Treatment of adult brainstem glioma with combined antiangiogenic therapy: a case report and literature review

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Abstract: Adult brainstem gliomas belong to a rare and heterogeneous group of brain tumors. The overall prognosis is poor; therapeutic options are limited, given the resistance to radiotherapy and the unclear role of chemotherapy/antiangiogenic therapy. Apatinib, a tyrosine kinase inhibitor that selectively inhibits the vascular endothelial growth factor receptor and mildly inhibits c-Kit, PDGFR-β, RET, and c-SRC, has been reported to show efficacy among some patients with malignant supratentorial gliomas. However, its effect on brainstem glioma has not been reported so far. Herein, a 66-year-old man with brainstem anaplastic astrocytoma isocitrate dehydrogenase (IDH) wild type was treated initially with combined radiotherapy, temozolomide, and apatinib. The patient achieved a complete response by MRI and continues to have an ongoing progression-free survival of over 8 months. To our knowledge, this is the first case report using apatinib to treat brainstem IDH wild-type anaplastic astrocytoma, displaying an excellent outcome. We also summarize cases of adult brainstem glioma treated with antiangiogenic therapy. Experiences using various regimens may improve understanding of this rare disease, and thus help physicians to seek more effective treatments for these patients.

Keywords: VEGFR, antiangiogenic therapy, apatinib, brainstem glioma

Introduction

Adult brainstem gliomas (BSGs) are uncommon and constitute only 2% of all adult brain tumors; as such, they are a poorly characterized cohort with variable prognosis.¹ ² ³ The available treatment strategies for BSG include surgery, radiotherapy, chemotherapy, and/or biological treatment. However, owing to the rarity of BSG in adults, no large clinical trials have been conducted, and as such, standard strategies are still controversial. Radiotherapy is considered the cornerstone of adult BSG, while the role of chemotherapeutic and biological agents has not been well studied. Herein, we report an adult with isocitrate dehydrogenase (IDH) wild-type anaplastic astrocytoma (WHO grade III) who experienced a remarkable response to the combination of external-beam fractionated radiation, concomitant temozolomide (TMZ), and the small molecule tyrosine kinase inhibitor (TKI) apatinib. Additionally, cases of adult BSG treated with antiangiogenic therapy are discussed in a review of the literature.

Ethics statement

Ethics approval of this case report was granted by the Institutional Ethics Review Board of Renmin Hospital of Wuhan University. Written informed consent was obtained from the patient for publication of this case report and any accompanying images. Institutional approval was not required to publish this manuscript.
Case description
Clinical presentation
A 66-year-old male was referred to the Department of Neurosurgery for 2 months of dizziness, weakness, and gait ataxia. Physical examination showed dysarthria and bilateral lower limb weakness with grade 2 muscle strength; the patient could not complete the heel–knee–tibia and left finger–nose–finger tests. The Karnofsky performance status (KPS) at diagnosis was 30.

T2-weighted (T2W) magnetic resonance imaging (MRI) showed a hyperintense signal alteration in the right side of the pons. Contrast-enhanced T1-weighted (T1W) MRI showed increased enhancement within the tumor. Three-dimensional arterial spin labeling (3D-ASL) showed increased cerebral blood flow (CBF) value in the right pons (Figure 1). A BSG was thus diagnosed radiologically.

Enhancing BSGs have a wide range of differential diagnoses on MRI, such as ependymoma, hemangioma, and metastasis. In order to rule out these image mimics, microsurgery was performed to obtain histologic diagnosis. The tumor was depicted by the surgeon as located in the pontine gray-white areas, ill-defined, and with a rich blood supply. Since resection was not possible, tissues were obtained for biopsy. No severe perioperative complications were observed. After operation, there was no improvement in neurological symptoms.

Histologic examination
H&E staining showed features of anaplastic astrocytoma (WHO grade III) with increased cellularity, nuclear atypia, and mitotic activity. Immunohistochemical staining for GFAP and oligodendrocyte transcription factor were positive, further supporting the diagnosis. High proliferative activity was present, with 60% positive cells staining for Ki-67. Vascular endothelial growth factor receptor-1 (VEGFR-1) and vascular endothelial growth factor receptor-2 (VEGFR-2) were positive; the tumor was negative for stem cell factor receptor (c-kit, CD117) and H3K27M (Figure 2).

The MGMT promoter region was methylated, and loss of heterozygosity was present in chromosome 1p and absent in chromosome 19q; no mutations were found in IDH1R132, IDH2R172, TERTC228T, TERTC250T, BRAFV600E, H3F3AK27M, and HIST1H3BK27M (Table 1).

Treatment
More than 1 month after microsurgery, the patient was referred for further treatment. Considering the poor location, history of rapid progression, as well as surgical and histopathologic findings, we decided to administer radiotherapy with concomitant chemotherapy (TMZ) and antiangiogenic therapy (apatinib). The patient’s hematologic, hepatic, renal, and cardiac functions were all within normal limits before commencing treatment.

Figure 1 Brain MRI scans.
Notes: MRI before treatment (A–D): (A) axial T2W shows a hyperintense signal alteration in the right side of the pons; (B) axial T1W post-contrast; (C) sagittal T1W post-contrast; and (B and C) increased enhancement within the tumor. (D) 3D-ASL CBF shows increased CBF value in the right pons. MRI after treatment (E–H): (E) axial T2W shows a decreased hyperintense signal in the right side of the pons compared to pretreatment; (F) axial T1W post-contrast; (G) sagittal T1W post-contrast; and (F and G) disappeared right pontine lesion. (H) 3D-ASL CBF shows decreased CBF value in the right pons compared to pretreatment.
Abbreviations: 3D-ASL, three-dimensional arterial spin labelling; CBF, cerebral blood flow; T1W, T1 weighted; T2W, T2 weighted.
From August 29 to October 11, 2017, the patient was treated with intensity-modulated radiation therapy (IMRT) and concomitant TMZ chemotherapy and apatinib-targeted therapy. The dose of IMRT was 56 Gy in 30 fractions, TMZ was 75 mg/m$^2$/day, and apatinib was 500 mg/day. Mannitol and nutritional support were administered during the treatment period, and headaches disappeared after ten fractions, on September 8.

During the period of combined-modality therapy, the patient experienced grade 1 leukopenia and thrombocytopenia, without other adverse events as defined by the National Cancer Institute Common Terminology Criteria for Adverse Events (version 4.0). When the combined-modality therapy got completed, on October 11, physical examination showed grade 4 muscle strength of bilateral lower limbs, which was greatly improved over that prior to treatment. However, his dysarthria persisted. After the concurrent therapy, we recommended continuation of apatinib (500 mg daily) following treatment. Unfortunately, 2 weeks later (October 25), apatinib was discontinued due to financial reasons. The patient began to receive adjuvant TMZ on November 18, according to the 5-day schedule, every 28 days. The dose of TMZ was 150–200 mg/m$^2$, and chemotherapy was continued if no hematologic toxicities occurred at the beginning of the next cycle, and a total of six cycles was planned.

The patient returned for reevaluation (after concurrent therapy) on November 16. At that time, the patient could walk slowly with assistance and dysarthria had ceased. The KPS was 60. T2W MRI showed a decreased hyperintense signal in the right side of the pons compared to pretreatment. T1W post-contrast MRI showed that the right pontine lesion had disappeared. 3D-ASL showed decreased CBF value in the right pons compared to pretreatment (Figure 1E–H). At the time of last follow-up, the patient was independent in all daily activities. MRI at the time (performed at outside facility) showed no signs of tumor recurrence. According to the Response Assessment in Neuro-Oncology criteria, he achieved a complete response (CR).^5

### Table 1 Results of molecular tests

| Item                                   | Result       |
|----------------------------------------|--------------|
| MGMT promoter methylation              | Positive     |
| Loss of heterozygosity of chromosome 1p| Positive     |
| Loss of heterozygosity of chromosome 19q| Negative     |
| IDH1 R132 mutation                     | Negative     |
| IDH2 R172 mutation                     | Negative     |
| TERT C228T mutation                    | Negative     |
| TERT C250T mutation                    | Negative     |
| BRAF V600E mutation                    | Negative     |
| H3F3A K27M mutation                    | Negative     |
| HIST1H3B K27M mutation                 | Negative     |

Abbreviation: VEGFR, vascular endothelial growth factor receptor.
**Discussion**

BSG in adults represents a heterogeneous group of tumors whose presentation and clinical course depend on pathologic and molecular features. Owing to the location of these tumors, biopsies are not routinely performed. An MRI-based radiological classification has been proposed to establish treatment strategies and to determine outcomes for 1) diffuse intrinsic low-grade gliomas, 2) enhancing malignant gliomas, 3) focal tectal gliomas, and 4) exophytic gliomas/other subtypes.²,⁶,⁷

Among the different subtypes above, enhancing malignant gliomas (WHO grade III–IV) carry the poorest prognosis.⁸,⁹ They (accounting for up to 30% of adult BSGs) comprise the majority of BSG in adults. Radiotherapy is reported to have limited efficacy, and only a small proportion of patients (<13%) exhibit clinical and radiographic improvement after radiation.² Even higher doses (up to 72 Gy) fail to improve efficacy and carry a higher risk of radiation-related complications.¹ The role of chemotherapy in the treatment of enhancing malignant adult BSG remains unclear; TMZ or combined procarbazine, lomustine, and vincristine added to radiation does lead to clinical improvement in some patients, but grade 3 and 4 myelosuppression may also occur.¹⁰

We summarize published reports of histologically diagnosed BSG in Table 2.⁴,⁸,¹¹–¹⁵ In these reports, the median overall survival for brainstem glioblastoma ranges from 6 to 16 months, along with a wide range of 12–77 months for brainstem anaplastic astrocytoma. The KPS score, age, duration of symptoms, pontine location, and tumor grade impact survival.¹,²,⁴,⁸,¹¹–¹⁶ In general, efficacious therapies are limited in malignant BSG. Therefore, novel therapies are required to avoid deterioration of clinical symptoms and prolong survival.

Antiangiogenic therapy can normalize the structure and function of tumor vessels, resulting in improved drug delivery and a normalized tumor microenvironment. This effect may underlie the therapeutic benefit of combined antiangiogenic and cytotoxic therapies.¹⁷,¹⁸ In recent years, there have been several reports regarding antiangiogenic therapy, such as bevacizumab (an anti-vascular endothelial growth factor monoclonal antibody, which has been intensively studied in recurrent glioblastoma), as an effective salvage therapy for progressive BSG.¹⁹ After administering bevacizumab, an improvement in clinical conditions has been shown, along with satisfactory radiologic responses and a progression-free survival (PFS) up to 2 years.²⁰–²⁴ However, the efficacy of first-line antiangiogenic therapy for adult BSG is unknown (Table 3). The cases we summarized include eleven patients with BSG, five diagnosed radiologically and the remainder with histologic diagnosis. For histologically diagnosed BSG, PFS with bevacizumab ranges from 4 to 21 months; for four

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**Table 2** Survival data of histologically diagnosed brainstem gliomas in adults

| References          | Tumor histology | No. of patients | Therapy | mPFS (months) | mOS (months) |
|---------------------|-----------------|-----------------|---------|---------------|--------------|
| Kesari et al (2008)² | WHO grade I     | 16              | Radiation, chemotherapy (PCV, lomustine, vincristine, carboplatin, carmustine, temozolomide, irinotecan) | 44 | 83 |
| Kesari et al (2008)² | WHO grade II    | 15              | Radiation, chemotherapy (PCV, lomustine, vincristine, carboplatin, carmustine, temozolomide, irinotecan) | 48 | 168 |
| Kesari et al (2008)² | WHO grade III   | 12              | Radiation, chemotherapy (PCV, lomustine, vincristine, carboplatin, carmustine, temozolomide, irinotecan) | 10 | 17 |
| Kesari et al (2008)² | WHO grade IV    | 3               | Radiation, chemotherapy (PCV, lomustine, vincristine, carboplatin, carmustine, temozolomide, irinotecan) | 10 | 16 |
| Babu et al (2013)²  | AA              | 3               | Radiotherapy and concurrent temozolomide, other salvage chemotherapy | 6.7 | 13.5 |
| Babu et al (2013)²  | GB              | 4               | Radiotherapy and concurrent temozolomide, other salvage chemotherapy (AA + GB) | 12.1 | 14 |
| Babu et al (2014)²  | AA              | 22              | Radiotherapy, temozolomide, other salvage chemotherapy | NA | 30.5 |
| Babu et al (2014)²  | GB              | 12              | Radiotherapy, temozolomide, other salvage chemotherapy | NA | 11.5 |
| Hundsberger et al (2014)² | LG-BSG       | 8               | Radiotherapy and/or chemotherapy, supportive care | 24.1 | 12 |
| Hundsberger et al (2014)² | HG-BSG       | 13              | Radiotherapy and/or chemotherapy, supportive care | 5.8 | 6 |
| Dey et al (2014)²  | AA              | 75              | Radiation, data regarding chemotherapy, and other treatment regimens NA | NA | 6 |
| Dey et al (2014)²  | GB              | 165             | Radiation, data regarding chemotherapy, and other treatment regimens NA | NA | 6 |
| Reithmeier et al (2014)² | WHO grade I  | 16              | Radiation, chemotherapy, interstitial radiosurgery (with l-125 seeds), combined radiation and chemotherapy, supportive care | NA | 26.2 |
| Reithmeier et al (2014)² | WHO grade II | 31              | Radiation, chemotherapy, interstitial radiosurgery (with l-125 seeds), combined radiation and chemotherapy, supportive care | NA | 12.9 |
| Reithmeier et al (2014)² | WHO grade III | 42              | Radiation, chemotherapy, interstitial radiosurgery (with l-125 seeds), combined radiation and chemotherapy, supportive care | NA | 9.8 |
| Reithmeier et al (2014)² | WHO grade IV  | 14              | Radiation, chemotherapy, interstitial radiosurgery (with l-125 seeds), combined radiation and chemotherapy, supportive care | NA | 77 |
| Theeler et al (2015)² | DA              | 4               | Radiation, chemotherapy (temozolomide, cytotoxic chemotherapy, and/or bevacizumab) | NA | 21.1 |
| Theeler et al (2015)² | AA              | 18              | Radiation, chemotherapy (temozolomide, cytotoxic chemotherapy, and/or bevacizumab) | NA | 14.8 |
| Theeler et al (2015)² | GB              | 21              | Radiation, chemotherapy (temozolomide, cytotoxic chemotherapy, and/or bevacizumab) | NA | 77 |

Notes:¹ The study contained patients who were diagnosed radiologically. In one patient, grading was not performed.
Abbreviations: AA, anaplastic astrocytoma; DA, diffuse astrocytoma; GB, glioblastoma; HG-BSG, high-grade brainstem glioma; LG-BSG, low-grade brainstem glioma; mOS, median overall survival; mPFS, median progression-free survival; NA, not available; PCV, procarbazine, lomustine, and vincristine.
| References          | Patient's age (years), gender | Tumor location       | Histology                      | Therapy                                      | PFS                                      |
|---------------------|-------------------------------|----------------------|---------------------------------|----------------------------------------------|------------------------------------------|
| Torcuator et al (2009)²⁰ | 43, female                    | Pons                 | NA (progressive diffuse BSG diagnosed by MRI) | Initial: radiotherapy (54 Gy)                | From bevacizumab: 1+ year               |
| Raza et al (2009)²¹  | Adult (age and sex not mentioned) | Pons and medulla    | NA (malignant infiltrative diffuse BSG diagnosed by MRI) | TMZ irinotecan and bevacizumab            | From bevacizumab: 30+ months           |
| Blesa et al (2012)²²  | 38, male                      | Brainstem            | Glioblastoma multiforme WHO grade IV | Initial: radiotherapy                       | From bevacizumab: 30+ months           |
| Reithmeier et al (2013)²³ | 32, female                    | Medulla oblongata    | Astrocytoma WHO grade II        | Initial: radiotherapy (54 Gy)                | From bevacizumab: 30+ months           |
|                     | 36, female                    | Cerebellar peduncle  | Initial: glioma WHO grade II    | Salvage: intensified TMZ, bevacizumab       | From bevacizumab: 97+ weeks            |
|                     | with infiltration of the      |                      | Recurrence: malignant astrocytoma WHO grade III | Initial: interstitial radiosurgery of iodine-125 seeds | From bevacizumab: 20 months           |
|                     | brain stem                    |                      |                                 | Salvage: radiotherapy (60 Gy)               | From bevacizumab: 35 weeks             |
|                     | 39, male                      | Pontine tegmentum    | Anaplastic astrocytoma WHO grade III | TMZ carmustine bevacizumab                  | From bevacizumab: 20 months           |
|                     | 36, male                      | Medulla              | Diffuse astrocytoma WHO grade II | Initial: radiotherapy (60 Gy)               | From bevacizumab: 39 weeks            |
|                     | 49, female                    | Right medulla        | Anaplastic astrocytoma WHO grade III | Initial: TMZ (concomitant)                  | From bevacizumab: 4 months            |
|                     | 21, male                      | Right pons           | NA (radiologically diagnosed)   | Changed to: carboplatin etoposide interferon-β | From bevacizumab: 10 months           |
|                     | 59, male                      | Right pons           | NA (radiologically diagnosed)   | Salvage: bevacizumab                        | From bevacizumab: 6 months            |
|                     | 66, male                      | Right pons           | NA (radiologically diagnosed)   | Initial: TMZ bevacizumab                    | From bevacizumab: 9 months in progression |

**Abbreviations:** BSG, brainstem glioma; NA, not available; PFS, progression-free survival; TMZ, temozolomide.
patients who were given bevacizumab as salvage therapy, the PFS was even much longer than the PFS in the initial radiotherapy setting.22,23

Despite the presence of the following poor prognostic factors such as declining performance status, age, tumor location, histology, as well as the limited efficacy of radiotherapy or chemotherapy for malignant BSG, we made the decision to treat the patient with combined-modality management. Because the tumor contained a rich blood supply as determined by the surgeon, and 3D-ASL showed increased CBF, we surmised that combined antiangiogenic therapy may achieve a meaningful benefit.

Receptor-specific antibodies and small molecular TKIs have been developed for antiangiogenic therapy. Because bevacizumab was most commonly used and reported to be effective in combined antiangiogenic therapy of BSG,20–24 bevacizumab was recommended initially, but the patient refused for financial reasons. Then, we suggested consideration of TKIs (such as sorafenib, pazopanib, sunitinib, or apatinib), which target the vascular endothelial growth factor receptor (VEGFR).25–29 Owing to the rarity of BSG in adults, reported experience of TKIs in this disease was also rare, and we were not able to find guidelines for our patient. In order to choose the best agent for our patient, we examined the expression of VEGFR-1 and VEGFR-2. Immunohistochemistry showed that both VEGFR-1 and VEGFR-2 were positive. As VEGFR-2 plays an important role in angiogenesis, agents targeting VEGFR-2 should be theoretically helpful for tumor suppression. Animal experiments showed that the value of IC50 to VEGFR-2 for apatinib was 2 nM, while the values for sorafenib, pazopanib, and sunitinib were 90, 30, and 10 nM, respectively,30 which indicated the high specificity of VEGFR-2 for apatinib. Apatinib can highly selectively compete for the ATP-binding site of intracellular VEGFR-2 to block downstream signal transduction and inhibit tumor angiogenesis. Moreover, besides antiangiogenic effects, apatinib can also mildly inhibit c-Kit, c-SRC, PDGFR-β, and MET, exerting antitumor effects directly.28,29 A case series reported that apatinib had outstanding efficacy for refractory malignant glioma; one patient achieved near CR as evaluated by MRI and had an overall survival of 27 weeks, and another achieved partial response and had a PFS of 12 months.31 Another pilot study of apatinib plus irinotecan in patients with recurrent high-grade glioma showed an objective response rate of 55% (5/9), disease control rate of 78% (7/9), and median PFS of 8.3 months. The most common adverse events were gastrointestinal reaction (31.8%), hypertension (22.7%), and myelosuppression (18.0%).32 In our case, positive VEGFR-2 staining observed on the histologic specimen, along with lower costs and ease of administration, led us to consider apatinib over intravenous bevacizumab and other TKIs.

To our knowledge, this is the first known case in which a malignant BSG was managed with apatinib and chemoradiation. The patient achieved a radiologic CR at 4 weeks after treatment. His KPS was 70 at last follow-up, the PFS was over 8 months, and only grade 1 leukopenia and thrombocytopenia were observed; other toxicities for apatinib, such as hypertension, proteinuria, or hand–foot skin reaction, were not developed in our patient. However, among cases that we summarized above in which malignant BSGs were managed with bevacizumab and chemoradiation,20–24 only one patient who was diagnosed by MRI showed disappeared contrast enhancement in the tumor lesion region after treatment.21 Compared to the therapeutic effect with bevacizumab, the treatment response with apatinib in our case was encouraging; although it is only one case, it provides an additional treatment option for malignant BSG.

H3K27M mutations are frequently observed in adult BSG and tend to be associated with a poorer prognosis.33 A publication indicated that the H3K27M mutation defined a distinct subgroup of IDH wild-type gliomas characterized by a constant midline location, low rate of MGMT promoter methylation, and poor prognosis in gliomas.34 In this patient, who was H3K27M negative, we also performed molecular analysis of histone H3 mutations (negative for H3F3A:K27M and HIST1H3B:K27M). Potentially, this may be a contributing factor to the excellent outcome of this patient.

In this case, owing to the limited biopsy tissue (except for c-Kit), we did not evaluate the expression of other potential target genes, such as PDGFR-β, RET, and c-SRC, which might also contribute to the potential therapeutic effect of apatinib. We consider apatinib to not just be an antiangiogenic agent, but it may also act on the tumor cells directly in this case, especially in the absence of an intact blood–brain barrier, as evidenced by the contrast enhancement on MRI.

Conclusion
To our knowledge, this is the first case report using apatinib to treat brainstem IDH wild-type anaplastic astrocytoma with an excellent outcome, and the toxicities are tolerable. The case describes a novel regimen for clinicians encountering this rare and difficult-to-treat brain tumor.

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