Adolescent and young adult glioma: systematic review of demographic, disease, and treatment influences on survival

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Abstract

Background. Prognostic factors in adolescent and young adult (AYA) glioma are not well understood. Though clinical and molecular differences between pediatric and adult glioma have been characterized, their application to AYA populations is less clear. There is a major need to develop more robust evidence-based practices for managing AYA glioma patients.

Methods. A systematic review using PRISMA methodology was conducted using multiple databases with the objective of identifying demographic, clinical, molecular and treatment factors influencing AYA glioma outcomes.

Results. 40 Studies met inclusion criteria. Overall survival was highly variable across studies depending on glioma grade, anatomic compartment and cohort characteristics. Thirty-five studies suffered from high risk of bias in at least one domain. Several studies included older adults within their cohorts; few captured purely AYA groups. Despite study heterogeneity, identified favorable prognosticators included younger age, higher functional status at diagnosis, low-grade pathology, oligodendroglioma histology and increased extent of surgical resection. Though isocitrate dehydrogenase (IDH) mutant status was associated with favorable prognosis, validity of this finding within AYA was compromised though may studies including older adults. The prognostic influence of chemotherapy and radiotherapy on overall survival varied across studies with conflicting evidence.

Conclusion. Existing literature is heterogenous, at high risk of bias, and rarely focused solely on AYA patients. Many included studies did not reflect updated pathological and molecular AYA glioma classification. The optimal role of chemotherapy, radiotherapy, and targeted agents cannot be determined from existing literature and should be the focus of future studies.

Key Points

• High-quality evidence on prognosticators in AYA glioma is lacking.
• Literature to date is heterogenous, rarely focused only on AYA, and prone to bias/ confounding.
• Optimal role of chemotherapy and radiation cannot be determined.
Gliomas represent a diverse histologic group of central nervous system tumors (CNS) with substantial molecular heterogeneity. Taken together, gliomas represent 29–35% of central nervous system tumors within the adolescent and young adult (AYA) demographic. Historically, AYA have been poorly represented in glioma research due to limited enrollment and representation in both pediatric- and adult-focused cohorts. This systematic review synthesizes available prognostic, treatment, and survival data for AYA glioma patients. We demonstrate the favorable impact of younger age and higher Karnofsky Performance Status (KPS) on overall survival (OS) and event-free survival (EFS). This review identified a positive association with OS and EFS with low-grade histology, oligodendrogial histology, isocitrate dehydrogenase (IDH) mutant molecular status and extent of surgical resection, though many included studies exhibited high bias risk and included older adults. It also highlights limited consensus on the role of adjuvant chemotherapy and radiotherapy in this population.

Glioma is a major contributor to oncologic morbidity and mortality in the adolescent and young adult (AYA) demographic. Historically, AYA have been poorly represented in glioma research due to limited enrollment and representation in both pediatric- and adult-focused cohorts. This systematic review synthesizes available prognostic, treatment, and survival data for AYA glioma patients. We demonstrate the favorable impact of younger age and higher Karnofsky Performance Status (KPS) on overall survival (OS) and event-free survival (EFS). This review identified a positive association with OS and EFS with low-grade histology, oligodendrogial histology, isocitrate dehydrogenase (IDH) mutant molecular status and extent of surgical resection, though many included studies exhibited high bias risk and included older adults. It also highlights limited consensus on the role of adjuvant chemotherapy and radiotherapy in this population.

**Importance of Study**

Glioma is a major contributor to oncologic morbidity and mortality in the adolescent and young adult (AYA) demographic. Historically, AYA have been poorly represented in glioma research due to limited enrollment and representation in both pediatric- and adult-focused cohorts. This systematic review synthesizes available prognostic, treatment, and survival data for AYA glioma patients. We demonstrate the favorable impact of younger age and higher Karnofsky Performance Status (KPS) on overall survival (OS) and event-free survival (EFS). This review identified a positive association with OS and EFS with low-grade histology, oligodendrogial histology, isocitrate dehydrogenase (IDH) mutant molecular status and extent of surgical resection, though many included studies exhibited high bias risk and included older adults. It also highlights limited consensus on the role of adjuvant chemotherapy and radiotherapy in this population.

**Methods**

Ethics approval was not required for this systematic review.

**Data Sources and Search Strategy**

The review followed the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines. Multiple databases including OVID MEDLINE, EMBASE and EBM Reviews-Cochrane library databases from inception to July 2020 were queried in collaboration with an academic librarian at the Hospital for Sick Children. A sample search strategy can be found in supplemental materials (Supplementary Table 1). Bibliographies of relevant reviews were further queried to ensure all relevant studies were captured.

**Screening and search strategy.**—Study inclusion criteria included: (1) original studies that reported predictors of cancer-related outcomes [e.g., PFS, time to malignant progression (TTP), OS]; (2) mean or median age at diagnosis within the AYA age range (15–39.9 years); (3) AYA patient sample size greater than 20; (4) diagnosis of glioma based on either WHO 2007 or WHO 2016 classification (Appendix 2); and (5) published in English between January 2010 and June 2020. Studies of pediatric and adult age groups were included if outcomes for AYA were reported separately, or if AYA patients represented more
than 50% of the entire group. Exclusion criteria included low- and middle-income country studies (World Bank Definition), reviews, commentaries, editorials, conference abstracts, articles published before 2010, case series fewer than 20 patients, and studies using population-based mortality statistics.14

Abstracts were screened and assessed to identify pertinent studies (VZ). Full text review was conducted by two independent authors (VZ and AM). Discrepancies were reviewed by a third author when required (VK). The kappa coefficient was calculated to determine agreement between reviewers.

Data extraction and analysis.—The Checklist for critical Appraisal and data extraction for systematic Reviews of prediction Modeling Studies-Prognostic Factors (CHARMS-PF) was used to extract data from included texts.16 The following data were extracted from each study: study type, country of origin, sample size, mean/median age at diagnosis, length of follow-up, and all factors included in univariate or multivariable models of outcomes. Study quality was evaluated independently by two reviewers (AM and VK) utilizing the Quality In Prognosis Studies (QUIPS) tool to assess risk of bias.16–18 Six domains of possible bias were assessed through QUIPS: study participation, study attrition, prognostic factor measurement, outcome measurement, adjustment for other prognostic factors, and statistical analysis and reporting. Meta-analysis was not possible due to significant study heterogeneity. When comparing outcomes across studies, “event-free survival” was used to describe any outcome which incorporated disease progression, such as malignant progression-free survival (MPFS) or PFS. Studies’ definitions of malignant transformation and disease progression were heterogenous. A common definition for malignant transformation was pathological diagnosis of grade 3 or 4 glioma or imaging consistent with malignant transformation based on new or increased contrast enhancement and or the lesional growth pattern. Progression was commonly defined in studies by previously described response assessment frameworks such as Response Assessment in Neuro-Oncology (RANO).39 In instances where a p-value was reported without a hazard ratio or risk ratio, the primary source was examined, and the directionality of the effect was included in parentheses. Several figures were generated using the R Studio version 1.4.1717 and the ggplot2 package.

Results

The search strategy yielded 12 294 studies; removal of duplicates resulted in 10 336 unique studies. After abstract screening, 261 studies were identified as possibly meeting inclusion criteria and their full texts reviewed. Following full text review, 40 studies met inclusion criteria. Supplementary Figure 1 depicts the PRISMA workflow identifying included studies and reasons for exclusion. The kappa measure of agreement between reviewers for final study inclusion was 94.6% (95% CI 89.5–99.8%), or excellent.

Study Characteristics

Forty studies met criteria for inclusion in the review: 39 studies were retrospective (single center, multi-center or national database studies) and 1 study was prospective. Countries of origin included: United States (n = 19), Germany (n = 8), France (n = 4), Italy (n = 2), Japan (n = 2), Poland (n = 1), Austria (n = 1), United Kingdom (n = 1), Norway (n = 1) and Korea (n = 1). There was substantial variability in sample size among studies, ranging from 25 to 3057 patients. Together, the studies represented 12 405 patients with an age range from 3 months to 86 years. Though greater than 50% of each study cohort was required to be AYA based on inclusion criteria, older adults and children were included in many studies as illustrated in Figure 1. There were three studies that specifically included spinal cord gliomas, 1 study that included both spinal cord and intracranial glioma and the remainder included intracranial glioma. Three studies did not provide OS for the overall cohort, while another 10 did not provide EFS. All studies included OS-based univariate or multivariable analyses.

Overall Survival and Event-Free Survival

Glioma outcomes are summarized in Table 1. Two studies reported only on intracranial grade 1 glioma in which one showed an OS of 80% at 5 years and the other showed a reduced survival in the cohort undergoing external beam radiation therapy (EBRT) (< 60% 5 year OS) compared to those not undergoing adjuvant EBRT (> 75% 5 year OS).20,21 Two studies included combined cohorts of both grade 1 and 2 glioma in which OS ranged from 75.7 to 91.0% at 5 years.22,23 Twenty-six studies included grade 2 glioma only and reported 5-year OS ranging from 84 to 98%, with one study reporting 5-year OS of 69.2% in a subset of patients with radiographic velocity of diametric expansion over 8 mm/year.24–49 Among studies of grade 2 glioma, 5-year EFS ranged from 30 to 94%. Several studies included glioma subgroups across multiple pathological grades. 2 studies grouped grade 2 and 3 pleomorphic xantho-astrocytoma (PXA) with combined OS 76.3–89.5% at 5-years, 3 studies grouped grade 2 and 3 glioma together, 2 studies included grade 3 and 4 glioma, and 3 studies reported varying grades of spinal cord glioma, with 5-year OS ranging from 85.4% in grade 1 cases to 36.4% in grades 2, 3 and 450–53 (Table 1).

Patient Factors

Several patient factors were associated with superior OS and EFS across glioma grade following adjusted multivariable analysis (Tables 2–4). Increased age was often associated with worse OS when age was evaluated as a continuous variable,20,23,34,38,59 including cohorts of pilocytic astrocytoma alone, combined grade 1 and 2 gliomas, combined grade 2 and 3 gliomas, and of peri-ventricular HGG. Within the AYA group, the following younger age clusters were associated with improved OS: age <18 years,51 age <30 years,53 and age <40 years.22,42 Only one study showed a negative impact of age younger than 40 on OS.48 Several studies in contrast did not find a significant association between age and OS in multivariable analysis.30–33,36,43,46,52,54 Three studies demonstrated that younger age was associated with improved EFS.49,41,53
The relationship between sex and OS and EFS was conflicting with no clear prognostic effect. Three studies showed no effect of patient sex on OS. Other patient-related factors associated with favorable OS included private health insurance in a United States cohort, median annual income greater than $38,000, and Charles-Deyo Comorbidity Index score of 0 vs. 2. KPS over 80 was associated with favorable EFS in 1 study following multivariable analysis, and though KPS was significantly associated with EFS in univariate analysis in three additional studies, it lost significance when adjusted for other factors.

Disease and Treatment-Related Factors

**Grade 1 glioma.**—Several disease and treatment-related factors were significantly associated with OS and EFS among patients with grade 1 glioma or studies combining grade 1 and 2 gliomas (Table 2). Pre-operative lesion size over 19 mm and grade 2 compared to grade 1 histology were associated with inferior OS, while location of tumor in the supratentorial compartment was favorable compared to spinal cord or infratentorial locations following univariate analysis, though non-significant after multivariable analysis (though brainstem lesion inclusion in the infratentorial category may have biased this finding). Symptom duration in spinal cord glioma was not significantly associated with OS after multivariable analysis. Treatment-related factors positively influencing OS included gross-total resection (GTR) in spinal cord glioma cases. Three studies found adjuvant radiation to be associated with inferior OS even after adjustment for other factors. The first study by Lee et al. examined a national cohort of patients with pilocytic astrocytoma and adjusted for age, median income, tumor volume and comorbidity scores. They found adjuvant external beam radiotherapy (EBRT) was associated with a significantly worsened OS compared to no radiotherapy (patients undergoing EBRT 5-year OS < 60% compared to ≥ 75% 5-year OS in patients receiving other therapies). The same study showed a trend towards inferior OS, though non-significant, when stereotactic radiotherapy was compared to no radiotherapy. The authors nonetheless attributed their finding to confounding by other important factors including eloquent location and tumor resectability. The second study examined the effect of pregnancy on LGG survival. They showed that post-operative radiation therapy was associated with significantly inferior OS in combined grade 1 and 2 gliomas as well as grade 2 gliomas alone following multivariable adjustment, though the authors did not provide a list of what variables were adjusted for. The third study, examining
| Glioma type | First author, year of Publication | Country | Study design | Sample size | Glioma pathological subtype | Age at diagnosis (years) | Length of follow-up (months) | Overall survival | Event-free survival |
|-------------|----------------------------------|---------|--------------|-------------|-----------------------------|-------------------------|---------------------------|----------------|------------------|
| Grade 1 glioma | Lee KJ 2018 | US | Retrospective national cohort | 3057 | WHO Grade 1 astrocytoma (includes spinal cord) | Median 32 | NOS | Patients undergoing EBRT 5 year < 60%, patients not undergoing EBRT 5 year > 75% |
| | Nelson AJ 2019 | UK | Retrospective single center | 50 | WHO Grade 1 glioma | Median 29 (16–76 range) | Median 3.5 years | 5 year—80% | Median 7 years (95% CI 4.5–9.5) |
| Grade 1 and 2 glioma | Bagley JH 2013 | US | Retrospective national cohort | 166 | Cerebellar WHO Grade 1 (n = 71) and Grade 2 (n = 95) astrocytoma | Median G1A 25.0 | Median G2A 34.0 | NOS | NOS |
| | Rønning PA 2016 | Norway | Retrospective national cohort | 346 | WHO Grade 1 and 2 glioma (female cohort) i) Pilocytic astrocytoma (n = 46) ii) Diffuse astrocytoma (n = 196) iii) Oligoastrocytoma (n = 26) iv) Oligodendroglioma (n = 78) | Median 26.4 (16–40 range) | Median 15.2 years | 5 year—75.7% 10 year—54.8% G2 glioma cases only: Median 12.2 years (95% CI 10.7–125) |
| Grade 2 glioma | Ahmadi R 2012 | Germany | Retrospective single center | 100 | Supratentorial WHO Grade 2 astrocytoma i) IDH1 mutant (n = 79) ii) IDH wt (n = 21) | Median 37.4 (19.8–72.1 range) | Median 81.1 (28–134.2 range) | Median 81.4 months (95% CI 5.5–247) | Median PFS 44.6 months (95% CI 1.0–267) MedianTtMP 74.9 (95% CI 1.6–236.2) |
| | Chang EF 2011 | US | Retrospective single center | 281 | Infiltrative WHO Grade 2 gliomas i) Astrocytoma (n = 81) ii) Oligoastrocytoma (n = 101) iii) Oligodendroglioma (n = 99) | Median 38 (15–72 range) | Median 62.4 (3–152 range) | 5 year 86% | 5 year 62% |
| | Coburger J 2016 | Germany | Retrospective multi-center | 288 | WHO Grade 2 gliomas i) Diffuse astrocytoma (n = 173) ii) Oligoastrocytoma (63) iii) Oligodendroglioma (n = 52) | Mean 39 (18–75 range) | Mean 52 | Mean 21 months (95% CI 17–25) | Mean 68 months (95% CI 58–77) 5 year 94% |
| | Eseonu CI 2017 | US | Retrospective single center | 109 | WHO Grade 2 gliomas i) Diffuse astrocytoma (n = 73) ii) Oligoastrocytoma (n = 36) | Median 37 (19–74 range) | Median 62.4 | 5 year 84% 8 year 65% | 5 year 70% 8 year 51% |
| | Gousias K 2014 | Germany | Retrospective single center | 148 | WHO Grade 2 supratentorial glioma i) Diffuse astrocytoma (n = 76) ii) Oligoastrocytoma (n = 54) iii) Oligodendroglioma (n = 18) | Median 38 (18–74 range) | Median 59 (1–196 range) | 5 year—86.1%* | Median PFS 70 months MedianTtMP 98 months |
| Glioma type | First author, year of Publication | Country | Study design | Sample size | Sample size | Glioma pathological subtype | Age at diagnosis (years) | Length of follow-up (months) | Overall survival | Event-free survival |
|-------------|---------------------------------|---------|--------------|-------------|-------------|-----------------------------|------------------------|-----------------------------|-----------------|------------------|
| Glioma type | First author, year of Publication | Country | Study design | Sample size | Sample size | Glioma pathological subtype | Age at diagnosis (years) | Length of follow-up (months) | Overall survival | Event-free survival |
| Goze C 2014 | France | Retrospective multi-center | 131 | WHO Grade 2 supratentorial glioma i) Diffuse astrocytoma (n = 25) ii) Oligoastrocytoma (n = 71) iii) Oligodendroglioma (n = 35) a) 1p19q co-deleted (n = 38 out of 119 tested) b) P53 over-expression (n = 65 out of 125 tested) c) IDH1 mutant (n = 107 out of 131 tested) | Median 38 (15–66 range) | Median 55 (3.6–262 range) | 82.4% survival at median observation period of 111 months | Median TiMP 51 months (42.7% of cohort in observed follow-up period) | |
| Harary M 2020 | US | Retrospective national cohort | 590 | WHO Grade 2 oligodendroglioma (1p/19q-co-deleted) | Median 39 (29–52 IQR) | Median 41.5 (23.8–61.6 IQR) | Biopsy only: 5 year—92.4% STR: 5 year—90.1% GTR: 5 year—96.5% | NOS | |
| Hartmann C 2011 | Germany | Retrospective multi-center | 89 | WHO Grade 2 glioma i) Diffuse astrocytoma (n = 40) ii) Oligoastrocytoma (n = 23) iii) Oligodendroglioma (n = 26) | Median 36.7 (17.4–75.7 range) | Median 75.6 | Median 15.5 years | Median 4.1 years (95% CI 3.1–5.1) | |
| Houillier C 2010 | France | Retrospective multi-center | 231 | WHO Grade 2 glioma i) Diffuse astrocytoma (n = 43) ii) Oligoastrocytoma (n = 58) iii) Oligodendroglioma (n = 130) | Median 39 (18–78 range) | Median 95.1 (95% CI 82.5–107.3) | Median 175.8 months (95% CI 150.1–261) | Median 39.6 months (95% CI 35.8–44.5) | |
| Houillier C 2010 | France | Retrospective single center | 271 | WHO Grade 2 glioma i) Diffuse astrocytoma (n = 47) ii) Oligoastrocytoma (n = 66) iii) Oligodendroglioma (n = 158) | Median 39 (18–78 range) | Median 69.2 (95% CI 60.3–78.7) | Median 133.3 months | Median 41.3 months | |
| Ius T 2012 | Italy | Retrospective single center | 190 | WHO Grade 2 glioma supratentorial eloquent location i) Diffuse astrocytoma (n = 98) ii) Oligoastrocytoma (n = 34) iii) Oligodendroglioma (n = 58) | Median 37 (18–75 range) | Median 56.4 (4–155 range) | 5 year—80% 8 year—66% 15 year—35% | 5 year—59% 8 year—35% | |
| Jairam V 2019 | US | Retrospective national cohort | 1032 | WHO Grade 2 glioma i) Diffuse astrocytoma (n = 433) ii) Oligoastrocytoma (n = 256) iii) Oligodendroglioma (n = 343) | Mean 29.8 ± 6 | Median 46.8 | 5 year—91.7% | NOS | |
| Jansen E 2019 | Germany | Retrospective multi-center | 110 | WHO Grade 2 glioma i) Diffuse astrocytoma IDH mutant (n = 53) ii) Diffuse astrocytoma IDH wt (n = 18) iii) Oligodendroglioma (n = 39) | Median 37 (18–79 range) | Median 126 (95% CI 109–143) | 5 year—88% 10 year—71% 15 year—57% | 5 year—38% 10 year—18% 15 year—1% | |
| Glioma type                        | First author, year | Country | Study design | Sample size | Sample pathological subtype | Age at diagnosis (years) | Length of follow-up (months) | Overall survival | Event-free survival |
|-----------------------------------|--------------------|---------|--------------|-------------|-----------------------------|-------------------------|----------------------------|----------------------|------------------|
| Glioma pathological subtype      |                    |         |              |             |                             |                         |                           |                      |                  |
| jGlioma grade 2 astrocytoma       | Jungk C 2016       | Germany | Retrospective single center | 46          | WHO Grade 2 astrocytoma | Median 35 (17–54 range) | Median 69 (127–164.6) | 5 year — 88.3% | 5 year — 80.8% |
|                                  |                    |         |              |             |                             |                         |                           |                      |                  |
| jGlioma grade 2 glioma with known radiologic progression (OS from date of progression) | Narang AK 2017     | US      | Retrospective single center | 108         | WHO Grade 2 glioma | Median 38 (18–62 range) | Median 36 (19–63 range) | 5 year — 91%    | 5 year — 81%    |
|                                  |                    |         |              |             |                             |                         |                           |                      |                  |
| jGlioma grade 2 diffuse glioma (IDH mutant only) | Nata M 2015        | Japan   | Retrospective single center | 144         | WHO Grade 2 diffuse glioma | Mean ± 11 | Median 13.1 | 5 year — 97.6% |                      |
|                                  |                    |         |              |             |                             |                         |                           |                      |                  |
| jGlioma grade 2 glioma with known radiologic progression (OS from date of progression) | Okita Y 2012       | Japan   | Retrospective single center | 72          | WHO Grade 2 glioma | Median 37.0 (15–76 range) | Median 34 (18–43 range) | 5 year — 96.4% | 10 year — 84.1% |
|                                  |                    |         |              |             |                             |                         |                           |                      |                  |
| jGlioma grade 2 oligodendroglioma | Pal'a A 2019       | Germany | Retrospective multi-center | 140         | WHO Grade 2 supratentorial glioma | Mean 39 ± 11 | Median 13.1 | 5 year — 97.6% |                      |
|                                  |                    |         |              |             |                             |                         |                           |                      |                  |
| jGlioma grade 2 glioma with known radiologic progression (OS from date of progression) | Pallud J 2013      | France  | Retrospective national cohort | 407         | WHO Grade 2 glioma | Mean 38.0 (18–70 range) | Median 38.0 (18–77 range) | 5 year — 97.6% |                      |
|                                  |                    |         |              |             |                             |                         |                           |                      |                  |
| jGlioma grade 2 oligodendroglioma | Scherer M 2020     | Germany | Retrospective multi-center | 486         | WHO Grade 2 glioma | Mean 38.0 (18–70 range) | Median 38.0 (18–77 range) | 5 year — 97.6% |                      |
|                                  |                    |         |              |             |                             |                         |                           |                      |                  |
| jGlioma grade 2 glioma with known radiologic progression (OS from date of progression) | Tom MC 2019        | US      | Retrospective single center | 77          | WHO Grade 2 glioma | Mean 39 ± 11 | Median 13.1 | 5 year — 97.6% |                      |
|                                  |                    |         |              |             |                             |                         |                           |                      |                  |
| Gioma type | First author, year of Publication | Country | Study design | Sample size | Glioma pathological subtype | Age at diagnosis (years) | Length of follow-up (months) | Overall survival | Event-free survival |
|------------|----------------------------------|---------|--------------|-------------|-----------------------------|---------------------|-----------------------------|----------------|-------------------|
| Tom MC 2019 | US Retrospective single center | 144 | WHO Grade 2 glioma i) Diffuse astrocytoma (n = 49) ii) Oligoastrocytoma (n = 36) iii) Oligodendroglioma (n = 59) | Median 29 (IQR 18–41) | Median 81 (IQR 36–132) | 5 year—98% 10 year—90% | 5 year—71% 10 year—53% |
| Wahl M 2017 | US Prospective single center | 120 | WHO Grade 2 glioma i) Diffuse astrocytoma (n = 43) ii) Oligoastrocytoma (n = 20) iii) Oligodendroglioma (n = 57) | Median 39 (19–71 range) | Median 75 years | Median 9.7 years (95% CI 7.2–11.3) | Median 3.8 years (95% CI 3.0–5.0) |
| Youland RS 2013 | US Retrospective single center | 852 | WHO Grade 2 glioma i) Diffuse astrocytoma (n = 293) ii) Oligoastrocytoma (n = 280) iii) Oligodendroglioma (n = 279) | Mean 39.1 (18.1–76.0) | Median 11.4 years (0.02–38.5) | Median 8.0 years | Median 4.4 years |
| Byun J 2018 | Korea Retrospective single center | 25 | WHO Grade 2 Pleomorphic xanthoastrocytoma (PXA) (n = 21) G3 PXA (n = 4) | Mean 29.9 (18–60 range) | Mean 51.4 (2–112 range) | G2 PXA: 5 year 89.5% 10 year 40.9% G3 PXA: 5 year 100% 10 year 0% | G2 PXA: 5 year 65.1% 7 year 52% G3 PXA: 5 year 0% 10 year 0% |
| Gallo P 2013 | Italy Retrospective single center | 40 | WHO Grade 2 PXA (n = 32) G3 PXA (n = 8) | Median 30.5 (12–65 range) | Median 74 | 5 year—76.3% 10 year—68.2% | 5 year—7.1% 10 year—58.0% |
| Hatanpaa KJ 2014 | US Retrospective single center | 50 | WHO Grade 2-III astrocytoma and oligoastrocytoma | Median 37.5 (20–66 range) | Median 51.6 | NOS | NOS |
| Miller JJ 2019 | US Retrospective single center | 275 | WHO Grade 2 (n = 134) and 3 glioma (n = 141) i) Oligodendroglioma (n = 95) ii) Astrocytoma (n = 180) | Median 38.0 (19–86 range) | Median 6.4 years | Median 18.7 years (95% CI 12.2–not reached) | Median 5.7 years (95% CI 4.7–6.4) |
| Olar A 2015 | US Retrospective multi-center | 558 | WHO Grade 2 and 3 diffuse glioma i) Grade 2 (n = 262) ii) Grade 3 (n = 296) | Median 38.2 (17.4–78.4 range) | Median 7.4 years | G2 glioma: median 12.41 years G3 glioma: Median 13.35 years | NOS |
| Yang W 2018 | US Retrospective national cohort | 353 | Peri-ventricular or subventricular zone Grade 3 and Grade 4 glioma i) Glioblastoma (n = 172) ii) Anaplastic ependymoma (n = 70) iii) Anaplastic astrocytoma (n = 65) iv) Other (n = 46) | Mean 38.77 ± 24.95 | NOS | Median n12 months (95% CI 10–15) | NOS |
| Leibetseder A 2013 | Austria Retrospective multi-center | 47 | WHO Grade 4 astrocytoma | Median 32 (18–39 range) | NOS | Median 28 months (95% CI 24–31.6) | Median 12 months (95% CI 9.5–14) |
| Glioma type                | First author, year of Publication | Country | Study design            | Sample size | Glioma pathological subtype                                                                 | Age at diagnosis (years) | Length of follow-up (months) | Overall survival | Event-free survival |
|----------------------------|----------------------------------|---------|-------------------------|-------------|-----------------------------------------------------------------------------------------------|-------------------------|-----------------------------|------------------|---------------------|
| Spinal cord glioma         | Diaz-Aguilar D                    | US      | Retrospective national cohort | 561         | WHO Grade 1 and 2 gliomas spinal cord  
  i) Pilocytic astrocytoma \( n = 247 \)  
  ii) Diffuse astrocytoma \( n = 64 \)  
  iii) Astrocytoma NOS \( n = 222 \)  
  iv) Glioma NOS \( n = 28 \) | Mean 28 (± 22) | NOS | NOS | NOS |
|                            | Fakhreddine MH                    | US      | Retrospective single center | 83          | Spinal cord astrocytoma  
  i) WHO Grade 1 \( n = 31 \)  
  ii) WHO Grade 2 \( n = 14 \)  
  iii) WHO Grade 3 \( n = 18 \)  
  iv) WHO Grade 4 \( n = 18 \)  
  v) Indeterminate either Grade 3 or IV \( n = 2 \) | Median 28.7 (0.25–77 range) | Median 49.2 | G1A: 5 year 85.4%  
  Infiltrative astrocytoma (G2A, G3A and G4A): 5 year 36.4% | G1A: Median 3.33 years  
  Infiltrative astrocytoma (G2A, G3A and G4A): Pooled median 0.89 years |
|                            | Liu J                             | US      | Retrospective national cohort | 158         | WHO Grade 3 and IV spinal cord glioma  
  i) Anaplastic astrocytoma \( n = 14 \)  
  ii) Anaplastic ependymoma \( n = 14 \)  
  iii) Glioblastoma \( n = 111 \) | Mean 36.23 (± 21.0) | NOS | Median 20 months (9–42.75) | NOS |

Pooled follow-up, median/mean age, OS and PFS when available unless reported separately in original article.  
NOS, not otherwise specified; G1A, Grade 1 astrocytoma; G2A, Grade 2 astrocytoma; G3A, Grade 3 astrocytoma; EBRT, external beam radiation therapy.  
*Limited number of patients died during follow-up therefore robust multivariate OS modeling was not possible.
| Imaging, treatment and tumor factors | Study | Overall survival | Event-free survival |
|-------------------------------------|-------|----------------|-------------------|
| **Demographic factors**             |       |                |                   |
| Age (continuous)                    | Rønning PA, 2016 | HR = 1.067, *P* < .001 | HR = 1.049, *P* < .001 |
| Age ≥ 40                            | Lee KJ, 2018 | *P* < .001 | HR = 1.050, *P* < .001 |
| Age 0–18 (ref.) vs. i) 18–65 ii) > 65 | Bagley JH, 2013 | | HR = 7.30, *P* < .0001 |
| Female sex                          | Bagley JH, 2013 | | HR = 0.28, *P* < .001 |
| Median annual income < $38 000 (ref.) vs. i) $38 000–$47 999 ii) $48 000–$62 999 iii) > $63 000 | Lee KJ, 2018 | *P* = .01 | i) *HR* = 0.621, *P* = .001 ii) *HR* = 0.543, *P* < .001 iii) *HR* = 0.600, *P* < .001 |
| Charlson-Deyo Comorbidity index = 0 (ref.) vs. i) 1 ii) 2 | Lee KJ, 2018 | *P* < .001 | i) NS ii) *HR* = 1.647, *P* = .009 |
| **Radiographic characteristics**    |       |                |                   |
| Tumor size 1–19 mm (ref.) vs. i) 20–39 mm ii) 40–59 mm iii) 60–79 mm iv) 80–99 mm v) 100+ mm | Lee KJ, 2018 | *P* < .001 | i) *HR* = 1.661, *P* = .010 ii) *HR* = 1.803, *P* = .006 iii) *HR* = 3.029, *P* < .001 iv) NS v) NS |
| Location of tumor supratentorial (ref.) vs. infratentorial and spinal cord | Lee KJ, 2018 | | Supratentorial superior *P* = .01 NS |
| **Tumor presentation**              |       |                |                   |
| Spinal astrocytoma motor deficit i) G1A cohort ii) Infiltrative cohort (G2A, G3A, G4A) | Fakhreddine MH, 2013 | | i) Motor deficit superior *P* = .040 ii) NS |
| Spinal astrocytoma symptoms ≥ 4.6 months i) G1A cohort ii) Infiltrative cohort (G2A, G3A, G4A) | Fakhreddine MH, 2013 | | ii) NS Symptoms ≥ 4.6 months superior *P* = .027 NS |
| Spinal astrocytoma motor deficit i) G1A cohort ii) Infiltrative cohort (G2A, G3A, G4A) | Fakhreddine MH, 2013 | | i) Motor deficit superior *P* = .040 ii) NS |
| **Histological factors**            |       |                |                   |
| G1 (ref.) vs G2 astrocytoma         | Bagley JH, 2013 | | *HR* = 2.76, *P* = .028 |
| Spinal cord G1 (ref.) vs. G2 astrocytoma | Díaz-Aguilar D, 2019 | *HR* = 2.34, *P* < .001 | NS |
| Diffuse astrocytoma (ref.) vs. i) Oligoastrocytoma ii) Oligodendroglioma iii) Pilocytic astrocytoma | Rønning PA, 2016 | i) NS ii) NS iii) *0.251, P* < .001 | i) NS ii) NS iii) *0.380, P* < .05 |
Table 2. Continued

| Imaging, treatment and tumor factors | Study | Overall survival | Event-free survival |
|-------------------------------------|-------|-----------------|--------------------|
|                                     |       | Univariate      | Multivariate       | Study                                    | Univariate | Multivariate |
| Chemotherapy                        |       |                 |                    |                                         |           |             |
| Spinal astrocytoma adjuvant chemotherapy | Fakhreddine MH, 2013 | ii) Adjuvant chemotherapy superior \(P = .032\) | ii) NS | Fakhreddine MH, 2013 | i) \(P = .023\) | ii) \(HR = 0.22\), \(P = .0075\) |
| Radiation therapy                  |       |                 |                    |                                          |           |             |
| G1 and G2 glioma post-operative radiotherapy | Rønning PA, 2016 | HR = 2.013, \(P < .001\) | HR = 1.808, \(P < .01\) | Lee KJ, 2018 | \(P < .001\) | ii) NS |
| Radiation technique no radiation (ref.) vs. | | | | | | |
| i) EBRT | P < .001 | \(i) HR = 3.370, P < .001\) | ii) NS | iii) NS |
| ii) Stereotactic radiosurgery | | | | | | |
| iii) Radiation NOS | | | | | | |
| Spinal cord G1 and G2 glioma post-operative radiotherapy | Diaz-Aguilar D, 2019 | \(P < .001\) | HR = 2.78, \(P < .001\) | |
| Spinal cord astrocytoma post-operative radiotherapy | | | | | | |
| i) G1A cohort | | | | | | |
| ii) Infiltrative cohort (G2A, G3A, G4A) | | | | | | |
| Radiation technique no radiation (ref.) vs. | Rønning PA, 2016 | HR = 3.370, \(P < .001\) | \(i) HR = 3.370, P < .001\) | Fakhreddine MH, 2013 | \(P = .047\) | ii) \(HR = 2.78, P < .001\) |
| i) EBRT | \(P < .001\) | \(i) HR = 3.370, P < .001\) | ii) NS | iii) NS |
| ii) Stereotactic radiosurgery | | | | | | |
| iii) Radiation NOS | | | | | | |
| Surgical factors                   |       |                 |                    |                                          |           |             |
| G1 and G2 spinal cord glioma no surgery (ref.) vs. | Diaz-Aguilar D, 2019 | \(P < .001\) | \(i) NS\) | Nelson AJ, 2019 | \(i) Biopsy inferior \(P = .002\) | ii) Biopsy inferior \(P = .005\) |
| i) STR | | | | | | |
| ii) GTR | | | | | | |
| G1 and G2 glioma biopsy (ref.) vs. resection | Rønning PA, 2016 | HR = 0.544, \(P < .01\) | NS | | | |
| Biopsy alone (ref.) vs. | | | | | | |
| i) < 25% residual following STR | | | | | | |
| ii) > 25% residual following STR | | | | | | |

Cases with pooled WHO Grade 2 gliomas were included if they included WHO Grade 1 lesions. NS, not significant; KPS, Karnofsky Performance Status; HR, Hazard ratio. Significant \(P\)-values without indication of effect directionality (absence of reported hazard ratio) contain a note about superior or inferior effect on OS or EFS. Bolded fields indicate statistical significance of the variable in cited study at an alpha of 0.05.
Table 3. Demographic and radiographic factors associated with AYA WHO Grade 2 glioma event-free survival (EFS) and overall survival (OS)

| Demographic and radiographic factors | Study | Overall survival | Event-free survival |
|-------------------------------------|-------|------------------|--------------------|
|                                     |       | Univariate       | Multivariate       |
|                                     |       |                  |                    |
| Age (continuous)                    | Eseonu CI, 2017 | HR = 1.098, \( P = .03 \) | Tom MC, 2019 | \( P = .005 \) |
|                                     | Ius T, 2012 | HR = 1.030, \( P = .011 \) | - | - |
|                                     | Kavouridis VK, 2020 | HR = 1.12, \( P = .032 \) | - | - |
|                                     | Majchrzak K, 2012 | HR = 1.035, \( P = .003 \) | - | - |
| Age ≤ 40                            | Okita Y, 2012 | HR = 0.400, \( P = .02 \) | Scherer M, 2020 | HR = 0.60, \( P = .03 \) |
| Age ≥ 40                            | Jansen E, 2019 | Age ≥ 40 inferior \( P = .048 \) | - | - |
|                                     | Youldand RS, 2013 | HR = 1.36, \( P = .001 \) | - | - |
|                                     | Tom MC, 2019 | \( P < .001 \) | - | - |
| Age ≥ 50                            | Nitta M, 2015 | HR = 5.43, \( P = .0089 \) | - | - |
| Age > 55                            | Houillier C, 2010 | Age > 55 inferior \( P = 0.001 \) | - | - |
| Sex                                 | Male sex | Goze C, 2014 | HR = 5.06, \( P = .002 \) | Tom MC, 2019 | \( P = .009 \) |
|                                     | Kavouridis VK, 2020 | HR = 2.02, \( P = .042 \) | - | - |
|                                     | Tom MC, 2019 | \( P = 0.003 \) | - | - |
|                                     | Houillier C, 2010 | HR = 0.45, \( P = .01 \) | - | - |
|                                     | Houillier C, 2010 | Female sex superior \( P = .01 \) | NS | NS |
| Female sex                          |        |                  |                    |
| Financial status                    | Non-insured (ref.) vs. | HR = 1.88, \( P = .043 \) | Harary M, 2020 | \( P = .04 \) |
|                                     | i) Private insurance | - | i) \( HR = 0.24, P = .04 \) |
|                                     | ii) Medicare | - | ii) NS |
|                                     | Median annual income < \$38 000 | - | - |
|                                     | Jairam V, 2019 | - | - |
| Functional status                   | KPS (continuous) | Ahmadi R, 2012 | Higher KPS superior \( P = .0004 \) | Ahmadi R, 2012 | Higher KPS superior \( P = .0009 \) |
|                                     | Gousias K, 2014 | HR = 0.136, \( P < .001 \) | - | - |
|                                     | Tom MC, 2019 | HR = 0.97, \( P = .045 \) | - | - |
| KPS ≥ 90                            | Gousias K, 2014 | HR = 0.136, \( P < .001 \) | - | - |
| KPS > 80                            | Houillier C, 2010 | HR = 0.40, \( P = .009 \) | - | - |
|                                     | Houillier C, 2010 | HR = 0.441, \( P = .001 \) | - | - |
|                                     | Okita Y, 2012 | HR = 0.045, \( P = .0002 \) | - | - |

Note: HR = Hazard Ratio, \( P \) = p-value, NS = Not Significant
Table 3. Continued

| Demographic and radiographic factors | Study | Overall survival | Event-free survival |
|-------------------------------------|-------|------------------|--------------------|
| | | Univariate | Multivariate | Univariate | Multivariate |
| Radiographic factors | | | | | |
| G2 glioma eloquent location | Chang EF, 2011 | $P < .0001$ | HR = 6.1, $P < .001$ | Chang EF, 2011 | $P < .0001$ | HR = 1.9, $P = .003$ |
| | Gousias K, 2014 | HR = 3.498, $P = .008$ | | Gousias K, 2014 | | Eloquent location inferior $P < .001$ |
| False eloquent group (ref.) vs. true eloquent group by intra-operative mapping* | Chang EF, 2011 | | False eloquent group superior $P < .001$ | | | |
| G2 glioma MRI contrast enhancement | Goze C, 2014 | HR = 1.79, $P = .001$ | NS | Gousias K, 2014 | HR = 2.335, $P = 0.013$ | HR = 2.441, $P = .012$ |
| | Narang AK, 2017 | Contrast enhancement inferior $P = .03$ (recurrent cases) | NS | Pallud J, 2013 | $P = .014$ | HR = 1.44, $P < .011$ |
| G2 glioma corpus callosum involvement | Goze C, 2014 | HR = 4.69, $P = .042$ | NS | Pallud J, 2013 | | HR = 1.73, $P = .003$ |
| G2 glioma tumor volume $\geq 100$ cm$^3$ | Goze C, 2014 | HR = 2.44, $P = .002$ | HR = 9.69, $P = .017$ | Goze C, 2014 | HR = 2.44, $P = .022$ | NS |
| | Pallud J, 2013 | HR = 2.31, $P = .002$ | HR = 2.92, $P = .001$ | Pallud J, 2013 | | $P = .001$ |
| G2 glioma tumor size/volume (continuous) | Ius T, 2012 | HR = 8.20, $P < .0001$ | HR = 1.01, $P = .016$ | Ius T, 2012 | HR = 3.256, $P = .001$ | |
| | Kavouridis VK, 2020 | | | | HR = 1.06, $P < .0001$ | HR = 1.07, $P < .0001$ |
| G2 glioma velocity of diametric expansion $\geq 8$ mm/year | Goze C, 2014 | HR = 6.61, $P < .0001$ | HR = 26.3, $P < .0001$ | Goze C, 2014 | HR = 4.18, $P < .0001$ | HR = 4.23, $P = .001$ |
| | Pallud J, 2013 | HR = 3.96, $P < .001$ | HR = 4.62, $P < .001$ | Pallud J, 2013 | HR = 3.50, $P < .001$ | HR = 3.87, $P < .001$ |
| G2 glioma > 5 cm | Jairam V, 2019 | HR = 2.27, $P = .010$ | HR = 1.95, $P = .03$ | Nitta M, 2015 | NS | HR = 1.89, $P = .0428$ |
| | Tom MC, 2019 | Gioma > 5 cm inferior $P = .05$ | | Tom MC, 2019 | $P < .001$ | HR = 3.5, $P < .001$ |
| | Youland RS, 2013 | | HR = 1.70, $P < .0001$ | Youland RS, 2013 | | HR = 1.85, $P < .0001$ |
| G2 glioma > 3 cm | | | | | | |
| G2 oligodendroglioma tumor size (ref. 2.1–4 cm) | Harary M, 2020 | i) $\leq 2$ cm | ii) $2$ cm | ii) NS | i) NS | ii) $HR = 4.56, P = .02$ |
| | i) $2$ cm | ii) $4.1$–$6$ cm | iii) $> 6$ cm | | | |
| G2 glioma relative cerebral blood volume measurements | Majchrzak K, 2012 | HR = 7.39, $P = .002$ | | Majchrzak K, 2012 | HR = 1.70, $P = .033$ | |
low-grade spinal cord glioma, demonstrated a negative association between adjuvant radiotherapy and OS following adjustment for grade, age and surgical history.51

**Grade 2 glioma.**—Radiographic factors associated with OS and EFS among patients with grade 2 gliomas are summarized in Table 3. Imaging-related factors negatively associated with OS following multivariable analysis included: eloquent location,25 tumor volume over 100 cm$^3$,29,44 larger tumor size as a continuous variable,38 velocity of diametric expansion over 8 mm/year,29,44 size greater than 5 cm35,49 and size greater than 6 cm.30 Factors initially significantly associated with OS in univariate analyses but which lost association in multivariable analyses included contrast enhancement on MR29,40 and corpus callosum involvement.29 There was significant negative influence of eloquent location,25 MRI contrast enhancement,28,44 tumor volume greater than 100 cm$^3$,44 tumor size as a continuous variable,38,45,47 diametric annual expansion greater than 8 mm,29,44 size greater than 5 cm$^3$,41,46,49 and parietal compared to frontal location29 on grade 2 glioma EFS following adjusted multivariable analysis.

Histological and molecular factors are shown in Table 5. Among patients with astrocytomas, grade 2 histology conferred significantly worse OS than grade 1 histology.22 Diffuse astrocytoma histology was associated with inferior OS compared to oligoastrocytoma or oligodendroglioma histology following multivariable analysis.34–36,41,42,49 Oligodendroglioma was variably defined either histologically or molecularly across articles. Oligodendroglioma showed significantly favorable OS compared to IDH mutant and IDH wildtype astrocytoma.38,46 IDH mutant status29,33,37,42 and 1p19q co-deletion32,33 were positively associated with longer EFS. In one cohort of diffuse supratentorial low-grade gliomas, 1p19q co-deletion status was non-significant after adjusted multivariable analysis.29 In multivariable analysis, EFS was significantly inferior among those with diffuse astrocytoma histology,34,49 IDH mutant status29,33,37,42 and 1p19q co-deletion32,33 compared to IDH wildtype gliomas. Diffuse astrocytic histology43,47 and p53 over-expression47 were significantly negatively associated with EFS in univariate analysis but after adjustment in multivariable analysis were no longer significant. Notably, the studies that described IDH mutational status and influence on prognosis all comprised of cohorts that despite meeting our inclusion criteria, included substantial numbers of older adults (Figure 1). For example, of the 26 studies that included AYA patients with grade 2 glioma, 24 had a mean or median age above 30.

Treatment-related variables are summarized in Table 6. The impact of adjuvant chemoradiotherapy on OS and EFS was mixed. Combined adjuvant chemotherapy and radiotherapy positively impacted OS and EFS among grade 2 glioma patients in one study compared to adjuvant radiotherapy alone following multivariable analysis.42 Within this study the effect of adjuvant chemoradiotherapy was most pronounced in cases of IDH 1/2 mutant cases. By contrast Pal’a et al43 examined only IDH mutant grade 2

### Table 3. Demographic and radiographic factors

| Study | Event-free survival | Univariate | Multivariate |
|-------|--------------------|------------|--------------|
| Goze C, 2014 | |
| ii) | NS                  | |
| iii) | NS                  | |
| Jung C, 2014 | |
| i) | NS                  | |
| ii) | HR = 4.20, P = 0.019 |
| iii) | NS                  | |
Table 4. Demographic, radiographic, tumor and treatment influences on AYA WHO Grade 3 and 4 glioma event-free survival (EFS) and overall survival (OS)

| Imaging, treatment and tumor factors | Study | overall survival | Event-free survival |
|-------------------------------------|-------|-----------------|--------------------|
|                                     |       | Univariate      | Multivariate       | Study | Univariate | Multivariate |
|                                     |       |                 |                    |       |            |              |
| **Demographic factors**             |       |                 |                    |       |            |              |
| Age (continuous)                    | Yang W, 2018 | P < .001         | HR = 1.19, P < .001 | Olar A, 2015 | HR = 1.03, P < .0001 |
| Age ≤ 30                            | Gallo P, 2013 | HR = 0.81, P = .024 | HR = 0.05, P = .01 | Gallo P, 2013 | NS          | HR = 0.15, P = .01 |
| Female sex                          | Leibetseder A, 2013 | Age ≤ 30 superior, P < .05 | Age ≤ 30 superior, P < .05 | Leibetseder A, 2013 | NS          | HR = 0.15, P = .01 |
|                                    | Hatanpaa KJ, 2014 | RR = 5.02, P = .022 | RR = 5.02, P = .022 | Hatanpaa KJ, 2014 | NS          | HR = 0.15, P = .01 |
| **Radiographic characteristics**    |       |                 |                    |       |            |              |
| G3 and G4 spinal cord glioma tumor extent (ref. localized) | Liu J, 2018 | i) NS           | ii) NS          | iii) HR = 1.68, P = .045 | Liu J, 2018 | i) NS           | ii) NS          | iii) HR = 1.68, P = .045 |
| i) Regional extension               |       |                 |                    |       |            |              |
| ii) Invasive/distal extension       |       |                 |                    |       |            |              |
| iii) Unknown                        |       |                 |                    |       |            |              |
| **Histological factors**            |       |                 |                    |       |            |              |
| G2 (ref.) vs G3 PXA                 | Gallo P, 2013 | HR = 12.58, P = .003 | HR = 12.58, P = .003 | Gallo P, 2013 | HR = 12.58, P = .003 |
| G2 and G3 glioma oligodendrogloma (ref.) vs. astrocytoma | Miller JJ, 2019 | Oligodendrogloma superior, P = .025 | Oligodendrogloma superior, P = .025 | Miller JJ, 2019 | Oligodendrogloma superior, P = .025 |
| Spinal astrocytoma G2A (ref.) vs. G3A vs. G4A | Fakhreddine MH, 2013 | P = .0004 | P = .0004 | Fakhreddine MH, 2013 | P = .0004 |
| Recurrent G2 glioma new Histological grade unchanged vs. malignant degeneration (G3 or G4 glioma) | Narang AK, 2017 | P < .001 | P < .001 | Narang AK, 2017 | P < .001 |
| **Molecular factors**               |       |                 |                    |       |            |              |
| G2 and G3 glioma IDH mutant 1p19q co-deletion (ref.) vs. other | Olar A, 2015 | 1p19q co-deletion superior, P < .0001 | 1p19q co-deletion superior, P < .0001 | Olar A, 2015 | HR = 0.53, P = .0265 |
| G2 and G3 glioma 1p19q status non co-deleted (ref.) vs. co-deleted | Olar A, 2015 | P < .0001 | P < .0001 | Olar A, 2015 | HR = 0.53, P = .0265 |
| G2 and G3 glioma IDH mutant (ref.) vs. wt | Hatanpaa KJ, 2014 | P = .0006 | IDH mutant superior, P = .015 | Hatanpaa KJ, 2014 | P = .0006 |
| G2 and G3 glioma wt (ref.) vs. mutant | Miller JJ, 2019 | P = .015 | P = .015 | Miller JJ, 2019 | P = .015 |
| G2 and G3 glioma nestin level (continuous) | Olar A, 2015 | NS | NS | Olar A, 2015 | NS |
| G2 and G3 glioma mitotic index >4% | Hatanpaa KJ, 2014 | P = .0022 | P = .0022 | Hatanpaa KJ, 2014 | P = .0022 |
| i) IDH mutant                       |       |                 |                    |       |            |              |
| ii) IDH wt                          |       |                 |                    |       |            |              |
| **Chemotherapy**                    |       |                 |                    |       |            |              |
| G2 and G3 glioma adjuvant chemotherapy only | Miller JJ, 2019 | HR = 1.6, P = .047 | NS | Miller JJ, 2019 | HR = 1.6, P = .047 |
| G2 and G3 glioma combined adjuvant chemoradiation | Miller JJ, 2019 | HR = 0.57, P = .0026 | HR = 0.38, P = .0002 | Miller JJ, 2019 | HR = 0.57, P = .0026 | HR = 0.38, P = .0002 |
| Study | Variable | Radiation therapy | Surgical factors | Event-free survival Multivariate | Event-free survival Univariate | Overall survival Multivariate | Overall survival Univariate |
|-------|----------|-------------------|------------------|-------------------------------|------------------------------|-------------------------------|-------------------------------|
| Olar A, 2015 | i) IDH mutant ii) IDH wt | G2 and G3 glioma adjuvant radiotherapy | G2 and G3 glioma GTR (ref.) vs. STR | HR = 0.58, P = .0020 | HR = 0.55, P = .0028 | NS | NS |
| Miller JJ, 2019 | i) IDH mutant ii) IDH wt | G2 and G3 glioma adjuvant radiotherapy | G2 and G3 glioma GTR (ref.) vs. STR | HR = 0.54, P = .013 | HR = 0.35, P < .001 | HR = 0.54, P = .00014 | HR = 0.35, P < .001 |
| Liu J, 2018 | | G2 and G3 glioma adjuvant radiotherapy | G3 and G4 spinal cord glioma post-radiotherapy | HR = 0.54, P = .031 | HR = 0.54, P = .031 | NS | NS |
| Yang W, 2018 | | G3 and G4 peri-ventricular glioma adjuvant radiotherapy | G3 and G4 peri-ventricular glioma no resection i) Biopsy ii) STR iii) GTR | HR = 0.55, P = .01 | HR = 0.50, P < .001 | HR = 0.50, P < .001 | HR = 0.50, P < .001 |
| Hatanpaa KJ, 2014 | | G2 and G3 glioma GTR (ref.) vs. STR | Recurrent G2 glioma with transformation to G3 or G4 vs. GTR or STR or biopsy (ref.) | RR = 3.97, P = .037 | HR = 0.55, P = .007 | HR = 0.36, P = .001 | HR = 0.36, P = .001 |
| Narang AK, 2017 | | G2 and G3 glioma GTR (ref.) vs. STR | Recurrent G2 glioma with transformation to G3 or G4 vs. GTR or STR or biopsy (ref.) | P = .02 | HR = 0.50, P < .001 | HR = 0.50, P < .001 | HR = 0.50, P < .001 |
| Yang W, 2018 | | G3 and G4 peri-ventricular glioma no resection i) Biopsy ii) STR iii) GTR | Recurrent G2 glioma with transformation to G3 or G4 vs. GTR or STR or biopsy (ref.) | HR = 0.62, P = .007 | HR = 0.45, P < .001 | HR = 0.45, P < .001 | HR = 0.45, P < .001 |
| Cases with pooled WHO Grade 2 gliomas were included if they included WHO Grade 3 lesions. Bolded fields indicate statistical significance of the variable in cited study at an alpha of 0.05. | | | | | | | |

Cases with pooled WHO Grade 2 gliomas were included if they included WHO Grade 3 lesions. Bolded fields indicate statistical significance of the variable in cited study at an alpha of 0.05.
| Table 5. Molecular and Histological influences on AYA WHO Grade 2 glioma event-free survival (EFS) and overall survival (OS) |
|---|
| **Histological and molecular factors** | **Study** | **Overall survival** | **Event-free survival** |
| | | **Univariate** | **Multivariate** | **Study** | **Univariate** | **Multivariate** |
| G1 (ref.) vs G2 astrocytoma | Bagley JH, 2013 | HR = 3.04, P < .001 | | |
| Non-oligodendroglia histology and tumor size > 5 cm after surgery (ref.) vs. all other groups | Jairam V, 2019 | | HR = 2.76, P = .028 |
| G2 glioma oligodendroglia or oligoastrocytoma (ref.) vs. diffuse astrocytoma | Wahl M, 2017 | Oligodendroglia superior P = .007 | Ius T, 2012 | HR = 2.273, P = .003 |
| | Ius T, 2012 | | | |
| | Jairam V, 2019 | HR = 2.69, P = .002 | | |
| | Youland RS, 2013 | HR = 1.60, P < .001 | | |
| | Ius T, 2012 | HR = 4.262, P = .001 | Nitta M, 2015 | HR = 4.98, P = .0143 |
| | | HR = 4.98, P = .0143 | | |
| G2 glioma diffuse astrocytoma (ref.) vs. oligodendroglia | Okita Y, 2012 | P = .04 | Houillier C, 2010 | Oligodendroglia superior P = .03 |
| | Jansen E, 2019 | P = .002 | Pal’a A, 2019 | Oligodendroglia superior P = .026 |
| G2 glioma oligodendroglia (ref.) vs. oligoastrocytoma | Tom MC, 2019 | HR = 2.28, P = .03 | | |
| G2 glioma oligodendroglia (ref.) vs. IDH mutant astrocytoma | Kavouridis VK, 2020 | | Kavouridis VK, 2020 | -- |
| | Wahl M, 2017 | Oligodendroglia superior P = .01 | Wahl M, 2017 | Oligodendroglia superior P < .001 |
| | Tom MC, 2019 | i) NS | i) HR = 7.76, P < .001 |
| | | ii) P = .001 | ii) HR = 20.6, P < .001 |
| G2 glioma IDH wt (ref.) vs. IDH1/2 mutant | Jungk C, 2016 | | Kavouridis VK, 2020 | -- |
| | Houillier C, 2010 | HR = 0.11, P = .0003 | | |
| | Okita Y, 2012 | HR = 0.091, P = .002 | | |
| | Goze C, 2014 | P = .002 | | |
| | | HR = 0.365, P = .01 | | |
| | | HR = 0.306*, P = .044 | | |
| | | HR = 0.306*, P = .044 | | |
| | | HR = 0.056*, P = .007 | | |
| | | HR = 0.056*, P = .007 | | |
| | | HR = 0.199*, P < .0001 | | |
| | | HR = 0.314, P = .025 | | |
| G2 glioma 1p19q co-deletion (ref. non co-deleted) | Houillier C, 2010 | P < .0001 | Houillier C, 2010 | P = .02 |
| | Houillier C, 2010 | P = .001 | Houillier C, 2010 | P = .002 |
| | Esenou CI, 2017 | HR = 0.291, P = .05 | Youland RS, 2013 | 1p19q co-deletion superior P < .0001 |
| | Pallud J, 2013 | HR = 0.45, P = .040 | | |
| | Youland RS, 2013 | 1p19q co-deletion superior P < .0001 | | |
| | Goze C, 2014 | HR = 0.256*, P = .031 | NS | |
glioma patients and found a negative impact of adjuvant chemoradiotherapy on EFS and OS after adjusting for age over 40 years, extent of resection, recurrent surgery and histology. Coburger et al. also showed a negative impact of adjuvant chemoradiotherapy compared to no adjuvant therapy on EFS in a cohort of grade 2 glioma after adjusting for age, recurrent surgery, histology and residual tumor in their multivariable model. One group showed in LGG that combined chemoradiotherapy (temozolomide) was superior in EFS compared to chemotherapy alone in a multivariable model with covariates gender, tumor size, molecular characteristics and adjuvant therapy regimen.

Several studies did not specify the adjuvant therapy regimen used, though showed chemoradiotherapy was associated with an unfavorable effect on OS following multivariable analysis. Gousias et al. showed a negative association between adjuvant therapy and OS, but did not conduct multivariable analyses for this outcome; only 5% of their cohort underwent either chemotherapy and or radiotherapy. In their multivariable analyses conducted for EFS however, including eloquent location as a covariate, adjuvant therapy had a favorable impact on EFS.

Conflicting results related to the role of adjuvant chemotherapy were observed; one group showed a positive association with both adjuvant chemotherapy and radiotherapy with increased EFS in multivariable analysis that included covariates age, histology, presenting symptoms, size and extent of resection. Another study showed increased EFS but no significant change in OS with adjuvant chemotherapy following LGG resection after multivariable analysis with covariates age, tumor diameter, pathology and adjuvant therapy.

Few studies analyzed the role of adjuvant radiotherapy alone upon OS, though one included study demonstrated a significant negative impact on OS after multivariable analysis including age at diagnosis, molecular class, eloquent location, and post-operative residual volume. Adjuvant radiotherapy significantly improved EFS in two studies and the effect was suggested to be greater with immediate as opposed to delayed radiotherapy following univariate analysis alone in two other reports.

Non-significant prognostic variables are shown in Supplementary Table 1. Following multivariable analysis, several studies found a non-significant association between OS and adjuvant chemotherapy, adjuvant radiotherapy and combined adjuvant chemoradiotherapy.

Several studies looked at the impact of surgery-related factors. Increased extent of resection compared to biopsy alone was associated with both OS and EFS in multivariable adjusted models. Extent of resection measured as either a continuous variable or lower magnitude of post-operative volumetric tumor residual correlated with prolonged OS and/or EFS. Several studies showed in adjusted multivariable analysis that GTR resulted in superior OS or EFS benefit compared to other resection categories, though one study showed negative effect on EFS in IDH mutant astrocymoma. One study found that first line surgical therapy compared to observation did not significantly influence OS though it favorably impacted EFS. Factors associated with positive impact on OS following univariate analysis (in absence of
### Table 6. Treatment-related influences on AYA WHO Grade 2 glioma event-free survival (EFS) and overall survival (OS)

| Treatment factors | Study | Overall survival | Event-free survival |
|-------------------|-------|------------------|---------------------|
|                   |       | Univariate | Multivariate | Univariate | Multivariate |
| Combined adjuvant therapy | G2 glioma post-operative radiotherapy alone (ref.) vs. chemoradiotherapy | Okita Y, 2012 | $P = .0002$ | $HR = 0.198, P = .002$ | Okita Y, 2012 | $P = .01$ | $HR = 0.408, P = .04$ |
|                   | G2 glioma IDH mutant adjuvant therapy (yes ref. vs. no) | Pal’a A, 2019 | No adjuvant therapy superior $P = .003$ | No adjuvant therapy superior $P = .009$ | Pal’a A, 2019 | No adjuvant therapy superior $P = .003$ | HR not stated | $P = .030$ | i) NS | ii) NS | iii) $HR = 2.745, P = .004$ |
|                   | i) No therapy vs. chemotherapy | Pal’a A, 2019 | i) NS | ii) NS | iii) $HR = 20.175, P = .001$ | Pal’a A, 2019 | i) NS | ii) NS | iii) $HR = 3.8, P = .008$ |
|                   | ii) No therapy vs. radiotherapy | Pal’a A, 2019 | i) NS | ii) NS | iii) $HR = 2.745, P = .004$ | Pal’a A, 2019 | i) NS | ii) NS | iii) $HR = 3.8, P = .008$ |
|                   | iii) No therapy vs. chemoradiotherapy | Pal’a A, 2019 | i) NS | ii) NS | iii) $HR = 3.8, P = .008$ | Pal’a A, 2019 | i) NS | ii) NS | iii) $HR = 3.8, P = .008$ |
|                   | G2 glioma temozolomide and radiotherapy (ref.) vs. i) Observation | Tom MC, 2019 | i) $HR = 0.3, P < .001$ | ii) NS | iii) $HR = 0.4, P = .004$ | Tom MC, 2019 | i) NS | ii) NS | iii) $HR = 0.4, P = .004$ |
|                   | ii) Radiation alone | Tom MC, 2019 | i) $HR = 0.3, P < .001$ | ii) NS | iii) $HR = 0.4, P = .004$ | Tom MC, 2019 | i) NS | ii) NS | iii) $HR = 0.4, P = .004$ |
|                   | iii) Temozolomide alone | Tom MC, 2019 | i) $HR = 0.3, P < .001$ | ii) NS | iii) $HR = 0.4, P = .004$ | Tom MC, 2019 | i) NS | ii) NS | iii) $HR = 0.4, P = .004$ |
|                   | G2 glioma post-operative tumor volume ≤ 68 cm³ prior to adjuvant therapy | Wahl M, 2017 | ≤ 68 cm³ superior $P < .001$ | | Wahl M, 2017 | ≤ 68 cm³ superior $P < .001$ | | | | |
|                   | G2 glioma adjuvant chemoradiation therapy | Coburger J, 2016 | – | | | | | | |
| Adjuvant therapy NOS | G2 glioma adjuvant therapy | Gousias K, 2014 | $HR = 8.115, P < .001$ | | Gousias K, 2014 | $HR = 2.449, P = .039$ | | | |
|                   | G2 astrocytoma adjuvant therapy following surgery at diagnosis (ref. is yes) | Jungk C, 2016 | $HR = 6.25, P = .0010$ | $HR = 7.13, P = .003$ | | Jansen E, 2019 | Adjuvant therapy and surgery superior $P = .0001$ | | | |
|                   | G2 glioma adjuvant therapy and surgery at first relapse vs surgery alone | Jansen E, 2019 | | | | | | |
| Chemotherapy | G2 glioma post-operative chemotherapy vs. no chemotherapeutic | Nitta M, 2015 | | | | | | |
| Radiation therapy | G2 glioma adjuvant radiotherapy (ref. no radiotherapy) | Kavouridis VK, 2020 | $HR = 2.99, P = .001$ | | Kavouridis VK, 2020 | | | | |
|                   | G2 glioma immediate (ref.) vs. delayed post-operative radiotherapy | Youland RS, 2013 | NS | | Youland RS, 2013 | | | | |
|                   | G2 glioma immediate (ref.) vs. delayed post-operative radiotherapy | Ius T, 2012 | | | | | | |
|                   | G2 glioma post-operative radiotherapy i) Diffuse astrocytoma ii) Oligodendroglioma | Houllier C, 2010 | Delayed radiotherapy inferior $P = .0001$ | Delayed radiotherapy inferior $P = .0001$ | Houllier C, 2010 | Delayed radiotherapy inferior $P = .0001$ | | | |
|                   | i) Diffuse astrocytoma ii) Oligodendroglioma | Nitta M, 2015 | i) NS | ii) Adjuvant radiotherapy superior $P = .02$ | | | | | |
| Treatment factors | Study | Overall survival | Event-free survival |
|-------------------|-------|------------------|--------------------|
| Surgical factors  |       |                  |                    |
| G2 glioma use of intra-operative electrical stimulation with or without addition of intra-op DTI/fMRI navigation | Ius T, 2012 | HR = 0.388, \( P < .016 \) |                    |
|                    |       |                  | Kavouridis V.K., 2020 | HR = 1.69, \( P = .007 \) |
| G2 glioma surgery (ref.) vs. biopsy alone | Gousias K, 2014 | HR = 0.132, \( P < .001 \) | Palud J, 2013 | Surgery superior, \( P < .001 \) |
|                    | Wahl M, 2017 | Surgery superior, \( P = .01 \) | Wahl M, 2017 | Surgery superior, \( P = .003 \) |
| G2 glioma EOR biopsy (ref.) | Goze C, 2014 | i) HR = 0.18, \( P = .031 \), ii) NS, iii) NS | Goze C, 2014 | i) NS, ii) HR = 0.34, \( P = .038 \), iii) NS | i) HR = 0.27, \( P = .021 \), ii) NS, iii) HR = 0.25, \( P = .025 \) |
| i) STR |                    |                    |                    |                    |
| ii) NTR |                    |                    |                    |                    |
| iii) GTR |                    |                    |                    |                    |
| G2 glioma % EOR (continuous) | Eseonu C.I, 2017 | HR = 0.994, \( P = .016 \) | Eseonu C.I, 2017 | HR = 0.983, \( P = .016 \) |
| i) ≥ 90% (ref) | Ius T, 2012 | HR = 0.933, \( P < .0001 \) | Ius T, 2012 | HR = 0.930, \( P < .0001 \) |
| ii) 70–90% | Majchrzak K, 2012 | HR = 0.96, \( P = .025 \) | Jung C, 2016 | HR = 0.23, \( P < .001 \) |
| iii) <70% | Majchrzak K, 2012 | HR = 0.96, \( P = .025 \) | Majchrzak K, 2012 | HR = 0.98, \( P = .004 \) |
|                | Scherer M, 2020 | Smaller tumor volume superior, \( P = .02 \) | Scherer M, 2020 | P < .001 |
| G2 glioma post-operative volume (cm\(^3\)) (continuous) | Kavouridis V.K., 2020 | HR = 1.02, \( P = .0001 \) | Kavouridis V.K., 2020 | HR = 1.01, \( P = .001 \) |
| i) Oligodendroglioma (9 vs. ≥9) | Kavouridis V.K., 2020 | Smaller tumor volume superior, i) \( P = .048 \), ii) \( P = .017 \), iii) \( P = .017 \) | Majchrzak K, 2012 | HR = 1.01, \( P = .008 \) |
| ii) IDH mutant astrocytoma (1 vs. ≥1) |                |                    |                    |                    |
| iii) IDH wt astrocytoma (1 vs. ≥1) |                |                    |                    |                    |
| G2 glioma % EOR | Ius T, 2012 | HR = 4.845, \( P = .002 \) | Ius T, 2012 | ii) HR = 3.402, \( P < .0001 \), iii) HR = 13.60, \( P < .0001 \) |
| i) ≥ 90% (ref) |                    |                    |                    |                    |
| ii) 70–90% |                    |                    |                    |                    |
| iii) <70% |                    |                    |                    |                    |
| Treatment factors | Study | Overall survival | Event-free survival |
|-------------------|-------|-----------------|--------------------|
|                   |       | Univariate      | Multivariate       | Study | Univariate      | Multivariate       |
|                   |       | G2 glioma non-GTR vs. GTR |                   |       |                   |                     |
| i) Oligodendrogioma |       |                 |                    |       |                   |                     |
| ii) Diffuse astrocytoma IDH wt IDH mutant |       |                 |                    |       |                   |                     |
|                   | Houiller C, 2010 | NS              | HR = 0.51, P = .03 |       |                   |                     |
|                   | Coburger J, 2016 | P < .05         |                   |       |                   |                     |
|                   | Houiller C, 2010 | GTR superior P = .0004 | NS              |       |                   |                     |
|                   | Youland RS, 2013 | GTR superior P < .0001 | HR = 0.51, P < .0001 |       |                   |                     |
|                   | Houiller C, 2010 |                   | GTR superior P = .02 |       |                   |                     |
|                   | Coburger J, 2016 |                   | P < .001          |       |                   |                     |
|                   | Scherer M, 2020 |                   | GTR superior P = .009 |       |                   |                     |
|                   | Jansen E, 2019 |                   | i) GTR superior P = .002 |       |                   |                     |
|                   |                   | i) GTR superior P = .037 | NS              |       |                   |                     |
|                   |                   | ii) NS           |                   |       |                   |                     |
|                   | Pal’a A, 2019 |                   | i) P = .035 |       |                   |                     |
|                   | Youland RS, 2013 |                   | ii) HR = 0.486, P = .019 |       |                   |                     |
|                   | Jansen E, 2019 | P = .003         | HR 2.6, P = .017 |       |                   |                     |
|                   | Jansen E, 2019 |                   | P = .001          |       |                   |                     |
|                   | Gallo P, 2013 |                   | HR 16.30, P = .004 |       |                   |                     |
|                   |                   |                   | HR = 4.60, P = .006 |       |                   |                     |
|                   |                   |                   | HR = 15.97, P = .001 |       |                   |                     |
|                   | Goze C, 2014 |                   | HR 0.41, P = .042 |       |                   |                     |
|                   | Goze C, 2014 |                   | NS              |       |                   |                     |
|                   | Pal’ud J, 2013 |                   | HR 0.42, P < .001 |       |                   |                     |
|                   |                   |                   | HR = 0.44, P < .001 |       |                   |                     |
|                   | Gousias K, 2014 |                   | i) HR = 0.306, P = .001 |       |                   |                     |
|                   |                   | i) HR = 0.045, P < .001 | ii) HR = 0.039, P < .001 |       |                   |                     |
|                   |                   | i) HR = 0.234, P < .001 |                   |       |                   |                     |
|                   |                   | ii) HR = 0.039, P < .001 |                   |       |                   |                     |
|                   | Harary M, 2020 |                   | i) P = .002 |       |                   |                     |
|                   | Tom MC, 2019 |                   | i) P = .002 |       |                   |                     |
|                   |                   | i) P = .003 |       |                   |                     |
|                   |                   | ii) P < .001 |       |                   |                     |
|                   |                   | ii) HR = 0.3, P < .001 |       |                   |                     |
|                   |                   | i) NS           |       |                   |                     |
|                   |                   | ii) GTR superior P = .002 |       |                   |                     |
|                   | Ius T, 2012 |                   | HR 1.040, P < .0001 |       |                   |                     |
|                   | Ius T, 2012 |                   | HR 1.034, P < .0001 |       |                   |                     |
|                   |                   |                   | HR 1.021, P < .001 |       |                   |                     |
|                   | Ius T, 2012 |                   | HR 3.699, P < .0001 |       |                   |                     |
|                   | Ius T, 2012 |                   | HR 1.035, P < .0001 |       |                   |                     |
|                   | Ius T, 2012 |                   | HR 3.427, P < .0001 |       |                   |                     |
|                   | Ius T, 2012 |                   | HR 1.022, P < .0001 |       |                   |                     |
|                   | Ius T, 2012 |                   | HR 1.023, P < .0001 |       |                   |                     |
Adjusted multivariable analysis) included: decreasing post-operative T2-weighted MRI signal volume, greater extent of resection across histological types, and smaller post-operative tumor volume.

**Grade 3 and 4 glioma.**—Groupings of Grade 3 and 4 glioma in included studies may not have reflected current classification schemes that include IDH mutational status. In addition, Grade 3 glioma may or may not be included in the definition of high-grade glioma. However, grouping Grade 3 and 4 glioma best reflected the categorization used by the papers identified in this systematic review.

**Table 4** summarizes disease and treatment-related factors influencing EFS and OS in HGG. Among high-grade spinal cord glioma, there was no significant influence on localized vs. regional or invasive location on OS. Oligodendroglioma histology showed superior influence on OS compared to astrocytic histology in pooled grade 2 and 3 cases following univariate analysis (no multivariable analysis reported). Grade 3 and 4 spinal cord glioma were negative influences on OS when compared to grade 2 histology. Grade 3 and 4 spinal cord glioma were negative influences on OS when compared to grade 2 histology. 1p19q co-deletion, IDH mutant status, low nestin level, and mitotic index less than 4% all positively impacted OS in combined grade 2 and 3 glioma cases.

Some studies included in this review showed adjuvant radiotherapy demonstrated favorable impact on OS in pooled grade 2 and 3 glioma, pooled grade 3 and 4 spinal cord glioma, and peri-ventricular HGG. STR or biopsy-only resulted in worse OS than GTR or near-total resection (NTR) in two studies. Though in peri-ventricular HGG STR and GTR were favorably associated with OS in univariate analysis compared to no surgery, they lost significance following adjusted multivariable analysis. Adjuvant chemoradiation positively impacted EFS in grade 2 and 3 glioma, though chemotherapy alone was not significant. One combined cohort of grade 2 and 3 glioma showed a non-significant influence of adjuvant chemoradiotherapy on OS following multivariable analysis.

Excluding spinal pilocytic astrocytoma, Fakrehddine et al showed adjuvant chemotherapy significantly improved EFS in infiltrative spinal cord glioma (grades 2, 3 and 4) after adjusting for treatment modality, age at diagnosis, grade, number of spinal levels, neurological deficits and symptom duration. In the same analysis, adjuvant radiotherapy did not significantly impact EFS nor did either chemotherapy or radiation contribute to OS benefit after multivariable analysis.

**Quality Assessment**

Given the absence of methodological limitation reporting across studies, the QUIPS assessment tool was utilized to provide a standardized risk of bias assessment (Supplementary Table 2). Most studies (35/40) had at least 1 domain that scored in the high risk of bias category. Among included studies only 1 was prospective. Common domains for high risk of bias include study participation and adjustment for other prognostic factors.

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**Table 6.** Continued

| Study | Overall survival | Event-free survival |
|-------|------------------|---------------------|
| Ius T , 2012 | ii) | i) HR = 3.281, \( P = .009 \) |
| | iii) | ii) NS |
| | iv) | iii) HR = 13.980, \( P < .0001 \) |

| Study | Overall survival | Event-free survival |
|-------|------------------|---------------------|
| Nitta M, 2015 | i) | i) \( P = .0096 \) |
| | ii) NS |
| | iii) | ii) NS |

| Study | Overall survival | Event-free survival |
|-------|------------------|---------------------|
| Pal’s A, 2019 | Recurrent surgery | Recurrent surgery |
| | superior | superior |
| | \( P = .012 \) | \( P = .012 \) |
Discussion

This systematic review identified 40 studies that reported on demographic, disease and treatment predictors of EFS and OS among AYA glioma patients in high income countries. Despite stringent definitions utilized to capture an adequately sized AYA cohort, several included studies captured a proportion of older adults (Figure 1). This points to a severe limitation in the existing AYA glioma literature, with all interpretation limited by the potential impact of older adult glioma biology in these cohorts. In contrast, only two studies included pediatric patients.52,53 Furthermore, many papers scored in the high-risk bias category in at least one domain. Despite this, several patient epidemiological, disease and treatment factors with prognostic impact on EFS and OS were identified.

Prognostication

There are important differences in glioma prognostication in adult and pediatric populations. In a national pediatric cohort study, lower tumor grade, GTR, non-brainstem location and age >1 year at diagnosis were all associated with longer OS.60 Recent clinical and molecular characterization has underscored the importance of single-nucleotide variant (SNV) and rearrangements in the pathobiology of pediatric LGG with SNV-driven tumors exhibiting inferior OS.5 Several molecular factors have important prognostic implications in pediatric LGG including mutations in BRAF V600E, KIAA1549-BRAF and NF-1 along with other less commonly encountered oncogenes. Identification of H3 K27M mutation in pediatric glioma portends a worse prognosis regardless of histologic diagnosis and modifies this clinical entity to WHO grade 4.51,52 Pathological and molecular favorable prognostic characteristics in adult glioma include IDH mutant, MGMT promoter methylation, non-astrocytoma histology or 1p/19q co-deletion and lower glioma grade when compared to IDH-WT glioma in older adults.63,64 Importantly, the influence of IDH mutation status in the AYA LGG is still not clear as this mutation does not portend the same prognostic importance in pediatric populations where it is encountered more rarely.5 Despite being highlighted as an important prognostic factor in this review, we are cognizant that this may reflect bias from inclusion of older adults, where IDH mutation is a known favorable molecular prognosticator (Figure 1). The role of IDH mutations in AYA, particularly younger AYA, remains uncertain.

Despite the AYA glioma demographic straddling the late pediatric and early adulthood age ranges, no studies in this systematic review comprehensively examined molecular prognostic markers. It is thus impossible to outline the specific prognostic impact of various molecular alterations in the AYA demographic. Instead, the literature could only confirm more the favorable impact of traditional adult prognosticators such as younger age at diagnosis, higher functional status, IDH mutant status (with limitations discussed above), lower glioma grade and 1p/19q co-deletion/ oligodendroglialoma histology with limited information on clinical behavior of tumors with other molecular alterations.

The effect of traditional functional status indicators such as KPS may reflect the older adults included in the review cohort. Furthermore, we have utilized previously described age parameters (15–39) for definition of AYA glioma patients; this is an assumption that will require future validation in this disease entity.5,55 Despite the widely accepted AYA age range, patients at the upper and lower end of the spectrum may be clinically distinct. Comprehensive molecular analyses among AYA cohort and their prognostic impact is a significant priority for future research.

Treatment

Several surgical factors were identified as important treatment-related factors for OS and EFS among AYA glioma patients. Extent of surgical resection was identified as an important positive factor associated with EFS and OS.26,27,29,30,32,34,36,38,40,45,46,49,51,53,54 The degree of resection and extent-of-resection categories within each study were not standardized nor was the definition of NTR and STR across studies. However, this favorable survival influence was present in several studies after multivariable analysis when GTR or NTR was compared to other resection categories in LGG or HGG cases.29,30,32,36,40,46,49,51,53,54 Furthermore, the impact of surgery was demonstrated in different anatomic compartments such as spinal cord glioma,51 in the setting of recurrent transformed LGG40 and different intracranial LGG pathological subtypes,52,49,53 though not in peri-ventricular HGG.59 This is in keeping with traditional surgical principles in glioma management across the age spectrum.

The role of adjuvant therapy and its influence on OS remains unclear in the current literature. One significant limitation is heterogeneous chemotherapy regimens in tumors with differing duration, agents and timing. Indeed, some studies did not provide any details of the regimen used. Radiotherapy doses ranged between 54 and 60 Gy. Secondly, despite attempts at adjustment for confounders through multivariable analyses, many studies could not fully account for patient, disease, surgical, or institutional factors that may influence the choice of chemotherapy and radiotherapy. For example, in several LGG studies, adjuvant radiotherapy conferred a negative survival benefit.20,23,38,43,51 The reasons for this disadvantage may include confounders such as residual tumor and radiographic or symptomatic progression or irradiation associated complications including secondary malignancies, transformation or vasculopathies.

Discussion about the role of chemotherapy and radiotherapy in AYA glioma raises several important points. First, AYA glioma patients have historically been under-represented in clinical trials that have established current chemotherapy and radiotherapy regimens.65–67 Our review shows that the current literature does not guide clinicians treating AYA with LGG on whether pediatric or adult approaches are more suitable, or indeed whether a tailored approach unique to AYA is required. In both groups, treatment approaches are informed by histopathological and molecular characteristics. Many pediatric patients treated with surgery alone despite post-surgical residual disease in an effort to avoid the long-term impacts of radiation or chemotherapy.5 In contrast, in older adults LGG or those with residual tumor following resection, combination
Chemotherapy and radiation therapy is usually considered. A major challenge is the lack of studies in this review including details about the presence of pediatric-type alterations in AYA glioma, thus limiting any meaningful molecularly informed conclusions about adjuvant chemoradiotherapy. Whether there is a role for adjuvant therapy among AYA with LGG either totally resected or with residual disease is a crucial question that should be prioritized.

Though HGG in pediatric and adult patients may share similarities in overall prognosis, there are important differences that exist between treatment regimens and biological considerations. At a molecular level, the profile of HGG is different with distinct copy number aberrations and driver mutations in pediatric HGG compared to adults. Furthermore, cancer predisposition syndromes are more common in pediatric populations compared to adults. The extent to which these pediatric-type alterations and predispositions exist in AYA demographics is not well known and was not clarified through this review, thus highlighting a major gap in understanding. Stupp et al showed that adults with HGG had improved OS with adjuvant temozolomide in combination with fractionated radiotherapy compared to radiotherapy alone. Radiotherapy typically begins 3–5 weeks following surgical resection and is typically administered at 50–60 Gy in 1.8–2 Gy fractions with limited evidence suggesting any added benefit at higher doses. For patients with MGMT methylated promoter glioblastoma, recurrent or progressive HGG, second line alkylating chemotherapeutics may be considered. By contrast, the benefit of adjuvant temozolomide in the treatment of pediatric HGG is debatable. This is highlighted by contrasting two prospective trials. Cohen et al. showed temozolomide administration during and after adjuvant radiotherapy in pediatric HGG did not improve outcomes. In contrast, Jakacki et al demonstrated that children with maximally resected non-metastatic HGG treated with radiotherapy and concomitant temozolomide followed by lomustine and temozolomide adjuvant chemotherapy experienced significantly improved outcomes. Despite the complexity in decision making surrounding HGG adjuvant therapy, our review highlights that AYA-specific data to guide clinicians is lacking.

Limitations stem from the predominance of retrospective studies included in this systematic review as well as the inclusion of older adults in many study cohorts. Despite intentions to identify and assess prognostic factors in AYA glioma, the inclusion of older adults skews the results and limits generalizability. However, stricter age-based inclusion criteria would have resulted in the exclusion of nearly all studies. Pediatric glioma mutational markers were rarely examined, precluding assessment of their prognostic value in AYA populations. Our review included all CNS gliomas, including spinal gliomas, though the latter may require different treatment approaches owing to differing biology anatomical considerations. Finally, the majority of studies were classified as at high risk of bias in at least one domain.

**Conclusion**

Although this study reveals some traditional factors that appear prognostically important in AYA glioma, most, including tumor grade, pathological subtype and genetic mutations such as IDH1/2, need to be considered with care given bias from the inclusion of older adults in many studies. Interestingly, the role of cytoreductive surgery remains an important prognostic factor in AYA gliomas and may not change until effective adjuvant medical therapies emerge. As such, the current literature does not provide clinicians with an evidence-based approach to treating AYA with gliomas, particularly regarding the role of adjuvant chemotherapy and radiotherapy. Available evidence is heterogeneous, of mixed quality, at high risk for confounding, and predominantly derived from older adult cohorts. Prospective studies of histopathological and molecularly-defined gliomas exposed to uniform treatment including both short- and long-term outcomes will allow the identification of optimal AYA-specific glioma management strategies.
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