Current diagnosis and treatment of oligodendroglioma

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Object. The strategies used to diagnose and treat oligodendrogliarial tumors have changed significantly over the past decade. The purpose of this paper is to review the topic of oligodendroglioma, emphasizing the new developments.

Methods. Information was obtained by conducting a Medline search in which the term oligodendroglioma was used. Recent editions of standard textbooks were also studied.

Because of tools such as magnetic resonance imaging, oligodendrogliomas are being diagnosed earlier, and they are being recognized more frequently histologically than in the past. Seizures are common in these patients. Functional mapping and image-guided surgery may now allow for a safer and more complete resection, especially when tumors are located in difficult areas. Genetic analysis and positron emission tomography may provide data that supplement the standard diagnostic tools. Unlike other low-grade gliomas, patients in whom residual or recurrent oligodendroglioma (World Health Organization Grade II) is present may respond to chemotherapy. Although postoperative radiotherapy prolongs survival of the patient, increasingly this therapeutic modality is being delayed until tumor recurrence, especially if a gross-total tumor resection has been achieved. Oligodendrogliomas are the first type of brain tumor for which “molecular” characterization gives important information. The most significant finding is that allelic losses on chromosomes 1p and 19q indicate a favorable response to chemotherapy.

Conclusions. Whereas surgery continues to be the primary treatment for oligodendroglioma, the scheme for postoperative therapy has shifted, primarily because of the lesion’s relative chemosensitivity. Molecular characterization of oligodendrogliomas may become a standard practice in the near future.

KEY WORDS • oligodendroglioma • brain neoplasm • chemotherapy • functional magnetic resonance imaging • positron emission tomography • radiation therapy
gliomatosis) have been reported to occur in up to 14% of cases.19,56,72,85 Oligodendrogliomas may metastasize outside the brain more frequently than other gliomas, possibly because of the longer periods of survival in these patients.56

Symptoms related to oligodendrogial tumors are non-specific—that is, they do not reliably distinguish this type of brain tumor from other types.31 Symptoms are caused by involvement of the central nervous system in general and/or because of the specific anatomical location of the tumor. In most series, seizure has been the most commonly reported presenting symptom, ranging in incidence from 35 to 85% of patients.12,31,36,57,82,92 Daumas-Duport, et al.23 have reported that 91% of their patients presented with seizures at some point in their clinical course. The seizures may be simple partial, complex partial, generalized, or a combination of these.60 Other reported presenting symptoms have included headache, mental status change, visual complaint, vertigo/nausea, and/or focal weakness.21

The duration of symptoms prior to diagnosis is highly variable; and a diagnosis may be reached after a new seizure, headache, or (rarely) hemorrhage, as well as after a decade of epilepsy.31,36

DIAGNOSTIC IMAGING

Findings on diagnostic imaging are often characteristic, but not pathognomonic.82,90 Usually oligodendrogliomas are mass lesions with fairly well-defined margins, located in the cortex and subcortical white matter.47 On CT scans, oligodendrogliomas appear hypodense or isodense and enhance poorly, if at all.51 On MR images, oligodendrogliomas are usually hypointense on T1-weighted sequences and hyperintense on T2-weighted sequences.23,46,47,51 Calcification is common; CT scanning demonstrates the calcium deposits better than plain radiography or MR imaging.47,50 Statistically, because of the higher incidence of astrocytomas, a glioma with calcium deposits is more likely to be an astrocytoma.31,46 Cystic changes and hemorrhage may be seen; peritumoral edema is usually mild or absent.27 Like astrocytomas, oligodendrogliomas may spread through the corpus callosum, the leptomeninges, and/or the ependyma.51,64 Magnetic resonance imaging is more sensitive than CT scanning in demonstrating contrast enhancement, which may be patchy or homogeneous. Contrast enhancement may indicate a more aggressive tumor.23,47,51,60,62 The appearance of ring enhancement has been associated with a poor prognosis.11 In children and adolescents with oligodendrogliomas, calcifications, enhancement, and edema are demonstrated less frequently than in adults.82 An example of a CT scan depicting an unusual oligodendroglioma is shown in Fig. 1; an example of an MR image is given in Fig. 2.

The authors of recent studies have indicated that PET scanning may provide useful information in patients with gliomas, including oligodendroglioma and anaplastic oligodendroglioma. Positron emission tomography can be performed to give quantitative information regarding the blood flow, glucose metabolism, and amino acid metabolism of the brain as well as brain tumors.26,27,55,81 Based on this information, PET has been used as a tool in distinguishing scar or gliotic tissue from tumor.88 With respect to gliomas, PET has been reported to be able to differentiate between low-grade astrocytoma and oligodendroglioma.27 In a study of 22 patients, Derlon, et al.27 have found that both of these tumor types exhibit glucose hypometabolism (slightly more pronounced in astrocytoma) but markedly different patterns of uptake of the labeled amino acid, MET. The uptake of MET was high in the oligodendrogliomas and decreased, normal, or only moderately increased in the astrocytomas.27 The authors speculated that this difference might be related to different cell densities in the tumor tissue. Determining MET uptake may also be more helpful than studying glucose metabolism in patients with oligodendroglioma who are being followed for residual or recurrent tumor.27

Positron emission tomography might also prove to be useful for grading oligodendrogliomas noninvasively. In a different study by Derlon, et al.,26 MET uptake and 18F-fluorodeoxyglucose uptake were used to determine amino acid metabolism and glycolysis in comparison of low- and high-grade oligodendrogliomas. The authors found that anaplastic oligodendrogliomas exhibited a higher 18F-fluorodeoxyglucose uptake than did the oligodendrogliomas but that the difference in MET uptake was even more pronounced. Positron emission tomography scans can be co-registered (that is, displayed together) with MR images to combine the metabolic data provided by PET with the superior anatomical resolution achieving using MR imaging.83 Positron emission tomography has been used to guide target selection for stereotactic biopsy sampling and to guide resection.88 Therefore, although confirmatory studies with large numbers of patients still need to be conducted, PET seems to hold promise for future use in the management of patients with oligodendroglioma.

Fig. 1. Unenhanced CT scan revealing a high density lesion extending from the midbrain to middle pons. Although this was believed likely to represent a cavernous hemangioma with hemorrhage, its appearance did not change over time, and examination of a biopsy sample later showed it to be an oligodendroglioma (WHO Grade II). Oligodendrogliomas may have a variable appearance and may occur in unusual locations.
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Fig. 2. A T2-weighted MR image of a Grade II oligodendroglioma, demonstrating the tumor as a bright signal in the anterior and medial temporal lobe. In this type of tumor, T1-weighted sequences may be more sensitive than T2-weighted sequences.

SURGICAL INTERVENTION

Surgery continues to be the primary treatment modality for most patients with gliomas, including oligodendroglioma.2,31,50,70,73 Until highly accurate noninvasive histological assessment becomes a reality, surgery (biopsy sampling or resection) will continue to be essential for establishing an accurate diagnosis.33,60 Tumor resection (when feasible) is also useful for reducing mass effect, which may be the cause of symptoms and/or neurological deficit. Tumor resection may also decrease the need for steroid therapy, allow a decrease of radiotherapeutic portal size, increase the effect of chemotherapy, and limit sampling error that may occur in cases in which a biopsy sample alone is obtained.33 A large cranial opening is typically made to facilitate optimum exposure and the use of multiple trajectories, as well as to aid in decompression of the brain. At the time of surgery, it may not be possible to distinguish reliably between oligodendroglial lesions and other intraxial tumors. Establishing an intraoperative diagnosis of oligodendroglioma can be difficult based on the frozen section histological specimen.32

The rate of surgery-related morbidity should be low but is dependent on the location of the tumor. Although these lesions are usually infiltrative and blend into normal brain, at times there may be an abrupt tumor–adjacent white matter interface.30 Staying within the confines of the tumor, as indicated by preoperative imaging studies, does not guarantee that a deficit will not occur. Functional MR imaging, functional cortical mapping, awake surgery, image-guided surgery (stereotactic craniotomy), and/or intraoperative MR imaging are modalities that may be used to increase the amount of tumor that can be safely resected.6,20,29,66 The value of radiosurgery, for instance in a patient in whom an oligodendroglioma has locally recurred, remains undetermined.

RESECTION AND SURVIVAL

Concerning extent of resection and survival, there are very little data available from the post–MR imaging era.68 In some clinical series addressing this issue the authors have not been able to confirm that a more complete resection is beneficial.30,80 In most reports, however, the authors have concluded that more complete resection is associated with increased patient survival.6,12,24,38,45,48,58,71,73 Interestingly, in a recent series reported by Giannini, et al.,35 gross-total resection or a biopsy procedure was associated with longer survival than subtotal resection. Most authors believe that the surgery-related goal for oligodendroglioma should be gross-total removal if the tumor can be safely resected.6,20,70,79 Aggressive resection in regions engendering significant risk of neurological damage should probably be avoided, because these tumors are likely to respond to other therapies.33

PATHOLOGICAL EXAMINATION

Oligodendrogliomas appear macroscopically to be soft masses of fleshy to pinkish-gray color, but they may be gelatinous due to mucoid degeneration. They may contain areas of firm, gritty calcifications and/or areas of soft cystic degeneration, and intratumoral hemorrhages can be present.10,32,64 Histological examination shows moderate cellularity, with tumor cells containing uniformly round, homogeneous nuclei and a swollen clear cytoplasm. This typical "honeycomb" or "fried-egg" appearance is actually an artifact of fixation but is a useful diagnostic feature if present.32 Additionally, oligodendrogliomas can display a dense network of branching capillaries in what has been termed a "chicken-wire" vascular pattern.32 Oligodendrogliomas may contain varying percentages of astrocytic cells. These are hypothesized to represent reactive astrocytes trapped by the invasive tumor, transitional forms of oligodendrogial cells, or differentiated neoplastic astrocytic cells. There is no specific immunocytochemical marker that allows for the recognition of human oligodendrogial tumor cells. Electron microscopy can help to discriminate oligodendrogliomas from other types of tumors.13,32 A photomicrograph of a hematoxylin and eosin stained oligodendroglioma specimen is shown in Fig. 3.

POSTOPERATIVE TREATMENT

Because oligodendrogliomas, like other gliomas, have an infiltrative growth pattern, and the true extent of tumor cells cannot be determined using current imaging modalities,30,31 an oligodendroglioma is very rarely completely resected, or cured.34 Recurrences are usually seen locally at the previous operative site.50 Patients with oligodendroglioma usually do succumb to progressive disease or to the effects of conversion of their tumor to a higher grade.79 Because of this, postoperative limited-field radiotherapy has been performed in the past as a treatment option in adults with low-grade oligodendrogliomas. The majority of the available literature does indicate that radiotherapy prolongs patient survival, especially if the tumor has been partially resected.2,24,34,40,71,73–75 In children, radiation is withheld whenever possible.2,50

Even in adults, because these patients experience a rel-
other infections, neutropenia, and thrombocytopenia. In alopathy, seizures, intracranial hemorrhage, pneumonia and nal pain, constipation, neuropathy, hepatotoxicity, enceph-egy, rash, numbness or paresthesias, weakness, abdominal pain, constipation, neuropathy, hepatotoxicity, encephalopathy, seizures, intracranial hemorrhage, pneumonia and other infections, neutropenia, and thrombocytopenia. In the series published by Olson, et al., 46% of patients treated with PCV chemotherapy developed significant myelosuppression. Given the potential benefit of chemotherapy, however, until new data become available, PCV chemotherapy continues to be an option for the treatment of patients with low-grade oligodendroglioma, either before or after radiotherapy, particularly if residual or recurrent tumor is present. Temozolomide is being tested in clinical trials.

A recurrent oligodendroglioma is often an anaplastic oligodendroglioma or glioblastoma multiforme. In patients with recurrent oligodendroglioma, either radiotherapy and/or PCV chemotherapy is undertaken, depending on which treatment has already been used. Patients in whom radiotherapy has been previously performed seem to be as likely to respond to PCV therapy as those who have not undergone irradiation therapy. Other salvage chemotherapy regimens have also been attempted. Veninga, et al., concluded that repeated radiotherapy can be considered as a treatment option in good-condition patients with recurrent gliomas, including oligodendrogliomas.

**CLINICAL PROGNOSTIC FACTORS AND SURVIVAL**

In addition to the aforementioned resection- and histology-related aspects, other prognostic factors have been reported for oligodendroglial tumors. Age has been found to be one of the strongest independent predictors of survival. In 2000 Olson, et al., reported 5- and 10-year survival rates of 73% and 49%, respectively, for patients with Grade II oligodendroglioma. In 2001 Henderson and Shaw reported a median survival time of 16.7 years in their series of patients with recurrent gliomas, including oligodendrogliomas.

Other favorable features include location in the frontal lobe (which may relate to the extent of resection possible) and higher initial or postoperative functional performance status. Additional studies are currently underway in which the authors may be able to confirm the validity of these clinical prognostic factors (H Engelhard, et al., unpublished data).

The survival times reported in recent studies are already significantly longer than those cited only a few years ago. In 2001 Henderson and Shaw reported 5- and 10-year survival rates of 73% and 49%, respectively, for patients with Grade II oligodendroglioma. In 2000 Olson, et al., reported a median survival time of 16.7 years in their series of patients with oligodendroglioma and mixed glioma. The authors of other studies have published similar results. Despite the information available, it is difficult to predict survival for the individual oligodendroglioma patient on clinical grounds, given the following factors: 1) earlier diagnosis and improved surveillance provided by MR imaging; 2) rapid evolution in treatment taking place; and 3) possibility of conversion to an anaplastic tumor that is refractory to all treatments.

**Fig. 3. Photomicrograph showing the histological features of a low-grade oligodendroglioma.** Note the round to oval-shaped nucleoli, with delicate chromatin and small nucleoli. The classic “fried-egg” appearance, which is an artifact of fixation, may not always be present. H & E, original magnification × 400.
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Molecular Biology and Tumor Markers

Significant advances have recently been made in our understanding of the molecular genetics of oligodendroglioma. Oligodendrogliomas generally demonstrate distinct genetic alterations that distinguish them from other types of gliomas. Their most frequent genetic alteration is the LOH on the long arm (designated “q”) of chromosome 19. In published studies, the incidence of LOH on 19q has varied from 50 to more than 80% of cases.\(^4,41,65,77,89\) The second most frequent genetic alteration is LOH on the short arm (designated “p”) of chromosome 1. The incidence of LOH on 1p has been reported to range from 40 to 92%.\(^4,41,65,77\) Such molecular markers have now been shown to have prognostic value in cases of oligodendrogliomas. Specifically, LOH on chromosome arm 1p (especially if accompanied by loss on 19q) appears to be strongly associated with the oligodendroglial phenotype and also an independent predictor of response to chemotherapy (with or without radiotherapy) and survival, in cases of high- and low-grade oligodendrogliomas.\(^3,11,39,77,83\) Ino, et al.,\(^41\) however, have recently reported that the loss of 1p does not identify all the chemosensitive tumors, nor was long survival demonstrated in all the patients with 1p loss. Therefore, further clarification of these findings is needed.

Cell proliferation markers such as Ki-67 and proliferative cell nuclear antigen have been studied in oligodendrogliomas to provide additional information regarding tumor biology.\(^63\) The Ki-67 antigen is recognized by the MIB-1 antibody. Typically, in WHO Grade II oligodendrogliomas, because mitotic activity is absent, labeling indices for proliferation markers such as Ki-67 are quite low. In some studies, Ki-67 staining has been reported to have prognostic significance (with higher staining implying poorer prognosis) and also to be higher in recurrent tumors than in the original tumors.\(^63,44,65,76\) Increased proliferative activity assessed by flow cytometry (as indicated by a higher percentage of cells in the S phase) has also been strongly associated with decreased survival time.\(^16\)

Enzyme markers have also been studied in oligodendroglioma. In a recent study, in which topoisomerase II-α (a molecular target for cytotoxic drugs) was evaluated, the authors found that it is associated with a higher proliferation rate and a poorer prognosis in patients with oligodendrogliomas. Topoisomerase was therefore suggested to be a useful marker for the selection of oligodendroglioma patients in whom a poorer prognosis exists and who would therefore be candidates for earlier adjuvant therapy.\(^53\) In another new immunohistochemical study, investigators evaluated the COX isoenzymes, COX-1 and COX-2.\(^25\) The COX-1 enzyme is expressed in macrophages/microglial cells. Patients with low COX-1 labeling scores were found to have a better survival than those with high scores. Because COX-2-expressing astrocytes were found to be present around areas of tumor necrosis, its expression was found to be significantly lower in the more benign oligodendrogliomas than in higher-grade oligodendrogliomas.

Uncommonly in oligodendrogliomas, there is decreased expression of the cell-cycle regulatory protein p16, which is encoded by a gene designated CDKN2A. If a CDKN2A deletion (or a decrease in p16 expression) does occur, it may be an important negative prognostic indicator.\(^7,11,54,69\) The p53 protein, nicknamed “the guardian of the genome,” has also been studied in oligodendrogliomas. Oligodendrogliomas have demonstrated p53 gene mutations in a small subset of cases.\(^3,19\) Growth factors that have been studied include vascular endothelial growth factor, the epidermal growth factor receptor, and platelet-derived growth factor. The expression of vascular endothelial growth factor in oligodendroglioma has been evaluated as a possible prognostic factor, and results have been mixed.\(^42\) Epidermal growth factor receptor does seem to be expressed in these tumors regardless of prognosis.\(^66,76\) Platelet-derived growth factors A and B and their receptors have also been found to be consistently expressed.\(^25,67\) Loss of the p18 tumor suppressor gene, which is located on chromosome 1p, may be involved in the progression of oligodendrogliomas.\(^37\)

**CONCLUSIONS**

Recent improvements in diagnostic imaging, surgical technique, histological diagnosis, and chemotherapy have significantly improved the prognosis for patients with oligodendroglioma.\(^14\) The observation that oligodendrogliomas (whether WHO Grade II or III) respond to PCV chemotherapy has been one of the more significant developments in medical neurooncology in recent years. Oligodendrogliomas seem to represent the first type of brain tumor for which genetic alterations and immunohistochemical findings may have significant prognostic value. Given the clear differences in biological features and response to treatment, investigators should no longer group oligodendrogliomas with other types of gliomas. With further advances in neuroimaging, molecular diagnostics, and chemotherapy, the treatment of patients with this tumor should be even more successful in the future.

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