CASE REPORT

TUBEROUS SCLEROSIS AND ACUTE HYDROCEPHALUS

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Tuberous sclerosis complex is a neurocutaneous and autosomal dominant disease characterized by multiple hamartomas in multiple viscera. It results from spontaneous mutation. The genetic anomaly is usually linked to the 9th chromosome. It may be accompanied by early childhood seizures, multiple brain tumors, skin lesions, angiomyolipomas in the kidneys and liver and rhabdomyomas. A careful physical examination, computerized tomography (CT) and magnetic resonance imaging (MRI) scans of the brain are essential in its diagnosis. In this study, we presented a 16 year old girl who was brought to our emergency service room due to acute loss of conscious and then underwent to ventriculo-peritoneal (V-P) shunt procedure after she had been diagnosed as acute hydrocephalus. The patient had been operated for intracranial mass when she was 2 years old and postoperative pathological diagnosis was established to be subependimal giant cell astrocytoma (SGCA).

Key words: Tuberous sclerosis, Subependimal giant cell astrocytoma

INTRODUCTION

Tuberous sclerosis complex (TSC) was first defined in 1862 by Von Recklinghausen. In 1880, Bourneville proposed the term “tuberous sclerosis” because of the relationship of the disease with mental retardation, epilepsy and cortical tubers. Its incidence has been estimated to be 1:10,000 to 1:70,000 in neonates. Its pathognomonic findings are angiofibroma on the face, cortical tuber, subependimal nodules or SGCA, multiple retinal astrocytoma and multiple subependimal nodules extending into the ventricle. SGCA develops from the growth of subependimal nodules. These tumors typically arise from terminal sulcus location adjacent to foramen monro. They usually lead to obstructive hydrocephalus and increased frequency of epileptic seizures. Cranial CT and MRI scanning are important in its diagnosis (1-3).

Our purpose in this case is to emphasize that SGCA in tuberosclerosis complex, especially due to involvement around the foramen magnum may constitute a potential pathology for acute hydrocephalus.

DISCUSSION

Tuberous sclerosis is a neurocutaneous syndrome characterized by hamartomas in a number of viscera. Takanashi et al. (1) proposed that half of the cases were autosomal dominant and the other half them were sporadic cases and suggested that the
genes TSC1 and TSC2 might be responsible for these cases. Hamartomas are benign lesions although they rarely may show malign progression. Subependimal giant cell astrocytoma is a slow-growing glial tumor (4). SGCAs are distinguished from other astrocytomas with their non-invasive nature and the fact that almost all of them locate in the ventricle (5,6). Spontaneous regression has not been observed in SGCAs. Curatolo et al. (3) reported that astrocytic astrocytoma regressed in a patient with Tuberosclerosis complex. Although rare, SGCAs may have such histological malignancy criteria as necrosis, mitosis and vascular proliferation. Kashiwagi et al. (2) reported a patient with SGCA only without any pathology in other systems.

Mortality in tuberous sclerosis is due to cardiac, renal and cerebral pathologies. Sudden deaths may be seen following cardiac arrhythmias, epilepsy, intracranial hemorrhages, obstructive hydrocephalus, aneurism rupture and spontaneous pneumothorax (7,8). SGCAs most commonly develop especially around the foramen magnum. Thus, they may lead to sudden deaths due to acute hydrocephalus (9). Hence, the patients with SGCA should be followed closely for hydrocephalus after the surgical procedure because of the importance of location of the tumor. The patient presented in this paper developed acute hydrocephalus due to growth of the mass 14 years after she had been operated.

In conclusion, in our opinion V-P shunt would be appropriate for hydrocephalus which may develop even though SGCT is not within surgical margins. We suggest that the patient should be followed closely or underwent V-P shunt for hydrocephalus if surgical treatment has been performed.

REFERENCE
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