CNS NEOPLASMS

PRESENTING SYMPTOMS OF PEDIATRIC BRAIN TUMORS

The presenting features of brain tumors in 200 children diagnosed between 1988 and 2001 were determined by retrospective chart review at the pediatric and neurosurgical services of the Wessex Neurology Centre and Southampton General Hospital, UK. Mean age at presentation was 7.4 years (range 15 weeks to 17 years). The male/female ratio was 4:3. The first symptoms in order of frequency were headache in 41%, vomiting (12%), ataxia (11%), visual problems (10%), educational or behavioral disorders (10%), and seizures (9%). Headaches were nocturnal or occurred early morning in 43 (61%) patients, and were continuous in 13 (18%). An early misdiagnosis of migraine had been made in 14 (13%), and tension headache in 8 (7%). Of 38 (19%) children with school problems, 12 had impairments in reading, writing, memory, concentration, and attendance. Visual disturbances in 76 (38%) patients included diplopia in 30, and blurred vision in 27. Seizures in 30 (15%) patients were focal in 19 and generalized in 11. Disturbances in growth and endocrine function were presenting features in 2% and were noted at some time in 18%.

Abnormal neurologic signs recorded in 175 (88%) children included papilledema in 66 (38%), cranial nerve abnormalities in 86 (49%), cerebellar signs in 84 (48%), long tract signs 47 (27%), and impaired consciousness in 21 (12%). Of 24 (12%) children with no neurologic findings at diagnosis, 17 had seizures (focal in 14), and 4 had headache. In 42 children (21%) aged 3 years or less, headache and seizures were less common than in older patients (12% vs 68% and 7% vs 17%, respectively), whereas behavioral problems were more common (48% vs 30%). The median duration of symptoms before diagnosis was 2.5 months (range 1 day to 120 months). Children aged 3 years or less had a significantly shorter symptom interval than older patients (1.0 vs 3.0 months). High grade tumors were diagnosed more rapidly than low grade tumors (median, 1.5 vs 4.0 months), and infratentorial tumors sooner than supratentorial tumors (2.0 vs 4.0 months). (Wilne SH, Ferris RC, Nathwani A,

COMMENT. Several similar studies of presenting symptoms of brain tumors in children are reported in the pre-medline literature, notably the classic monograph by Chicago authors, Bailey P, Buchanan DN, and Bucy PC. Intracranial Tumors of Infancy and Childhood. Chicago and London, University of Chicago Press, 1939. In the present report, the earliest reference cited was 1986, and the majority was published in the late 1990s and after 2000. Many of the authors’ findings corroborate those of earlier studies: 1) the incidence of seizures of 15% in children of all ages, with lesser frequency of 7% in children less than 3 years of age, and 2) delay in diagnosis of low grade vs high grade tumors and supratentorial vs infratentorial tumors. Relatively less recognized early manifestations of brain tumor emphasized in the present study include visual symptoms and learning and behavioral disorders, including lethargy.

In a study of 291 consecutive children treated for intracranial tumor at the Mayo Clinic from 1950-59, seizures occurred in 17% of the total group (in 25% of patients with supratentorial tumors and 12% of those with infratentorial tumors). (Backus RE, Millichap JG. Pediatrics 1962;29:978-984). The diagnosis of supratentorial tumors was delayed for an average of 2 years, whereas infratentorial tumors were diagnosed within 3 months of the initial seizure. Seizures were more common in patients with slowly growing grades 1 and 2 than rapidly expanding grades 3 and 4 astrocytomas (67% vs 10% incidence). Diagnosis was facilitated by symptoms of increased intracranial pressure associated with the first seizure in 79% of infratentorial vs 20% of supratentorial tumors. In a report of presenting features of brain tumors in 21 infants (Gordon GS et al. Arch Dis Child 1995;73:345-347), vomiting in 9 (43%) and large head in 16 (76%) were the most common findings. Meningitis was suspected initially in 5 (24%) infants with brain tumor, one with nuchal rigidity, and a CSF pleocytosis and elevated protein on premature spinal tap led to incorrect treatment with antibiotics and delay in CT scan and diagnosis.

POST-SURGICAL SPONTANEOUS REGRESSION OF RESIDUAL CEREBELLAR ASTROCYTOMA

Spontaneous regression of residual cerebellar astrocytoma following subtotal resection is reported in a 2-year-old boy followed for 11 years at the University of British Columbia, Canada. Tumor tissue left adjacent to the lateral wall of the fourth ventricle because of intraoperative bradycardia remained stable for 2 years. Between 2 and 7 years, repeated CT scans showed progressive decrease in size of the tumor, which became more cystic in appearance. The most recent scan at 11 years post-surgery showed complete regression of the tumor. Clinically, the child remained stable. The time course of regression is thought to rule out ischemic necrosis caused by surgery, and apoptosis is a more likely mechanism. (Steinbok P, Poskitt K, Henderson G. Spontaneous regression of cerebellar astrocytoma after subtotal resection. Childs Nerv Syst June 2006;22:572-576). (Respond: Dr P Steinbok, Division of Neurosurgery, Department of Surgery, University of British Columbia, Vancouver, BC, Canada).