Age-stratified clinical performance and survival of patients with IDH-wildtype glioblastoma homogeneously treated by radiotherapy with concomitant and maintenance temozolomide

Kerstin Berger1 · Bernd Turowski2 · Jörg Felsberg3 · Bastian Malzkorn3 · Guido Reifenberger3 · Hans-Jakob Steiger1 · Wilfried Budach4 · Jan Haussmann4 · Johannes Knipps1 · Marion Rapp1 · Daniel Hänggi1 · Michael Sabel1 · Hendrik-Jan Mijderwijk1 · Marcel A. Kamp1

Received: 9 April 2020 / Accepted: 22 July 2020 / Published online: 3 August 2020
© The Author(s) 2020

Abstract
Objective Isocitrate dehydrogenase (IDH)-wildtype glioblastomas are the most malignant glial tumours. Median survival is only 14–16 months after diagnosis, with patients aged ≥ 65 years reportedly showing worse outcome. This study aimed to further evaluate the prognostic role of age in a homogenously treated patient cohort.

Methods The study includes 132 IDH-wildtype glioblastoma patients treated between 2013 and 2017 with open resection followed by radiotherapy with concomitant and maintenance temozolomide. Patients were dichotomized into a non-elderly (< 65 years) and an elderly (≥ 65 years) group. Extent of resection and the O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation status were determined for each tumour. Clinical and radiological follow-up data were obtained at 6 weeks after the end of radiation therapy and thereafter in 3-month intervals. Progression-free survival (PFS) and overall survival (OS) were evaluated in univariate and multivariate cox regression analyses.

Results The elderly group consisted of 58 patients (median age: 70.5 years) and the non-elderly group of 74 patients (median age: 55 years). Median pre- and postoperative operative Karnofsky Performance Scale (KPS), Eastern Cooperative Oncology Group (ECOG) score and National Institutes of Stroke Scale (NIHSS) were not significantly different between the groups, but KPS and ECOG scores became significantly worse in the elderly group at 6 weeks after termination of radiation therapy. Neither PFS nor OS differed significantly between the age groups. Patients with MGMT promoter-methylated tumours survived longer.

Conclusion Elderly patients in good pre- and postoperative clinical conditions may show similar outcome as younger patients when treated according to standard of care. However, elderly patients may suffer more frequently from clinical deterioration following chemoradiotherapy. In both age groups, MGMT promoter methylation was linked to longer PFS and OS.

Keywords Age at diagnosis · Glioblastoma · MGMT promoter methylation · Chemoradiotherapy · Survival

Introduction
Isocitrate dehydrogenase (IDH)-wildtype glioblastomas are the most frequent and most malignant intrinsic brain tumours that lead to an average survival of only three months without treatment (Malmstrom A et al. 2012; Tamimi and Juweid 2017). The incidence of IDH-wildtype glioblastoma increases with patient age and peaks between 65 and 75 years of age (Brodbelt et al. (2015); Thakkar et al. 2014). More than half of the patients are older than 64 years at the time of diagnosis (Brodbelt et al. (2015); Thakkar et al. 2014).
Standard treatment of glioblastoma patients is, if feasible, a gross total resection of the tumour followed by fractionated tumour irradiation up to 60 Gy with concomitant systemic chemotherapy with temozolomide (TMZ) followed by six cycles of maintenance TMZ therapy (Stummer and Kamp 2009; Weller et al. 2017a). This combined chemoradiotherapy prolonged overall survival as compared to radiation therapy alone in glioblastoma patients younger than 70 years in the European Organization for Research and Treatment of Cancer (EORTC)-22,981/26,981/National Cancer Institute of Canada (NCIC) CE3 trial (Stupp et al. 2005, 2009). The benefit from combined chemoradiotherapy has been reported to decrease with increasing age of the patients (Stupp et al. 2005, 2009), a finding that might be related to an increased toxicity of the therapy in elderly patients who commonly suffer from comorbidities (Sijben et al. 2008; Tabatabai et al. 2013). In addition, differences in tumour biology, such as higher prevalence of IDH mutations in glioblastomas of younger as compared to older patients, might contribute to prognostic differences, in particular in studies that did not stratify according to IDH mutation status (Eckel-Passow et al. 2015; Hartmann et al. 2010; Houillier et al. 2010; Parsons et al. 2008). Two prospective phase III clinical trials have analysed the impact of either chemotherapy with TMZ or radiotherapy alone in the treatment of elderly glioblastoma patients (Hartmann et al. 2010; Perry et al. 2017). The NOA-08 trial compared TMZ chemotherapy (100 mg/m² TMZ, one week on/1 week off protocol) with radiotherapy (30 × 1.8–2.0 Gy ad 60 Gy) in glioblastoma patients with a Karnofsky Performance Score (KPS) > 50 and an age > 65 years. Extent of resection was identified as an independent prognostic factor. Adjuvant chemotherapy was not inferior to radiotherapy but its effectiveness was related to the O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation status (Wick et al. 2012). In elderly glioblastoma patients with MGMT promoter-methylated glioblastoma, TMZ chemotherapy was associated with a longer event-free survival and a trend towards longer overall survival (OS) as compared to radiation therapy alone. The opposite was true for elderly glioblastoma with MGMT promoter- unmethylated glioblastoma (Wick et al. 2012). In the Nordic trial (Perry et al. 2017), TMZ chemotherapy alone (200 mg/m²; 5/28 day cycle), radiation therapy (30 × 2 Gy ad 60 Gy) and hypofractionated radiotherapy (10 × 3.4 Gy ad 34 Gy) were compared in newly diagnosed glioblastoma patients who were older than 60 years at diagnosis. Again, chemotherapy alone was not inferior to radiation therapy alone in terms of overall survival and the MGMT promoter methylation status was a valuable marker for response to TMZ chemotherapy. For patients older than 70 years, OS was prolonged in the chemotherapy and hypofractionated radiotherapy groups when compared to the standard radiotherapy group (Perry et al. 2017). The CCTG CE.6/EORTC 26,062–22,061 phase III trial randomized newly diagnosed glioblastoma patients aged 65 years or older to hypofractionated radiotherapy (40 Gy/15 fractions) alone versus hypofractionated radiotherapy with concomitant and adjuvant temozolomide (Perry et al. 2017). The results of this study suggested that the addition of temozolomide to short-course radiotherapy resulted in longer survival than short-course radiotherapy alone. According to the European Association for Neuro-Oncology (EANO) guideline of 2017 (Hegi et al. 2005), treatment of glioblastoma patients should consider molecular biomarkers like IDH mutation and MGMT promoter methylation. Patients diagnosed with IDH-mutant glioblastoma should be treated with surgery and radiotherapy with or without concomitant TMZ followed by TMZ regardless of age. In case of IDH-wildtype glioblastoma, patients aged ≥ 70 years should be treated with surgery and hypofractionated radiotherapy in case of an MGMT promoter unmethylated tumour, while elderly patients with an MGMT promoter-methylated tumour should be treated by surgery and TMZ plus hypofractionated radiotherapy or surgery and TMZ alone (Weller et al. 2017b). As elderly patients more often suffer from other comorbidities and higher frailty when compared to younger patients, therapeutic decisions need to be adopted to their general health status. On the other hand, elderly patients in good general condition and without any relevant comorbidities may be suitable to standard treatment according to the EORTC-22981/NCIC CE3 protocol (Stupp et al. 2005).

The aim of the present study was to analyse survival, perioperative neurologic functioning and complication rates following multimodal initial therapy in an institutional cohort of non-elderly (< 65 years) and elderly (≥ 65 years) glioblastoma patients who otherwise were in good clinical condition and were treated uniformly with open resection followed by chemoradiotherapy according to the EORTC 26,981 NCIC CE3 protocol (Felsberg et al. 2010; Glaser et al. 2017).

Materials and methods

Study design, inclusion and exclusion criteria

Patients fulfilling the following inclusion criteria were considered for this study and their data were retrospectively analysed: (1) Surgical treatment at the Department of Neurosurgery, Heinrich Heine University Düsseldorf, between 01/2013 and 12/2017, (2) neuropathologically confirmed diagnosis of an IDH-wildtype glioblastoma, World Health Organization (WHO) grade IV (Louis et al. 2016), (3) primary surgical resection or surgical resection within four weeks following initial biopsy, (4) preoperative Karnofsky performance scale (KPS) ≥ 70% and (5) postsurgical
therapy according to the EORTC-22981/NCIC CE3 protocol. Exclusion criteria included (1) other histopathological diagnoses than glioblastoma, IDH-wildtype (2) glioblastoma in patients with a preceding diagnosis of a WHO grade II or III glioma, (3) primary biopsy without resection within 4 weeks, (4) preoperative KPS < 70%, (5) postsurgical therapy different from the EORTC-22981/NCIC CE3 protocol (e.g., chemotherapy with TMZ and lomustine (CCNU), application of tumour-treating fields, either chemotherapy or radiation therapy alone), (6) treatment at another institution (e.g. first biopsy or surgery at another neurosurgical department).

**Surgical and postoperative treatment**

All patients underwent surgical resection with standard 5-aminolevulinic acid (5-ALA) fluorescence-guided resection and intraoperative neuro-navigation. For tumours located in eloquent brain regions, surgery was planned with intraoperative monitoring and as awake surgery in an asleep–awake–asleep protocol (Kamp et al. 2012,2015b). Eloquent brain regions were defined as cortical or subcortical brain areas for which intraoperative stimulation was expected to elicit changes in neurologic condition (particularly regarding speech, movement and tactile sensation) or to elicit a response in electrophysiological recordings in corresponding areas (Kamp et al. 2015b). Extent of surgical resection was postoperatively determined by contrast-enhanced magnetic resonance imaging (MRI) within 72 h after surgery (Kamp et al. 2015a). All patients received TMZ chemoradiotherapy according to the EORTC protocol (Kocher et al. 2011) as initial treatment following resection, with at least one cycle of maintenance TMZ chemotherapy (median: six cycles, range 1–6 cycles). Radiotherapy was administrated as standard fractionated therapy (30 × 2 Gy ad 60 Gy). Follow-up consisted of regular clinical and radiological re-assessment 6 weeks after end of radiotherapy and thereafter every 3 months after diagnosis.

**Histopathological and molecular analyses**

All tumours were neuropathologically classified as glioblastoma, IDH-wildtype, WHO grade IV according to the WHO classification of central nervous system tumors 2016 (Louis et al. 2016). Tumours from patients diagnosed before 2016 were neuropathologically re-evaluated and reclassified according to the WHO 2016 criteria. The IDH mutation status was assessed by immunohistochemistry for IDH1-R132H as reported (Felsberg et al. 2010; Hartmann et al. 2010). Tumours of patients younger than 55 years of age were additionally investigated for less common mutations at codon 132 of IDH1 and codon 172 of IDH2 by Sanger sequencing or pyrosequencing as reported (Felsberg et al. 2010). The MGMT promoter methylation status was determined by methylation-specific PCR and pyrosequencing of sodium bisulphite-treated DNA as reported (Felsberg et al. 2009,2011).

**Study variables and neuroimaging**

Preoperative Karnofsky Performance Scale (KPS), Eastern Co-operative Oncology Group Score (ECOG) and the National Institute of Health Stroke Scale (NIHSS) were determined at initial admission and postoperative KPS at discharge from the Department of Neurosurgery (Karnofsky et al. 1949) (Goldstein et al. 1989; Verger et al. 1992). In addition, KPS was assessed at follow-up visits at the department’s outpatient’s office.

All MRI images were obtained by contrast-enhanced 1.5 T MRI (Avanto; Siemens, Erlangen, Germany). For detection of residual tumour tissue after surgery or for diagnosis of disease progression/tumour recurrence, non-contrast-enhanced and contrast-enhanced T1- and T2-weighted, diffusion and fluid attenuated inversion recovery sequences were evaluated. Evaluations of the MR images were performed by an attending neurosurgeon and a neuroradiologist. The extent of surgical resection was assessed by MRI within 72 h after surgery. A complete surgical resection (CR) was defined as a complete resection of the contrast-enhancing tumour tissue and was distinguished from subtotal resection (SR), in which residual contrast-enhancing tumour tissue was present in postoperative MRI. A progression/tumour recurrence was diagnosed when Response Assessment in Neuro-Oncology (RANO) criteria were fulfilled (Kamp et al. 2015a). In 81 patients (61%), additional investigations by FET-PET were performed to confirm evidence of tumour recurrence (Kamp et al. 2015b). Moreover, 77 patients were re-operated for recurrent disease and neuropathological analysis confirmed presence of recurrent tumour in 70 of these patients. 47 patients received an additional adjuvant therapy after diagnosis of tumour recurrence. Adjuvant therapy was only radiation therapy in ten patients, only chemotherapy in 21 (5/23 TMZ chemotherapy in 12 patients, weekly TMZ in seven patients, procarbazine / CCNU in two patients, only bevacizumab in one patient). One patient got only Tumour Treating Fields and 14 patients a combination of radiation and TMZ chemotherapy.

**Outcome variables**

Progression-free survival was defined as the time span between initial surgery and diagnosis of tumour progression on MRI. Overall survival was defined as the time span between initial surgery and tumour-related death. Patients with unknown date of death were censored at the time of last follow-up.
Data management

Demographic data including information on age at diagnosis and gender, KPS pre- and post-surgery as well as at 6-week follow-up following radiation, extent of resection, PFS and OS were collected retrospectively from patients’ charts. The follow-up ended on January 29, 2019 and all patients who were still alive were censored at the date of last follow-up. Continuous variables are presented as mean ± standard error of mean, ordinal values were presented as median values and minimum–maximum ranges.

All patients that fulfilled the inclusion criteria were divided into two groups, those < 65 years and those ≥ 65 years of age. The two groups were statistically compared.

Statistical analysis

For continuous data, median and interquartile range are presented. For categorical data, frequencies and percentages are presented. Kaplan–Meier survival analysis including log-rank test and Cox regression analyses are used for statistical significance testing. Sidak’s correction was applied to adjust for multiplicity. Therefore, for statistical significance, P values were considered significant at a level of <0.003. A tendency towards a correlation was defined for p values between 0.05 and 0.003. All statistical analyses were performed with SPSS software (Version 25.0,—IBM-, USA) and the Graph Pad Prism 5 package 3.3.2 (GraphPad Software, Inc., La Jolla, USA).

Results

Patient cohort

Between 2013 and 2017, 132 patients among a total of 683 glioblastoma patients (including recurrent glioblastoma) treated at our centre during this time period met the inclusion criteria for this study. The median age of the 132 patients at diagnosis was 61 (range 22–83 years). 74 patients were younger than 65 years and 58 patients were 65 years and older. In the group of the elderly patients, the median age was 70.5 years (range: 65–83 years, interquartile range, IQR: 10). Median age in the patients aged < 65 years was 55 years (range: 22–64 years, IQR: 10). In both groups, males dominated the study population: 45 (61%) in the non-elderly group and 34 (58%) in the elderly group. All patients were diagnosed with glioblastoma, IDH-wildtype, WHO grade IV according to the 2016 WHO classification of central nervous system tumours (Louis et al. 2016). MGMT promoter methylation was detected in 54/132 tumours (41%), including 23/74 tumours (31%) in the group of patients aged < 65y and 31/58 tumours (53%) in the elderly patient group.

Clinical performance and adverse effects of therapy

The median preoperative KPS was 90% for both groups (IQR: 10) and was unaffected at the day of hospital discharge after initial resection. The pre- and postoperative ECOG and NIHSS were 0 and 1 without any significant differences, respectively (each median 90, IQR: 10). While the NIHSS values 6 weeks after the end of the radiation therapy were not significantly different, KPS and ECOG values were significantly worse in the group of elderly patients at this time point. More detailed information on the neurologic performance is summarized in Table 1; Fig. 1.

All patients received at least one cycle of maintenance TMZ following the concomitant chemoradiotherapy. The median number of administered TMZ cycles in the entire cohort was 6 in both elderly and non-elderly patients (interquartile range 4–6, for the elderly patients: 4–6 and 6–6 for the non-elderly patients). TMZ had been stopped prematurely in 23 elderly (39%) and 18 non-elderly patients (24.7%). Reasons for prematurely stopped chemotherapy were adverse side effects. In total, 32 patients suffered from relevant adverse effects of the adjuvant therapy (23.5%). Among these, 20 patients were ≥ 65 years of age (20/58 elderly patients, 33.9%) and 12 patients were < 64 years of age (12/74 patients < 65 y, 16.2%). Most commonly, patients exhibited an impairment of their general health condition as assessed by an impairment of the KPS of more than 20 points (13/58 patients ≥ 65 y, 24.1%; 6/74 patients; 8.1%, $X^2=6.5, p=0.01$). Nine patients showed haematologic toxicities with severe thrombocytopenia and/or leucocytopenia according to the common toxicity criteria grade three or four (five patients < 65 years, four patients ≥ 65 years). Two patients suffered from grade-four hepatic toxicities (CTC criteria; one patient in each age group). The incidence of haematological and hepatic toxicities did not differ significantly between age groups.

Hazard ratios for progression-free and overall survival

A total of 56 non-elderly patients (76.7%) and 41 elderly patients (70.7%) developed tumour progression within the observation period. The median PFS was 17 months and not significantly different for both age groups (11 months vs. 19 months; $X^2=1.7; p=0.2$). The median OS in the present cohort was 21 months as depicted in the Kaplan–Meier survival curves. OS did not significantly differ between the two age groups (20 months vs. 24 months, Log-Rank $X^2=0.002; p=0.96$). Univariate hazard ratios for overall survival and
progression-free survival are given in Table 2. In the univariate analyses, the degree of surgical resection and MGMT promoter methylation tended to correlate with the progression-free survival (for the degree of surgical resection: hazard ration, HR: 1.8; 95%-CI 1.04–3.16; p = 0.04; for the MGMT promoter methylation HR: 0.5; 95%-CI 0.28–0.9; p = 0.02) and the overall survival (for the degree of surgical resection: HR: 1.3; 95%-CI 0.75–2.28; p = 0.34; for the MGMT promoter methylation HR: 0.46; 95%-CI 0.26–0.83; p = 0.01). Multivariate hazard ratios revealed a tendency towards significance for the effect of the degree of surgical resection on the progression-free survival (for the degree of surgical resection: HR: 2.1; 95%-CI 1.09–4.09; p = 0.03). However, the MGMT promoter methylation was the only analysed variable that had a significant effect on overall survival. In the present cohort, age had neither an effect on the progression-free nor on the overall survival in both statistical analyses. Univariate and multivariate hazard ratios are given in (Table 2, 3).

**Discussion**

The present single-centre study analysed neurological performance and survival of patients diagnosed with IDH-wildtype glioblastoma and treated by surgical resection followed by concomitant and maintenance
chemoradiotherapy according to the Stupp protocol (Felsberg et al. 2010; Glaser et al. 2017). A particular focus was placed on the comparison of groups of patients stratified according to age at diagnosis into elderly (≥ 65 years of age) and non-elderly (< 65 years of age) patients. There are three key findings in this study: (1) Elderly patients suffer more frequently from a deterioration of their general health condition than younger patients following initial multimodal therapy according to Stupp protocol. However, hematologic and hepatic complications of TMZ chemotherapy were similar in both groups. (2) Survival of elderly patients in good clinical condition is similar to survival of non-elderly patients. (3) MGMT promoter methylation was the only analyzed variable with a significant influence on OS in the multivariate analysis (Table 4).

The optimal treatment of elderly patients with glioblastoma is still part of an on-going debate and has been addressed in both retrospective studies as well as prospective clinical trials (Glaser et al. 2017; Harrison and Groot 2018; Karsy et al. 2018; Morgan et al. 2017; Okada et al. 2017; Palmer et al. 2018; Pretanvil et al. 2017; Putz et al. 2016; Socha et al. 2016; Victor et al. 2019; Zhang et al. 2016). The EORTC-22981/26,981 / NCIC CE3 trial led to the establishment of concomitant TMZ chemotherapy followed by TMZ maintenance therapy as standard of care for adult glioblastoma patients aged less than 70 years (Stupp et al. 2005, 2009). The benefit of this multimodal therapy concept, however, decreased with increasing patient age (Sijben et al. 2008; Tabatabai et al. 2013). Two prospective randomized trials, therefore, analysed less-toxic therapy concepts based

### Table 2: Univariate hazard ratios for overall survival and progression free survival

|          | OAS HR 95% CI | PFS HR 95% CI |
|----------|---------------|---------------|
|          | Lower | Upper | p value | Lower | Upper | p value |
| Age      |        |       |         |        |       |         |
| Dichotomized | 1.013 | 0.587 | 1.747 | 0.963 | 0.702 | 1.208 | 0.202 |
| Continuous | 1.003 | 0.98  | 1.026 | 0.805 | 0.99  | 1.011 | 0.351 |
| Gender   | 1.32  | 0.771 | 2.258 | 0.312 | 1.552 | 0.635 | 3.797 | 0.335 |
| KPS      |        |       |         |        |       |         |
| Preoperative | 0.999 | 0.965 | 1.035 | 0.967 | 1.012 | 0.979 | 1.047 | 0.474 |
| Postoperative | 0.981 | 0.95  | 1.014 | 0.257 | 0.995 | 0.964 | 1.028 | 0.774 |
| 3rd Month of follow-up | 0.975 | 0.95  | 1.002 | 0.067 | 0.981 | 0.954 | 1.008 | 0.173 |
| GTR      | 1.308 | 0.751 | 2.277 | 0.343 | 1.808 | 1.036 | 3.156 | 0.037 |
| MGMT    | 0.46  | 0.256 | 0.826 | 0.009 | 0.497 | 0.276 | 0.895 | 0.02 |

OAS overall survival; PFS progression free survival; CI confidence interval; HR hazard ratio; GTR gross total resection (0 = yes, 1 = no); MGMT O6-methylguanine-DNA methyltransferase (0 = unmethylated, 1 = methylated)

### Table 3: Multivariate hazard ratios for overall survival and progression free survival

|          | OAS HR 95% CI | PFS HR 95% CI |
|----------|---------------|---------------|
|          | Lower | Upper | p value | Lower | Upper | p value |
| Age      |        |       |         |        |       |         |
| Dichotomized | 1.023 | 0.979 | 1.068 | 0.319 | 1.013 | 0.975 | 1.052 | 0.511 |
| Continuous | 0.691 | 0.244 | 1.955 | 0.486 | 0.524 | 0.198 | 1.392 | 0.195 |
| Gender   | 1.169 | 0.637 | 2.147 | 0.614 | 1.106 | 0.57  | 2.145 | 0.766 |
| KPS      |        |       |         |        |       |         |
| Preoperative | 1.001 | 0.955 | 1.05  | 0.957 | 1.034 | 0.98  | 1.092 | 0.224 |
| Postoperative | 0.983 | 0.937 | 1.031 | 0.474 | 0.986 | 0.934 | 1.04  | 0.601 |
| 3rd Month of follow-up | 0.956 | 0.926 | 0.988 | 0.006 | 0.964 | 0.932 | 0.997 | 0.033 |
| GTR      | 1.28  | 0.681 | 2.406 | 0.442 | 2.107 | 1.086 | 4.087 | 0.027 |
| MGMT    | 0.353 | 0.18  | 0.689 | 0.002 | 0.53  | 0.273 | 1.03  | 0.061 |

OAS overall survival; PFS progression free survival; CI confidence interval; HR hazard ratio; GTR gross total resection (0 = yes, 1 = no); MGMT O6-methylguanine-DNA methyltransferase (0 = unmethylated, 1 = methylated)
on either TMZ, standard radiotherapy or hypofractionated radiotherapy (Hartmann et al. 2010; Perry et al. 2017). Both studies found that the effectiveness of adjuvant chemotherapy or radiotherapy is closely linked to the MGMT promoter methylation status (Wick et al. 2012). Similarly, a prospective cohort study of the German Glioma Network of 233 glioblastoma patients aged ≥ 70 years revealed longer PFS (5.2 vs. 4.7 months) and OS (8.4 vs. 6.4 months) in patients with MGMT promoter-methylated tumours (Reifenberger et al. 2012). The NORDIC trial also reported that standard radiotherapy may be less favorable compared to chemotherapy or hypofractionated radiotherapy in patients older than 70 years (Malmstrom A, et al. 2012). In a further prospective cohort study derived from the Norwegian Cancer Registry, the median overall survival was 7.4 month in the radiotherapy alone and 13.4 month in the group treated with radiotherapy and TMZ in patients older than 70 years of age (Ronning et al. 2012). In a further prospective cohort study derived from the Norwegian Cancer Registry, the median overall survival was 7.4 month in the radiotherapy alone and 13.4 month in the group treated with radiotherapy and TMZ in patients older than 70 years of age (Ronning et al. 2012). The recent CCTG CE.6/EORTC 26,062–22,061 phase III trial on glioblastoma patients aged ≥ 65 years or older similarly revealed longer survival of patients treated with hypofractionated radiotherapy and TMZ compared to hypofractionated radiotherapy alone (Perry et al. 2017).

Several neuro-oncologic centres offer elderly patients in a good general health condition adjuvant chemoradiotherapy according to the EORTC-22981/26,981/NCIC CE3 protocol (Felsberg et al. 2010; Glaser et al. 2017). Data regarding toxicity and efficacy of this approach in elderly patients with IDH-wildtype glioblastoma are still limited. The present study provides a retrospective analysis of a homogenous cohort of IDH-wildtype glioblastoma patients who were in good clinical condition at the time of diagnosis and were treated homogeneously according to the EORTC protocol. In line with the data in the original data in the EORTC-22981/26,981/NCIC CE3 trial (Stupp et al. 2005, 2009), we observed a higher frequency of deterioration of the general health status in elderly patients and subsequently a higher complication rate following multimodal initial treatment. In contrast, the frequency of haematological and hepatic complication was comparable in elderly patients and non-elderly patients of our cohort. The extent of surgical resection and the MGMT promoter methylation status were found to be prognostic, as documented in previous studies (Hegi et al. 2005, 2008, 2019; Kamp et al. 2018; Reifenberger et al. 2012; Senft et al. 2011; Stummer and Kamp 2009; Stummer et al. 2006). Interestingly, in the present study, the higher complication rate of initial therapy in elderly patients had no effect on PFS and OS, which were comparable in the groups of elderly and non-elderly patients.

The present study, thus, suggests that elderly IDH-wildtype glioblastoma patients in a good clinical condition might benefit from a similarly aggressive therapy as applied to younger patients. Although clinical deterioration following initial therapy was more common in elderly patients, PFS and OS were statistically not affected by this observation. Furthermore, our data confirm MGMT promoter methylation as a powerful predictor of outcome in both elderly and non-elderly glioblastoma patients. As the incidence of glioblastomas and also cerebral metastases increases with age, treatment of elderly brain tumour patients becomes increasingly more relevant in neuro-oncology and novel therapy concepts need to be developed that consider the special needs of the large geronto–neuro–oncological population of patients (Mason et al. 2016; Munoz-Bendix et al. 2019; Tabatabai et al. 2013; Wick et al. 2012; Wirsching et al. 2015).

**Limitations**

We acknowledge that our present study has several limitations: (1) All data are derived from a retrospective, single-centre study. (2) By defining strict inclusion criteria (IDH-wildtype glioblastoma treated by surgical resection

---

**Table 4** TMZ chemotherapy and adverse events

|                          | Patients < 65 years | Patients ≥ 65 years | p value** |
|--------------------------|---------------------|---------------------|-----------|
| TMZ cycles (median (IQR))| 6 (6–6)             | 6 (4–6)             | 0.1713    |
| Number of patients in which TMZ had to be stopped prematurely (%) | X (x) | X (x) |          |
| Adverse effects* (%)     | 12 (16.4)           | 20 (33.9)           |           |
| Impairment of the general health condition (%) radiation therapy | 6 (8.1) | 13 (24.1) | X²=6.5, p=0.01 |
| Haematologic toxicities (%) | 5 (6.8) | 4 (6.8%) | n.a       |
| Hepatic toxicities (%)   | 2 (2.7)             | 2 (3.4)             | n.a       |

KPS, Karnofsky performance score; ECOG Eastern Cooperative Oncology Group scale of performance; NIHSS National Institutes of Health Stroke Scale

*Adverse effects included AZV, leukopenia, leukocytopenia, and thrombocytopenia

**p Values according to the Mann–Whitney test
and standard chemoradiotherapy according to the EORTC-NCIC-protocol, presurgical KPS > 70), we aimed to minimize potential confounders and to constitute a homogenous patient cohort. However, the analysed cohort is highly selected and, therefore, not representative of the entire patient cohort of glioblastoma patients. Furthermore, we cannot exclude potential influences related to a selection bias in this relatively small patient cohort. (3) Not all patients received the same cycles of chemotherapy due to side effects, such as hematotoxicity and other reasons. We have not analysed a potential correlation between the administered amount of temozolomide and the progression-free and overall survival. (4) This cohort includes all patients who were treated in our neuro-oncologic centre between 2013 and 2017 and who met the inclusion criteria. During and after this period, neuro-oncological treatment of glioblastomas has progressed, e.g. the advent of chemotherapy with TMZ and CCNU for patients with MGMT promoter-methylated glioblastoma (Herrlinger et al. 2019) as well as the introduction of tumour-treating fields (Stupp et al. 2017). These concepts were not considered in the current study. “(5) In the present analysis, KPS and ECOG scores were significantly worse in elderly patients 6 weeks after the end of radiotherapy while NIHSS scores did not significantly changed. This difference might in part be related to the different metrics of the respective scales. (6) We did not perform a quality-of-life assessment. Possibly, a more aggressive therapy might lead to an impairment of the patient’s quality of life and some patients, particularly among the elderly patients, might prefer a less aggressive post-surgical therapy associated with lower life expectancy but a higher quality of life. Evaluation of the individual patient’s therapy goals by shared decision-making is additionally essential.

Conclusion

In a retrospective analysis of 132 patients with IDH-wildtype glioblastoma treated by surgical resection and TMZ chemoradiotherapy, patients aged ≥ 65 years showed comparable PFS and OS as patients aged < 65 years. Elderly patients, however, suffered more frequently from a deterioration of their general health condition following aggressive tumour therapy than younger patients. Incidences of hematologic and hepatic toxicities of TMZ were similar in both groups. MGMT promoter methylation had a significant influence on PFS and OS independent from patient age.

Acknowledgements Open Access funding provided by Projekt DEAL.

Funding The study was supported by the institutional budgets from the participating institutions. There was no support by external funding.

Compliance with ethical standards

Conflicts of interest M. Sabel and M. Rapp work as consultants for Johnson and Johnson Company and Integra Company. G. Reifenberger received research grant from Roche and Merck as well as honoraria for advisory boards from AbbVie. All other authors certify that they have no affiliations with or involvement in any organisation or entity with any financial interest (such as honoraria; educational grants; participation in speakers’ bureaus; membership, employment, consultancies, stock ownership, or other equity interest; and expert testimony or patent-licensing arrangements), or non-financial interest (such as personal or professional relationships, affiliations, knowledge or beliefs) in the subject matter or materials discussed in this manuscript.

Ethics approval The present retrospective analysis was performed in accordance with the Declaration of Helsinki and with approval by the institutional review board of the medical Faculty, Heinrich Heine University Düsseldorf (study number: 2018-156-RetroDEuA).

Informed consent Patients had provided informed consent for the use of their tissue samples and clinical data for research purposes.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

Brodbelt A, Greenberg D, Winters T, Williams M, Vernon S, Collins VP, NCINB (2015) Glioblastoma in England: 2007–2011. Eur J Cancer 51:533–542. https://doi.org/10.1016/j.ejca.2014.12.014

Eckel-Passow JE, Lachance DH, Molinaro AM, Walsh KM, Decker PA, Sicotte H, Pekmezci M, Rice T, Kosel ML, Smirnov IV, Sarkar G, Caron AA, Kollmeyer TM, Praska CE, Chada AR, Halder C, Hansen HM, McCoy LS, Bracci PM, Marshall R, Zheng S, Reis GF, Pico AR, O’Neill BP, Buckner JC, Giannini C, Huse JT, Perry A, Tihan T, Berger MS, Chang SM, Prados MD, Wiemels J, Wiebecke JK, Wrensch MR, Jenkins RB (2015) Glioma groups based on 1p/19q, IDH, and TERT promoter mutations in tumors. N Engl J Med 372:2499–2508. https://doi.org/10.1056/NEJMoa1407297

Felsberg J, Rapp M, Loeser S, Fimmers R, Stummer W, Goepfert M, Steiger HJ, Friedensdorf B, Reifenberger G, Sabel MC (2009) Prognostic significance of molecular markers and extent of resection in primary glioblastoma patients. Clin Cancer Res 15:6683–6693. https://doi.org/10.1158/1078-0432.CCR-08-2801

Felsberg J, Thon N, Eigenbrod S, Hentschel B, Sabel MC, Westphal M, Schackert G, Kretz FW, Pietsch T, Loffler M, Weller M, Reifenberger G, Tonn JC, German Glioma N (2011) Promoter methylation and expression of MGMT and the DNA mismatch repair genes MLH1, MSH2, MSH6 and PMS2 in paired primary and recurrent glioblastomas. Int J Cancer 129:659–670. https://doi.org/10.1002/ijc.26083
Felsberg J, Wolter M, Seul H, Friedensdorf B, Goppert M, Sabel MC, Reifenberger G (2010) Rapid and sensitive assessment of the IDH1 and IDH2 mutation status in cerebral gliomas based on DNA pyrosequencing. Acta Neuropathol 119:501–507. https://doi.org/10.1007/s00401-010-0647-4

Glaser SM, Dobopolski MJ, Balasubramani GK, Flickinger JC, Beriwal S (2017) Glioblastoma multiforme (GBM) in the elderly: initial treatment strategy and overall survival. J Neurooncol 134:107–118. https://doi.org/10.1007/s11060-017-2493-x

Goldstein LB, Bertels C, Davis JN (1989) Interrater reliability of the NIH stroke scale. Arch Neurol 46:660–662. https://doi.org/10.1001/archneur.1989.0052042000026

Harrison RA, de Groot JF (2018) Treatment of glioblastoma in the elderly. Drugs Aging 35:707–718. https://doi.org/10.1007/s40266-018-0568-9

Hartmann C, Hentschel B, Wick W, Capper D, Felsberg J, Simon M, Harrison RA, de Groot JF (2018) Treatment of glioblastoma in the elderly. Drugs Aging 35:707–718. https://doi.org/10.1007/s40266-018-0568-9

Kamp MA, Dibue M, Niemann L, Reichelt DC, Felsberg J, Steiger HJ, Tzaridis T, Mack F et al (2019) MGMT promoter methylation cutoff with safety margin: implications for classification of gliomas. Acta Neuropathol 138:123–132. https://doi.org/10.1007/s00701-018-2777-9

Kocher M, Soffietti R, Abacioglu U, Villa S, Fauchon F, Baumert BG, Farselli L, Trzk-Shina T, Kortmann RD, Carrie C, Ben Hassel M, Kouri M, Valeinis E, van den Berge D, Collette S, Collette L, Mueller RP (2011) Adjuvant whole-brain radiotherapy versus observation after radiosurgery or surgical resection of one to three cerebral metastases: results of the EORTC 22952–26001 study. J Clin Oncol 29:134–141. https://doi.org/10.1200/JCO.2010.30.1655

Koeller M, Reifenberger G, von Deimling A (2010) Patients with IDH1 wild type anaplastic astrocytomas exhibit worse prognosis than IDH1-mutated glioblastomas, and IDH1 mutation status accounts for the unfavorable prognostic effect of higher age: implications for classification of gliomas. Acta Neuropathol 120:707–718. https://doi.org/10.1007/s00401-010-0781-z

Hegi ME, Dierens AC, Gorlia T, Hamou MF, de Tribolet N, Weller M, Kros JM, Hainfellner JA, Mason W, Mariani L, Bromberg JE, Hau P, Mirimanoff RO, Cairncross JG, Janzer RC, Stupp R (2005) MGMT gene silencing and benefit from temozolomide in glioblastoma. N Engl J Med 352:997–1003. https://doi.org/10.3745/COR.2017-000343

Hegi ME, Genbrugge E, Gorlia T, Stupp R, Gilbert MR, Chinot OL, Nabors LB, Jones G, Van Crickening W, Staub J, Weller M (2019) MGMT promoter methylation cutoff with safety margin for selecting glioblastoma patients into trials omitting temozolomide: a pooled analysis of four clinical trials. Clin Cancer Res 25:1809–1816. https://doi.org/10.1158/1078-0432.CCR-18-3181

Hegi ME, Liu L, Herman JG, Stupp R, Wick W, Weller M, Mehta MP, Gilbert MR (2008) Correlation of O6-methylguanine methylationtransferase (MGMT) promoter methylation with clinical outcomes in glioblastoma and clinical strategies to modulate MGMT activity. J Clin Oncol 26:4189–4199. https://doi.org/10.1200/JCO.2007.11.3964

Herrlinger U, Tarzidis T, Mack F et al (2019) Lomustine-temozolomide combination therapy versus standard temozolomide therapy in patients with newly diagnosed glioblastoma with methylated MGMT promoter (CeTeG/NOA-09): a randomised, open-label, phase 3 trial. Lancet Oncol 13:916–926. https://doi.org/10.1016/S1470-2045(12)70265-6

Mason M, Lapierriere N, Wick W, Reardon DA, Malmstrom A, Hovey E, Weller M, Perry JR (2016) Glioblastoma in the elderly: making sense of the evidence. Neurooncol Pract 3:77–86. https://doi.org/10.1093/nop/npw027

Morgan ER, Norman A, Laing K, Seal MD (2017) Treatment and outcomes for glioblastoma in elderly compared with non-elderly patients: a population-based study. Curr Oncol 24:e92–e98. https://doi.org/10.3747/co.24.3424

Munoz-Bendix C, Rapp M, Mjördiskwjik HJ, von Sass C, Dibue-Adjei M, Cornelius JF, Steiger HJ, Turowski B, Sabel M, Kamp MA (2019) Risk factors for in-brain local progression in elderly patients after resection of cerebral metastases. Sci Rep 9:7431. https://doi.org/10.1038/s41598-019-43942-9

Okada M, Miyake K, Tamiya T (2017) Glioblastoma treatment in the elderly. Neurol Med Chir (Tokyo) 57:667–676. https://doi.org/10.2176/nmc.ra.2017-0009

Palmer JD, Bhamidipati D, Mehta M, Williams NL, Dicker AP, Wernir-Wasik M, Shi W (2018) Treatment recommendations for elderly patients with newly diagnosed glioblastoma lack worldwide consensus. J Neurooncol 140:421–426. https://doi.org/10.1007/s00780-018-0699-3

Parsons DW, Jones S, Zhang X, Lin JC, Leary RJ, Angenendt P, Mankoo P, Carter H, Sui IM, Gallia GL, Olivi A, McLendon R, Rasheed BA, Keir S, Nikolaouky T, Nikolsky Y, Busam DA, Tekleah H, Diaz LA Jr, Hartigan J, Smith DR, Strausberg RL, Marie SK, Shinjo SM, Yan H, Riggins GI, Bigner DD, Karchin R, Papadopoulos N, Parmigiani G, Vogelstein B, Velculescu VE, Kinzler KW (2008) An integrated genomic analysis of human glioblastoma multiforme. Science 321:1807–1812. https://doi.org/10.1126/science.1164382

Perry JR, Laperriere N, O’Callaghan CJ, Brandes AA, Menten J, Phillips C, Fay M, Nishikawa R, Cairncross JG, Roa W, Osoba D,
Putz F, Putz T, Goerig N, Knippen S, Gryc T, Eyupoglu I, Rossler K, Pretanvil JA, Salinas IQ, Piccioni DE (2017) Glioblastoma in the elderly: treatment patterns and survival. CNS Oncol 6:19–28. https://doi.org/10.2217/cncs-2016-0023

Putz F, Putz T, Goerig N, Knippen S, Gryc T, Eyupoglu I, Rossler K, Pretanvil JA, Salinas IQ, Piccioni DE (2017) Glioblastoma in the elderly: treatment patterns and survival. CNS Oncol 6:19–28. https://doi.org/10.2217/cncs-2016-0023

Stupp R, Hegi ME, Mason WP, van den Bent MJ, Taphoorn MJ, Bendszus M, Reijneveld JC, Dhermain F, French P, Marosi C, Bogdahn U, Reifenberger G, Soffietti R, Cohen-Jonathan-Moyal E, Henriksson R, Le Ruhn E, Balana C, Chinot O, Bendszus M, Reijneveld JC, Dhermain F, French P, Marosi C, Bogdahn U, Reifenberger G, Soffietti R, Wick W (2017a) European Association for Neuro-Oncology (EANO) guideline on the diagnosis and treatment of adult astrocytic and oligodendrogial gliomas. Lancet Oncol 18:e315–e329. https://doi.org/10.1016/S1470-2045(17)30194-8

Wick W, Platten M, Meisner C, Fidler J, Stupp R, Tabatabai G, Simon M, Nikkhah H, Papsdorf K, Steinbach JP, Sabel M, Combs SE, Vecht CJ, Watts C, Oberg I, Pilkington G, Baumert BG, Hau J, Janzer RC, Ludwin SK, Allgeier A, Fishner B, Belanger K, Hau P, Brandes AA, Gjengebek J, Marosi C, Vecht CJ, Mokhtari K, Wesseling P, Villa S, Eisenhauer E, Gorlia T, Weller M, Lacombe D, Cairncross JG (2008) Toxicity from chemoradiotherapy in older patients with glioblastoma multiforme. Neuro Oncol 10:1178–1184. https://doi.org/10.1093/neuonc/nos153

Senft C, Bink A, Franz K, Vatter H, Gasser T, Seifert V (2011) Intraoperative MRI guidance and extent of resection in glioma surgery: a randomised, controlled trial. Lancet Oncol 12:997–1003. https://doi.org/10.1016/S1470-2045(11)70196-6

Sijben AE, McIntyre JB, Roldan GB, Easaw JC, Yan E, Forsyth PA, Parney IF, Magliocca AM, Bernsen H, Cairncross JG (2008) Toxicity from chemoradiotherapy in older patients with glioblastoma multiforme. J Neurosurg 89:97–103. https://doi.org/10.1093/jnci/dsl106

Socha J, Kepka L, Ghosh S, Roa W, Kumar N, Sinaika V, Mattiello J, Lomizde D, de Castro DG, Bentati D, Fidarova E (2016) Outcome of treatment of recurrent glioblastoma multiforme in elderly and/or frail patients. J Neurosurg 126:493–498. https://doi.org/10.3171/2015.10.JNS151121

Stummer W, Kamp MA (2009) The importance of surgical resection in malignant glioma. Curr Opin Neurol 22:645–649. https://doi.org/10.1097/WCO.0b013e3283320165

Stummer W, Pichlmier U, Meinel T, Wiestler OD, Zanella F, Reulen HJ, Group AL-GS (2006) Fluorescence-guided surgery with 5-aminolevulinic acid for resection of malignant glioma: a randomised controlled multicentre phase III trial. Lancet Oncol 7:392–401. https://doi.org/10.1016/S1470-2045(06)70665-9

Stupp R, Hegi ME, Mason WP, van den Bent MJ, Taphoorn MJ, Janzer RC, Lanzi PD, Allgeier A, Fisher B, Belanger K, Hau P, Brandes AA, Gjengerbeek J, Marosi C, Vecht CJ, Wesseling P, Villa S, Eisenhauer E, Gorlia T, Weller M, Lacombe D, Cairncross JG, Mirimanoff RO (2005) Radiotherapy plus concomitant and adjuvant temozolomide for glioblastoma. N Engl J Med 352:987–996. https://doi.org/10.1056/NEJMa043330

Stupp R, Taillibert S, Kanner A et al (2017) Effect of tumor-treating fields plus maintenance temozolomide vs maintenance temozolomide alone on survival in patients with glioblastoma: a randomized clinical trial. JAMA. 318(23):2306–2316. https://doi.org/10.1001/jama.2017.18718

Tabatabai G, Stupp R, Wick W, Weller M (2013) Malignant astrocytoma in elderly patients: where do we stand? Curr Opin Neurol 26:693–700. https://doi.org/10.1097/WCO.0b013e3283320037

Tammari AF, Juweid M (2017) Epidemiology and Outcome of Glioblastoma. In: De Vleeschouwer S (ed) Glioblastoma. Brisbane, AU

Thakkar JP, Dolecek TA, Horbinski C, Ostrom QT, Lightner DB, Barnholtz-Sloan JS, Villano JL (2014) Epidemiologic and molecular prognostic review of glioblastoma. Cancer Epidemiol Biomarkers Prev 23:1985–1996. https://doi.org/10.1158/1055-9965.EPI-14-0275

Verger E, Salamero M, Conill C (1992) Can Karnofsky performance status be transformed to the eastern cooperative oncology group scoring scale and vice versa? Eur J Cancer 28A:1328–1330. https://doi.org/10.1016/0959-8409(92)90510-9

Victor M, Lu PK, Brown DA, Burns TC, Quinones-Hinojosa A, Chai-chana KL (2019) Hypofractionated versus standard radiation therapy in combination with temozolomide for glioblastoma in the elderly: a meta-analysis. J Neurooncol 143(2):177–185

Weller M, van den Bent M, Tonn JC, Stupp R, Preusser M, Cohen-Jonathan-Moyal E, Henriksson R, Le Ruhn E, Balana C, Chinot O, Renders M, Reinefeld JC, Dhermain F, French P, Marosi C, Watts C, Oberg I, Pilkington G, Baumert BG, Taphoorn MJ, Hegi M, Westphal M, Reifenberger G, Soffietti R, Wick W (2017a) European Association for Neuro-Oncology (EANO) guideline on the diagnosis and treatment of adult astrocytic and oligodendrogial gliomas. Lancet Oncol 18:e315–e329. https://doi.org/10.1016/S1470-2045(17)30194-8

Wick W, Platten M, Meisner C, Fidler J, Tabatabai G, Simon M, Nikkhah H, Papsdorf K, Steinbach JP, Sabel M, Combs SE, Vesper J, Braun C, Meixensberger J, Ketter R, Mayer-Steinacker R, Reifenberger G, Weller M (2012) Temozolomide chemotherapy alone versus radiotherapy alone for malignant astrocytoma in the elderly: the NOA-08 randomised, phase 3 trial. Lancet Oncol 13:707–715. https://doi.org/10.1016/S1470-2045(12)70164-x

Wirsching HG, Happold C, Roth P, Weller M (2015) Management of diffusely infiltrating glioma in the elderly. Curr Opin Oncol 27:502–509. https://doi.org/10.1097/CCO.0000000000000236

Zhang C, Wang X, Hao S, Su Z, Zhang P, Li Y, Song G, Yu L, Wang J, Ji N, Xie J, Gao Z (2016) Analysis of treatment tolerance and factors associated with overall survival in elderly patients with glioblastoma. World Neurosurg 95:77–84. https://doi.org/10.1016/j.wneu.2016.07.079

Publisher’s Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.