Cervical lipomyelomeningocele: case Illustration

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Summary

Cervical lipomyelomeningocele is a rare congenital spinal pathology. Lipomyelomeningocele is the commonest cause of congenital tethering, which causes neurological deterioration due to the conus medullaris and root ischemia. Early intervention is recommended even in cases with normal neurological examinations in order to prevent deterioration but our patient with cervical lipomyelomeningocele had a normal neurological examination despite his age (22 year-old) and had no urodynamic dysfunction.

KEY WORDS: Lipomyelomeningocele. Cervical. Occult spinal dysraphism.

Lipomielomeningoceles cervical. Caso clínico

Resumen

El lipomielomeningocele cervical es una patología raquídea congénita rara. El lipomielomeningocele es la causa más frecuente de anclaje medular, que da lugar a deterioro neurológico, debido a isquemia del cono medular y de las raíces. Se recomienda la intervención precoz, incluso en casos con examen neurológico normal, con el fin de prevenir un deterioro, pero nuestro paciente con lipomielomeningocele cervical tenía un examen neurológico normal, a pesar de su edad (22 años) y no tenía ninguna disfunción urinaria.


Introduction

Lipomyelomeningocele is a type of congenital occult spinal dysraphism consisting of the presence of lipomatous tissue attached to the dorsal spinal cord, which protrudes though a spinal defect along with the menings or spinal cord to form a posterior mass under the skin, usually in the lumbosacral region. Neural ectoderm separates from the cutaneous ectoderm and periaxial mesoderm comes in intact with the unfused ventral neural ectoderm. The mesoderm then differentiates into fatty tissue, thus preventing the neural canal and the posterior aspect of the spine from fusing4,5.

Lipomyelomeningocele is the commonest cause of congenital tethering and causes neurological deterioration due to the conus medullaris and root ischemia1.

Here we presented a 22 year-old patient harboring a lipomyelomeningocel in the cervical region without any neurological deficit and tethered cord.

Case

A 22 year-old man suffering from a posterior mass in the cervical region was admitted to our out-patient clinic (Figure 1). Neurological examination of the patient was unremarkable and he had no urinary dysfunction. Plain x-ray films of the cervical vertebrae revealed C5-6 posterior fusion defect (Figure 2). The magnetic resonance imaging (MRI) of the cervical spine revealed a lipomyelomeningocele at C6 level (Figure 3).

The patient was operated under general anesthesia in prone position and the lipomyelomeningocele was corrected without any complication. Postoperative neurological examination of the patient was unremarkable.

Discussion

The lipomyelomeningocele rate has been estimated to be 2,5 per 10000 births3. Their occurrence in the cervical region is even rarer4. The defect is more commonly found in females2. An autosomal inheritance is also suggested for lipomyelomeningocele3.

Cervical and upper thoracic myelomeningoceles account for only 1-5% of all spinal dysraphism and lipomyelomeningocele as an additional congenital spinal pathology is very rare6. Although lipomyelomeningocele
Figure 1. Our patient with cervical lipomyelomeningocele.

Figure 2. Cervical plain x-ray film showing C5 and C6 posterior fusion defect.

Figure 3. MRI revealing lipomyelomeningocele at C6 level.
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is the commonest cause of tethering, our patient was not harboring a tethered cord and had no urologic dysfunction\(^1\). Unless other significant spinal anomalies exist, such as a tethered cord, the risk of voiding disorders and upper tract deterioration is minimal\(^7\). Although early intervention is recommended even in cases with normal neurological examinations in order to prevent deterioration\(^8\), our patient was 22 years old and still had a normal neurological examination probably due to the lack of an additional significant spinal pathology.

It has been 5 years since we have operated the patient, and we are still following the patient's urodynamic functions because deterioration can occur up to 8 years postoperatively\(^8\).

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References


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References


Comentario al trabajo Cervical lipomyelomeningocele: case Illustration de Gürkanlar and col.

We read with great interest the article by Gurkanlar and al, entitled ‘Cervical lipomyelomeningocele’. The author describes a fatty mass starting under the skin of the child’s back, in the cervical midline line. The mass extended inward to the spinal canal and was covered by skin and visible from outside. The two biggest problems lipomyelomeninocoeles have in children are: The spinal cord is stuck (fixed) to the fatty mass and the fatty mass puts pressure on the spinal cord. The defect happens early in the mother’s pregnancy, (about the fourth to sixth week) and has no known cause. Unlike a mielomeningocele, it does not have a genetic component nor it is due to a lack of folic acid during pregnancy. The authors describe very well a rare cause of cervical lipomyelomeningocele.

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