Surgical Timing of the Subependymal Giant Cell Astrocytoma (SEGA) with the Patients of Tuberous Sclerosis Complex

Tüberus Sklerozis Kompleksli Subependimal Dev Hücreli Astrositomlu Hastalarda Cerrahi Zamanlama

ABSTRACT

AIM: Tuberous sclerosis complex has shown a wide variety of clinical, pathologic and radiologic manifestations. Many tumor types are found in tuberous sclerosis, which includes subependimal giant cell astrocytoma. The aim of this study is to focus on surgical timing of the tumor.

MATERIAL and METHODS: This study included 37 children with tuberous sclerosis presenting to Erciyes Univercity Medical School, whose hospital record were retrospectively evaluated between 1995 and 2010. Of the 5 patients had diagnosed with the subependimal giant cell astrocytoma and three patients were opereted on.

RESULTS: In the 27 of the patients had subependimal nodules (73%), cortical tubers were in the 19 patients (51.4%), giant cell astroitoma (SEGA) were in the 5 patients (13.5%). Mental retardation in different level was detected in the 18 patients (48.6%). The other clinical findings of the patients were angiomyolipomas (37.8%), hypomelanotic macules (91.9%), Convulsion (54.1%), adenoma sebaceum (32.4%), West syndrome (16.2%), shagreen patch (16.2%), intracardiac tumor (37.8%), subungual fibroma (2.7%), fibroadenom in the neck (2.7%).

CONCLUSION: A multidisciplinary approach is essential for an early, accurate diagnosis and proper management of affected individuals. The early surgical menagement for subependimal giant cell astrocytoma are recommended, and also periodic monitoring even for asymptomatic patients with subependymal nodules.

KEYWORDS: Tuberous sclerosis complex, Subependimal giant cell astrocytoma, Surgical timing

ÖZ


BULGULAR: 27 hastada (%73) subependimal nodul, 19 hastada (%51.4) kortikal tüber, 5 hastada (%13.5) dev hücreli astroitom vardır. 18 hastada (%48.6) farklı seviyelerde mental retardasyon tespit edildi. Hastaların diğer klinik bulguları; anjiyomyolipom (%37.8), hypomelanotik makul (%91.9), convulsion (%54.1), adenoma sebaseum (%32.4), west syndrome (%16.2), shagreen lekeleri (%16.2), intrakardıak tümör (%37.8), subungual fibroma (%2.7), boynuda fibro adenom (%2.7).

SONUÇ: Etki列enmiş bireylerin erken, doğru ve uygun tedavisi için multidisipliner yaklaşım esasdır. Subependimal dev hücreli astroitomada erken cerrahi önerilir ve de subependimal nodullü hastalar için peryodik izleme önerilir.

ANAHTAR SÖZCÜKLER: Tüberus sklerozis kompleks, Subependimal dev hücreli astroitom, Cerrahi zamanlama

Correspondence address: Mehmet Ali EkICI / E-mail: mehmetali.ekici@gmail.com

1Bozok University, Faculty of Medicine, Department of Neurosurgery, Yozgat, Turkey
2Erciyes University, Faculty of Medicine, Department of Pediatric Neurology, Kayseri, Turkey
3Osmanlı University, Faculty of Medicine, Department of Pediatric Neurology, Eskişehir, Turkey
4Erciyes University, Faculty of Medicine, Department of Pathology, Kayseri, Turkey
5Erciyes University, Faculty of Medicine, Department of Neurosurgery, Kayseri, Turkey

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INTRODUCTION

Tuberous sclerosis complex (TSC) is a multisystem genetic disorder that occurs in approximately 1 in 5,800 to 10,000 individuals (24,35). TSC may cause hamartomatous lesions of all organs but primarily affects tissues of the skin, central nervous system, kidneys, eyes, heart, and lungs. The disease is characterized by a broad physical and neuroimaging spectrum with epilepsy, mental retardation, renal dysfunction, and dermatologic abnormalities (26, 27, 32). The pathogenesis of TSC lies in the expression and function of the gene products, tuberin in TSC2 and hamartin in TSC1, in tissue (7, 21, 36).

The classic TSC central nervous system findings of cortical tubers, subependymal nodules (SENs), subependymal giant cell astrocytomas (SEGAs), and white-matter abnormalities are now easily identified by modern neuroimaging techniques (3, 4, 17, 19, 20). SEGAs are benign tumors that occur in approximately 10% of patients with TSC. They are associated with TSC and believed to originate from subependymal TS nodules. SEGAs are clinically significant because enlargement of these lesions may obstruct the flow of cerebrospinal fluid through the foramen of monro, causing obstructive hydrocephalus (5, 6, 13, 31).

We report 37 patients with tuberous sclerosis complex, which include 5 patients with subependymal giant cell astrocytomas. Of these patients with SEGA, three have been treated with surgical resection.

MATERIAL and METHODS

This study included 37 children with tuberous sclerosis presenting to Erciyes University Medical School during the 15-year period from 1995 to 2010. The Patients’ hospital records for this period were scanned retrospectively. The diagnosis of TSC was made according to the criteria given by Roach et al (Table I). The patients with the diagnosis of tuberous sclerosis were observed retrospectively. The study group was chosen from among the patients who had come to the pediatric neurology department with the complaints of epilepsy and mental motor retardation (MMR). MRI and CT demonstrated neorotubers, as well as allowing us to determine the involvement, if any, of other organs. For this purpose, renal, cardiac, ophthalmic and cutaneous scans were obtained. Controls and scannings were repeated every three months especially for the patients having the brain and renal involvement. In order to determine the patients’ intelligence level, Denver II development test for patients under, and WISC-R test for those above, 6 years of age, were performed. The data obtained were saved at each control. Because of the obstructive hydrocephaly development risk due to foramen of monro compression, MRI and CT were performed each year for the patients with SEGA. Of the 5 patients with the diagnosis of subependymal giant cell astrocytoma, three were operated on with the anterior transcicallosal approach.

Patient Reports

Patient 1

A 14-year-old boy presented with headache and hyperactivity and seizures. The patient had been diagnosed with tuberousclerosis earlier when he was only 5 months old. He had cortical tuber and subependymal giant cell astrocytoma (SEGA) in the brain and hypomelanotic macules on the skin (Table III). MR disclosed a mass in the left and right ventricular anterior horn at the level of the foramen of monro (Figure 1 A-D). The patient was operated on. The navigation was used intraoperatively for planning and during the surgery. Under general anesthesia and after Mayfield’s spiky heading binding, the position of the patient was matched using neuronavigation. Anterior interhemispheric transcicallosal approach was performed. The pericallosal artery was anomalous as azygos shape when the corpus callosum was reached. Subsequently callosotomy was performed, and passed over above the pericallosal artery and the right lateral ventricle was reached. The corpus and genu of the lateral ventricle were filled with the mass. The lesion was macroscopically consistent with astrocytomas, and firstly debulking was performed and then subtotal resection was done because the lesion had no favorable cleavage plane with adhesion to the head of the caudate nucleus. The lateral side of the mass was resected subtotally to control bleeding. Secondly, septum pellicidum was opened to reach the left lateral ventricle followed by the total resection of the mass that was moderately hemorrhagic and infiltrative into the neural tissue near the foramen monro. Although no acute hydrocephaly occurred in the early postoperative period, a burrhole was opened on the right side, 1cm away from the anterior of the coronal suture, to save time. The postoperative course was uneventful. Histological examination disclosed a highly cellular neoplasm with marked pleomorphism. In this case, the clustering of tumor cells and perivascular pseudopalisading was striking (x 100, H-E). Prominent giant cells and numerous atypical mitotic figures were found (Figure 2). The picture was consistent with SEGA.

Patient 2

A 15-year-old girl presented with headache and seizures and vomiting. The patient had been diagnosed with tuberousclerosis when she was 15 months old. She had cortical tuber and subependymal giant cell astrocytoma (SEGA) in the brain and hypomelanotic macules on the skin and renal angiomyolipoma (Table III). MR disclosed a mass in the right ventricular anterior horn at the level of the foramen monro (Figure 3 A-D). The patient had pupil edema on fundoscopy. The patient was operated on owing to the symptoms of increased intracranial pressure. Navigation was used intraoperatively for planning and during the surgery. Under general anesthesia, and after Mayfield’s spiky heading binding, the position of the patient was matched using neuronavigation. Anterior interhemispheric transcicallosal approach was performed. The pericallosal artery was passed over to reach the corpus callosum. Subsequently, callosotomy was performed and the right lateral ventricle was reached. The anterior and posterior margin of the tumor was found and debulked with ultrasonic surgical aspirator (CUSA). The tumor was resected completely and the foramen monro was opened. The lesion was macroscopically consistent with astrocytomas, and was

The postoperative course was uneventful. Histological examination disclosed a highly cellular neoplasm with marked pleomorphism. Reticulin was abundant in the stroma. Tumor cells demonstrated GFAP positivity (x100, Immunoperoxidase) (Figure 4). The picture was consistent with SEGA.

Patient 3

An 11-year-old girl presented with headache and seizures and vomiting. The patient was diagnosed with tuberous sclerosis when she was admitted the hospital. She had cortical tuber and subependymal giant cell astrocytoma (SEGA) in the brain and hypomelanotic macules on the skin of the lumbar and left knee and renal angiomyolipoma (Table III). MR disclosed a mass in the right ventricle anterior horn and at the level of the foramen of monro (Figure 5A-D). The patient had pupil edema on fundoscopy. She was operated on owing to the symptoms of elevated intracranial pressure. The navigation was used intraoperatively for planning and during the surgery. Under general anesthesia, after Mayfield’s spiky heading binding, the position of the patient was matched using neuronavigation. Anterior interhemispheric transcallosal approach was performed. The pericallosal artery were passed over and the corpus callosum was reached. Following callosotomy,
Table I: Diagnostic Criteria for Tuberous Sclerosis Complex (TSC)

<table>
<thead>
<tr>
<th>Major features</th>
<th>Minor features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Facial angiofibromas or forehead plaque</td>
<td>Dental pits (more than 14), randomly distributed</td>
</tr>
<tr>
<td>Nontraumatic ungual or peringual fibroma</td>
<td>Hamartomatous rectal polyps</td>
</tr>
<tr>
<td>Hypopigmented macules (more than 3)</td>
<td>Bone cysts</td>
</tr>
<tr>
<td>Shagreen patch (connective tissue nevus)</td>
<td>Cerebral white matter radial migration lines</td>
</tr>
<tr>
<td>Cortical tuber</td>
<td>Gingival fibromas</td>
</tr>
<tr>
<td>Subependymal nodule</td>
<td>Nonrenal hamartomas</td>
</tr>
<tr>
<td>Subependymal giant cell astrocytoma</td>
<td>Retinal achromic patch</td>
</tr>
<tr>
<td>Multible retinal nodular hamartomas</td>
<td>Confetti skin lesions</td>
</tr>
<tr>
<td>Cardiac rhabdomyoma, single or multible</td>
<td>Multible renal cysts</td>
</tr>
<tr>
<td>Lymphangiomyomatosis</td>
<td></td>
</tr>
<tr>
<td>Renal angiomyolipoma</td>
<td></td>
</tr>
</tbody>
</table>

**Definite TSC**

**Probable TSC**

**Possible TSC**

2 major features or 1 major + 2 minor features

1 major + 1 minor features

1 major features or ≥2 minor features

the right lateral ventricle was reached. The anterior and posterior margin of the tumor was found and debulked with ultrasonic surgical aspirator (CUSA). The tumor was resected completely and the foramen monro was opened. The lesion was macroscopically consistent with astrocytomas, and was moderately hemorrhagic and infiltrative to the neural tissue level near of the foramen of monro. The postoperative course was uneventful. Histological examination disclosed a highly cellular neoplasm with marked pleomorphism. Pleomorphic multinucleated eosinophilic tumor cells were found (x200, H-E) (Figure 6). The picture was consistent with SEGA.

Patient 5
A 9-year-old girl presented with seizures. The patient had been diagnosed with tuberous sclerosis when she was 14 months old. Because of the seizures MRI and computed tomography (CT) were performed, which revealed hyperintense left ventricular lesion and periventricular neurotubers (Figure 8 A-C). He had mental retardation and adenoma sebaceum, and Shagreen patch, hypomelanotic macules on the skin besides the MRI and CT findings (Table III). The patient had no sign and symptom of increased intracranial pressure on the examination. Thus, She was diagnosed with tuberosclerosis were given was treated with two antiepileptic drugs (vigabatrin and sodium valproate). Since the patient has had no complaints of the seizures and increased intracranial pressure symptoms so far, she has not been operated.

Patient 4
A 16-year-old girl presented with seizures. The patient had been diagnosed with tuberous sclerosis when she was 3 months old. MRI and computed tomography (CT), performed for seizures, revealed hyperintense right ventricular lesions and periventricular neurotubers (Figure 7A-C). She had mental retardation and adenoma sebaceum, and a single intracardiac tumor, hypomelanotic macules on the skin besides the MRI and CT findings (Table III). Having no signs and symptoms of increased intracranial pressure symptoms on the examination, the patient was diagnosed with tuberosclerosis and treated with two antiepileptic drugs (vigabatrin and sodium valproate). Since the patient has had no complaints

Table II: Tuberosclerosis Clinical Findings of the 37 Patietns

<table>
<thead>
<tr>
<th>Features</th>
<th>N=37</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age</td>
<td>63.6 month</td>
<td></td>
</tr>
<tr>
<td>Gender</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>16</td>
<td>43.2</td>
</tr>
<tr>
<td>Female</td>
<td>21</td>
<td>56.8</td>
</tr>
<tr>
<td>Subependimal nodules</td>
<td>27</td>
<td>73</td>
</tr>
<tr>
<td>SEGA</td>
<td>5</td>
<td>13.5</td>
</tr>
<tr>
<td>Convulsion</td>
<td>20</td>
<td>54.1</td>
</tr>
<tr>
<td>Mental retardation</td>
<td>18</td>
<td>48.6</td>
</tr>
<tr>
<td>West syndroma</td>
<td>6</td>
<td>16.22</td>
</tr>
<tr>
<td>hypomelanotic macules</td>
<td>34</td>
<td>91.9</td>
</tr>
<tr>
<td>adenoma sebaceum</td>
<td>12</td>
<td>32.4</td>
</tr>
<tr>
<td>“shagreen” patch</td>
<td>6</td>
<td>16.2</td>
</tr>
<tr>
<td>subungual fibroma</td>
<td>1</td>
<td>2.7</td>
</tr>
<tr>
<td>fibroadenom in the neck</td>
<td>1</td>
<td>2.7</td>
</tr>
<tr>
<td>Angiomyolipomas</td>
<td>14</td>
<td>37.8</td>
</tr>
<tr>
<td>Polycyst</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Simple cysts</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Renal-cell carcinoma</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>intracardiac tumor</td>
<td>14</td>
<td>37.8</td>
</tr>
<tr>
<td>Retinal hamartomas</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Bone cyst</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
Neurological findings:

Our patients were admitted to hospital primarily because of convulsion. There were 20 (54.1%) patients with convulsion in our study. Subependymal nodule was the most frequent finding in cranial imaging. Subependimal nodules were detected in 27 (73%), cortical tubers in 19 (51, 4%), and giant cell astrocytomas in 5 (13, 5%) patients. Mental retardation at varying degrees was detected in 18 of the patients (48.6).

West syndrome was in 6 patients (16.2%). There were 3 female and 2 male patients with SEGA. The age range at the time of surgery was 10–18 years, with a mean of 15 years. Four children who did not show any progressive neurological deficit related to the intraventricular lesion were admitted because sequential neuroimaging studies revealed the growth of “subependymal nodules” into the intraventricular tumours, and these four patients with epileptic seizures resistant to a combination of anticonvulsant agents, were diagnosed as having TS by pathognomonic features. In the remaining one patient, headache was the first and only sign of a CNS disease. This patient was admitted because of the symptoms of increased intracranial pressure, such as progressive headaches, nausea, vomiting.

Cutaneous finding:

Of the patients, 34 (91, 9%) had hypomelanotic macules in our study. There were hypopigmented lesions on the skin of 34 patients (91.9%). In 12 of the patients (32, 4%) adenoma sebaceum, in 6 of the patients (16,2%) “shagreen” patch, in one of the patients (2.7%) subungual fibroma and in one of them (2.7%) fibroadenoma in the neck were detected.

Cardiac, renal and eye findings

Cardiological assessment revealed an intracardiac mass in 14 of the patients (37.8%). Renal USG of these patients demonstrated angiomyolipoma. The ophthalmic and lung examination revealed no pathological finding.
DISCUSSION

The first reason for morbidity and mortality is neurological complications and the second one is renal involvement. The neurological complications of TSC are the most common and often the most impairing aspect of this disease. In brain tissue, mutations in the TSC1 and TSC2 genes cause abnormal neuronal and glial differentiation and proliferation as well as abnormal cellular migration. TSC derives its name from cortical tubers, so called for their gross pathologic “potato-like” appearance (9). They occur in as many as 95% of patients (3, 17). Tubers most commonly involve surface cortical and
Subcortical areas but may also be seen in the cerebellum. Their architecture expands the gyri and disrupts the gray-white matter junction. Tumors vary in size from millimeters to centimeters and are frequently multiple (4, 10, 17, 19, 27).

Subependymal nodules are also seen in the majority of TSC patients. They can be identified early in infancy and increase in number through the first decade of life. Histologically, they are hamartomatous lesions formed from dysplastic astrocytic and neuronal cells. Radiographically, SENs appear as small protrusions into the walls of the lateral ventricles, giving the pathologic appearance of “candle guttering.” SENs are most commonly found around the area of the foramen Monro. They tend to calcify with time and for this reason are often more easily identified on CT imaging (1, 4, 16, 17, 27). Literature proposes that subependymal nodules a turn into tumor as a risk factor when they are greater than 5mm, with incomplete calcification and contrast enhancement. Therefore, if the nodules are located near the foramen monro, MRI should be performed every year up to 10 years of age (22).

Research has shown that tubers and SEGAs express similar neuroglial markers and thus likely to originate from a defect in progenitor cell differentiation during brain development. SEGAs are clinically significant because enlargement of these lesions may obstruct the flow of cerebrospinal fluid at the foramen of Monro, causing obstructive hydrocephalus. These lesions are benign slow-growing grade I tumors containing giant cells with morphologic features of glia. Although traditionally classified as an astrocytic tumor, many SEGAs also exhibit a mixed glioneuronal phenotype and express both neuronal as well as astrocytic markers. In addition, SEGAs may develop directly from a subependymal nodule or possibly originate de novo from an unidentified cell type. Surgical resection of the lesion has been accepted as the treatment of choice but the timing of surgery remains debatable. Some surgeons advocate early surgery, while others offer waiting for symptomatic lesions or hydrocephalus. In children, 110 cases of subependymal giant cell astrocytomas have been reported in the literature. They were usually discovered after 10 years, at a mean age of 13 years (5,9,16,17,20,22). SEGAs were detected in five of the patients. The age range at the time of surgery was 10–18 years, with a mean of 15 years in our patients.

In this study, 3 patients were operated on in elective conditions. In one case there were signs of high intracranial pressure on fundoscopic examination for pupil edema, but no signs of transformonial herniation. In another case the patients’ parents did not consent to the surgery and the other patient had no high intracranial pressure sign on fundoscopy and no tumor growing on the MRI.

According to literature, SEGAs are seen by 2-14% (22, 29, 30). The nature of the SEGAs is benign and slowly growing but they can lead to sudden deterioration in the patient’s clinic condition due to foramen monro obstruction related with their localization.

Causes of foramen monro obstruction are described as intratumoral bleeding and tumor volume increasing. Foramen of monro obstruction results in high intracranial pressure. It is reported that if the characteristics of the nodule have a noncalcification, contrast enhancement and near localization to the foramen of monro, these characteristics might be described as high risk factor for tumor transformation (22).

When the patients underwent emergency surgery, mortality rate reached approximately 10% and when the tumoral resection is done in selective conditions, patients outcome were improved (23).

Ribapiere et al (8) reported that if the radiological characteristics of the SEGA have an incomplete or noncalcification, with a location near the foramen of monro and a size greater than 5mm, the lesions should be excised as early as possible. Their numbers were larger or symptomatic lesions tend to have a higher morbidity.

Patients with SEGAs might be able to present different neurological symptoms to the clinic, which include not only increased ICP symptoms, epilepsy, behavioral modification but also acute hydrocephalus due to foramen of monro obstruction or intraventricular hemorrhage (2,12).

Moncef Berhouma (2) reports that if there is evidence of growth on the successive MRI, tumors should be removed early as soon as possible.

According to the literature, early surgical approach is advised (2,6,13,33).

<table>
<thead>
<tr>
<th>Features</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
<th>Case 4</th>
<th>Case 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Giant cell astrocytoma</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Cortical tuber</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Hypomelanotic macules</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Adenoma sebaceum</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Renal Angiomyolipomas</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Cardiac rhabdomyoma</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Shagreen patch</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Subependymal nodule</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>
In this study, the tumor in patient 1 was resected subtotally because of the adhesion to the head of the caudate nucleus. It is reported that large tumors have had some drawbacks during surgery due to the deformation of the foramen of monro like difficulties in operative hemostasis and dissection from surrounding structures (2).

SEGA is diagnosed in adolescents as a part of the tuberous sclerosis (TSC) complex, but it can be seen without any associated stigmata of TSC. Surgery is the preferred treatment and should be done as soon as SEGA is diagnosed, because early diagnosis and treatment can reduce surgical morbidity and risk of tumor recurrence. MRI should be performed at the time of diagnosis of tuberous sclerosis when there are nodules located near the foramen of Monro and if they measure >5 mm in diameter, are not or incompletely calcified, and are enhanced by gadolinium, MRI should be repeated every year until the age of 15 years. (1,3,5,9,11,17,22,27,31,32,34). In our study, subependymal nodule was the most frequent symptom detected in the cranial imaging. In 27 of the patients (73%) subependymal nodules, in 19 of them (51,4%) cortical tubers, in 5 of them (13,5%) giant cell astrocytoma were detected. Four children who did not show any progressive neurological deficit related to the intraventricular lesion were admitted because sequential neuroimaging studies revealed the growth of ‘subependymal nodules’ into the intraventricular area, and these four patients who had epileptic seizures resistant to a combination of anticonvulsant agents, were diagnosed as having TS by pathognomonic features. In the remaining one patient, headache was the first and only sign of a CNS disease.

More than 80% of patients experience seizures, and 50 to 80% have mild to severe cognitive impairment. Sudden deaths in TS patients have been described. Seizures are the presenting symptom in 92% of individuals with TSC, making epilepsy a significant cause of morbidity associated with this disease. Seizure onset usually occurs during infancy or early childhood but may happen at any age (18,27,28). There was no mortality in our series. Twenty patients (54%) were admitted to hospital with the complaints of having convulsion in our study. Tuberous sclerosis was detected in the etiology of 6 of our patients being followed with the diagnosis of west syndrome.

Cognitive dysfunction has long been recognized as a common neurological complication of TSC. It was previously assumed that mental retardation was inevitable, but more recent studies have shown that many affected individuals may have normal intelligence. Cognitive abnormalities have been found in about 50% to 65% of patients in earlier population-based studies, but the results in these studies were not based on standardized psychometric evaluations (15,27). In our study, mental retardation was detected by 48.6%, a rate that was in various levels (mild 19%, medium 22%, serious 8%). This rate is similar to that which is in the literature.

The renal manifestations of TSC are found in a majority of patients. They include angiomyolipomas (AMLs), simple cysts, polycystic kidney disease, and renal-cell carcinoma. AMLs are found in as many as 80% of patients with TSC, but are typically asymptomatic (14,26,27). After the renal USG, performed to the patients, angiomyolipoma was detected in the 14 of the patients (37,8%).

The cutaneous manifestations in TSC are common, and are typically the first clue to the diagnosis. They include hypomelanotic macules, facial angiofibroma, forehead plaques, shagreen patches, and ungual fibromas. Hypomelanotic macules or ash leaf spots occur in 61% to 97.2% of individuals with TSC. Facial angiofibromas (adenoma sebaceum) are pathognomonic for TSC and are seen in over 70% of patients. They first appear during the preschool years and with time develop into discrete, papular, pink, or erythematous lesions in a symmetrical malar distribution on the face (26, 27).

Cardiac rhabdomyomas are the most common intracardiac tumor of infancy and childhood, and are highly associated with TSC. They are found in 30% to 50% of affected individuals (25-28). In this study, after the cardiological assessment intracardiac mass has been detected in the 14 of the patients (37,8%).

In conclusion, a multidisciplinary approach is essential for an early, accurate diagnosis and proper management of affected individuals. In the light of the literature, we concluded that early and planned surgery was suggested although we operated on the patients in chosen conditions.

REFERENCES


