We describe a case of anaplastic astrocytoma in a 14-year-old boy arising at the site of a dysembryoplastic neuroepithelial tumor (DNT) 3 years after combined radiation and chemotherapy. The subtotally excised superficial right temporoparietal tumor was originally diagnosed as mixed oligoastrocytoma in 1974; the patient was treated with radiation therapy postoperatively. One year later he underwent a craniotomy to remove cyst fluid and no change was reported in the size of the residual tumor. Postoperatively, he received a 6-week course of chemotherapy (lovustine, CCNU). He remained clinically and radiographically stable until 3 years later, when seizure activity returned and imaging studies were consistent with tumor recurrence. He was lost to follow-up until 1986, when records showed that he had died. Review of the initial biopsy showed cortical fragments containing abundant calcifications and multinodular structures typical of the complex form of DNT, in addition to specific glioneuronal elements. The Ki-67 labeling index ranged from 0.1% to 3% focally. The specimen from the third surgery showed an anaplastic astrocytoma (Ki-67 up to 12%) and morphologic features characteristic of radiation effect. This is the first documented case of malignant transformation of DNT following radiation and adjuvant chemotherapy. The implications of malignant transformation in subtotally excised complex DNTs and the intriguing issue of the contribution of radiation/chemotherapy are discussed.

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a right frontoparietal delta focus. Radiographic studies including brain scan and arteriogram, which confirmed a large superficial lesion in the right posterior parietal area with neovascularization and a posterior temporal component. The patient underwent a right temporal craniotomy, and an extensive cortical-based tumor in the mid-posterior temporoparietal lobe was subtotally resected. The pathologic diagnosis was mixed low-grade oligoastrocytoma. From February 27th to April 5th, 1975, the patient was referred for cobalt radiation therapy, which consisted of a course of 3,000 rads whole brain, followed by an additional 2,500 rads to the primary tumor site. He was apparently free of neurologic symptoms until February 1976, about 1 year after the initial surgery, when he experienced a generalized seizure. Examination of the surgical site revealed a mass pulsating through a 1 inch section of bone flap. A brain scan and arteriogram were abnormal and a second craniotomy was performed. At surgery, a large amount of fluid was drained from the site and the residual tumor, thought to be unchanged, was noted. Postoperatively, the patient received a 6-week course of oral CCNU chemotherapy. In March 1977, 3 years after the initial surgery, the patient experienced a recurrence of headaches and vomiting. Computerized tomography with intravenous contrast material revealed a mass with a small area of enhancement at the site of the two previous surgeries. A third craniotomy was performed and the tumor was grossly excised and interpreted as an anaplastic astrocytoma. Follow-up magnetic resonance imaging in September 1977 showed no evidence of residual tumor. The patient was lost to follow-up until 1986, when review of social security records showed that he had died. Additional information could not be obtained.

Microscopic sections from the initial resection were stained with hematoxylin-eosin and reviewed and compared with slides from the third surgery by the authors. Additional immunohistochemical stains including neurofilament proteins, Ki-67, synaptophysin, and glial fibrillary acidic protein were prepared from the accompanying paraffin blocks. Examination of tissue samples obtained at the time of the first surgical procedure showed multiple discrete cortical tumor nodules (Fig 1A) composed of a mixture of glial fibrillary acidic protein-positive astrocytic and glial fibrillary acidic protein-negative oligodendroglial-like components with foci of microcalcifications (Fig 1B). In addition, the neoplastic nodules contained several glioneuronal elements consisting of isolated neurons suspended in a pale eosinophilic matrix. Synaptophysin immunolabeling highlighted the neuronal cytoplasm and processes within the glioneuronal component. Mitotic activity, vascular changes, or necrosis were not identified. The Ki-67 labeling index varied from 0.1% to 3.0% focally. These morphologic features represented the complex form of DNT. Review of sections from the recurrent tumor obtained at the time of the third surgery revealed an infiltrating glial neoplasm composed of moderately pleomorphic cells, mostly of astrocytic lineage, accompanied by vascular endothelial hyperplasia and conspicuous mitotic activity (Fig 1C). There were foci of tumor with morphologic features similar to the original biopsy, including collections of oligodendroglial-like areas. Necrosis was not identified. Histopathologic features of radiation injury included areas of vascular hyalinization and hyperchromatic bizarre-appearing nuclei. Sections stained for Ki-67 revealed a labeling index of up to 12% in the more cellular areas. These histologic findings were diagnostic of a high-grade glioma, namely an anaplastic astrocytoma.

Discussion

The induction of central nervous system tumors within the irradiated field and the malignant transformation of a previously low-grade tumor are among the recognized adverse effects of radiotherapy.7,8,15-17 According to Huang et al,15 the criteria that define radiation-induced tumors are as follows: the tumor has a long quiescent or “latency period,” the location is in a previously irradiated field, and there is a verified histologic difference from the primary condition. In the present case, the anaplastic astrocytoma identified at the third surgery fulfilled the latter two criteria. Moreover, histopathologic features of radiation injury were identified in the tissue samples. The latency period in the present case was only 3 years; however, the combined mutagenic effects of radiation and chemotherapy may have accelerated the malignant transformation.18 The precise mechanisms that correlate irradiation with oncogenesis are not completely understood. A number of studies clearly implicate radiation and chemotherapy-induced deletions of critical DNA sequences as an important
event in triggering radiation-induced carcinogenesis.\textsuperscript{8,15} An alternative explanation is that the malignant transformation within the residual neoplastic tissue occurred independent of radiation or chemotherapy. In the original series of 39 cases by Daumas-Duport,\textsuperscript{1} 17 patients underwent subtotal resection with no clinical or radiologic evidence of recurrence on long-term follow-up. In the 13 patients in the original study who had undergone postoperative radiotherapy compared with the 26 subjects who had not received radiotherapy, there was no difference in survival or recurrence.\textsuperscript{1} In another series of patients with DNT who underwent temporal lobectomy for seizure control, there was adequate seizure control and no recurrences despite subtotal resection.\textsuperscript{19} Conversely, the recent case report by Hammond et al\textsuperscript{20} presents compelling evidence of the malignant potential of an incompletely resected DNT. In that case, the pathologic specimen obtained during subtotal removal of the tumor was originally diagnosed as a microcystic astrocytoma and the patient received no additional therapy. The patient remained well for 11 years, at which time progressive seizure activity recurred and imaging studies showed a recurrent enhancing partially cystic lesion. A glioblastoma multiforme, which contained a single focus reminiscent of DNT, was resected. Review of the original biopsy confirmed the diagnosis of DNT of the complex form.\textsuperscript{20} In another recent report, Nakatsuka\textsuperscript{21} published a case of an unclassified tumor closely resembling DNT with rapid growth in a 3-year-old girl who presented with a single generalized seizure. The tumor was restricted to the cortex, but lacked the specific glioneuronal element. These two cases and the present case challenge the widespread belief that all DNTs are slow-growing neoplasms and that incompletely excised DNTs have a uniformly favorable prognosis.

Although our case antedated the debut of DNT

Figure 1. (A) Photomicrograph from the original biopsy showing a representative cortical nodule typical of the complex form of DNT. (Hematoxylin-eosin stain, magnification $\times$120.) (B) Original biopsy with focus of calcification adjacent to the sharply demarcated neoplasm with microcystic features (Hematoxylin-eosin stain, magnification $\times$230.) (C) Histologic features of the third biopsy specimen, dominated by microvascular proliferation and astrocytic pleomorphism (Hematoxylin-eosin stain, magnification $\times$330.)
in the scientific literature, which underscores the potential pitfalls that persist in diagnosing this uncommon entity. In particular, stereotactic biopsies need to be interpreted with caution because of sampling limitations. The original diagnosis in our patient was oligoastrocytoma. On the basis of this diagnosis and the presence of residual neoplasm, the patient received adjuvant therapy. Indeed, the differential diagnosis of DNT includes oligodendroglioma, mixed oligoastrocytoma, ganglioglioma, and hamartoma. There are reliable morphologic clues to help distinguish these similar entities. It is notable that although oligodendrogliomas may include nodular foci, they lack the neuronal component of DNT. Hamartomas are characterized by the presence of disordered neuronal elements, without evidence of astrocytic or oligodendroglial components. Gangliogliomas are not intracortical and do not exhibit multi-nodular architecture. Gangliogliomas have an abundant reticulin-rich connective tissue stroma and often contain perivascular inflammatory infiltrates, which are absent in DNT. Of note, mixed or transitional neoplasms containing both discrete DNT and ganglioglioma components have been cited in the literature. The heterogeneous histopathologic features of DNTs and the recognition of the increasing number of morphologic variants of this entity continue to present a diagnostic challenge. The distinction of DNT from more aggressive neoplasms can be facilitated if viewed as a clinicopathologic entity. Daumas-Duport emphasizes that whatever the histologic appearance of a glial tumor, the diagnosis of DNT must be considered when the following criteria are associated: (1) partial seizures, with or without secondary generalization, beginning before the age of 20 years; (2) no neurologic deficit or stable congenital deficit; (3) cortical topography of the lesion as better demonstrated by magnetic resonance imaging; and (4) no mass effect on imaging. An increasing number of reports have described DNTs in extracortical sites. For example, a recent report by Baisden et al identifies 10 DNT-like neoplasms in the septum pellicundum. Another series of 11 cases of glioneuronal tumors with histologic features resembling DNT were found in the fourth ventricle. These reports provide strong evidence that DNTs may occur in other sites than the cerebral cortex.

In summary, we have presented the case of malignant transformation of a DNT following radiation and chemotherapy. Although it is not currently feasible to confirm the mutagenic role of these agents in an individual case, in the present case, there is convincing circumstantial evidence linking the two events. This case illustrates that DNTs may demonstrate a somewhat unpredictable biological behavior that warrants close monitoring.

References