



CyberKnife for Recurrent Malignant Gliomas: A Systematic Review and Meta-Analysis

Lucio De Maria^{1*}, Lodovico Terzi di Bergamo², Alfredo Conti^{3†}, Kazuhiko Hayashi^{4†}, Valentina Pinzi^{5†}, Taro Murai^{6†}, Rachelle Lanciano^{7†}, Sigita Burneikiene^{8†}, Michela Buglione di Monale⁹, Stefano Maria Magrini⁹ and Marco Maria Fontanella¹

¹ Unit of Neurosurgery, University of Brescia and ASST Spedali Civili, Brescia, Italy, ² Institute of Oncology Research, Bellinzona, Switzerland, ³ Unit of Neurosurgery, Alma Mater Studiorum University of Bologna and IRCCS Istituto delle Scienze Neurologiche, Bologna, Italy, ⁴ Unit of Radiation Oncology, Osaka University Graduate School of Medicine, Suita, Japan, ⁵ Unit of Neurosurgery, Fondazione IRCCS Istituto Neurologico Carlo Besta, Milan, Italy, ⁶ Unit of Radiology, Nagoya City University Graduate School of Medical Sciences, Nagoya, Japan, ⁷ Philadelphia CyberKnife/Crozer Health, Havertown, PA, United States, ⁸ Boulder Neurosurgical Associates, Boulder, CO, United States, ⁹ Unit of Radiation Oncology, University of Brescia and ASST Spedali Civili, Brescia, Italy

OPEN ACCESS

Edited by:

Christine Marosi, Medical University of Vienna, Austria

Reviewed by:

Birgit Flechl, MedAustron, Austria Pierina Navarria, Humanitas Research Hospital, Italy

*Correspondence:

Lucio De Maria I.demaria@unibs.it

[†]These authors have contributed equally to this work

Specialty section:

This article was submitted to Neuro-Oncology and Neurosurgical Oncology, a section of the journal Frontiers in Oncology

> Received: 12 January 2021 Accepted: 15 March 2021 Published: 29 March 2021

Citation:

De Maria L, Terzi di Bergamo L, Conti A, Hayashi K, Pinzi V, Murai T, Lanciano R, Burneikiene S, Buglione di Monale M, Magrini SM and Fontanella MM (2021) CyberKnife for Recurrent Malignant Gliomas: A Systematic Review and Meta-Analysis. Front. Oncol. 11:652646. doi: 10.3389/fonc.2021.652646 **Background and Objective:** Possible treatment strategies for recurrent malignant gliomas include surgery, chemotherapy, radiotherapy, and combined treatments. Among different reirradiation modalities, the CyberKnife System has shown promising results. We conducted a systematic review of the literature and a meta-analysis to establish the efficacy and safety of CyberKnife treatment for recurrent malignant gliomas.

Methods: We searched PubMed, MEDLINE, and EMBASE from 2000 to 2021 for studies evaluating the safety and efficacy of CyberKnife treatment for recurrent WHO grade III and grade IV gliomas of the brain. Two independent reviewers selected studies and abstracted data. Missing information was requested from the authors via email correspondence. The primary outcomes were median Overall Survival, median Time To Progression, and median Progression-Free Survival. We performed subgroup analyses regarding WHO grade and chemotherapy. Besides, we analyzed the relationship between median Time To Recurrence and median Overall Survival from CyberKnife treatment. The secondary outcomes were complications, local response, and recurrence. Data were analyzed using random-effects meta-analysis.

Results: Thirteen studies reporting on 398 patients were included. Median Overall Survival from initial diagnosis and CyberKnife treatment was 22.6 months and 8.6 months. Median Time To Progression and median Progression-Free Survival from CyberKnife treatment were 6.7 months and 7.1 months. Median Overall Survival from CyberKnife treatment was 8.4 months for WHO grade IV gliomas, compared to 11 months for WHO grade III gliomas. Median Overall Survival from CyberKnife treatments who underwent CyberKnife treatment alone, compared to 9.5 months for patients who underwent CyberKnife treatment plus chemotherapy. We did not observe a correlation between median Time To Recurrence and median Overall Survival from CyberKnife. Rates of acute neurological and acute non-neurological side effects were

March 2021 | Volume 11 | Article 652646

1

3.6% and 13%. Rates of corticosteroid dependency and radiation necrosis were 18.8% and 4.3%.

Conclusions: Reirradiation of recurrent malignant gliomas with the CyberKnife System provides encouraging survival rates. There is a better survival trend for WHO grade III gliomas and for patients who undergo combined treatment with CyberKnife plus chemotherapy. Rates of complications are low. Larger prospective studies are warranted to provide more accurate results.

Keywords: CyberKnife, stereotactic radiosurgery, malignant gliomas, recurrence, HGG, high-grade gliomas, glioblastoma, anaplastic astrocytoma

INTRODUCTION

The majority of malignant brain tumors are represented by gliomas (70%) (1). The standard management of newly diagnosed malignant gliomas (MGs) is maximal resection followed by radiotherapy (RT) with concomitant and adjuvant chemotherapy (CMT) (2). Although a solid treatment strategy has been established for MGs, recurrence still occurs in almost all patients within 2 years after initial treatment (3-5). Possible treatment strategies for recurrent malignant gliomas (rMGs) include second-line CMTs, surgery with or without adjuvant therapies, and RT (2, 6, 7). Reirradiation appears to be an efficacious and safe treatment modality, providing survival benefits with acceptable risk (8, 9). Among different reirradiation modalities, hypofractionated stereotactic radiotherapy (HFSRT) has shown promising results as it allows delivery of a large total dose, in a precise target volume and short treatment duration (10, 11). Nowadays, various HFSRT and stereotactic radiosurgery (SRS) machines are available and their usage has been gradually increasing. All systems have excellent accuracy with targeting areas close to 1 mm (12-14). Among those, the CyberKnife[®] (CK) is a frameless image-guided radiotherapy system mounting a 6-MV linear accelerator on a highly maneuverable robotic arm (15). The CK System is a non-invasive and pain-free treatment strategy that requires a customized thermoplastic face mask, reducing patient discomfort associated with other frame-based radiosurgical systems. Unlike other SRS techniques, the CK does not require general or local anesthesia still ensuring a comparable level of accuracy (12). Particularly, the CK was found to have clinically relevant accuracy of 0.7 +/- 0.3 mm, minimizing normal brain radiation exposure and allowing for high doses of radiation to targeted areas (12, 16). Given its recent development, few case series have been reported on CK for rMGs of the brain, and indications are still debated. We hereby conducted a systematic review of the literature and a meta-analysis to provide physicians awareness about the efficacy and safety of CK treatment for rMGs.

MATERIALS AND METHODS

Literature Search

The systematic review was performed according to the Preferred Reporting Items for Systematic Reviews and Meta-Analysis (PRISMA) guidelines (17). A comprehensive literature search of the databases PubMed, Ovid MEDLINE, and Ovid EMBASE databases was designed and conducted by an experienced librarian with input from the authors. The keywords "glioblastoma", "anaplastic astrocytoma", "malignant glioma", "high-grade glioma", "HHG", "recurrence", "recurrent malignant glioma", "brain", "CyberKnife", "CK", "stereotactic radiosurgery", "SRS", and "stereotactic radiotherapy" were used in "AND" and "OR" combinations. The search was limited to articles published between 2000 and 2021.

The following inclusion criteria were used: 1) English language, 2) case series reporting greater than 5 patients 3) studies reporting exclusively histologically proven World Health Organization (WHO) grade IV gliomas or WHO grade III gliomas of the brain (18), 4) studies reporting recurrence, and 5) studies reporting retreatment with the CK System at recurrence. The exclusion criteria were: 1) case series reporting fewer than 10 patients and case reports, 2) brain lesions other than MGs, 3) lesions not located in the brain (e.g. gliomas of the spinal cord), 4) studies reporting only newly diagnosed MGs, 5) studies reporting on irradiation techniques other than the CK System, 6) studies not reporting survival data.

Two authors determined the inclusion and exclusion criteria for the studies in the literature search. In studies with overlapping patient populations written by the same author/institution, we only included the largest or most complete dataset. In cases where outcomes were separated by WHO grade or CMT at recurrence, we abstracted outcomes separately to perform our subgroup analyses. Missing baseline data and outcomes information was requested from the authors via email correspondence. The authors of six included studies replied and the information provided was integrated into the data abstraction process.

Data Extraction

For each study, we abstracted the following baseline information: number of patients; median age at CK treatment; gender; median Karnofsky Performance Status (KPS) at CK treatment; WHO grade and histotype at recurrence. Regarding treatment at initial diagnosis we collected information about: the extent of resection (EOR), i.e. gross total resection (GTR, resection of more than 99% of the preoperative tumor volume), subtotal resection (STR, 95%–99% resection); partial resection (PR, < 95% resection), and biopsy (B) (19); the number of patients who underwent conventional radiation therapy (CRT); and the number of patients who underwent CMT. About the recurrence interval, we abstracted the Time To Recurrence (TTR, the time span between initial treatment and CK) (20, 21). As for treatment at recurrence, we gathered the following data: median planned target volume (PTV); the median number of fractions; total radiation dose in Gray (Gy); the number of patients who underwent CMT.

Objectives

Our primary endpoints were median Overall Survival (OS), median Time To Progression (TTP), and median Progression-Free Survival (PFS). As for OS, we extracted data from initial diagnosis (i.e. time-length from the date of initial diagnosis to death from any cause) and from CK (i.e. time-length from the date of the start of CK treatment to death from any cause) (22). Concerning TTP and PFS, we abstracted data from CK. The former was defined as the time elapsed between the start of CK treatment to Beside Recurrence (BR, new lesion developed after 4 weeks beside or inside the prescribed marginal isodose line of previous CK treatment) or Progressive Disease (PD, more than 25% growth of Gd-enhanced area within 4 weeks after CK treatment) (21). The latter was defined as the time elapsed between the start of CK treatment to any disease recurrence or death from any cause (23). For our subgroup analysis, we were able to abstract median OS from initial diagnosis and from CK treatment for WHO grade IV gliomas versus WHO grade III gliomas separately and for CK plus CMT versus CK treatment alone separately. Besides, we analyzed the relationship between median TTR and median OS from CK treatment.

The secondary endpoints were Local Response (LR), New Lesion (NL), and complications. The LR was assessed with Gdenhanced Magnetic Resonance Imaging (MRI) at 1 month after CK treatment and was classified into the following categories: Complete Response (CR, Gd-enhanced area disappears and no regrowth is recognized for at least 4 weeks after treatment), Partial Response (PR, Gd-enhanced area is reduced by more than 50% and maintains this state for at least 4 weeks after treatment), No Change (NC, less than 50% reduction or less than 25% growth of Gd-enhanced area, maintained for at least 4 weeks after treatment) and PD (24). The development of NLs following initially controlled disease (i.e. CR, PR, NC), was divided into BR and Remote Recurrence (RR, lesion located remotely from the prescribed marginal isodose line of previous CK treatment) (25). Regarding complications, we extracted the number of acute neurological and non-neurological side effects, corticosteroid dependency (the onset of neurological deficits and/or cephalalgia requiring daily doses of dexamethasone > 4 mg for more than 8 weeks), radiation necrosis, and other toxicities (26).

Study Risk of Bias Assessment

We modified the Newcastle-Ottawa Quality Assessment Scale to assess the methodologic quality of the studies included in this meta-analysis (27). This tool is designed for use in comparative studies; however, our analyzed studies did not have control groups, therefore, we assessed the study risk of bias based on selected items from the scale, focusing on the following questions: 1) Did the study include all patients or consecutive patients versus a selected sample? 2) Was the study retrospective or prospective? 3) Was clinical follow-up satisfactory, thus allowing ascertainment of all outcomes? 4) Were outcomes clearly reported? 5) Were there clearly defined inclusion and exclusion criteria?

Statistical Analysis

We estimated each cohort's cumulative prevalence and 95% confidence interval for each outcome. Event rates were pooled across studies using a random-effects meta-analysis. Heterogeneity across studies was evaluated using the I² statistic. An I² value of >50% suggests substantial heterogeneity. Meta-regression was not used in this study. For some outcomes it was not possible to estimate the standard errors, therefore a standard error of 0 was used in the meta-analysis. Pearson's correlation was used to correlate median TTR and median OS from CK treatment. Statistical analyses were performed using OpenMeta [Analyst] (http://www.cebm.brown.edu/openmeta/) and R statistical package v3.4.1 (http://www.r-project.org).

RESULTS

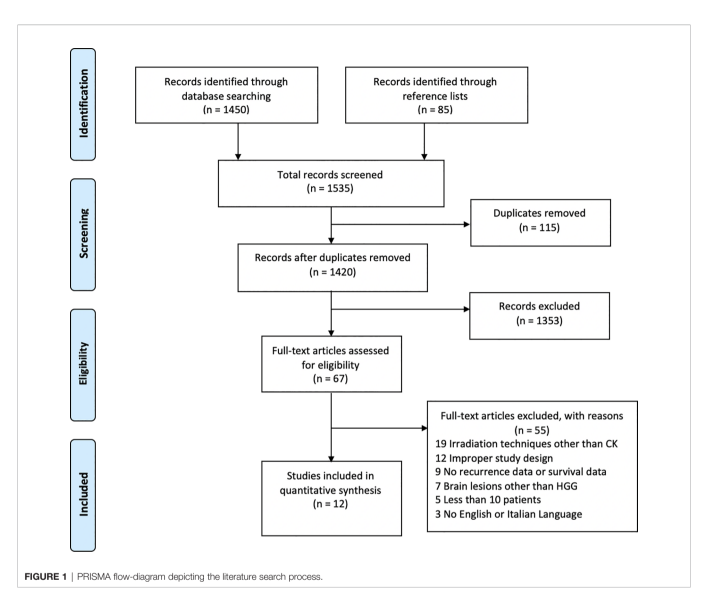
Literature Review

A total of 1420 papers were identified after duplicates removal. After title and abstract analysis, 67 articles were identified for full-text analysis. Eligibility was ascertained for 12 articles (20, 21, 24–26, 28–34). The remaining 55 articles were excluded for the following reasons: 1) irradiation techniques other than the CK System (19 articles), 2) improper study design (12 articles), 3) studies reporting only on newly diagnosed MGs or not reporting survival data (9 articles), 4) studies reporting on brain lesions other than MGs (7 articles), 5) case series reporting fewer than 10 patients (5 articles), and 6) studies in other languages (3 articles). All studies included in the analysis had at least one or more outcome measures available for one or more of the patients' groups analyzed. **Figure 1** shows the flow chart according to the PRISMA statement (17).

Study and Patients Characteristics

Our meta-analysis included a total of 398 patients. The smallest study included 13 patients (32) and the largest included 128 patients (33). The median age at CK treatment was 54.5 years. There was a male predominance (1.6:1). The median KPS at CK treatment was 80. Six studies (50%) reported on WHO grade IV and III gliomas and other 6 studies (50%) reported on WHO grade IV gliomas. The histotype was available in 9 studies (75%): six studies (67%) reported on glioblastomas (GBMs) and 3 studies (33%) reported on GBMs and anaplastic astrocytomas (AAs).

At the time of initial diagnosis, most of the patients underwent STR (131, 33%), followed by GTR in 112 patients (28%), B in 30 patients (8%), and PR in 12 patients (3%). Post-operative CRT was undertaken in 343 patients (86%) and the median dose was 60 Gy.



Post-operative CMT was undertaken in 275 patients (69%) and Temozolomide (TMZ) was the CMT regimen reported in most studies (193 patients, 48%). Patients were followed-up with a Gd-enhanced MRI performed every 1 to 3 months. The median TTR was 14 months (range 1-171).

At recurrence, the GTV was defined as the MRI Gd-enhanced area and the PTV was reconstructed adding 0 to 3 mm margin to the GTV. The median target volume (PTV) was 12.1 ml. The median number of fractions was 3 (range 1-6) and the median dose was 24.5 Gy (range 13.9-48.8). The prescribed marginal isodose ranged from 78% to 91%. Half of the patients (203, 51%) underwent CMT at recurrence. Although TMZ was the most reported CMT regimen (66 patients, 17%), other therapies were undertaken, particularly Bevacizumab (BEV) in 22 patients (5%) and Interferon in 16 patients (4%). Administration of CMT was concomitant and/or after CK treatment in 199 patients (98%) and before CK treatment in 4 patients (2%). The latter received BEV-based salvage therapy prior to CK treatment. A summary of the included studies is provided in **Table 1**.

Primary Outcomes

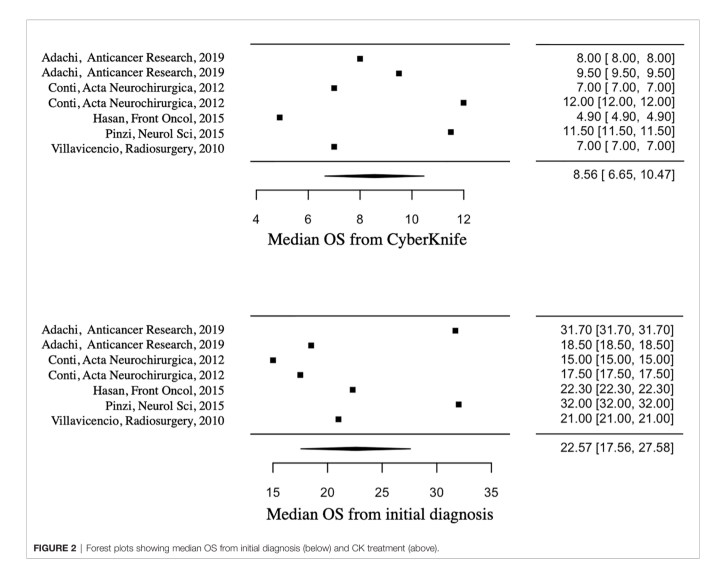
Median OS from initial diagnosis and CK treatment was 22.57 months (95%CI=17.56-27.58) and 8.56 months (95%CI=6.65-10.47) respectively. **Figure 2** shows the median OS forest plots. Median TTP and median PFS from CK treatment were 6.68 months (95%CI=2.13-11.22) and 7.05 months (95%CI=1.30-12.79) respectively.

Concerning the WHO grade, the median OS from initial diagnosis was 19.88 months (95%CI=17-22.76) for WHO grade IV gliomas, compared to 48.35 months (95%CI=15.72-80.98) for WHO grade III gliomas. Median OS from CK treatment was 8.4 months (95%CI=6.35-10.45) for WHO grade IV gliomas, compared to 11 months (95%CI=5.12-16.88) for WHO grade III gliomas.

About the treatment, median OS from initial diagnosis was 25.4 months (95%CI=16.97-33.83) for patients who underwent CK plus CMT treatment, compared to 16.05 months (95% CI=13.99-18.11) for patients who underwent CK treatment alone; median OS from CK treatment was 9.52 months (95%

Study			Baseline Data						1st Treatment			Recurrence Interval	2nd Treatment			
No.	Author, Journal, Year	Prospective/ Retrospective	No. of Patients	Median Age; range	M:F	Median KPS; range	WHO Grade	Histotype	GTR/STR/ PR/B (%)	CRT (%)	СМТ (%)	Median TTR (months); range	Median PTV (ml)	CK Median No. of Fractions	Median Dose (Gy)	СМТ (%)
1	Adachi, Anticancer Research, 2019 (28)	R	29	53	19:10	75	IV-III	NA	NA	29 (100)	29 (100)	NA	4.1	1	25.5	29 (100)
2	Conti, Acta Neurochirurgica, 2012 (26)	Ρ	23	58	13:10	80	IV	GBM	NA	23 (100)	23 (100)	NA	15.7	2	20	12 (52)
3	Ekici, J BUON, 2014 (29)	R	27	NA	NA	70-100	IV-III	NA	NA	NA	NA	NA	NA	NA	NA	NA
4	Glavatskyi, Neuro-Oncology, 2014 (30)	R	26	NA	NA	NA	IV-III	GBM-AA	8 GTR (30), 9 STR (35), 9 B (35)	9 (35)	17 (65)	NA	NA	NA	NA	0 (0)
5	Greenspoon, Onco Targets Ther, 2014 (20)	Ρ	31	53; 36-75	NA	80; 60-90	IV	GBM	NA	31 (100)	0 (0)	NA	12.1	5	NA	31 (100)
6	Hasan, Front Oncol, 2015 (31)	R	19	56; 29-79	13:6	80; 40-100	IV	GBM	7 GTR (37), 8 STR (42), 4 NA (21)	19 (100)	19 (100)	16; 2-122	20.9	5	25	15 (79)
7	Lévy, Cancer/Radiothérapie, 2017 (32)	R	13	55	11:02	85; 65-100	IV-III	NA	6 GTR (46), 3 STR (23), 4 B (31)	13 (100)	13 (100)	11	23.6	NA	30	11 (85)
8	Pinzi, Neurol Sci, 2015 (33)	R	128	51; 18-79	80:48	60-100	IV-III	GBM-AA	38 GTR (30), 77 STR (60), 13 B (10)	128 (100)	84 (67)	15; 6-171	6.5	2	19	61 (48)
9	Torok, Technol Cancer Res Treat, 2011 (25)	R	14	58; 39-76	7:7	NA	IV	GBM	11 GTR (79),	14 (100)	14 (100)	8; 1-28	7	3	24	12 (86)
10	Villavicencio, Radiosurgery, 2010 (34)	R	26	55.5; 36-82	18:8	90; 50-100	IV	GBM	3 PR (21) 15 GTR (58),	26 (100)	25 (96)	13; 5-89	7	2	NA	5 (19)
11	Yazici, J Neurooncol, 2014 (21)	R	37	37; 22-69	18:19	60-100	IV	GBM	9 STR (34), 2 B (8) 23 GTR (62),	37 (100)	37 (100)	15; 5-45	NA	5	30	11 (30)
12	Yoshikawa, Minim Invasive	Ρ	25	54; 28-78	13:12	77.3; 30-90	IV-III	GBM-AA	13 STR (35), 1 B (3) 3 GTR (12), 12	. ,	14	NA	19.1	3	20.3	(30)
	Neurosurg, 2006 (24)		-	,		-,	·		STR (48),	(56)	(56)		-	-		(64)
									9 PR (36), 1 B (4)							

De Maria et al.



CI=7.78-11.25) for patients who underwent combined treatment, compared to 4.44 months (95%CI=0-9.46) for patients who underwent CK treatment alone. Primary outcomes are reported in **Table 2**. We did not observe a positive correlation between median TTR and median OS from CK.

Secondary Outcomes

Rates of acute neurological and non-neurological side effects after CK treatment at recurrence were reported in 287 patients. The overall rate of the former was 3.6% (95%CI=1.5-5.7), while 13% for the latter (95%CI=0-26.1). Acute neurological effects included worsening of pre-existing symptoms, dizziness, nausea/ vomiting, and neurological deterioration. Acute non-neurological effects included alopecia, fatigue, asthenia, and clinical deterioration. **Figure 3** shows the acute neurological side effects forest plot. Rates of corticosteroid dependency and radiation necrosis were reported in 271 patients and 306 patients respectively. The overall rate of corticosteroid dependency was

18.8% (95%CI=10.0-27.6), while the overall rate of radiation necrosis was 4.3% (95%CI=2.1-6.6). Rates of other toxicities were reported in 267 patients. The overall rate was 1.1% (95%CI=0.4-2.7) and these were hematological toxicities.

Rates of LR after CK treatment at recurrence were reported in 84 patients. The overall rate of PD was 37.9% (95%CI=26.5-49.3), followed by overall rates of 29.2% for NC (95%CI=15.7-42.7), 27.7% for PR (95%CI=18.2-37.2), and 2% for CR (95% CI=1.0-4.9).

Rates of NLs developed following CK treatment were reported in 61 patients, and the overall rate was 88.4% (95% CI=80.5-96.4). Rates of BR or RR were reported in 50 patients. The overall rate of BR was 75.9% (95%CI=64.3-87.6), compared to 17.7% for RR (95%CI=1.5-33.8). The secondary outcomes are summarized in **Table 3**.

Study Heterogeneity

 $\rm I^2$ values were <50% indicating a lack of substantial heterogeneity for all the outcomes.

TABLE 2 | Primary outcomes.

	Months (95%CI)
Median Overall Survival	
From initial diagnosis	22.57 (17.56-27.58)
WHO grade IV	19.88 (17.00-22.76)
WHO grade III	48.35 (15.72-80.98)
CK alone	16.05 (13.99-18.11)
CK plus CMT	25.40 (16.97-33.83)
From CK treatment	8.56 (6.65-10.47)
WHO grade IV	8.40 (6.35-10.45)
WHO grade III	11.00 (5.12-16.88)
CK alone	4.44 (0-9.46)
CK plus CMT	9.52 (7.78-11.25)
Median Time To Progression	
From CK treatment	6.68 (2.13-11.22)
Median Progression Free Survival	
From CK treatment	7.05 (1.30-12.79)

DISCUSSION

Findings

The treatment strategy for patients harboring rMGs is still debated and no clear consensus has been achieved yet. Treatment modalities include surgery, CMT, RT, and combined treatments. Reirradiation with SRS can provide survival benefits with acceptable risks. Among diverse SRS machines currently available, we focused on the CK System. Our study's primary aim was to establish the efficacy of CK treatment for rMGs, concerning survival and time to disease progression. Our secondary aims were to establish the local disease response, recurrence of disease, and toxicities. We performed a systematic review and meta-analysis of published studies on CK for rMGs and found several interesting findings.

Patients Characteristics

In our meta-analysis, we observed a male predominance (1.6:1). Recent evidence suggested that sex-associated biological features can play a role in MGs incidence, regardless of the age, race, and geographic location of patients (1, 35, 36). An average male-to-

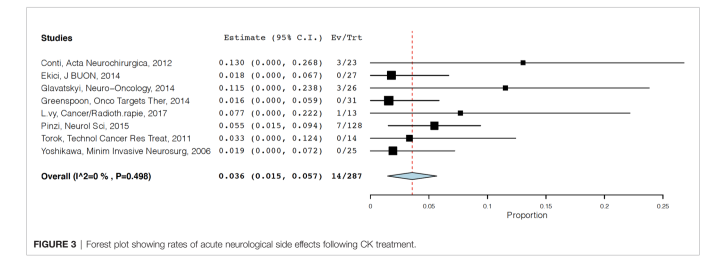
TABLE 3 | Secondary outcomes.

	Overall % (95%CI)
Local Response	
Complete Response	2.0 (1.0-4.9)
Partial Response	27.7 (18.2-37.2)
No Change	29.2 (15.7-42.7)
Progressive Disease	37.9 (26.5-49.3)
Recurrence	
New Lesion	88.4 (80.5-96.4)
Beside Recurrence	75.9 (64.3-87.6)
Remote Recurrence	17.7 (1.5-33.8)
Complications	
Acute Neurological Side Effects	3.6 (1.5-5.7)
Acute non-Neurological Side Effects	13.0 (0-26.1)
Corticosteroid Dependency	18.8 (10.0-27.6)
Radiation Necrosis	4.3 (2.1-6.6)
Other Toxicities	1.1 (0.4-2.7)

female ratio of 1.6:1 has been previously reported for MGs, with greater incidence in men (1, 37). The prevalence of MGs in males appeared to be related mainly to genetic dissimilarities and not only to the presence of sex hormones (38). Gender differences can be pivotal for developing tailored approaches to MGs and pursuing studies are taking into account sex differences for innovative treatment strategies (37).

Primary Outcomes

Median OS of rMGs without any treatment has been reported to range between 3 and 6 months (5). Reoperation of recurrent GBMs provides 3 to 5 months median survival, without a significant increase in morbidity and mortality, and is still limited to 10-30% of patients due to the infiltrative nature of the disease and the involvement of eloquent areas (39–42). Over the past years, reirradiation has been increasingly proposed as an alternative treatment strategy with successful results (43, 44). Among different reirradiation modalities, HFSRT and SRS have been reported to provide a median OS ranging from 8.6 to 18 months with acceptable side effects (45). Our meta-analysis on CK System revealed a median OS of 8.6 months (95%CI=6.65-10.47) from SRS treatment and 22.6 months (95%CI=17.56-27.58)



considering survival from initial diagnosis. Median TTP and median PFS after CK treatment were comparable (6.7 vs 7.1 months), with a slightly longer median PFS as this outcome only differs for the inclusion of remote recurrence or death from any cause (23). Barbagallo et al. reported a similar mean PFS for patients with rMGs undergoing second surgery (7.7 months) (46). Randomized controlled trials (RCTs) are needed to provide more definitive answers about differences in particular treatment strategies for rMGs.

Regarding the grade of the disease, WHO grade IV rMGs showed a shorter median survival from CK treatment (8.4 months), compared to WHO grade III rMGs (11 months). Notably, Murai et al. reported a 3-year survival rate of 38% for re-irradiated patients with recurrent WHO grade III anaplastic ependymomas (AEs) and a median OS from CK treatment of 31.5 months (47). Therefore, treatment of recurrent AEs with CK System is a promising alternative, especially for deep-seated lesions or lesions located adjacent to eloquent areas (47–50).

The subgroup analysis of treatment strategy revealed a longer survival for patients undergoing CK plus CMT treatment (9.5 months) compare with patients undergoing CK treatment alone (4.4 months). Hu et al. previously reported that HFSRT combined with CMT confers a slight survival improvement for patients with rMGs compared with HFSRT alone (8.23-23.0 months vs 3.9-12.0 months) (51). In their meta-analysis including 388 patients, 3 out of the 7 selected studies presented statistically significant differences (P < 0.05) between these two treatment approaches, and 3 out of the 4 remaining studies showed a favorable survival for patients treated with combined therapy rather than HFSRT alone. Likewise, our meta-analysis suggests a longer survival for patients who undergo combined treatment, but we cannot ascertain the absence of confounding bias between the two groups and stratified RCTs would be needed for ultimate conclusions. Moreover, we were unable to perform qualitative subgroup analyses of the systemic agents used and the time of systemic therapy sessions with respect to CK treatment. Among the different agents used in the included studies, TMZ was the most reported CMT regimen (66 patients, 16%), followed by BEV in 22 patients (5%) and Interferon in 16 patients (4%). Administration of CMT was concomitant and/or after CK treatment in 199 patients (98%) and before CK treatment in 4 patients (2%). The latter received BEV-based salvage therapy prior to CK treatment (31). The most commonly used systemic therapies for rMGs include TMZ, nitrosoureas, and BEV (52-54). The combination of lomustine with BEV has shown improved PFS but not OS, and a higher toxicity rate compared with lomustine alone (55). Bevacizumab alone or in combination with chemotherapy agents such as lomustine or irinotecan has demonstrated a median survival time from recurrence around 9 months and radiographic response rates of approximately 30 to 40 percent (55, 56). Few reports described the combination of bevacizumab with HFSRT for recurrent GBMs with safe and effective results (57-59). This treatment strategy is under study in an ongoing larger randomized trial (60). Among the studies included in our meta-analysis, Hasan et al. showed a better survival for patients with recurrent GBMs treated with BEV

either before or after CK treatment (31). Palmer et al. reported a slightly higher survival for patients with recurrent GBMs treated with HFSRT before BEV rather than BEV before HFSRT (13.9 vs 13.3 months) but stressed the importance of a randomized multiinstitutional trial for more definite conclusions (61).

We did not observe a positive correlation between median TTR and median OS from CK. Likewise, Greenspoon et al. did not find a statistical difference in OS or PFS when stratifying by TTR (<12 months or >12 months) (20). Conversely, Yazici et al. reported improved survival for patients with a TTR of more than 12 months (21).

Secondary Outcomes

Our meta-analysis shows that CK is a relatively safe and effective treatment modality for rMGs. Rates of complications were relatively low. Corticosteroid dependency had the highest rate among the complications (18.8%), followed by acute nonneurological side effects (13%, including fatigue, alopecia, and clinical deterioration), and by radiation necrosis (4.3%). Notably, the authors of the included studies included steroid use among side effects only when requiring daily doses of dexamethasone > 4 mg for more than 8 weeks. However, we must acknowledge that current guidelines mention steroid use as a side effect from basic prescription (62). Larger re-irradiated tumors (maximum diameter greater than 4 cm) are more inclined to develop radiation necrosis (33, 63). Indeed, a crucial factor in developing radiation necrosis is the volume of the irradiated normal brain, which is relative to the tumor volume (64, 65). Radiation necrosis is known to occur in the normal brain when the normalized total dose (NTD) is greater than 100 Gy (66). Other authors reported that using a fractionated scheme aimed to maintain a normalized total dose (NTD)<100 Gy can reduce the risk of radionecrosis in larger tumors (26, 33). Conversely, rates of acute neurological effects (3.6%) such as worsening of pre-existing symptoms, dizziness, nausea/vomiting, neurological deterioration, and rates of hematological toxicities (1.1%) were the lowest. Acute side effects were higher in patients treated with large single fraction volumes, supporting the hypothesis that fractioned schemes may be safer for tumors larger than 4 cm in maximum diameter or proximal to eloquent areas (33). Hematological toxicities such as leukopenia and thrombocytopenia were mainly reported for patients who underwent CK treatment plus CMT (26). Although we meta-analyzed the side effects reported by the authors, it was not possible to grade toxicity because of a lack of uniformity among studies. Future trials should report the side effects according to standardized grading systems to enhance uniformity and facilitate interpretation of results (62, 67).

The analysis of LR at 4 weeks after CK treatment showed disease progression in 37.9% of cases, stability in 29.2%, reduction in 27.7%, and complete disappearance in 2%. Yoshikawa et al. reported a higher control rate (i.e. CR, PR, NC) for GBM patients than AA patients (63.6% vs 45.5%) (24). However, LR after CK was reported in a small overall cohort (84 patients), and this outcome should be validated by more extensive analyses. Moreover, true progression may often be indistinguishable from pseudoprogression (21). Pseudoprogression is a subacute effect of

radiotherapy observed in the first 12 weeks after treatment, first described by Hoffman et al (68). It was pathologically defined by Chamberlain et al. as necrosis without evidence of tumor and appears as increased contrast enhancement following radiotherapy (69). These imaging findings are consequences of disruption of the blood-brain barrier and represent proof of radiation's efficacy rather than progression or toxicity, indeed correlate with longer OS (21, 70, 71). Diagnosis of pseudoprogression is made during follow-up when stabilization or improvement of clinical and radiographic findings is observed (21). Instead, true progression within the first 12 weeks after radiotherapy, can only be defined if the majority of new enhancement is outside the radiation field or if there is pathological confirmation of PD (72).

The overall rate of NLs was considerably high (88.4%), with a greater rate of BR rather than RR (75.9% vs 17.7%). However, the rate of NLs was reported in only 61 patients, and the location of recurrence in only 50 patients overall. Therefore, this outcome needs to be corroborated by larger studies as well. Although it is known that recurrence of MGs appears mainly within 2 cm of the enhancing edge of the original tumor, Yoshikawa et al. reported the development of BR despite an initial high control rate (63.2% for GBM and 42.9% for AA controlled patients) (73). Despite surgery plays a key role in GBM recurrence, most of all for large volumes, CK radiosurgery has shown good results with a low rate of toxicity. Some aspects though remain unclear, such as radiation dose and fractionation.

A focus on the quality of life (QoL) is imperative given the poor prognosis and short life expectancy of patients with a diagnosis of rMGs. The QoL of MG patients is most often affected by the development of CMT/RT side effects, changes in physical functioning, and global health status (74). Unlike surgery and other SRS techniques, the CK treatment can be delivered without sedation and as an outpatient, which would help maximize the QoL. The primary and secondary end-points of our meta-analysis were based on outcomes reported by authors of the included studies. Therefore, we were unable to meta-analyze the effect of CK treatment on KPS, cognitive function, and QoL. However, Greenspoon et al. reported on the benefit of BEV in preventing toxicity and improving QoL of patients undergoing CK plus TMZ (20). Quality of Life after HFSRT for rMGs patients has been previously reported to remain stable for a median follow-up of 9 months (75). A subsequent study on high-dose reirradiation in selected patients with recurrent/progressive MGs found a stable QoL and improvement of activities of daily living (ADL) over a 1-year time period (76). Future studies should include KPS and QoL among their primary outcomes to evaluate the impact of CK treatment in life-limiting diseases such as rMGs.

Limitations

Despite the significant number of patients included in our study, this meta-analysis was based primarily on a few single-center case series and thus has limitations inherent to single-center retrospective studies. Based on the data abstracted from the articles and provided by the authors of the included studies, we could not ascertain the number of patients undergoing repeat surgery and the EOR at recurrence. The different ways in which each study provided the confidence intervals and/or standard deviations did not allow the use of the standard errors for some of the outcomes in the meta-analysis. In such cases, a standard error of 0 was adopted for each study. This led to an imperfect approximation of the meta-analyzed outcome and its confidence interval. While we were able to perform subgroup analyses based on WHO grade and CMT at recurrence as well as analyzing the relationship between TTR and OS, we were unable to perform more granular analyses stratifying outcomes by other relevant variables such as the histotype, the PTV, irradiation dose, the number of fractions, the patients' age and KPS. Moreover, we were unable to perform qualitative subgroup analyses of the CMT agents used and the time of CMT sessions with respect to CK treatment at recurrence. The assessment of LR was reported in a small cohort and differential diagnosis of lesions developed post-CK treatment can be misleading. Therefore, this outcome should be validated by more extensive analyses and future studies should focus on discrimination of lesions developed following CK treatment.

Nonetheless, to the best of our knowledge, this is the first meta-analysis providing helpful conclusions on the treatment of rMGs with the CK System and a potential start point for future studies.

CONCLUSIONS

Reirradiation of rMGs with the CK System has reasonable efficacy and provides encouraging survival rates. There is a better survival trend for WHO grade III lesions and for patients who undergo combined treatment with CK plus CMT. Treatment of rMGs with CK is a safe alternative, considering the low rates of complications. Larger and well-designed prospective studies are warranted to provide more accurate results.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/supplementary material. Further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

LDM: Writing - Original Draft, Term, Methodology, Validation, Resources, Project administration, Formal analysis, Investigation, Data Curation. LTB: Investigation, Resources, Data Curation, Formal analysis. AC: Resources, Writing - Review & Editing. KH: Resources, Writing - Review and Editing. VP: Resources, Writing - Review and Editing. TM: Resources, Writing - Review and Editing. RL: Resources, Writing - Review and Editing. SB: Resources, Writing - Review and Editing. MBM: Writing - Review and Editing. SMM: Writing - Review and Editing. MMF: Conceptualization, Validation, Writing - Review and Editing, Supervision, Project administration. All authors contributed to the article and approved the submitted version

REFERENCES

- Ostrom QT, Gittleman H, Truitt G, Boscia A, Kruchko C, Barnholtz-Sloan JS. CBTRUS Statistical Report: Primary Brain and Other Central Nervous System Tumors Diagnosed in the United States in 2011-2015. *Neuro Oncol* (2018) 20: iv1–iv86. doi: 10.1093/neuonc/noy131
- Nayak L, Reardon DA. High-grade Gliomas. Continuum (Minneap Minn) (2017) 23:1548–63. doi: 10.1212/CON.00000000000554
- Kortmann R-D, Jeremic B, Weller M, Plasswilm L, Bamberg M. Radiochemotherapy of malignant glioma in adults. Clinical experiences. *Strahlenther Onkol* (2003) 179:219–32. doi: 10.1007/s00066-003-1027-y
- Kirkpatrick JP, Sampson JH. Recurrent malignant gliomas. Semin Radiat Oncol (2014) 24:289–98. doi: 10.1016/j.semradonc.2014.06.006
- Martínez-Carrillo M, Tovar-Martín I, Zurita-Herrera M, Del Moral-Ávila R, Guerrero-Tejada R, Saura-Rojas E, et al. Salvage radiosurgery for selected patients with recurrent malignant gliomas. *BioMed Res Int* (2014) 2014:657953. doi: 10.1155/2014/657953
- Seystahl K, Wick W, Weller M. Therapeutic options in recurrent glioblastoma-An update. Crit Rev Oncol Hematol (2016) 99:389-408. doi: 10.1016/j.critrevonc.2016.01.018
- Birk HS, Han SJ, Butowski NA. Treatment options for recurrent high-grade gliomas. CNS Oncol (2017) 6:61–70. doi: 10.2217/cns-2016-0013
- Rades D, Witteler J, Leppert J, Schild SE. Re-Irradiation for Recurrent Glioblastoma Multiforme. Anticancer Res (2020) 40:7077–81. doi: 10.21873/ anticanres.14735
- Kaul D, Pudlitz V, Böhmer D, Wust P, Budach V, Grün A. Reirradiation of High-Grade Gliomas: A Retrospective Analysis of 198 Patients Based on the Charité Data Set. Adv Radiat Oncol (2020) 5:959–64. doi: 10.1016/ j.adro.2020.06.005
- Gzell C, Back M, Wheeler H, Bailey D, Foote M. Radiotherapy in Glioblastoma: the Past, the Present and the Future. *Clin Oncol (R Coll Radiol)* (2017) 29:15–25. doi: 10.1016/j.clon.2016.09.015
- Vordermark D, Kölbl O, Ruprecht K, Vince GH, Bratengeier K, Flentje M. Hypofractionated stereotactic re-irradiation: treatment option in recurrent malignant glioma. *BMC Cancer* (2005) 5:55. doi: 10.1186/1471-2407-5-55
- Chang SD, Main W, Martin DP, Gibbs IC, Heilbrun MP. An analysis of the accuracy of the CyberKnife: a robotic frameless stereotactic radiosurgical system. *Neurosurgery* (2003) 52:140–6; discussion 146-7. doi: 10.1097/ 00006123-200301000-00018
- Yu C, Main W, Taylor D, Kuduvalli G, Apuzzo MLJ, Adler JR. An anthropomorphic phantom study of the accuracy of Cyberknife spinal radiosurgery. *Neurosurgery* (2004) 55:1138–49. doi: 10.1227/01.neu. 0000141080.54647.11
- Yan H, Yin F-F, Kim JH. A phantom study on the positioning accuracy of the Novalis Body system. *Med Phys* (2003) 30:3052–60. doi: 10.1118/1.1626122
- Home. CyberKnife. Available at: https://cyberknife.com/ (Accessed October 11, 2020).
- Pantelis E, Moutsatsos A, Antypas C, Zoros E, Pantelakos P, Lekas L, et al. On the total system error of a robotic radiosurgery system: phantom measurements, clinical evaluation and long-term analysis. *Phys Med Biol* (2018) 63:165015. doi: 10.1088/1361-6560/aad516
- Moher D, Shamseer L, Clarke M, Ghersi D, Liberati A, Petticrew M, et al. Preferred reporting items for systematic review and meta-analysis protocols (PRISMA-P) 2015 statement. *Systematic Rev* (2015) 4:1. doi: 10.1186/2046-4053-4-1
- Louis DN, Ohgaki H, Wiestler OD, Cavenee WK. WHO Classification of Tumours of the Central Nervous System. Available at: https://publications.iarc. fr/Book-And-Report-Series/Who-Classification-Of-Tumours/WHO-Classification-Of-Tumours-Of-The-Central-Nervous-System-2016 (Accessed July 19, 2020).
- Southwell DG, Birk HS, Han SJ, Li J, Sall JW, Berger MS. Resection of gliomas deemed inoperable by neurosurgeons based on preoperative imaging studies. *J Neurosurg* (2017) 129:567–75. doi: 10.3171/2017.5.JNS17166
- Greenspoon JN, Sharieff W, Hirte H, Overholt A, Devillers R, Gunnarsson T, et al. Fractionated stereotactic radiosurgery with concurrent temozolomide chemotherapy for locally recurrent glioblastoma multiforme: a prospective cohort study. Onco Targets Ther (2014) 7:485–90. doi: 10.2147/OTT.S60358

- Yazici G, Cengiz M, Ozyigit G, Eren G, Yildiz F, Akyol F, et al. Hypofractionated stereotactic reirradiation for recurrent glioblastoma. J Neurooncol (2014) 120:117–23. doi: 10.1007/s11060-014-1524-0
- 22. Fetcko K, Lukas RV, Watson GA, Zhang L, Dey M. Survival and complications of stereotactic radiosurgery: A systematic review of stereotactic radiosurgery for newly diagnosed and recurrent high-grade gliomas. *Medicine* (2017) 96:e8293. doi: 10.1097/MD.00000000008293
- Lamborn KR, Yung WKA, Chang SM, Wen PY, Cloughesy TF, DeAngelis LM, et al. Progression-free survival: An important end point in evaluating therapy for recurrent high-grade gliomas. *Neuro Oncol* (2008) 10:162–70. doi: 10.1215/15228517-2007-062
- Yoshikawa K, Saito K, Kajiwara K, Nomura S, Ishihara H, Suzuki M. CyberKnife Stereotactic Radiotherapy for Patients with Malignant Glioma. *Minim Invasive Neurosurg* (2006) 49:110–5. doi: 10.1055/s-2006-932183
- Torok JA, Wegner RE, Mintz AH, Heron DE, Burton SA. Re-irradiation with Radiosurgery for Recurrent Glioblastoma Multiforme. *Technol Cancer Res Treat* (2011) 10:253–8. doi: 10.7785/tcrt.2012.500200
- Conti A, Pontoriero A, Arpa D, Siragusa C, Tomasello C, Romanelli P, et al. Efficacy and toxicity of CyberKnife re-irradiation and "dose dense" temozolomide for recurrent gliomas. *Acta Neurochir* (2012) 154:203–9. doi: 10.1007/s00701-011-1184-1
- Ottawa Hospital Research Institute. Available at: http://www.ohri.ca/ programs/clinical_epidemiology/oxford.asp (Accessed December 8, 2020).
- Adachi K, Hayashi K, Kagawa N, Kinoshita M, Sumida I, Akino Y, et al. Feasibility of Salvage Re-irradiation With Stereotactic Radiotherapy for Recurrent Glioma Using CyberKnife. *Anticancer Res* (2019) 39:2935–40. doi: 10.21873/anticanres.13423
- Ekici K, Ozseker N, Mayadagli A, Erdogan Kocak M, Olmezoglu A. Efficacy of stereotactic radiotherapy as salvage treatment for recurrent malignant gliomas. J BUON (2014) 19:1029–34.
- Glavatskyi O, Buryk VM, Kardash KA, Pylypas OP, Chebotaryova TI. P13.11
 * USAGE OF CYBER KNIFE HYPOFRACTIONATED RADIOSURGERY IN HIGH GRADE GLIOMAS COMPLEX TREATMENT. *Neuro Oncol* (2014) 16:ii68–8. doi: 10.1093/neuonc/nou174.257
- 31. Hasan S, Chen E, Lanciano R, Yang J, Hanlon A, Lamond J, et al. Salvage Fractionated Stereotactic Radiotherapy with or without Chemotherapy and Immunotherapy for Recurrent Glioblastoma Multiforme: A Single Institution Experience. Front Oncol (2015) 5:1–11. doi: 10.3389/fonc.2015.00106
- 32. Lévy S, Chapet S, Scher N, Debbi K, Ruffier A, Bernadou G, et al. Reirradiation of gliomas under stereotactic conditions: Prognostic factors for survival without relapse or side effects, a retrospective study at Tours regional university hospital (France). *Cancer Radiother* (2017) 21:759–65. doi: 10.1016/j.canrad.2017.05.006
- Pinzi V, Orsi C, Marchetti M, Milanesi IM, Bianchi LC, DiMeco F, et al. Radiosurgery reirradiation for high-grade glioma recurrence: a retrospective analysis. *Neurol Sci* (2015) 36:1431–40. doi: 10.1007/s10072-015-2172-7
- Villavicencio AT, Burneikienė S, Romanelli P, McNeely L, Lipani JD, Fariselli L, et al. Survival following Stereotactic Radiosurgery for Newly Diagnosed and Recurrent Glioblastoma Multiforme: A Multicenter Experience. *Radiosurgery* (2010) 7:288–99. doi: 10.1159/000288740
- Ostrom QT, Gittleman H, Stetson L, Virk SM, Barnholtz-Sloan JS. Epidemiology of gliomas. *Cancer Treat Res* (2015) 163:1–14. doi: 10.1007/ 978-3-319-12048-5_1
- Gould J. Breaking down the epidemiology of brain cancer. *Nature* (2018) 561: S40–1. doi: 10.1038/d41586-018-06704-7
- Matteoni S, Abbruzzese C, Villani V, Malorni W, Pace A, Matarrese P, et al. The influence of patient sex on clinical approaches to malignant glioma. *Cancer Lett* (2020) 468:41–7. doi: 10.1016/j.canlet.2019.10.012
- Dunford A, Weinstock DM, Savova V, Schumacher SE, Cleary JP, Yoda A, et al. Tumor-suppressor genes that escape from X-inactivation contribute to cancer sex bias. *Nat Genet* (2017) 49:10–6. doi: 10.1038/ng.3726
- Barbagallo GMV, Jenkinson MD, Brodbelt AR. 'Recurrent' glioblastoma multiforme, when should we reoperate? Br J Neurosurg (2008) 22:452–5. doi: 10.1080/02688690802182256
- Montemurro N, Perrini P, Blanco MO, Vannozzi R. Second surgery for recurrent glioblastoma: A concise overview of the current literature. *Clin Neurol Neurosurg* (2016) 142:60–4. doi: 10.1016/j.clineuro.2016.01.010

- Park JK, Hodges T, Arko L, Shen M, Dello Iacono D, McNabb A, et al. Scale to predict survival after surgery for recurrent glioblastoma multiforme. J Clin Oncol (2010) 28:3838–43. doi: 10.1200/JCO.2010.30.0582
- Park C-K, Kim JH, Nam D-H, Kim C-Y, Chung S-B, Kim Y-H, et al. A practical scoring system to determine whether to proceed with surgical resection in recurrent glioblastoma. *Neuro Oncol* (2013) 15:1096–101. doi: 10.1093/neuonc/not069
- Nieder C, Andratschke NH, Grosu AL. Re-irradiation for Recurrent Primary Brain Tumors. Anticancer Res (2016) 36:4985–95. doi: 10.21873/anticanres. 11067
- 44. Clarke J, Neil E, Terziev R, Gutin P, Barani I, Kaley T, et al. Multicenter, Phase 1, Dose Escalation Study of Hypofractionated Stereotactic Radiation Therapy With Bevacizumab for Recurrent Glioblastoma and Anaplastic Astrocytoma. *Int J Radiat Oncol Biol Phys* (2017) 99:797–804. doi: 10.1016/ j.ijrobp.2017.06.2466
- Hu Y, Chen D, Zhang L, Chen J. Efficacy and Safety of Hypofractionated Stereotactic Radiotherapy for Recurrent Malignant Gliomas: A Systematic Review and Meta-analysis. World Neurosurg (2019) 127:176–85. doi: 10.1016/ j.wneu.2019.03.297
- 46. Vincenzo Barbagallo GM, Certo F, Di Gregorio S, Maione M, Garozzo M, Peschillo S, et al. Recurrent high-grade glioma surgery: a multimodal intraoperative protocol to safely increase extent of tumor resection and analysis of its impact on patient outcome. *Neurosurg Focus* (2021) 50:E20. doi: 10.3171/2020.10.FOCUS20744
- 47. Murai T, Sato K, Iwabuchi M, Manabe Y, Ogino H, Iwata H, et al. Reirradiation of recurrent anaplastic ependymoma using radiosurgery or fractionated stereotactic radiotherapy. *Jpn J Radiol* (2016) 34:211–8. doi: 10.1007/s11604-015-0511-5
- Iqbal MS, Lewis J. An overview of the management of adult ependymomas with emphasis on relapsed disease. *Clin Oncol (R Coll Radiol)* (2013) 25:726– 33. doi: 10.1016/j.clon.2013.07.009
- Rodríguez D, Cheung MC, Housri N, Quinones-Hinojosa A, Camphausen K, Koniaris LG. Outcomes of malignant CNS ependymomas: an examination of 2408 cases through the Surveillance, Epidemiology, and End Results (SEER) database (1973-2005). J Surg Res (2009) 156:340–51. doi: 10.1016/ j.jss.2009.04.024
- Bouffet E, Hawkins CE, Ballourah W, Taylor MD, Bartels UK, Schoenhoff N, et al. Survival benefit for pediatric patients with recurrent ependymoma treated with reirradiation. *Int J Radiat Oncol Biol Phys* (2012) 83:1541–8. doi: 10.1016/j.ijrobp.2011.10.039
- Hu Y, Zhang L, Ding C, Chen D, Chen J. Hypofractionated stereotactic radiotherapy combined with chemotherapy or not in the management of recurrent malignant gliomas: A systematic review and meta-analysis. *Clin Neurol Neurosurg* (2019) 183:105401. doi: 10.1016/j.clineuro.2019. 105401
- Chamberlain MC. Salvage therapy with lomustine for temozolomide refractory recurrent anaplastic astrocytoma: a retrospective study. *J Neurooncol* (2015) 122:329–38. doi: 10.1007/s11060-014-1714-9
- 53. Taal W, Oosterkamp HM, Walenkamp AME, Dubbink HJ, Beerepoot LV, Hanse MCJ, et al. Single-agent bevacizumab or lomustine versus a combination of bevacizumab plus lomustine in patients with recurrent glioblastoma (BELOB trial): a randomised controlled phase 2 trial. *Lancet* Oncol (2014) 15:943–53. doi: 10.1016/S1470-2045(14)70314-6
- 54. Stupp R, Taillibert S, Kanner AA, Kesari S, Steinberg DM, Toms SA, et al. Maintenance Therapy With Tumor-Treating Fields Plus Temozolomide vs Temozolomide Alone for Glioblastoma: A Randomized Clinical Trial. JAMA (2015) 314:2535–43. doi: 10.1001/jama.2015.16669
- Wick W, Gorlia T, Bendszus M, Taphoorn M, Sahm F, Harting I, et al. Lomustine and Bevacizumab in Progressive Glioblastoma. N Engl J Med (2017) 377:1954–63. doi: 10.1056/NEJMoa1707358
- 56. Kreisl TN, Kim L, Moore K, Duic P, Royce C, Stroud I, et al. Phase II trial of single-agent bevacizumab followed by bevacizumab plus irinotecan at tumor progression in recurrent glioblastoma. J Clin Oncol (2009) 27:740–5. doi: 10.1200/JCO.2008.16.3055
- Gutin PH, Iwamoto FM, Beal K, Mohile NA, Karimi S, Hou BL, et al. Safety and efficacy of bevacizumab with hypofractionated stereotactic irradiation for recurrent malignant gliomas. *Int J Radiat Oncol Biol Phys* (2009) 75:156–63. doi: 10.1016/j.ijrobp.2008.10.043

- Cabrera AR, Cuneo KC, Desjardins A, Sampson JH, McSherry F, Herndon JE, et al. Concurrent Stereotactic Radiosurgery and Bevacizumab in Recurrent Malignant Gliomas: A Prospective Trial. *Int J Radiat Oncol Biol Phys* (2013) 86:873–9. doi: 10.1016/j.ijrobp.2013.04.029
- Shapiro LQ, Beal K, Goenka A, Karimi S, Iwamoto FM, Yamada Y, et al. Patterns of Failure After Concurrent Bevacizumab and Hypofractionated Stereotactic Radiation Therapy for Recurrent High-Grade Glioma. *Int J Radiat Oncol Biol Phys* (2013) 85:636–42. doi: 10.1016/j.ijrobp.2012.05.031
- 60. Radiation Therapy Oncology Group. Randomized Phase II Trial of Concurrent Bevacizumab and Re-Irradiation Versus Bevacizumab Alone as Treatment for Recurrent Glioblastoma(2020). Available at: https:// clinicaltrials.gov/ct2/show/NCT01730950 (Accessed February 24, 2021). clinicaltrials.gov.
- Palmer JD, Bhamidipati D, Song A, Eldredge-Hindy HB, Siglin J, Dan TD, et al. Bevacizumab and re-irradiation for recurrent high grade gliomas: does sequence matter? *J Neurooncol* (2018) 140:623–8. doi: 10.1007/s11060-018-2989-z
- 62. RTOG Foundation. Available at: https://www.rtog.org/ (Accessed March 2, 2021).
- Shaw E, Scott C, Souhami L, Dinapoli R, Kline R, Loeffler J, et al. Single dose radiosurgical treatment of recurrent previously irradiated primary brain tumors and brain metastases: final report of RTOG protocol 90-05. *Int J Radiat Oncol Biol Phys* (2000) 47:291–8. doi: 10.1016/s0360-3016(99) 00507-6
- Combs SE, Thilmann C, Edler L, Debus J, Schulz-Ertner D. Efficacy of fractionated stereotactic reirradiation in recurrent gliomas: long-term results in 172 patients treated in a single institution. J Clin Oncol (2005) 23:8863–9. doi: 10.1200/JCO.2005.03.4157
- Hall WA, Djalilian HR, Sperduto PW, Cho KH, Gerbi BJ, Gibbons JP, et al. Stereotactic radiosurgery for recurrent malignant gliomas. J Clin Oncol (1995) 13:1642–8. doi: 10.1200/JCO.1995.13.7.1642
- Mayer R, Sminia P. Reirradiation tolerance of the human brain. Int J Radiat Oncol Biol Phys (2008) 70:1350–60. doi: 10.1016/j.ijrobp.2007.08.015
- Common Terminology Criteria for Adverse Events (CTCAE) | Protocol Development | CTEP. Available at: https://ctep.cancer.gov/protocol development/electronic_applications/ctc.htm (Accessed March 2, 2021).
- Hoffman WF, Levin VA, Wilson CB. Evaluation of malignant glioma patients during the postirradiation period. *J Neurosurg* (1979) 50:624–8. doi: 10.3171/ jns.1979.50.5.0624
- Chamberlain MC, Glantz MJ, Chalmers L, Van Horn A, Sloan AE. Early necrosis following concurrent Temodar and radiotherapy in patients with glioblastoma. *J Neurooncol* (2007) 82:81–3. doi: 10.1007/s11060-006-9241-y
- de Wit MCY, de Bruin HG, Eijkenboom W, Sillevis Smitt P a E, van den Bent MJ. Immediate post-radiotherapy changes in malignant glioma can mimic tumor progression. *Neurology* (2004) 63:535–7. doi: 10.1212/ 01.wnl.0000133398.11870.9a
- Brandes AA, Bartolotti M, Franceschi E. Second surgery for recurrent glioblastoma: advantages and pitfalls. *Expert Rev Anticancer Ther* (2013) 13:583–7. doi: 10.1586/era.13.32
- Ellingson BM, Wen PY, Cloughesy TF. Modified Criteria for Radiographic Response Assessment in Glioblastoma Clinical Trials. *Neurotherapeutics* (2017) 14:307–20. doi: 10.1007/s13311-016-0507-6
- Brady LW, Perez CA, Wazer DE. Perez & Brady"s Principles and Practice of Radiation Oncology. Alphen aan den Rijn Netherlands: Lippincott Williams & Wilkins (2013).
- Weitzner MA, Meyers CA. Cognitive functioning and quality of life in malignant glioma patients: a review of the literature. *Psychooncology* (1997) 6:169–77. doi: 10.1002/(SICI)1099-1611(199709)6:3<169::AID-PON269>3.0.CO;2-
- Ernst-Stecken A, Ganslandt O, Lambrecht U, Sauer R, Grabenbauer G. Survival and quality of life after hypofractionated stereotactic radiotherapy for recurrent malignant glioma. *J Neurooncol* (2007) 81:287–94. doi: 10.1007/ s11060-006-9231-0
- 76. Maitre P, Gupta T, Maitre M, Goda J, Krishnatry R, Chatterjee A, et al. Prospective Longitudinal Assessment of Quality of Life and Activities of Daily Living as Patient-Reported Outcome Measures in Recurrent/Progressive Glioma Treated with High-dose Salvage Re-irradiation. *Clin Oncol (Royal Coll Radiol)* (2021) 33:1–4. doi: 10.1016/j.clon.2020.08.011

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2021 De Maria, Terzi di Bergamo, Conti, Hayashi, Pinzi, Murai, Lanciano, Burneikiene, Buglione di Monale, Magrini and Fontanella. This is an open-

access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.