AGENESIS AND LIPOMA OF CORPUS CALLOSUM

CASE REPORT


SUMMARY - The agenesis and lipoma of the corpus callosum is a very rare association. We report the case of a 18-years old woman with rare epileptic seizures since the age of 6 years, normal neurological examination, as well as normal electroencephalogram. The brain computed tomography scanning and the magnetic resonance showed the lipoma and the agenesis of the corpus callosum.

KEY WORDS: corpus callosum, lipoma, agenesis.

Corpus callosal lipoma (CCL) is a very rare benign tumor of questionable etiology. Most of the reported cases have been found at postmortem examination. Its incidence is around 0.06% and is often associated with other malformations, most commonly partial or complete callosum agenesis (CA), which occurs in as many as 50% of CCL patients. Since the first report in 1856 by Rokitansky, an extensive review of the literature revealed a total of less than 85 reported cases of CCL with CA. We describe one case of this unusual association.

REPORT OF CASE

JR, a 18-year-old woman was referred for investigation of epileptic seizures. She had a history of tonic-clonic generalized seizures since 6 years of age. Their seizures lasts no longer than a few minutes, occurred up to 4 per year, mainly during the sleep and complaint of headache as the only postictal change. Pregnancy and delivery were normal, as was the child's early development. The antenatal history had been unremarkable. Physical and neurological examination were normal. Phenobarbital monotherapy was started with 100 mg/day. A conventional EEG performed for 30 min was normal during the awake and sleep state. CT as well as the MR showed the lipoma and the agenesis of CC. The CT scans were characterized by regular low density image filling the place of the CC (lipoma) an bordered on each side by two shell-like calcifications. The MR was carried out with T1 and T2 weighted images and showed the following findings: (a) indirect signs of callosal
agenesis: dilatation of the trigone of lateral ventricles and ascension of the third ventricle; (b) partial (nearly total) agenesis of the body and splenium of the CC; (c) a slice of adipose tissue over the region of the genu and the rostrum that extended as a voluminous mass (lipoma) into the ventricles with high intensity sign at T1 (Fig 1a, b, and c). During the follow-up of 4 years the EEG remained normal and the CT and MR images showed the same findings without new changes. The patient is still under treatment with phenobarbital 100 mg per day and presents rare epileptic seizures.

COMMENTS

The rarity of intracranial lipomas is indicated by the fact that in a series occurred only four times in 5000 postmortem examinations\(^3\). The first case discovered at necropsy occurred after 23813 consecutive necropsies\(^2\). In a CT department only four lipomas of the CC were found in nearly 13000 patients, a frequency of 0.03%\(^7\). The commonest associated anomaly is agenesis of the CC, as in our patient, occurring in 48% of cases with this type of tumor\(^24,34\), and may affect part or all of this structure\(^25\). Other deviations include agenesis of the cerebellar vermis, cleft lip, funnel chest, cardiac ventricular septal defect\(^34\), spina bifida and myelomeningocele\(^25\), and cortical heterotopia\(^10\). The lipoma of the CC is a congenital\(^14\), hamartomatous condition\(^4\), probably due to meningeal maldifferentiation, and the agenesis has been regarded as an example of cerebral dysraphism, the term dysraphism referring to the midline defects caused by imperfect closure of neural tube\(^1,2,27,28\).

The commonest presentation of the CCL is epilepsy, either partial motor or complex partial seizures\(^7\). Some of these seizures may evolve to secondary generalization and might terminate in generalized tonic-clonic seizures. In our patient the generalized convulsive seizures could have been secondarily generalized seizure in which the partial beginning was not appreciated. The origin of the epilepsy in this condition is not clear. One theory advanced is that the invasion of cerebral cortex by the collagenous capsule sets up a focus or because of the interhemispheric disconnection\(^7,34\). Other nonspecific symptoms have been described as headache, paresthesias, vomiting, vertigo and emotional lability\(^16,18,21,23,30\). On the other hand, approximately 50% of cases are asymptomatic\(^3,32\).

CT and MR are the diagnostic procedures of choice\(^1,2,6,9,21,27\). The CT appearance of lipomas of the CC was first described by New and Scott in 1975\(^22\) and the diagnosis depends on the characteristic low attenuation values which are characteristic of fat tissue only. When calcification occurs, it is curvilinear and mural\(^8\). On contrast-enhanced CT, the tumor does not show any change in density\(^13\). MR provides highly visible images of the CC and is the procedure of choice in the evaluation of this structure\(^6,23\). In our patient the diagnosis of partial agenesis (nearly total) of CC was established with the MR images (Fig 1a, b, and c). The differential diagnosis must be made with

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Fig 1. MR evaluation with T1 weighted images demonstrates the agenesis and the lipoma of corpus callosum. Sagittal (a), coronal (b), and axial MR images shows a high-intensity signal (white) of lipoma circumscribing the region of corpus callosum and extending into the ventricles.
dermoid cyst, teratoma, craniopharyngioma, and epidermoid tumors\textsuperscript{17,19,30}. The cystic component of craniopharyngiomas may be fat dense but its complex appearance and bizarre calcifications are usually suggestive of this tumor\textsuperscript{26}. Teratomas and dermoid cysts are frequently inhomogenous because they contain tissues other than fat\textsuperscript{13,33}. Epidermoid tumors are generally located within the cerebellopontine angle, parapituitary region or in the fourth ventricle and have CT appearance that usually fall in the range of cerebrospinal fluid density\textsuperscript{20}.

Surgery has little place in the management of this condition\textsuperscript{13}. The necessity for removal of the CCL must be seriously questioned, since it is a non-malignant, non-progressive lesion, and some patients live to old age, trouble-free, untreated. Firm adhesion between the collagenous capsule and brain together with the high vascularity of the tumour add to the surgical hazard that may end with death\textsuperscript{25,30,34}. Laser techniques may make these lipomas more amendable to surgery\textsuperscript{4}. Surgery has been performed in the face of a rapidly growing tumor, signs of increased intracranial pressure, progressive neurologic signs, or severe uncontrolled seizures\textsuperscript{11,15}. Even when surgery is performed, a patient's epilepsy will unlikely be resolved\textsuperscript{9}. Tahmouresie et al.\textsuperscript{30} reported that of 21 surgical patients, 10 (47.6\%) died during the postoperative period, 4 were no better, 1 was worse, and 5 (23.7\%) improved. Our patient had a history of 12 years of rare seizures, was under control with anticonvulsant, had not progression of their symptoms and the CT and MR images did not change during the follow-up.

**REFERENCES**