months following discontinuation of dabrafenib or until initiation of another anticancer therapy. 187 The primary endpoint was objective response rate (ORR) as determined using RANO 188 criteria. Responses were determined both by the investigator and by an independent 189 pediatric neuro-radiologist. Imaging was performed using MRI. Radiographic tumor 190 191 assessment occurred at baseline and every 8 weeks thereafter through 56 weeks; subsequent scans were performed every 12 weeks or as per the standard of care. 192 Clinical activity was assessed based on the Response Assessment in Neuro-Oncology 193 (RANO) criteria (19). Adverse events (AEs) were graded according to the National 194 Cancer Institute Common Terminology Criteria for Adverse Events (NCI-CTCAE) 195 version 4.0 criteria. 196

197 Outcomes

The RP2D of 5.25 mg/kg/day in patients <12 years of age and 4.5 mg/kg/day in patients
≥12 years of age was determined in part 1 of the study and is described in the
companion paper (16). Further evaluation of the safety, tolerability, pharmacokinetics,
and clinical activity of dabrafenib in tumor-specific pediatric populations was performed
in part 2 of the study.

203 Statistical analysis

Data were summarized using descriptive statistical methods. For the part 1 doseescalation phase (16), a minimum of three patients per dose level were evaluated to determine the RP2D. For the part 2 expansion, \geq 10 patients per disease cohort were enrolled. Planned analysis populations for statistical considerations included the alltreated population of patients who received \geq 1 dose of study treatment. The all-treated population was used for the safety and efficacy analyses, and for summarizing the baseline and disease characteristics.

Safety data were based on the initial dose of dabrafenib assigned and were
summarized for AEs and laboratory abnormalities based on the maximum toxicity
grade. The extent of exposure was summarized for all patients. For the results
described here, data were grouped for all pediatric patients with LGG enrolled across
parts 1 and 2.

Efficacy analyses were conducted in the all-treated population. In addition, sensitivity
analysis was performed on the response-evaluable population, which was defined as
the proportion of all-treated patients with a pre-dose and ≥1 post-dose efficacy
assessment. For efficacy analyses, assessments of response were based on the RANO

criteria for all pediatric patients with LGG (17,18). Per RANO criteria, a patient must 220 have measurable disease at baseline in order to qualify for a complete response (CR) 221 or partial response (PR) determination. The ORR was defined as the proportion of all 222 treated patients with the best overall response of CR or PR (95% confidence intervals 223 [CIs] were calculated). Both CR and PR were confirmed by repeat assessments not less 224 225 than 4 weeks after the criteria for response were first met (17). The duration of response was defined as the time from the initial response (CR or PR) to the first documented 226 disease progression or death. 227

228 **Results**

From December 2013 through July 2015, 32 pediatric patients with investigator-229 230 determined BRAF V600 mutation-positive, refractory or recurrent LGG were enrolled across 15 centers in Australia, Canada, France, Spain, and the United States across 231 three dose levels, and were included in the efficacy and safety analyses 232 233 (Supplementary Fig. S1). Fifteen patients were enrolled in part 1 (dose finding) and 17 were enrolled in part 2 (at the RP2D). In part 1, the RP2D was determined (16). Patients 234 enrolled in part 2 were treated at the RP2D, determined as dabrafenib 5.25 mg/kg/day 235 for patients <12 years of age (n = 9) and 4.5 mg/kg/day for patients ≥12 years of age (n236 = 8). There was no correlation between dose and response in this relatively small study. 237 Overall, 24 patients were treated at the age-defined RP2D (seven in part 1 and 17 in 238 part 2). Demographics and baseline disease characteristics of pediatric patients with 239 LGG are summarized in Table 1. The median age was 8.5 years (range 2–17), and 22 240 241 of 32 patients (69%) were <12 years of age. Pilocytic astrocytoma (n = 13; 41%), ganglioglioma (n = 7; 22%), and pleomorphic xanthoastrocytoma (n = 3; 9%) were the 242

most common LGG diagnoses; other tumors are reported in Table 1. All 32 patients had 243 a documented progression. The median time since initial LGG diagnosis (in 26 patients 244 with available data) was 32 months (range 6–190). Ten of 32 patients did not have 245 progressive disease within the previous 4 months and were eligible for enrolment with 246 indolent disease as per the phase I part of the study. Twenty-two patients (69%) were 247 248 initially diagnosed with grade 1 disease, nine patients (28%) with grade 2 disease, and one patient had undetermined grade 1 or grade 2 disease. Most patients had a good 249 Karnofsky/Lansky performance status; only 13% of the patients had a performance 250 251 status below 80 at baseline. Prior treatments were predominantly chemotherapy (n =28; 88%) and radiotherapy (n = 6; 19%). Five patients had a best overall response of 252 PR to the most recent therapy received before starting dabrafenib treatment; no prior 253 CRs were reported. 254

As of the data cutoff date (September 12, 2017) with a minimum follow-up of 26.2 months, 15 patients (47%) were continuing treatment with dabrafenib (Table 1).

257 The study set no minimum treatment duration and the most common reason for treatment discontinuation was physician and/or parent decision. Four patients (13%) 258 discontinued due to disease progression, including one patient who discontinued at 259 week 8 of the treatment and underwent subsequent therapy, but later died due to 260 disease progression. This patient was enrolled into this study 11 years after the initial 261 diagnosis of LGG (pilocytic astrocytoma). At autopsy, this patient's tumor was found to 262 have transformed into a BRAF V600-mutated, PDGFRA-amplified glioblastoma (World 263 264 Health Organization grade 4). Two patients (6%) discontinued dabrafenib due to AEs. The median duration of dabrafenib exposure was 108 weeks (range <1-185; Table 1 265

and Fig. 1A), and 17 patients (53%) were on treatment for >2 years. Ten patients (31%)
had dose reductions and/or interruptions.

The confirmed ORR with dabrafenib by independent radiological review was 44% 268 (14/32, 95% CI 26–62), and included one patient with CR and 13 with PR (Table 2). 269 Five of these 32 patients were not evaluable for CR or PR per RANO criteria due to 270 non-measurable but evaluable disease at study entry; these five patients were 271 evaluable for and met the definition of stable disease (SD). An example of PR (ongoing 272 at data cutoff) achieved after 8 weeks of dabrafenib therapy in an 11-year-old male 273 patient with BRAF V600-mutant ganglioglioma is shown by MRI scan (Fig. 2). Eight of 274 275 32 patients (25%) had a first response within 4 months of dabrafenib initiation. The median time to first response was 3.8 months (range 1.7–24.0; Fig. 1A). The median 276 duration of response was 26 months (95% CI 9-not estimable). Eight out of 14 patients 277 278 had an ongoing response at the time of data cutoff, and six out of 14 patients who relapsed had a duration of response of more than 2 years to dabrafenib. The disease 279 control rate (CR+PR+SD) by independent review was 78% (95% CI 60–91). Among the 280 27 patients with measurable disease as determined by independent radiological review, 281 19 (73%) had at least one occurrence of a maximum reduction in lesion size of at least 282 50% from baseline (Fig. 1B). 283

The disease control rate (CR+PR+SD) by investigator assessment was 88% (95% Cl 71–97). Among the 31 patients with measurable lesions as per investigator assessment, 22 (71%) achieved a maximum reduction in lesion size of at least 50% from baseline (Supplementary Fig. S2). Eleven of the patients with a best overall response of SD by independent review had significant tumor reductions that were categorized by

investigators as PRs, accounting for most of the observed discordance between theindependent and investigator assessment of response.

A total of 11 disease progression events were determined by independent review, three 291 of which occurred after ending dabrafenib treatment. Five of the eight patients 292 determined as disease progression on treatment with independent review were 293 continuing treatment at the data cutoff; these patients did not have progression per 294 investigator assessment. The median progression-free survival (independent review) 295 was 35.0 months (interguartile range 12.9-not estimable), and the Kaplan-Meier 296 estimate of the proportion of patients with progression-free survival at 1 year of 297 298 dabrafenib treatment was 85% (95% CI 64–94; Fig. 3). One survival event occurred after treatment discontinuation. 299

Treatment-related AEs of any grade occurred in 29 patients (91%); the most common 300 were fatique (34%), rash (31%), dry skin (28%), pyrexia (28%), and maculopapular rash 301 302 (28%; Table 3). Grade 3/4 treatment-related AEs were reported in nine patients (28%) and included maculopapular rash (n = 3), arthralgia, lymphocytopenia, increased 303 weight, thrombocytopenia, back pain, increased blood alkaline phosphatase, 304 hypotension, neutropenia, and migraine (n = 1 each). In this pediatric population, there 305 were no cases of squamous cell carcinoma of the skin or keratoacanthoma, as have 306 been reported commonly in adult patients treated with dabrafenib. Note that new or 307 increased size of melanocytic nevi was reported in 8 of 32 patients (25%), all grade 1 or 308 2. AEs were well managed by supportive care, dose interruption, and dose reduction. 309 310 Ten patients had AEs that led to dose interruptions and/or reductions. AEs of allergic reaction/hypersensitivity (n = 1) and hip pain/arthralgia with erythema nodosum (n = 1)311

led to treatment discontinuation in two patients (6%). Treatment-related serious AEs of any grade occurred in five patients (16%) and were reported as grade 3/4 in three patients (9%), which included arthralgia, disseminated intravascular coagulation with hypotension, and maculopapular rash (n = 1 each). No treatment-related deaths occurred in the study; one patient died due to disease progression 2 weeks after discontinuing the therapy.

318 **Discussion**

This study represents the largest report of successful outcomes from a clinical trial of a 319 BRAF V600-targeted therapy in a pediatric population selected based on a specific 320 driver mutation. Previous reports have been limited to case study observations (19-22) 321 322 and the report of an adult glioma subset from the vemurafenib basket trial that included 9 adult patients with BRAF V600-mutant LGG (23). In this study, we demonstrated 323 clinical activity of dabrafenib in pediatric patients with BRAF V600-mutant relapsed or 324 325 refractory LGG in a clinical trial setting; a high proportion of these patients had a radiographic response. Dabrafenib was tolerable and demonstrated a manageable 326 safety profile with a minimum follow-up of >2 years. These results in pediatric patients 327 add to those previously reported for adult patients with other BRAF V600 mutation-328 positive tumors, including melanoma, NSCLC, anaplastic thyroid cancer, and gliomas 329 (24-27). Taken together, these data clearly demonstrate the clinical benefit of targeting 330 the V600 mutation with dabrafenib in pediatric patients with relapsed refractory BRAF 331 V600 mutation-positive LGG. 332

Current treatment options for pediatric patients with progressive or recurrent LGG are 333 limited to radiotherapy and chemotherapy. These are associated with various clinically 334 significant long-term adverse effects, including risk of secondary malignancies, cognitive 335 impairment, hormonal deficiencies, vasculopathies and infertility (5), which are of 336 particular concern in a pediatric patient population. Standard chemotherapy treatments 337 338 appear to have worse efficacy in patients with BRAF V600-mutant LGG than in those with non-BRAF V600 LGG (13), including a 10-year progression-free survival of 27% vs 339 60%. The apparent ORR (CR+PR at the 6-month milestone) observed in historical 340 341 cohorts of this population treated with chemotherapy is approximately 10% (13). In this study, an ORR of 44% and a 1-year progression-free survival rate of 85% were reported 342 by independent review using the RANO criteria. Approximately half of responders by 343 independent review had an ongoing response at the time of data cutoff. Notably, among 344 patients assessed by independent review, only two had a best response of progressive 345 disease. 346

The most common reason for discontinuation of treatment in this study was physician and/or parent decision with the majority having at least one year of treatment. It is likely that the typical duration of standard chemotherapy for pLGG of 12-24 months had an impact on the decision to stop therapy in patients with SD or better. Further data generation is needed to determine the optimal duration of treatment, and if patients can be retreated successfully. Anecdotal reports from investigators of this clinical trial, showed that retreatment with dabrafenib can result in tumor control.

354 Observations from experienced neuro-oncologists and neuro-radiologists involved in the 355 study suggest that *BRAF* V600–mutant LGG tumors may have some unique

characteristics on MRI imaging, which can prove challenging in recording tumor 356 response consistently and accurately as illustrated by the disconcordance seen 357 between the local and central independent review in this study. Generally, LGG tumors 358 are monitored for response by T2/FLAIR MRI sequences, and these T2/FLAIR images 359 are recommended for the observation of tumor size changes in LGG assessment (28). 360 361 However, some of the LGG tumors on this study appeared more like typical HGG tumors and displayed enhancement in post-gadolinium T1-weighted images ("T1 362 enhancement"). Further, this enhancement can decrease quickly upon initiation of 363 364 treatment with dabrafenib and occurs before changes in tumor size are observed on T2/FLAIR sequences. There are a few reports of rapid increase in T1 enhancement 365 upon elective cessation of treatment, with subsequent decrease upon rechallenge with 366 dabrafenib. The mechanism of this rapid change in T1 enhancement is not well 367 understood, nor is its biologic significance. Until more experience is gained, caution 368 should be exercised, as these rapid changes in the size of apparent T1-enhancing 369 BRAF V600-mutant LGG tumors on starting or stopping dabrafenib treatment may not 370 accurately represent true changes in tumor size. 371

The safety profile of dabrafenib in pediatric patients with LGG was manageable and was consistent with that observed in adult patients across other indications, except for the absence of observations of squamous cell carcinoma (as of April 2019). Similar to the observations in patients with melanoma and NSCLC (24,25), fatigue and pyrexia were among the most common treatment-related AEs observed in pediatric patients with LGG treated with dabrafenib; these AEs and others were manageable and did not lead to discontinuation.

Recent research from several different groups led to the identification of multiple 379 molecular aberrations in pLGG tumors (20,21,29), including a BRAF V600-mutation 380 rate of 15%–20% across LGG histologies (12,13). A recent study of gene expression 381 profiling of 151 LGG biopsies from pediatric patients demonstrated that BRAF gene 382 383 abnormalities were observed across a variety of histological subtypes, with BRAF: KIAA1549 fusions occurring most frequently in pilocytic astrocytomas and BRAF 384 385 V600 point mutations occurring most frequently in pleomorphic xanthoastrocytomas and 386 gangliogliomas (29). Taken together with the results of the current report, these data 387 suggest that only specific patient subgroups may be more likely to derive benefit from dabrafenib therapy. It is important to note that patients with the BRAF gene fusion or 388 389 duplications should not receive BRAF inhibitor therapy, as preclinical data demonstrate that BRAF inhibition activates the MAPK signaling pathway in cells with wild-type BRAF 390 at V600 (30). Furthermore, a phase 2 study of the multikinase inhibitor sorafenib, which 391 targets BRAF, VEGFR, PDGFR, and c-kit, in pediatric patients with recurrent low-grade 392 astrocytomas—some of whom harbored BRAF duplications—indicated that sorafenib 393 treatment was associated with accelerated tumor growth (31). The authors concluded 394 that sorafenib may have led to paradoxical ERK activation that caused rapid tumor 395 progression. These data underscore the importance of detailed molecular profiling prior 396 397 to treatment with BRAF inhibitors in pLGG patients.

The results presented here provide additional rationale for increased efforts worldwide to molecularly characterize newly diagnosed tumors in children, with the intent to identify targetable aberrations for each patient. Indeed, efforts ongoing at centers around the world, including INFORM (German Cancer Research Center), MAPPYACTS

(NCT02613962; Gustave Roussy, France), PEDS-MIONCOSEQ (University of 402 Michigan), BASIC3 (Baylor College of Medicine), iCat (NCT01853345; Dana-Farber 403 Cancer Institute), SMPaeds - Stratified Medicine Pediatrics (ISRCTN21731605; UK) 404 and the Pediatric MATCH program (US NCI) among others, are showing good promise 405 in the ability to provide targeted therapies for pediatric cancer patients who may 406 407 otherwise have limited treatment options (32-35). Although the tumors of patients enrolled in this study were already determined to harbor the BRAF V600 mutation, it is 408 apparent that broad molecular profiling of LGG tumors (as well as other pediatric tumor 409 410 types) at diagnosis may lead to enhanced treatment options for an increasing number of pediatric cancer patients (36). 411

Overall, these results demonstrate a distinct clinical benefit and favorable tolerability for 412 dabrafenib in pediatric patients with BRAF V600 mutation-positive relapsed or 413 414 refractory LGG and provide support for further evaluation in this population. Determination of optimal duration of treatment and biological correlates of response to 415 dabrafenib remains important areas of study. As has been demonstrated in several 416 BRAF V600–mutant adult tumor types, the addition of trametinib to dabrafenib therapy 417 may provide improved outcomes in pediatric patients with BRAF V600-mutant LGG. A 418 phase 2 study of dabrafenib in combination with the MEK inhibitor trametinib in pediatric 419 patients with BRAF V600 mutation-positive newly diagnosed LGG or recurrent HGG 420 (NCT02684058) is ongoing. 421

423 Authors'	Contributions
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435	Administrative, technical, or material support (i.e., reporting or organizing data,
436	constructing databases): L. Tseng
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Table 1. Patient demographics, baseline characteristic	s, prior treatments, disposition, and dabrafenib exposure ^a
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	Part 1			Part 2	All patients	All nationts
	Dabrafenib	Dabrafenib	Dabrafenib	Dabrafenib	treated with	with LGG
	3.75 mg/kg	4.5 mg/kg	5.25 mg/kg	RP2D	dabrafenib at	with LGG
	(<i>n</i> = 3)	(<i>n</i> = 6)	(<i>n</i> = 6)	(<i>n</i> = 17)	RP2D	(N - 22)
Characteristic					(<i>n</i> = 24)	(14 = 32)
Median age (range), years	8 (4–13)	8.5 (2–16)	7.5 (3–11)	11 (2–17)	9.5 (2–17)	8.5 (2–17)
<2 years, <i>n</i>	0	0	0	0	0	0
2 to <6 years, <i>n</i>	1	2	2	5	7	10
6 to <12 years, <i>n</i>	1	3	4	4	8	12
12 to ≤18 years, <i>n</i>	1	1	0	8	9	10
Sex, <i>n</i> (%)						
Male	2 (67)	5 (83)	3 (50)	9 (53)	13 (54)	19 (59)
Female	1 (33)	1 (17)	3 (50)	8 (47)	11 (46)	13 (41)
Race, <i>n</i> (%)						
White	3 (100)	5 (83)	6 (100)	13 (76)	19 (79)	27 (84)
Black	0	1 (17)	0	2 (12)	3 (13)	3 (9)
Asian	0	0	0	2 (12)	2 (8)	2 (6)
Performance status, n (%) ^b						
100	2 (67)	3 (50)	3 (50)	9 (53)	12 (50)	17 (53)
80–90	1 (33)	1 (17)	2 (33)	7 (41)	10 (42)	11 (34)
<80	0	2 (33)	1 (17)	1 (6)	2 (8)	4 (13)
Histology at initial diagnosis, n (%)						
Pilocytic astrocytoma	1 (33)	3 (50)	1 (17)	8 (47)	10 (42)	13 (41)

Ganglioglioma	0	1 (17)	1 (17)	5 (29)	6 (25)	7 (22)
Pleomorphic xanthoastrocytoma	0	0	1 (17)	2 (12)	3 (13)	3 (9)
Pilomyxoid astrocytoma	1 (33)	0	0	1 (6)	1 (4)	2 (6)
Other ^c	1 (33)	2 (33)	3 (50)	1 (6)	4 (17)	7 (22)
Histological grade at initial diagnosis, n						
(%) ^d	2 (67)	4 (67)	4 (67)	12 (71)	16 (67)	22 (69)
Grade I	1 (33)	2 (33)	2 (33)	4 (24)	7 (29)	9 (28)
Grade II						
Median time since initial diagnosis						
(range), months	36 (32–39)	15 (11–90)	39 (18–83)	26 (6–190)	31 (6–190)	32 (6–190)
Prior treatments, $n(\%)^{e}$						
Chemotherapy	3 (100)	5 (83)	6 (100)	14 (82)	20 (83)	28 (88)
Radiotherapy	1 (33)	1 (17)	1 (17)	3 (18)	5 (21)	6 (19)
Small-molecule therapy	0	0	1 (17)	1 (6)	2 (8)	2 (6)
Immunotherapy	0	0	0	1 (6)	1 (4)	1 (3)
Other	0	0	0	3 (18)	3 (13)	3 (9)
Median time from last recurrence to	NA	NA	0.8 (0.2–1.3)	1.1 (0.1–81.5)	1.1 (0.1–81.5)	1.1 (0.1–81.5)
dabrafenib start (range), months ^f						
Median time from last progression to	7.6 (0.5–14.7)	0.8 (0.5–1.1)	1.8 (0.2–26.2)	1.6 (0.1–10.3)	1.5 (0.1–26.2)	1.1 (0.1–26.2)
dabrafenib start (range), months ^g						
Continuing treatment, n (%)	2 (67)	3 (50)	2 (33)	8 (47)	10 (42)	15 (47)
Discontinued treatment, n (%)	1 (33)	3 (50)	4 (67)	9 (53)	14 (58)	17 (53)
Reasons for discontinuation						
Investigator discretion	1 (33)	1 (17)	4 (67)	5 (29)	10 (42)	11 (34)

Disease progression	0	2 (33)	0	2 (12)	2 (8)	4 (13)
Adverse event	0	0	0	2 (12)	2 (8)	2 (6)
Median duration of exposure to dabrafenib (range), weeks	157 (62–159)	120 (8–185)	96 (25–152)	105 (<1–149)	104 (<1–152)	108 (<1–185)
Patients with dose reductions and/or interruptions, <i>n</i> (%)	1 (33)	3 (50)	1 (17)	5 (29)	6 (25)	10 (31)

^aAs of data cutoff (September 12, 2017); ^bUsing Karnofsky (\geq 16 years of age; *n* = 28) or Lansky (<16 years of age; *n* = 4) performance status, as appropriate; ^cDesmoplastic neuroepithelial neoplasm, cervicomedullary tumor, glioneuronal brain stem tumor, posterior fossa brain tumor, optic pathway glioma, gliomatosis cerebri, and other low-grade glioma; ^dOne patient had missing data for disease grade at initial diagnosis but was confirmed to have LGG; ^ePatients may have had multiple therapies and prior therapy type was undetermined in 2 patients; best response to last therapy received included five patients with a partial response, 13 patients with stable disease, and nine patients with progressive disease (response to last therapy was undetermined in five patients); ^fIn 11 patients with recurrence; ^gIn 25 patients with disease progression.

Table 2. Dabrafenib efficacy

		Part 1			All patients	All nationts
	Dabrafenib	Dabrafenib	Dabrafenib	Dabrafenib	treated with	with LCC
	3.75 mg/kg	4.5 mg/kg	5.25 mg/kg	RP2D	dabrafenib at	with LGG
	(<i>n</i> = 3)	(<i>n</i> = 6)	(<i>n</i> = 6)	(<i>n</i> = 17)	RP2D	(N - 22)
Characteristic					(<i>n</i> = 24)	(N = 32)
Independent review ^a						
Best overall response, <i>n</i> (%)						
Complete response	0	1 (17)	0	0	0	1 (3)
Partial response	2 (67)	2 (33)	2 (33)	7 (41)	9 (38)	13 (41)
Stable disease ^b	1 (33)	3 (50)	4 (67)	8 (47)	13 (54)	16 (50)
Progressive disease	0	0	0	2 (12)	2 (8)	2 (6)
Objective response, n (%)	2 (67)	3 (50)	2 (33)	7 (41)	9 (38)	14 (44)
[95% CI]	[9–99]	[12–88]	[4–78]	[18–67]	[19–59]	[26–62]
Median duration of response						
(range), months	-	-	-	-	11.0 (3.7–14.5)	11.0 (7.4–14.5)
Disease control, <i>n</i> (%)	3 (100)	5 (83)	5 (83)	12 (71)	18 (75)	25 (78)
[95% CI]	[29–100]	[36–100]	[36–100]	[44–90]	[53–90]	[60–91]
Median progression-free	25 (15 NE)		12 (12 NE)		14 (12 NE)	25 (12 NE)
survival (95% CI), months ^c	35 (15-INE)		13 (13-INE)		14 (13-INE)	55 (13-INE)
1-year progression-free	100 (100-100)	80 (20-97)	100 (100-100)	78 (46-92)	79 (53-92)	85 (64-94)
survival rate (95% CI), % ^c	100 (100-100)	00 (20-37)	100 (100-100)	10 (1 0-32)	19 (00-92)	00 (04-04)

Abbreviation: NE, not estimable.

^aUsing RANO criteria; ^bIncludes five patients with independent review of stable disease but lacking any confirmation scan results; ^cKaplan-Meier estimate.

Table 3.	Safety sumr	nary and treati	ment-related AEs
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	All patients wit	h LGG (<i>N</i> = 32)
	All grade	Grade 3/4
Patients with a treatment-related AE, n (%)	29 (91)	9 (28)
Treatment-related AEs (in >20% of patients), <i>n</i> (%)		
Fatigue	11 (34)	0
Rash	10 (31)	0
Dry skin	9 (28)	0
Pyrexia	9 (28)	0
Rash maculopapular	9 (28)	3 (9)
Arthralgia	8 (25)	1 (3)
Headache	7 (22)	0
Vomiting	7 (22)	0
AEs leading to discontinuation, n (%)	2 (6)	2 (6)
Treatment-related deaths, n (%)	0	0
Patients with a treatment-related serious AE, n (%)	5 (16)	3 (9)
Treatment-related serious AEs, <i>n</i> (%)		
Arthralgia	1 (3)	1 (3)
Disseminated intravascular coagulation	1 (3)	1 (3)
Ejection fraction decreased	1 (3)	0
Febrile neutropenia	1 (3)	0
Hypotension	1 (3)	1 (3)
Pyrexia	1 (3)	0
Rash maculopapular	1 (3)	1 (3)

Figure Legends

Figure 1. Dabrafenib treatment duration and best response.

- **A.** Duration of exposure to dabrafenib analyzed by best overall response assessed by independent review using the RANO criteria.
- **B.** Best reduction in tumor size analyzed by best overall response, assessed by independent review using the RANO criteria, for the subset of patients with measurable disease. Dashed line represents a 50% reduction from baseline, which corresponds to the threshold for partial response per the RANO criteria.

^aIncludes only patients with measurable disease and \geq 1 post-baseline evaluation. Five of these patients had the best overall response as stable disease, with no confirmation from the scan results; one patient was not evaluable.

Figure 2. MRI of a partial response (ongoing) achieved after 8 weeks of dabrafenib therapy in an 11-year-old male patient with *BRAF* V600–mutant ganglioglioma determined using coronal T1 post-gadolinium contrast sequence.

Figure 3. Kaplan-Meier progression-free survival.

Kaplan-Meier estimates of progression-free survival. Eleven disease progression events occurred; eight were on-treatment and three were off-treatment. Tumor assessments were by independent review using the RANO criteria.

Figure 1



Figure 2





Pre-treatment

Dabrafenib week 8

Figure 3

