Adult Central Nervous System Tumors Treatment (PDQ®)

Health Professional Version

PDQ Adult Treatment Editorial Board.

Published online: October 27, 2015.

This PDQ cancer information summary for health professionals provides comprehensive, peer-reviewed, evidence-based information about the treatment of adult central nervous system tumors. It is intended as a resource to inform and assist clinicians who care for cancer patients. It does not provide formal guidelines or recommendations for making health care decisions.

This summary is reviewed regularly and updated as necessary by the PDQ Adult Treatment Editorial Board, which is editorially independent of the National Cancer Institute (NCI). The summary reflects an independent review of the literature and does not represent a policy statement of NCI or the National Institutes of Health (NIH).

General Information About Adult Primary Central Nervous System (CNS) Tumors

Incidence and Mortality

Brain tumors account for 85% to 90% of all primary central nervous system (CNS) tumors.[1] Estimated new cases and deaths from brain tumors and other nervous system tumors in the United States in 2015:[2]

- New cases: 22,850.
- Deaths: 15,320.

Available registry data from the Surveillance, Epidemiology, and End Results (SEER) database for 2011 indicate that the combined incidence of primary invasive CNS tumors in the United States is 6.4 per 100,000 persons per year, with an estimated mortality of 4.3 per 100,000 persons per year.[3] Worldwide, approximately 256,213 new cases of brain and other CNS tumors were diagnosed in the year 2012, with an estimated 189,382 deaths.[4]

In general, the incidence of primary CNS tumors is higher in whites than in blacks, and mortality is higher in males than in females.[1]

Primary brain tumors include the following in decreasing order of frequency:[1]

- Anaplastic astrocytomas and glioblastomas (38% of primary brain tumors).
- Meningiomas and other mesenchymal tumors (27% of primary brain tumors).
- Pituitary tumors.
- Schwannomas.
- CNS lymphomas.
- Oligodendroglialomas.
- Ependymomas.
- Low-grade astrocytomas.
- Medulloblastomas.

Primary spinal tumors include the following in decreasing order of frequency:

- Schwannomas, menigiomas, and ependymomas (79% of primary spinal tumors).
- Sarcomas.
Primary brain tumors rarely spread to other areas of the body, but they can spread to other parts of the brain and to the spinal axis.

Anatomy

Anatomy of the inside of the brain. The supratentorium contains the cerebrum, ventricles (with cerebrospinal fluid shown in blue), choroid plexus, hypothalamus, pineal gland, pituitary gland, and optic nerve. The infratentorium contains the cerebellum and brain stem.

Risk Factors

Few definitive observations have been made about environmental or occupational causes of primary CNS tumors.[1]

The following potential risk factors have been considered:
Exposure to vinyl chloride may be a risk factor for glioma.

Epstein-Barr virus infection has been implicated in the etiology of primary CNS lymphoma.

Transplant recipients and patients with AIDS have a substantially increased risk of primary CNS lymphoma.[1,5] (Refer to the PDQ summary on Primary CNS Lymphoma Treatment for more information.)

The familial tumor syndromes and related chromosomal abnormalities that are associated with CNS neoplasms include the following:[6,7]

- Neurofibromatosis type I (17q11).
- Neurofibromatosis type II (22q12).
- von Hippel-Lindau disease (3p25-26).
- Tuberous sclerosis (9q34, 16p13).
- Li-Fraumeni syndrome (17p13).
- Turcotte syndrome type 1 (3p21, 7p22).
- Turcotte syndrome type 2 (5q21).
- Nevoid basal cell carcinoma syndrome (9q22.3).

Clinical Features

The clinical presentation of various brain tumors is best appreciated by considering the relationship of signs and symptoms to anatomy.[1]

General signs and symptoms include the following:

- Headaches.
- Seizures.
- Visual changes.
- Gastrointestinal symptoms such as loss of appetite, nausea, and vomiting.
- Changes in personality, mood, mental capacity, and concentration.

Seizures are a presenting symptom in approximately 20% of patients with supratentorial brain tumors and may antedate the clinical diagnosis by months to years in patients with slow-growing tumors. Among all patients with brain tumors, 70% with primary parenchymal tumors and 40% with metastatic brain tumors develop seizures at some time during the clinical course.[8]

Diagnostic Evaluation

All brain tumors, whether primary, metastatic, malignant, or benign, must be differentiated from other space-occupying lesions that can have similar clinical presentations, such as abscesses, arteriovenous malformations, and infarctions.[9]

Imaging tests

Contrast-enhanced computed tomography (CT) and magnetic resonance imaging (MRI) have complementary roles in the diagnosis of CNS neoplasms.[1,9,10]

- The speed of CT is desirable for evaluating clinically unstable patients. CT is superior for detecting calcifications, skull lesions, and hyperacute hemorrhages (bleeding less than 24 hours old) and helps direct differential diagnosis and immediate management.
- MRI has superior soft-tissue resolution. MRI can better detect isodense lesions, tumor enhancements, and associated findings such as edema, all phases of hemorrhagic states (except hyperacute), and infarctions.
High-quality MRI is the diagnostic study of choice in the evaluation of intramedullary and extramedullary spinal cord lesions.[1]

In posttherapy imaging, single-photon emission computed tomography (SPECT) and positron emission tomography (PET) may be useful in differentiating tumor recurrence from radiation necrosis.[9]

Biopsy

Biopsy confirmation to corroborate the suspected diagnosis of a primary brain tumor is critical, whether before surgery by needle biopsy or at the time of surgical resection. Cases in which the clinical and radiologic picture clearly point to a benign tumor, which could potentially be managed with active surveillance without biopsy or treatment, are the exception. For other cases, radiologic patterns may be misleading, and a definitive biopsy is needed to rule out other causes of space-occupying lesions, such as metastatic cancer or infection.

CT- or MRI-guided stereotactic techniques can be used to place a needle safely and accurately into almost all locations in the brain.

Prognostic Factors

Several genetic alterations have emerged in recent years as powerful prognostic factors in diffuse glioma (astrocytoma, oligodendroglioma, mixed glioma, and glioblastoma), and these alterations may guide patient management. Specific alterations include the following:

- DNA methylation of the O6-methylguanine-DNA methyltransferase (MGMT) gene promoter.
- Mutation of isocitrate dehydrogenase (IDH) 1 (IDH1) or IDH2 genes.
- Codeletion of chromosomes 1p and 19q.

(Refer to the Treatment of Primary Central Nervous System Tumors by Tumor Type section of this summary for more information.)

Related Summaries

Refer to the following PDQ summaries for more information:

- Primary CNS Lymphoma Treatment.
- Childhood Brain and Spinal Cord Tumors Treatment Overview (treatment of brain tumors in children).

References

World Health Organization (WHO) Classification of Adult Primary CNS Tumors

This classification is based on the World Health Organization (WHO) classification of central nervous system (CNS) tumors.[1] The WHO approach incorporates and interrelates morphology, cytogenetics, molecular genetics, and immunologic markers in an attempt to construct a cellular classification that is universally applicable and prognostically valid. Earlier attempts to develop a TNM-based classification were dropped for the following reasons:[2]

- Tumor size (T) is less relevant than are tumor histology and location.
- Nodal status (N) does not apply because the brain and spinal cord have no lymphatics.
- Metastatic spread (M) rarely applies because most patients with CNS neoplasms do not live long enough to develop metastatic disease.

The WHO grading of CNS tumors establishes a malignancy scale based on histologic features of the tumor.[3] The histologic grades are as follows:

- **WHO grade I** includes lesions with low proliferative potential, a frequently discrete nature, and the possibility of cure following surgical resection alone.
- **WHO grade II** includes lesions that are generally infiltrating and low in mitotic activity but recur more frequently than do grade I malignant tumors after local therapy. Some tumor types tend to progress to higher grades of malignancy.
- **WHO grade III** includes lesions with histologic evidence of malignancy, including nuclear atypia and increased mitotic activity. These lesions have anaplastic histology and infiltrative capacity. They are usually treated with aggressive adjuvant therapy.
- **WHO grade IV** includes lesions that are mitotically active, necrosis prone, and generally associated with a rapid preoperative and postoperative progression and fatal outcomes. The lesions are usually treated with aggressive adjuvant therapy.

Table 1 lists the tumor types and grades.[4] Tumors limited to the peripheral nervous system are not included. Histopathology, grading methods, incidence, and what is known about etiology specific to each tumor type have been described in detail elsewhere.[4,5]

Table 1. WHO Grades of CNS Tumors

<table>
<thead>
<tr>
<th>Tumor Type</th>
<th>Grade I</th>
<th>Grade II</th>
<th>Grade III</th>
<th>Grade IV</th>
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<tr>
<td>Astrocytic tumors</td>
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<td>Subependymal giant cell astrocytoma</td>
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<td>Pilocytic astrocytoma</td>
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<td>Pilomyxoid astrocytoma</td>
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<td>Diffuse astrocytoma</td>
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<td>Pleomorphic xanthoastrocytoma</td>
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<td>Anaplastic astrocytoma</td>
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<td>Glioblastoma</td>
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<tr>
<td>Giant cell glioblastoma</td>
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<td>Gliosarcoma</td>
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<tr>
<td>Oligodendrogial tumors</td>
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<th>Tumor Type</th>
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<tr>
<td>Oligodendroglioma</td>
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<tr>
<td>Anaplastic oligodendroglioma</td>
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<tr>
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<td>Oligoastrocytoma</td>
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<td>Anaplastic oligoastrocytoma</td>
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<td><strong>Ependymal tumors</strong></td>
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<td>Subependymoma</td>
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<tr>
<td>Myxopapillary ependymoma</td>
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<td>Ependymoma</td>
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<td>Anaplastic ependymoma</td>
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<tr>
<td><strong>Choroid plexus tumors</strong></td>
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<td>Choroid plexus papilloma</td>
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<tr>
<td>Atypical choroid plexus papilloma</td>
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<tr>
<td>Choroid plexus carcinoma</td>
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<td>X</td>
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<tr>
<td><strong>Other neuroepithelial tumors</strong></td>
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<tr>
<td>Angiocentric glioma</td>
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<td>Chordoid glioma of the third ventricle</td>
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<tr>
<td><strong>Neuronal and mixed neuronal-glial tumors</strong></td>
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<tr>
<td>Gangliocytoma</td>
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<tr>
<td>Ganglioglioma</td>
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<tr>
<td>Anaplastic ganglioma</td>
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<tr>
<td>Desmoplastic infantile astrocytoma and ganglioglioma</td>
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<tr>
<td>Dysembryoplastic neuroepithelial tumor</td>
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<tr>
<td>Central neurocytoma</td>
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<td>Extraventricular neurocytoma</td>
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<tr>
<td>Cerebellar liponeurocytoma</td>
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<td>Paragangioma of the spinal cord</td>
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<td>Papillary glioneuronal tumor</td>
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<td>Rosette-forming glioneural tumor of the fourth ventricle</td>
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<td><strong>Pineal tumors</strong></td>
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<tr>
<td>Pineocytoma</td>
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<td>Pineal parenchymal tumor of intermediate differentiation</td>
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<td>Pineoblastoma</td>
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<td>Papillary tumor of the pineal region</td>
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<td><strong>Embryonal tumors</strong></td>
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<tr>
<td>Medulloblastoma</td>
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<tr>
<td>CNS primitive neuroectodermal tumor</td>
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<tr>
<td>Atypical teratoid/rhabdoid tumor</td>
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<td><strong>Tumors of the cranial and paraspinal nerves</strong></td>
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<tr>
<td>Schwannoma</td>
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<td>Neurofibroma</td>
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<td>Perineurioma</td>
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<td>Malignant peripheral nerve sheath tumor</td>
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### Meningeal tumors

| Meningioma | X |
| Atypical meningioma | X |
| Anaplastic/malignant meningioma | X |
| Hemangiopericytoma | X |
| Anaplastic hemangiopericytoma | X |
| Hemangioblastoma | X |

### Tumors of the sellar region

| Craniopharyngioma | X |
| Granular cell tumor of the neurohypophysis | X |
| Pituicytoma | X |
| Spindle cell oncocytoma of the adenohypophysis | X |

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Genomic Alterations

Recently discovered alterations in the \(BRAF\) and \(isoctirate dehydrogenase (IDH) 1\) and \(IDH2\) genes, and genomic 1p/19q codeletion, appear to be hallmark aberrations in particular glioma subtypes. Assessment for the presence of these mutations aids diagnosis and prognosis and, with regard to 1p/19q codeletion, predicts for response to chemotherapy.

In pilocytic astrocytomas (WHO grade I), tandem duplication at 7q34 leading to a fusion between \(KIAA1549\) and \(BRAF\) is found in approximately 70% of pilocytic astrocytomas.[6-8] An activating point mutation in \(BRAF\) (V600E) is found in an additional 5% to 9% of these tumors and in general, RAF alterations occur in approximately 80% of pilocytic astrocytomas.

\(BRAF\) V600E mutations are observed (in about 60%) of other benign glioma variants, including pleomorphic xanthoastrocytoma and ganglioglioma, while \(BRAF\) tandem duplications are not found in these variant glioma tumors.[9-11]

The majority of WHO grade II and III diffuse gliomas (astrocytomas, oligodendrogliomas, and oligoastrocytomas) and 5% to 10% of glioblastomas (WHO grade IV) harbor point mutations in the \(R132\) position of \(IDH1\) or, rarely, the analogous codon in \(IDH2\) (R172).[12-16] The presence of an \(IDH1\) or \(IDH2\) mutation is a strong prognostic factor. Patients with these mutant tumors have significantly longer survival independent of WHO grade or histologic subtype.

Deletion of chromosomes 1p and 19q occurs through a translocation event [17] and is common in oligodendrogliomas. 1p/19q codeletion is a powerful prognostic factor and may predict for response to chemotherapy. (Refer to the Anaplastic oligodendrogliomas treatment section of this summary for more information.)

These genetic alterations have potential diagnostic utility. Presence of the \(IDH1\) and \(IDH2\) mutations may distinguish diffuse gliomas from other glioma variants, which often have \(BRAF\) genetic alterations, and non-neoplastic reactive astrocytosis.[18] Most (90%) \(IDH\) mutations in gliomas result in an \(R132H\) substitution, which can be detected with a highly sensitive and specific monoclonal antibody. A rapid immunohistochemical analysis using the mutant-specific \(IDH1\) antibody can aid diagnostic analysis.[19]
Other CNS tumors are associated with characteristic patterns of altered oncogenes, altered tumor suppressor genes, and chromosomal abnormalities. Familial tumor syndromes with defined chromosomal abnormalities are associated with gliomas.

References

Treatment Option Overview for Adult Primary CNS Tumors

Primary CNS Tumors

This section discusses general treatment modalities for primary central nervous system (CNS) tumors. (Refer to the Treatment of Primary Central Nervous System Tumors by Tumor Type section of this summary for a description of specific treatment options for each tumor type.)

Radiation therapy and chemotherapy options vary according to histology and anatomic site of the CNS tumor. For glioblastoma, combined modality therapy with resection, radiation, and chemotherapy is standard. Anaplastic astrocytomas, anaplastic oligodendrogliomas, and anaplastic oligoastrocytomas represent only a small proportion of CNS gliomas; therefore, phase III randomized trials restricted to these tumor types are not generally practical. The natural histories of these tumors are variable, depending on histological and molecular factors; therefore, treatment guidelines are evolving. Therapy involving surgically implanted carmustine-impregnated polymer wafers combined with postoperative external-beam radiation therapy (EBRT) may play a role in the treatment of high-grade (grades III and IV) gliomas in some patients.[1]

Standard treatment options for primary CNS tumors include the following:

2. Radiation therapy.
3. Chemotherapy.
4. Active surveillance.
5. Supportive therapy.

Surgery

For most types of CNS tumors in most locations, complete or near-complete surgical removal is generally attempted, within the constraints of preserving neurologic function and the patient's underlying health. This practice is based on observational evidence that survival is better in patients who undergo tumor resection than in those who have closed biopsy alone.[2,3] The benefit of resection has not been tested in randomized trials. Selection bias can enter into observational studies despite attempts to adjust for patient differences that guide the decision to resect the tumor; therefore, the actual difference in outcome between radical surgery and biopsy alone may not be as large as noted in the retrospective studies.[3]

An exception to the use of resection is the case of deep-seated tumors such as pontine gliomas, which are diagnosed on clinical evidence and treated without initial surgery approximately 50% of the time. In most cases, however, diagnosis by biopsy is preferred. Stereotactic biopsy can be used for lesions that are difficult to reach and resect.

The primary goals of surgical resection include the following:[4]

- To establish a histologic diagnosis.
- To reduce intracranial pressure by removing as much tumor as is safely possible to preserve neurological function.

Total elimination of primary malignant intraparenchymal tumors by surgery alone is rarely achievable. Therefore, intraoperative techniques have been developed to reach a balance between removing as much tumor as is practical and preserving functional status. For example, craniotomies with stereotactic resections of primary gliomas can be performed in cooperative patients while they are awake, with real-time assessment of neurologic function.[5]

Examples of intraoperative neurologic assessment include the following:

- Resection proceeds until either the magnetic resonance imaging (MRI) signal abnormality being used to monitor the extent of surgery is completely removed or subtle neurologic dysfunction appears (e.g., a slight decrease in rapid alternating motor movement or anomia).
- When the tumor is located in or near language centers in the cortex, intraoperative language mapping can be
performed by electrode discharge-induced speech arrest while the patient is asked to count or read.[6]

As is the case with several other specialized operations [7,8] in which postoperative mortality has been associated with the number of procedures performed, postoperative mortality after surgery for primary brain tumors may be associated with hospital and/or surgeon volume.[9] Using the Nationwide Inpatient Sample hospital discharge database for the years 1988 to 2000, which represented 20% of inpatient admissions to nonfederal U.S. hospitals, investigators observed the following:[9]

- Large-volume hospitals had lower in-hospital mortality rates after craniotomies for primary brain tumors (odds ratio [OR], 0.75 for a tenfold higher caseload; 95% confidence interval [CI], 0.62–0.90) and after needle biopsies (OR, 0.54; 95% CI, 0.35–0.83).

- Although there was no specific sharp threshold in all-cause mortality outcomes between low-volume hospitals and high-volume hospitals, craniotomy-associated in-hospital mortality was 4.5% for hospitals with 5 or fewer procedures per year and 1.5% for hospitals with at least 42 procedures per year.

- In-hospital mortality rates decreased over the study years (perhaps because the proportion of elective nonemergent operations increased from 45% to 57%), but the decrease was more rapid in high-volume hospitals than in low-volume hospitals.

- High-volume surgeons had lower in-hospital patient mortality rates after craniotomy (OR, 0.60; 95% CI, 0.45–0.79).

As with any study of volume-outcome associations, these results may not be causal because of residual confounding factors such as referral patterns, private insurance, and patient selection, despite multivariable adjustment.

**Radiation therapy**

**High-grade tumors**

Radiation therapy has a major role in the treatment of patients with high-grade gliomas.

Evidence (postoperative radiation therapy [PORT]):

1. A systematic review and meta-analysis of five randomized trials (plus one trial with allocation by birth date) comparing PORT with no radiation therapy showed a statistically significant survival advantage with radiation (risk ratio, 0.81; 95% CI, 0.74–0.88).[10][Level of evidence: 1iiA]

2. A randomized trial comparing 60 Gy (in 30 fractions over 6 weeks) with 45 Gy (in 25 fractions over 4 weeks) showed superior survival in the first group (12 months vs. 9 months median survival; hazard ratio [HR], 0.81; 95% CI, 0.66–0.99). The accepted standard dose of EBRT for malignant gliomas is 60 Gy.[11][Level of evidence: 1iiA]

EBRT using either 3-dimensional conformal radiation therapy (3D-CRT) or intensity-modulated radiation therapy (IMRT) is considered an acceptable technique in radiation therapy delivery. Typically used are 2- to 3-cm margins on the MRI-based volumes (T1-weighted and fluid-attenuated inversion recovery [FLAIR]) to create the planning target volume.

Dose escalation using radiosurgery has not improved outcomes. A randomized trial tested radiosurgery as a boost added to standard EBRT, but the trial found no improvement in survival, quality of life, or patterns of relapse compared with EBRT without the boost.[12,13]

Brachytherapy has been used to deliver high doses of radiation locally to the tumor while sparing normal brain tissue. However, this approach is technically demanding and has fallen out of favor with the advent of 3D-CRT and IMRT.

**Low-grade tumors**

The role of immediate PORT for low-grade gliomas (i.e., low-grade astrocytomas, oligodendrogliomas, and mixed oligoastrocytomas) is not as clear as in the case of high-grade tumors.

Evidence (PORT versus observation):
The European Organisation for Research and Treatment of Cancer (EORTC) randomly assigned 311 patients with low-grade gliomas to undergo either radiation or observation in the EORTC-22845 (MRC BR04) trial. On review of central pathology, about 25% of patients in the trial were reported to have high-grade tumors. Most of the control patients received radiation therapy at the time of progression. After a median follow-up of 93 months, median progression-free survival was 5.3 years in the radiation arm versus 3.4 years in the control arm (HR, 0.59; 95% CI, 0.45–0.77).

There was no difference in the overall survival (OS) rate (median survival = 7.4 years in the radiation arm vs. 7.2 years in the control arm; HR, 0.97; 95% CI, 0.71–1.34; P = .87). This was caused by a longer survival after progression in the control arm (3.4 years) than in the radiation arm (1.0 year) (P < .0001).

The investigators did not collect reliable quality-of-life measurements, so it is not clear whether the delay in initial relapse in the radiation therapy arm translated into improved function or quality of life.

Disease progression, subsequent neoplasms, or recurrences

There are no randomized trials to delineate the role of repeat radiation after disease progression or the development of radiation-induced cancers. The literature is limited to small retrospective case series, which makes interpretation difficult. The decision to repeat radiation must be made carefully because of the risk of neurocognitive deficits and radiation-induced necrosis. One advantage of radiosurgery is the ability to deliver therapeutic doses to recurrent tumors that may require the re-irradiation of previously irradiated brain tissue beyond tolerable dose limits.

Chemotherapy

Systemic chemotherapy

For many years, the nitrosourea carmustine (bis-chloroethylnitrosourea) BCNU was the standard chemotherapy agent added to surgery and radiation therapy for malignant gliomas, based on the Radiation Therapy Oncology Group's randomized trial (RTOG-8302). A modest impact on survival with the use of nitrosourea-containing chemotherapy regimens for malignant gliomas was confirmed in a patient-level meta-analysis of 12 randomized trials (combined HR, 0.85; 95% CI, 0.78–0.91).

A large multicenter trial (NCT00006353) of glioblastoma patients conducted by the EORTC-National Cancer Institute of Canada reported a survival advantage with the use of temozolomide in addition to radiation therapy. On the basis of these results, the oral agent temozolomide has replaced BCNU as the standard systemic chemotherapy for malignant gliomas. (Refer to the Glioblastomas treatment section of this summary for more information.)

Long-term results of randomized trials in high-risk, low-grade (World Health Organization [WHO] grade II) gliomas and anaplastic (WHO grade III) oligodendroglial tumors have demonstrated that the addition of procarbazine, lomustine (CCNU), and vincristine (PCV) chemotherapy to radiation therapy after surgery extends survival. Radiation and PCV chemotherapy should be considered for patients deemed appropriate for therapy. (Refer to the Treatment of Primary Central Nervous System Tumors by Tumor Type section of this summary for more information.)

Localized chemotherapy (carmustine wafer)

The concept of delivering high doses of chemotherapy while avoiding systemic toxicity is attractive because malignant glioma–related deaths are nearly always the result of an inability to control intracranial disease rather than the result of distant metastases. A biodegradable carmustine wafer has been developed for that purpose. The wafers contain 3.85% carmustine, and up to eight wafers are implanted into the tumor bed lining at the time of open resection, with an intended total dose of about 7.7 mg per wafer (61.6 mg maximum per patient) over a period of 2 to 3 weeks.

Two randomized, placebo-controlled trials of this focal drug-delivery method have shown an OS advantage associated with the carmustine wafers versus radiation therapy alone. In both trials, the upper age limit for patients was 65 years.

Evidence (carmustine wafer):
1. A small trial was closed because of a lack of continued availability of the carmustine wafers after 32 patients with high-grade gliomas had been entered.[24]

   - Although OS was better in the carmustine-wafer group (median 58.1 vs. 39.9 weeks; \(P = .012\)), there was an imbalance in the study arms (only 11 of 16 patients in the carmustine-wafer group vs. 16 of the 16 patients in the placebo-wafer group had grade IV glioblastoma tumors).

2. A multicenter study of 240 patients with primary malignant gliomas, 207 of whom had glioblastoma, was more informative.[25,26] At initial surgery, patients received either carmustine wafers or placebo wafers, followed by radiation therapy (55–60 Gy). Systemic therapy was not allowed until recurrence, except in the case of anaplastic oligodendrogliomas (n = 9). Unlike the initial trial, patient characteristics were well balanced between the study arms.

   - Median survival in the two groups was 13.8 months in patients treated with carmustine wafers versus 11.6 months in placebo-treated patients (HR, 0.73; 95% CI, 0.56–0.96; \(P = .017\)).

3. A systematic review combining both studies [24-26] estimated an HR for overall mortality of 0.65; 95% CI, 0.48–0.86; \(P = .003\).[27][Level of evidence: 1iA]

**Active surveillance**

Active surveillance is appropriate in some circumstances. With the increasing use of sensitive neuroimaging tools, detection of asymptomatic low-grade meningiomas has increased; most appear to show minimal growth and can often be safely observed, with therapy deferred until the detection of tumor growth or the development of symptoms.[28,29]

**Supportive therapy**

Dexamethasone, mannitol, and furosemide are used to treat the peritumoral edema associated with brain tumors. The use of anticonvulsants is mandatory for patients with seizures.[4]

**References**


Treatment of Primary Central Nervous System Tumors by Tumor Type

Table 2. Treatment of Primary Central Nervous System Tumors by Tumor Type
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### Astrocytic Tumors Treatment

#### Brain stem gliomas treatment

Patients with brain stem gliomas have relatively poor prognoses that correlate with histology (when biopsies are performed), location, and extent of tumor. The overall median survival time of patients in studies has been 44 to 74 weeks.

Standard treatment options for brain stem gliomas include the following:

1. Radiation therapy.

#### Pineal astrocytic tumors treatment

Depending on the degree of anaplasia, patients with pineal astrocytomas have variable prognoses. Patients with higher-grade tumors have worse prognoses.

Standard treatment options for pineal astrocytic tumors include the following:

1. Surgery plus radiation therapy for pineal astrocytoma.
2. Surgery plus radiation therapy and chemotherapy for higher-grade tumors.

#### Pilocytic astrocytomas treatment

This astrocytic tumor is classified as a World Health Organization (WHO) grade I tumor and is often curable.

Standard treatment options for pilocytic astrocytomas include the following:

1. Surgery alone if the tumor is totally resectable.
2. Surgery followed by radiation therapy to known or suspected residual tumor.

#### Diffuse astrocytomas treatment

This WHO grade II astrocytic tumor is less often curable than is a pilocytic astrocytoma.

Standard treatment options for diffuse astrocytomas (WHO grade II) include the following:

1. Surgery with or without radiation therapy.
2. Surgery followed by radiation therapy and chemotherapy.

Controversy exists about the timing of radiation therapy after surgery. (Refer to Low-grade tumors in the Radiation therapy section of the Treatment Option Overview for Adult Central Nervous System Tumors Treatment section in this summary for more information.)

- Radiation therapy improved progression-free survival (PFS) in patients who received early radiation therapy in the European Organisation for Research and Treatment of Cancer (EORTC) (EORTC-22845) trial. (Refer to Oligodendrogliomas treatment in the Oligodendroglial Tumors Treatment section of the Treatment of Primary Central Nervous System Tumors by Tumor Type section of this summary for more information.)[1][Level of evidence: 1iiA]

- In the same trial, there was no difference in overall survival (OS) between patients who had radiation therapy after surgery and those who were treated with radiation therapy at the time of progression.[1][Level of evidence: 1iiA]

Some physicians use surgery alone if a patient has clinical factors that are considered low risk, such as age less than 40 years and the lack of contrast enhancement on a computed tomographic scan.[2]
Evidence (surgery followed by radiation therapy and chemotherapy):

1. For patients with low-grade (WHO grade II) tumors, which are considered high risk, radiation therapy followed by six cycles of vincristine (PCV) chemotherapy is a recommended option based on the long-term follow-up results of the Radiation Therapy Oncology Group (RTOG)'s 1986-initiated randomized trial (RTOG 9802 [NCT00003375]).[3][Level of evidence: 1iiA] In this trial, high-risk, low-grade glioma patients, defined as patients aged 18 to 39 years with biopsy or subtotal resection, or patients aged 40 years or older, were randomly assigned to either 54 Gy of radiation therapy or radiation therapy followed by six cycles of PCV chemotherapy.

   a. The addition of PCV to radiation therapy increased median PFS from 4.0 years to 10.4 years (P = .002; hazard ratio [HR], 0.50) and median OS from 7.8 years to 13.3 years (P = .03; HR, 0.59).

   b. Notably, the RTOG 9802 study enrolled patients with a variety of tumors, including astrocytomas, oligodendrogliomas, and mixed oligoastrocytomas.

      - In a risk-adjusted multivariate analysis, patients treated with PCV and patients with an oligodendroglial histology had better survival outcomes. A subset analysis of histologic type suggested that the addition of PCV mainly benefited patients with oligodendroglial tumors, although this data is yet to be validated.[4]

      - Median OS for PCV versus the control arm was not reached versus 10.8 years for oligodendrogliomas (P = .008), 11.4 years versus 5.9 years for oligoastrocytomas (P = .05), and 7.7 years versus 4.4 years for astrocytomas (P = .31).

The discovery of the isocitrate dehydrogenase (IDH) 1 and IDH2 mutations in diffuse gliomas has greatly helped to identify patients who are considered high risk. A number of large, retrospective studies has demonstrated that the IDH1 and IDH2 mutation is a powerful independent prognostic factor for improved survival.[5-9] The majority of WHO grade II and III gliomas harbor the IDH1 and IDH2 mutation,[6,10,11] and, therefore, the presence of the IDH1 and IDH2 mutation should be included in the assessment of high risk. Molecular correlative data from the RTOG 98-02 trial, which would be informative about which patients benefited the most from the addition of PCV, have not yet been reported.

Anaplastic astrocytomas treatment

Patients with anaplastic astrocytomas (WHO grade III) have a low cure rate with standard local treatment.

Standard treatment options for anaplastic astrocytomas include the following:

1. Surgery plus radiation therapy with or without chemotherapy.
2. Surgery plus chemotherapy.

A subset of anaplastic astrocytomas is aggressive; these tumors are frequently managed in the same way as glioblastomas, with surgery and radiation, and often with chemotherapy. However, the optimal treatment for these tumors is not established. Two phase III randomized trials restricted to patients with anaplastic gliomas (NCT00626990 and NCT00887146) are currently enrolling patients, but efficacy data are not available. It is not known whether the improved survival of patients with chemotherapy-treated glioblastoma can be extrapolated to patients with anaplastic astrocytomas.

The IDH1 and IDH2 mutation is present in 50% to 70% of anaplastic astrocytomas and is independently associated with significantly improved survival.[6,9] Assessment of the IDH1 and IDH2 mutation status may guide decisions about treatment options.

Evidence (surgery plus radiation therapy or chemotherapy):

1. Postoperative radiation alone has been compared with postoperative chemotherapy alone in patients with anaplastic gliomas (i.e., 144 astrocytomas, 91 oligoastrocytomas, and 39 oligodendrogliomas), with crossover to the other modality at the time of tumor progression. Of the 139 patients randomly assigned to undergo radiation therapy, 135 were randomly assigned to receive chemotherapy, with a 32-week course of either PCV or single-agent temozolomide (2:1:1 randomization).[12][Levels of evidence: 1iiA and 1iiD]
The order of the modalities did not affect time-to-treatment failure (TTF) or OS.

Neither TTF nor OS differed across the treatment arms.

Patients with anaplastic astrocytomas are appropriate candidates for clinical trials designed to improve local control by adding newer forms of treatment to standard treatment. Information about ongoing clinical trials is available from the NCI website.

Glioblastomas treatment

For patients with glioblastoma (WHO grade IV), the cure rate is very low with standard local treatment. Methylation of the promoter of the O6-methylguanine-DNA methyltransferase (\textit{MGMT}) DNA repair enzyme gene is an independent prognostic factor for improved survival in newly diagnosed glioblastoma.\cite{13,14} \textit{MGMT} promoter methylation and concomitant inactivation of the DNA repair enzyme activities may also predict for response to temozolomide chemotherapy.\cite{13} However, the clinical data that \textit{MGMT} promoter methylation is a predictive marker is less certain. (Refer to Glioblastomas treatment in the Astrocytic Tumors Treatment section of the Treatment of Primary Central Nervous System Tumors by Tumor Type section of this summary for more information about the RTOG-0525 [NCT00304031] trial.)

Standard treatment options for patients with newly diagnosed glioblastoma include the following:

1. Surgery plus radiation therapy and chemotherapy.
2. Surgery plus radiation therapy.
3. Carmustine-impregnated polymer implanted during initial surgery.
4. Radiation therapy and concurrent chemotherapy.

The standard treatment for patients with newly diagnosed glioblastoma is surgery followed by concurrent radiation therapy and daily temozolomide, and then followed by six cycles of temozolomide. The addition of bevacizumab to radiation therapy and temozolomide did not improve OS.

Evidence (Surgery plus radiation therapy and chemotherapy):

1. Standard therapy is based on a large, multicenter, randomized trial (NCT00006353) conducted by the EORTC and National Cancer Institute of Canada (NCIC). This trial reported a survival benefit with concurrent radiation therapy and temozolomide, compared with radiation therapy alone.\cite{15,16} [Level of evidence: 1iiA] In this study, 573 patients with glioblastoma were randomly assigned to receive standard radiation therapy to the tumor volume with a 2- to 3-cm margin (60 Gy, 2 Gy per fraction, over 6 weeks) alone or with temozolomide (75 mg/m\textsuperscript{2} orally per day during radiation therapy for up to 49 days, followed by a 4-week break and then up to six cycles of five daily doses every 28 days at a dose of 150 mg/m\textsuperscript{2}, increasing to 200 mg/m\textsuperscript{2} after the first cycle).

   a. OS was statistically significantly better in the combined radiation therapy–temozolomide group (HR\textsubscript{death} = 0.6; 95% confidence interval [CI], 0.5–0.7; OS at 3 years was 16.0% for the radiation therapy–temozolomide group vs. 4.4% radiation therapy–alone group).

   b. A companion molecular correlation subset study to the EORTC-NCIC trial provided strong evidence that epigenetic silencing of the \textit{MGMT} DNA-repair gene by promoter DNA methylation was associated with increased OS in patients with newly diagnosed glioblastoma.\cite{13}

      - \textit{MGMT} promoter methylation was an independent favorable prognostic factor ($P < .001$ by the log-rank test; HR, 0.45; 95% CI, 0.32–0.61).

      - The median OS for patients with \textit{MGMT} methylation was 18.2 months (95% CI, 15.5–22.0), compared with 12.2 months (95% CI, 11.4–13.5) for patients without \textit{MGMT} methylation.

2. To test whether protracted (dose-dense) temozolomide enhances treatment response in patients with newly diagnosed glioblastoma, a multicenter, randomized, phase III trial conducted by the RTOG, EORTC, and the North Central Cancer Therapy Group, RTOG 0525 (NCT00304031), compared standard adjuvant temozolomide
treatment (days 1–5 of a 28-day cycle) with a dose-dense schedule (days 1–21 of a 28-day cycle). All patients were treated with surgery followed by radiation therapy and concurrent daily temozolomide. Patients were then randomly assigned to receive either standard adjuvant temozolomide or dose-dense temozolomide. [14][Level of evidence: 1iiA]

- Among 833 randomly assigned patients, no statistically significant difference between standard and dose-dense temozolomide was observed for median OS (16.6 months for standard temozolomide vs. 14.9 months for dose-dense temozolomide; HR, 1.03; \( P = .63 \)) or for median PFS (5.5 vs. 6.7 months; HR, 0.87; \( P = .06 \)).

- Protracted temozolomide, which depletes intracellular MGMT, was predicted to have greater efficacy in tumors with MGMT-promoter methylation. To test this retrospectively, MGMT status was determined in 86% of randomly assigned patients. No difference in efficacy was observed in either the MGMT-methylated or MGMT-unmethylated subsets. There was no survival advantage for the use of dose-dense temozolomide versus standard-dose temozolomide in newly diagnosed glioblastoma patients, regardless of MGMT status. However, this study confirmed the strong prognostic effect of MGMT methylation because the median OS was 21.2 months (95% CI, 17.9–24.8) for patients with methylation versus 14 months (HR, 1.74; 95% CI, 12.9–14.7; \( P < .001 \)) for patients without methylation.

- The efficacy of dose-dense temozolomide for patients who have recurrent glioblastoma, however, is yet to be determined.

Evidence (surgery and chemoradiation with or without bevacizumab):

In 2013, final data from two multicenter, phase III, randomized, double-blind, placebo-controlled trials of bevacizumab in patients who had newly diagnosed glioblastoma were reported: RTOG 0825 (NCT00884741) and the Roche-sponsored AVAglio (NCT00943826).[17,18][Level of evidence: 1iA] Bevacizumab did not improve OS in either trial.

There was significant crossover in both trials. Approximately 40% of RTOG 0825 patients and approximately 30% of AVAglio patients received bevacizumab at the first sign of disease progression.

1. RTOG 0825 (NCT00884741): Patients were randomly assigned to receive standard therapy (chemoradiation with temozolomide) or standard therapy plus bevacizumab. OS and PFS were coprimary endpoints.[17][Level of evidence: 1iA]

   - Bevacizumab did not improve OS (median OS was 16–17 months for each arm); however, it increased median PFS (10.7 months in the bevacizumab arm vs. 7.3 months in the placebo arm; HR, 0.79; \( P = .007 \)).

   - The PFS result in the RTOG 0825 trial did not meet the prespecified significance level (\( P = .004 \)).

2. AVAglio (NCT00943826): Patients were randomly assigned to receive standard therapy (chemoradiation with temozolomide) or standard therapy plus bevacizumab. OS and PFS were coprimary endpoints.[18][Level of evidence: 1iA]

   - Bevacizumab did not improve OS (median OS was 16–17 months for each arm); however, it increased median PFS (10.6 months in the bevacizumab arm vs. 6.2 months in the placebo arm; HR, 0.64; \( P < .0001 \)).

   - The PFS result was statistically significant and associated with clinical benefit because bevacizumab-treated patients remained functionally independent longer (9.0 months in the bevacizumab arm vs. 6.0 months in the standard therapy arm) and had a longer time until their Karnofsky Performance status deteriorated (HR, 0.65; \( P < .0001 \)).

   - Bevacizumab-treated patients also had delayed initiation of corticosteroids (12.3 months vs. 3.7 months; HR, 0.71; \( P = .002 \)), and more patients were able to discontinue corticosteroids if they were already taking them (66% in the bevacizumab arm vs. 47% in the standard therapy arm).

The two trials had contradictory results in health-related quality of life (HRQoL) and neurocognitive outcomes studies. In the mandatory HRQoL studies in the AVAglio trial, bevacizumab-treated patients experienced improved HRQoL,
but bevacizumab-treated patients in the elective RTOG 0825 studies showed more decline in patient-reported HRQoL and neurocognitive function. The reasons for these discrepancies are unclear.

On the basis of these results, there is no definite evidence that the addition of bevacizumab to standard therapy is beneficial for all newly diagnosed glioblastoma patients. Certain subgroups may benefit from the addition of bevacizumab, but this is not yet known.

Glioblastoma patients are appropriate candidates for clinical trials designed to improve local control by adding newer forms of treatment to standard treatment. Information about ongoing clinical trials is available from the NCI website.

**Oligodendroglial Tumors Treatment**

**Oligodendrogliomas treatment**

Patients who have oligodendrogliomas (WHO grade II) generally have better prognoses than do patients who have diffuse astrocytomas. In particular, patients who have oligodendrogliomas with 1p/19q codeletion have a much longer survival.[3] Most of the oligodendrogliomas eventually progress.

Standard treatment options for oligodendrogliomas include the following:

1. Surgery with or without radiation therapy.
2. Surgery with radiation therapy and chemotherapy.

Controversy exists concerning the timing of radiation therapy after surgery. A study (EORTC-22845) of 300 patients with low-grade gliomas who had surgery and were randomly assigned to either radiation therapy or watchful waiting, did not show a difference in OS between the two groups.[1][Level of evidence: 1iiA] (Refer to Low-grade tumors in the Radiation Therapy section of the Treatment Overview for Adult Central Nervous System Tumors Treatment section of this summary for more information.)

For low-grade (WHO grade II) tumors that are considered high risk, radiation therapy followed by six cycles of PCV chemotherapy is a recommended option based on the long-term follow-up results of RTOG-9802, a randomized trial for high-risk, low-grade gliomas.[3][Level of evidence: 1iiA] (Refer to Diffuse astrocytomas treatment in the Astrocytic Tumors Treatment section of the Treatment of Primary Central Nervous System Tumors by Tumor Type section of this summary for more information.)

Notably, the RTOG-9802 study enrolled patients with a mixed variety of tumors, including astrocytomas, oligodendrogliomas, and mixed oligoastrocytomas; in a retrospective subset analysis, only the oligodendrogial tumors appeared to benefit from the addition of PCV.[4] (Refer to Diffuse astrocytomas treatment in the Astrocytic Tumors Treatment section of the Treatment of Primary Central Nervous System Tumors by Tumor Type section of this summary for more information.)

The discovery of the IDH1 and IDH2 mutations, which are independent prognostic factors for significantly improved survival in diffuse gliomas, has greatly helped to identify patients who are considered high risk. (Refer to Diffuse astrocytomas treatment in the Astrocytic Tumors Treatment section of the Treatment of Primary Central Nervous System Tumors by Tumor Type section of this summary for more information.) In addition, a high proportion of WHO grade II oligodendrogliomas have 1p/19q codeletion, which is a powerful prognostic factor for improved survival. [19-21] Therefore, the presence of the IDH1 and IDH2 mutation and 1p/19q codeletion should be included in the assessment of high-risk patients. Molecular correlative data from the RTOG-9802 trial, which would be informative about which patients benefited most from the addition of PCV, have not yet been reported.

**Anaplastic oligodendrogliomas treatment**

Patients with anaplastic oligodendrogliomas (WHO grade III) have a low cure rate with standard local treatment, but their prognoses are generally better than are the prognoses of patients with anaplastic astrocytomas. Prognoses are particularly better for patients with 1p/19q codeletion, which occurs in a majority of these tumors. Two phase III randomized trials restricted to patients with anaplastic gliomas (NCT00626990 and NCT00887146) are currently enrolling patients; however, efficacy data are not yet available. (Refer to Anaplastic astrocytomas treatment in the Astrocytic Tumors Treatment section of the Treatment of Primary Central Nervous System Tumors by Tumor Type section of this summary for more information.)
section of this summary for more information.) These patients are appropriate candidates for clinical trials designed to improve local control by adding newer forms of treatment.

Information about ongoing clinical trials is available from the NCI website.

Standard treatment options for anaplastic oligodendrogliomas include the following:

1. Surgery plus radiation with or without chemotherapy.[22]

Evidence (surgery followed by radiation therapy with or without chemotherapy):

1. Mature results from the EORTC Brain Tumor Group Study 26951 (NCT00002840), a phase III, randomized study with 11.7 years of follow-up, demonstrated increased OS and PFS in patients with anaplastic oligodendrogial tumors with six cycles of adjuvant PCV chemotherapy after radiation therapy, compared with radiation therapy alone.[23][Level of evidence: 1iiA]
   - OS was significantly longer in the radiation therapy and PCV arm (42.3 months vs. 30.6 months; HR, 0.75; 95% CI, 0.60–0.95).
   - Patients with 1p/19q-codeleted tumors derived more benefit from adjuvant PCV chemotherapy than did those with non–1p/19q-deleted tumors.[23]

2. In contrast, the RTOG trial (RTOG-9402 [NCT00002569]) demonstrated no differences in median survival by treatment arm between an 8-week, intensive PCV chemotherapy regimen followed by immediate involved-field-plus-radiation therapy and radiation therapy alone.[24]
   - In an unplanned subgroup analysis, patients with 1p/19q-codeleted anaplastic oligodendrogliomas and mixed anaplastic astrocytomas demonstrated a median survival of 14.7 years versus 7.3 years (HR, 0.59; 95% CI, 0.37–0.95; \( P = .03 \)).
   - For patients with non-codeleted tumors, there was no difference in median survival by treatment arm (2.6 vs. 2.7 years; HR, 0.85; 95% CI, 0.58–1.23; \( P = .39 \)).[24][Level of evidence: 1iiA]

3. Postoperative radiation therapy alone has been compared with postoperative chemotherapy alone in patients with anaplastic gliomas (including 144 astrocytomas, 91 oligoastrocytomas, and 39 oligodendrogliomas) with crossover to the other modality at the time of tumor progression. Of the 139 patients randomly assigned to undergo radiation therapy, 135 were randomly assigned to receive chemotherapy, with a 32-week course of either PCV or single-agent temozolomide (2:1:1 randomization).[12][Levels of evidence: 1iiA and 1iiD]
   - TTF or OS did not differ across the treatment arms and were not affected by the order of the modalities.

On the basis of these data, CODEL (NCT00887146), a study that randomly assigned patients to receive radiation therapy alone (control arm), radiation therapy with temozolomide, and temozolomide alone (exploratory arm), was halted because radiation therapy alone was no longer considered adequate treatment in patients with anaplastic oligodendroglioma with 1p/19q-codeletions.[25] Temozolomide and PCV chemotherapy in anaplastic oligodendroglioma have not been compared, although in the setting of grade 3 anaplastic gliomas, no survival difference was seen between PCV chemotherapy and temozolomide.[12,26]

The combination of radiation and chemotherapy is not known to be superior in outcome to sequential modality therapy.

A high proportion of anaplastic oligodendrogliomas have the \textit{IDH1} and \textit{IDH2} mutation and 1p/19q codeletion, both powerful prognostic factors for improved survival. (Refer to Diffuse astrocytomas treatment in the Astrocytic Tumors Treatment section of the Treatment of Primary Central Nervous System Tumors by Tumor Type of this summary for more information.)[23,24] In addition, PCV chemotherapy has been shown to be predictive in a retrospective analysis of the phase III trials described earlier. Therefore, assessment of these molecular markers may aid management decisions for anaplastic oligodendrogliomas.

**Mixed Gliomas Treatment**

Patients with mixed glial tumors, which include oligoastrocytoma (WHO grade II) and anaplastic oligoastrocytoma
(WHO grade III), have highly variable prognoses based upon their status of the IDH1 and IDH2 genes and 1p/19q chromosomes.[27-29] Therefore, the optimal treatment for these tumors as a group is uncertain. Often, they are treated similarly to astrocytic tumors because a subset of tumors may have outcomes similar to WHO grade III astrocytic or WHO grade IV glioblastoma tumors. Testing for these known, strong, prognostic molecular markers should be performed, which may help to guide the assessment of risk and subsequent management.

Standard treatment options for mixed gliomas include the following:

1. Surgery plus radiation therapy with or without chemotherapy.

(Refer to the Astrocytic Tumors section in the Treatment of Primary Central Nervous System Tumors by Tumor Type section of this summary for more information about astrocytic tumors.)

**Ependymal Tumors Treatment**

Ependymal tumors (WHO grade I) and ependymomas (WHO grade II)—i.e., subependymomas and myxopapillary ependymomas—are often curable.

Standard treatment options for grades I and II ependymal tumors include the following:

1. Surgery alone if the tumor is totally resectable.
2. Surgery followed by radiation therapy to known or suspected residual tumor.

Patients with anaplastic ependymomas (WHO grade III) have variable prognoses that depend on the location and extent of disease. Frequently, but not invariably, patients with anaplastic ependymomas have worse prognoses than do those patients with lower-grade ependymal tumors.

Standard treatment options for anaplastic ependymomas include the following:

1. Surgery plus radiation therapy.[30]

**Embryonal Cell Tumors (Medulloblastomas) Treatment**

Medulloblastoma occurs primarily in children, but may also occur in adults.[31] (Refer to the PDQ summary on Childhood Central Nervous System Embryonal Tumors Treatment for more information.)

Standard treatment options for medulloblastomas include the following:

1. Surgery plus craniospinal radiation therapy for good-risk patients.[32]

**Treatment options under clinical evaluation for medulloblastomas**

Treatment options under clinical evaluation include the following:

1. Surgery plus craniospinal radiation therapy and various chemotherapy regimens are being evaluated for poor-risk patients.[32]

**Pineal Parenchymal Tumors Treatment**

Pineocytomas (WHO grade II), pineoblastomas (WHO grade IV), and pineal parenchymal tumors of intermediate differentiation are diverse tumors that require special consideration. Pineocytomas are slow-growing tumors and prognosis varies.

Pineoblastomas grow more rapidly and patients with these tumors have worse prognoses. Pineal parenchymal tumors of intermediate differentiation have unpredictable growth and clinical behavior.

Standard treatment options for pineal parenchymal tumors include the following:

2. Surgery plus radiation therapy and chemotherapy for pineoblastoma.
Meningeal Tumors Treatment

WHO grade I meningiomas are usually curable when they are resectable. With the increasing use of sensitive neuroimaging tools, there has been more detection of asymptomatic low-grade meningiomas. Most appear to show minimal growth and can often be safely observed while therapy is deferred until growth or the development of symptoms.[33,34]

Standard treatment options for meningeal tumors include the following:

1. Active surveillance with deferred treatment, especially for incidentally discovered asymptomatic tumors.[33,34].
2. Surgery.
3. Stereotactic radiosurgery for tumors smaller than 3 cm.
4. Surgery plus radiation therapy in selected cases, such as for patients with known or suspected residual disease or with recurrence after previous surgery.
5. Fractionated radiation therapy for patients with unresectable tumors.[35]

The prognoses for patients with WHO grade II meningiomas (atypical, clear cell, and chordoid), WHO grade III meningiomas (anaplastic/malignant, rhabdoid, and papillary), and hemangiopericytomas are worse than the prognoses for patients with low-grade meningiomas because complete resections are less commonly feasible, and the proliferative capacity is greater.

Standard treatment options for grades II and III meningiomas and hemangiopericytomas include the following:


Germ Cell Tumors Treatment

The prognoses and treatment of patients with germ cell tumors—which include germinomas, embryonal carcinomas, choriocarcinomas, and teratomas—depend on tumor histology, tumor location, presence and amount of biological markers, and surgical resectability.

Treatment of Tumors of the Sellar Region

Craniopharyngiomas (WHO grade I) are often curable.

Standard treatment options for craniopharyngiomas include the following:

1. Surgery alone if the tumor is totally resectable.
2. Debulking surgery plus radiation therapy if the tumor is unresectable.

Treatment Options Under Clinical Evaluation for Primary CNS Tumors

Patients who have CNS tumors that are either infrequently curable or unresectable should be considered candidates for clinical trials. Information about ongoing clinical trials is available from the NCI website.

Heavy-particle radiation, such as proton-beam therapy, carries the theoretical advantage of delivering high doses of ionizing radiation to the tumor bed while sparing surrounding brain tissue. The data are preliminary for this investigational technique, and are not widely available.

Novel biologic therapies under clinical evaluation for patients with CNS tumors include the following:[36]

- Dendritic cell vaccination.[37]
- Tyrosine kinase receptor inhibitors.[38]
- Farnesyl transferase inhibitors.
- Viral-based gene therapy.[39,40]
- Oncolytic viruses.
• Epidermal growth factor-receptor inhibitors.
• Vascular endothelial growth factor inhibitors.[36]
• Other antiangiogenesis agents.

Current Clinical Trials

Check the list of NCI-supported cancer clinical trials that are now accepting patients with adult brain tumor. The list of clinical trials can be further narrowed by location, drug, intervention, and other criteria.

General information about clinical trials is also available from the NCI website.

References


Treatment of Primary Tumors of the Spinal Axis

Surgery and radiation therapy are the primary modalities used to treat tumors of the spinal axis; therapeutic options vary according to the histology of the tumor.[1] The experience with chemotherapy for primary spinal cord tumors is limited; no reports of controlled clinical trials are available for these types of tumors.[1,2] Chemotherapy is indicated for most patients with leptomeningeal involvement from a primary or metastatic tumor and positive cerebrospinal fluid cytology.[1] Most patients require treatment with corticosteroids, particularly if they are receiving radiation therapy.

Patients who have spinal axis tumors that are either infrequently curable or unresectable should be considered candidates for clinical trials. Information about ongoing clinical trials is available from the NCI website.

References


Metastatic Brain Tumors

General Information About Metastatic Brain Tumors

Brain metastases outnumber primary neoplasms by at least 10 to 1, and they occur in 20% to 40% of cancer patients, with subsequent median survival generally less than 6 months.[1] The exact incidence is unknown because no national cancer registry documents brain metastases, but it has been estimated that 98,000 to 170,000 new cases are diagnosed in the United States each year.[2,3] This number may be increasing because of the capacity of magnetic resonance imaging (MRI) to detect small metastases and because of prolonged survival resulting from improved systemic therapy.[1,2]

The most common primary tumors with brain metastases and the percentage of patients effected are as follows:[1,2]

- Lung (18%-64%).
- Breast (2%-21%).
- Cancer of unknown primary (1%-18%).
- Melanoma (4%-16%).
- Colorectal (2%-12%).
- Kidney (1%-8%).

Eighty percent of brain metastases occur in the cerebral hemispheres, 15% occur in the cerebellum, and 5% occur in the brain stem.[2] Metastases to the brain are multiple in more than 70% of cases, but solitary metastases also occur.[1] Brain involvement can occur with cancers of the nasopharyngeal region by direct extension along the cranial nerves or through the foramina at the base of the skull. Dural metastases may constitute as much as 9% of total brain metastases.

Clinical Features

The diagnosis of brain metastases in cancer patients is based on the following:

- Patient history.
Neurological examination.

Diagnostic procedures, including a contrast MRI of the brain.

Patients may describe any of the following:

- Headaches.
- Weakness.
- Seizures.
- Sensory defects.
- Gait problems.

Often, family members or friends may notice the following:

- Lethargy.
- Emotional lability.
- Personality change.

**Diagnostic Evaluation**

A physical examination may show objective neurological findings or only minor cognitive changes. The presence of multiple lesions and a high predilection of primary tumor metastasis may be sufficient to make the diagnosis of brain metastasis.

A lesion in the brain should not be assumed to be a metastasis just because a patient has had a previous cancer; such an assumption could result in overlooking appropriate treatment of a curable tumor.

**Imaging tests**

Computed tomography scans with contrast or MRIs with gadolinium are quite sensitive in diagnosing the presence of metastases. Positron emission tomography scanning and spectroscopic evaluation are new strategies to diagnose cerebral metastases and to differentiate the metastases from other intracranial lesions.[4]

**Biopsy**

In the case of a solitary lesion or a questionable relationship to the primary tumor, a brain biopsy (via resection or stereotactic biopsy) may be necessary.

**Treatment of Metastatic Brain Tumors**

The optimal therapy for patients with brain metastases continues to evolve.[1,2,5] The following treatments have been used in the management of metastatic brain tumors:

- Radiation therapy.
- Radiosurgery.
- Surgical resection.
- Corticosteroids.
- Anticonvulsants.

Because most cases of brain metastases involve multiple metastases, a mainstay of therapy has historically been whole-brain radiation therapy (WBRT); however, stereotactic radiosurgery has come into increasingly common use. The role of radiosurgery continues to be defined. Stereotactic radiosurgery in combination with WBRT has been assessed.

Surgery is indicated to obtain tissue from a metastasis with an unknown primary tumor or to decompress a
symptomatic dominant lesion that is causing significant mass effect.

Chemotherapy is usually not the primary therapy for most patients; however, it may have a role in the treatment of patients with brain metastases from chemosensitive tumors and can even be curative when combined with radiation for metastatic testicular germ cell tumors.[1,6] Intrathecal chemotherapy is also used for meningeal spread of metastatic tumors.

**Treatment for patients with one to four metastases**

*Standard treatment options for patients with one to four metastases*

About 10% to 15% of patients with cancer will have a single brain metastasis. Radiation therapy is the mainstay of palliation for these patients. The extent of extracranial disease can influence treatment of the brain lesions. In the presence of extensive active systemic disease, surgery provides little benefit for overall survival (OS). In patients with stable minimal extracranial disease, combined modality treatment may be considered, using surgical resection followed by radiation therapy. However, the published literature does not provide clear guidance.

Standard treatment options for patients with one to four metastases include the following:

1. WBRT with or without surgical resection.
2. WBRT with or without stereotactic radiosurgery.
3. Focal therapy alone (surgical resection or stereotactic radiosurgery).

**Evidence (treatment for one to four metastases):**

1. Three randomized trials examined resection of solitary brain metastases followed by WBRT versus WBRT alone, totaling 195 randomly assigned patients.[7-9] The process that necessarily goes into selecting appropriate patients for surgical resection may account for the small numbers in each trial. In the first trial,[7][Level of evidence: 1iiD] performed at a single center, all patients were selected and operated upon by one surgeon.
   a. The first two trials showed an improvement in survival in the surgery group,[7,8] but the third trial showed a trend in favor of the WBRT–only group.[9]
   b. The three trials were combined in a trial-level meta-analysis.[10] The combined analysis showed the following:
      - The combined analysis did not show a statistically significant difference in OS (hazard ratio [HR] = 0.72; 95% confidence interval [CI], 0.34–1.53; $P = .4$); or in death from neurologic causes (relative risk of death = 0.68; 95% CI, 0.43–1.09; $P = .11$).[10]
      - One of the trials reported that combined therapy increased the duration of functionally independent survival.[7][Level of evidence: 1iiD]
      - None of the trials assessed or reported quality of life.

2. The need for WBRT after resection of solitary brain metastases has been studied.[11] Patients were randomly assigned to either undergo postoperative WBRT or receive no further treatment after resection.
   - Patients in the WBRT group were less likely to have tumor progression in the brain and were significantly less likely to die of neurologic causes.
   - OS was the same in each group, and there was no difference in duration of functional independence.

3. One additional randomized study of observation versus WBRT after either surgery or stereotactic radiosurgery for solitary brain metastases was closed after 19 patients had been entered because of slow accrual; therefore, little can be deduced from the trial.[12]

4. A Radiation Therapy Oncology Group (RTOG) study (RTOG-9508) randomly assigned 333 patients with one to three metastases with a maximum diameter of 4 cm to WBRT (37.5 Gy over 3 weeks) with or without a stereotactic boost.[13] Patients with active systemic disease requiring therapy were excluded. The primary
endpoint was OS with predefined hypotheses in both the full study population and the 186 patients with a solitary metastasis (and no statistical adjustment of \( P \) values for the two separate hypotheses).[13][Levels of evidence: 1iiDii for the full study population and 1iIA for patients with solitary metastases]

a. Mean OS in the combined-therapy group was 5.7 months, and mean OS in the WBRT–alone group was 6.5 months (\( P = .14 \)).

- In the subgroup with solitary metastases, OS was better in the combined-therapy group (6.5 months vs. 4.9 months; \( P = .039 \) in univariate analysis; \( P = .053 \) in a multivariable analysis adjusting for baseline prognostic factors).
- In patients with multiple metastases, survival was 5.8 months in the combined-therapy group versus 6.7 months in the WBRT–only group (\( P = .98 \)).
- The combined-treatment group had a survival advantage of 2.5 months in patients with a single metastasis but not in patients with multiple lesions.

b. Local control was better in the full population with combined therapy.

c. At the 6-month follow-up, Karnofsky Performance status (considered a soft endpoint because of its imprecision and subjectivity) was better in the combined-therapy group, but there was no difference in mental status between the treatment groups. Acute and late toxicities were similar in both treatment arms. Quality of life was not assessed.

5. A phase III randomized trial compared adjuvant WBRT with observation after surgery or radiosurgery for a limited number of brain metastases in patients with stable solid tumors.[14][Level of evidence: 1iIC]

- Health-related quality of life was improved in the observation-only arm, compared with WBRT.
- Patients in the observation arm had better mean scores in physical, role, and cognitive functioning at 9 months.
- In an exploratory analysis, statistically significant worse scores for bladder control, communication deficit, drowsiness, hair loss, motor dysfunction, leg weakness, appetite loss, constipation, nausea/vomiting, pain, and social functioning were observed in patients who underwent WBRT, compared with those who underwent observation only.

6. A meta-analysis of two trials with a total of 358 participants found no statistically significant difference in OS between the WBRT plus stereotactic radiosurgery group and the WBRT–alone group (HR, 0.82; 95% CI, 0.65–1.02).[15][Level of evidence: 1iIDiii]

- Patients in the WBRT plus stereotactic radiosurgery group had decreased local failure, compared with patients who received WBRT alone (HR, 0.27; 95% CI, 0.14–0.52).
- Unchanged or improved Karnofsky Performance status at 6 months was seen in 43% of patients in the combined-therapy group versus only 28% in the WBRT-alone group (\( P = .03 \)).

A study that had a primary endpoint of learning and neurocognition, using a standardized test for total recall, was stopped by the Data and Safety Monitoring Board because of worse outcomes in the WBRT group.[16][Level of evidence: 1iiD]

Given this body of information, focal therapy plus WBRT or focal therapy alone, with close follow-up with serial MRIs and initiation of salvage therapy when clinically indicated, appear to be reasonable treatment options. The pros and cons of each approach should be discussed with the patient.

Several randomized trials have been performed that were designed with varying primary endpoints to address whether WBRT is necessary after focal treatment. The results can be summarized as follows:[16–18]

1. Studies consistently show that the addition of WBRT to focal therapy decreases the risk of progression and new metastases in the brain.

2. The addition of WBRT does not improve OS.
3. The decrease in risk of intracranial disease progression does not translate into improved functional or neurologic status, nor does it appear to decrease the risk of death from neurologic deterioration.

4. About half or more of the patients who receive focal therapy alone ultimately require salvage therapy, such as WBRT or radiosurgery, compared with about a quarter of the patients who are given up-front WBRT.

5. The impact of better local control associated with WBRT on quality of life has not been reported and remains an open question.

**Leptomeningeal carcinomatosis (LC)**

LC occurs in about 5% of all cancer patients. The most common types of cancer to spread to the leptomeninges are:

- Breast tumors (35%).
- Lung tumors (24%).
- Hematologic malignancies (16%).

Diagnosis includes a combination of neurospinal axis imaging and cerebrospinal fluid (CSF) cytology. Median OS is in the range of 10 to 12 weeks.

The management of LC includes the following:

- Intrathecal chemotherapy.
- Intrathecal chemotherapy and systemic chemotherapy.
- Intrathecal chemotherapy and radiation therapy.
- Supportive care.

In a series of 149 patients with metastatic non-small cell lung carcinoma, cytologically proven LC, poor performance status, high protein level in the CSF, and a high initial CSF white blood cell count were significant poor prognostic factors for survival. Patients received active treatment, including intrathecal chemotherapy, WBRT, or epidermal growth factor receptor-thymidine kinase-1, or underwent a ventriculoperitoneal shunt procedure. In a retrospective series of 38 patients with metastatic breast cancer and LC, the proportion of LC cases varied by breast cancer subtype:

- Luminal A (18.4%).
- Luminal B (31.6%).
- Human epidermal growth factor receptor 2 (HER2)-positive (26.3%).
- Triple-negative breast cancer subtype (23.7%).

Patients with triple-negative breast cancer had a shorter interval between metastatic breast cancer diagnosis and the development of LC. Median survival did not differ across breast cancer subtypes. Consideration of intrathecal administration of trastuzumab in patients with HER2-positive LC has also been described in case reports.

**References**

Recurrent Adult CNS Tumors Treatment

Patients who have recurrent CNS tumors are rarely curable and should be considered candidates for clinical trials. Information about ongoing clinical trials is available from the NCI website.

Standard treatment options for recurrent CNS tumors include the following:

1. Chemotherapy.
2. Antiangiogenesis therapy.
3. Radiation therapy.

Chemotherapy

Localized chemotherapy (carmustine wafer)

Carmustine wafers have been investigated for the treatment of recurrent malignant gliomas, but the impact on survival is less clear than at the time of initial diagnosis and resection.

Evidence (localized chemotherapy):

1. In a multicenter randomized, placebo-controlled trial, 222 patients with recurrent malignant primary brain tumors requiring reoperation were randomly assigned to receive implanted carmustine wafers or placebo biodegradable wafers.[1][Level of evidence: 1iA] Approximately half of the patients had received prior systemic chemotherapy. The two treatment groups were well balanced at baseline.
   - Median survival was 31 weeks in the group receiving carmustine wafers versus 23 weeks in the group receiving placebo wafers. The statistical significance between the two overall survival curves depended on the method of analysis.
   - The hazard ratio (HR) for risk of dying in the direct intention-to-treat comparison between the two groups was 0.83 (95% confidence interval [CI], 0.63–1.10; \( P = .19 \)). The baseline characteristics were similar in the two groups, but the investigators performed an additional analysis, adjusting for prognostic factors, because they felt that even small differences in baseline characteristics could have a powerful influence on outcomes. In the adjusted proportional hazards model, the HR for risk of death was 0.67 (95% CI, 0.51–0.90, \( P = .006 \)). The investigators emphasized this latter analysis and reported this as a positive trial. [1][Level of evidence: 1iA]

2. A Cochrane Collaboration systematic review of chemotherapeutic wafers for high-grade glioma focused on the unadjusted analysis and reported the same trial as negative.[2]

Systemic chemotherapy

Systemic therapy (e.g., temozolomide, lomustine, or the combination of procarbazine, a nitrosourea, and PCV in patients who have not previously received the drugs) has been used at the time of recurrence of primary malignant brain tumors. However, this approach has not been tested in controlled studies. Patient-selection factors likely play a strong role in determining outcomes, so the impact of therapy on survival is not clear.

Antiangiogenesis Therapy

In 2009, the U.S. Food and Drug Administration (FDA) granted accelerated approval of bevacizumab monotherapy for patients with progressive glioblastoma. The indication was granted under the FDA's accelerated approval program that permits the use of certain surrogate endpoints or an effect on a clinical endpoint other than survival or irreversible morbidity as bases for approvals of products intended for serious or life-threatening illnesses or conditions.

The approval was based on the demonstration of improved objective response rates observed in two historically controlled, single-arm, or noncomparative phase II trials.[3,4][Level of evidence: 3iiiDiv] On the basis of these data and FDA approval, bevacizumab monotherapy has become standard therapy for recurrent glioblastoma.

Evidence (antiangiogenesis therapy):

1. The FDA independently reviewed an open-label, multicenter, noncomparative phase II study that randomly assigned 167 recurrent glioblastoma multiforme (GBM) patients to receive bevacizumab alone or bevacizumab in combination with irinotecan,[3] although only efficacy data from the bevacizumab monotherapy arm (\( n = 85 \)) were used to support drug approval.
   - Tumor responses were observed in 26% of patients treated with bevacizumab alone, and the median duration of response in these patients was 4.2 months.
   - On the basis of this externally controlled trial, the incidence of adverse events associated with bevacizumab did not appear to be significantly increased in GBM patients.
2. The FDA independently assessed another single-arm, single-institution trial in which 56 recurrent glioblastoma patients were treated with bevacizumab alone.[4]

- Responses were observed in 20% of patients, and the median duration of response was 3.9 months.

Currently, however, no data are available from prospective, randomized controlled trials demonstrating improvement in health outcomes, such as disease-related symptoms or increased survival with the use of bevacizumab to treat glioblastoma.

Radiation Therapy

Because there are no randomized trials, the role of repeat radiation after disease progression or the development of radiation-induced cancers is also ill defined. Interpretation is difficult because the literature is limited to small retrospective case series.[5] The decision must be made carefully because of the risk of neurocognitive deficits and radiation necrosis.

Surgery

Re-resection of recurrent CNS tumors is used for some patients. However, most patients do not qualify because of a deteriorating condition or technically inoperable tumors. The evidence is limited to noncontrolled studies and case series of patients who are healthy enough and have tumors that are small enough to technically debulk. The impact on survival of reoperation versus patient selection is not known.

Current Clinical Trials

Check the list of NCI-supported cancer clinical trials that are now accepting patients with recurrent adult brain tumor. The list of clinical trials can be further narrowed by location, drug, intervention, and other criteria.

General information about clinical trials is also available from the NCI website.

References


Changes to This Summary (10/27/2015)

The PDQ cancer information summaries are reviewed regularly and updated as new information becomes available. This section describes the latest changes made to this summary as of the date above.

General Information About Adult Primary Central Nervous System (CNS) Tumors

An editorial change was made to this section.

This summary is written and maintained by the PDQ Adult Treatment Editorial Board, which is editorially independent of NCI. The summary reflects an independent review of the literature and does not represent a policy statement of NCI or NIH. More information about summary policies and the role of the PDQ Editorial Boards in maintaining the PDQ summaries can be found on the About This PDQ Summary and PDQ® - NCI's Comprehensive Cancer Database pages.

About This PDQ Summary
Purpose of This Summary

This PDQ cancer information summary for health professionals provides comprehensive, peer-reviewed, evidence-based information about the treatment of adult central nervous system tumors. It is intended as a resource to inform and assist clinicians who care for cancer patients. It does not provide formal guidelines or recommendations for making health care decisions.

Reviewers and Updates

This summary is reviewed regularly and updated as necessary by the PDQ Adult Treatment Editorial Board, which is editorially independent of the National Cancer Institute (NCI). The summary reflects an independent review of the literature and does not represent a policy statement of NCI or the National Institutes of Health (NIH).

Board members review recently published articles each month to determine whether an article should:

- be discussed at a meeting,
- be cited with text, or
- replace or update an existing article that is already cited.

Changes to the summaries are made through a consensus process in which Board members evaluate the strength of the evidence in the published articles and determine how the article should be included in the summary.

The lead reviewers for Adult Central Nervous System Tumors Treatment are:

- Andrew S. Chi, MD, PhD (Massachusetts General Hospital)
- Minh Tam Truong, MD (Boston University Medical Center)

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