Computed Tomography of a False Postoperative Meningocele

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False meningoceles following laminectomy have been reported; however, calcified postoperative meningoceles have not been demonstrated by computed tomography (CT). These cystic lesions may be considered as postoperative false (or spurious) meningoceles. The development of these changes occurs because a collection of cerebrospinal fluid (CSF) accumulates in the soft tissue through a rent in the dura. The surrounding soft tissues can absorb only a finite amount of fluid; then a connective tissue reaction occurs that wall off the CSF. In rare instances, as seen in our case, the connective tissue can calcify, thus forming a false postoperative meningocele.

Case Report

A 26-year-old man originally was seen after 3 months of low back pain that radiated down the left leg. Myelography demonstrated an intradural defect (fig. 1A) at the L5 level with displacement of the nerve roots of the cauda equina to the right side. There was also a large extradural component of the mass on the left side that displaced the entire lumbar cul-de-sac to the contralateral side. Close examination revealed that there were also additional focal areas of widening of the individual nerve roots of the cauda equina.

The patient was operated through a wide laminectomy at the L4–L5 level and a plexiform neurofibroma was removed. At the operative site, after removal of the neurofibroma, a small posterior hole in the dura was packed with Gelfoam. A dural patch was not put in place at the operative site.

The patient did well for several months and then returned with increasing low back pain. At this time the pain did not radiate down either leg. Myelography revealed a complete block to the flow of contrast material at the L4 level. The block suggested an extradural lesion that dissected superiorly along the left lateral aspect of the canal at the L3 level, displacing the contrast column to the right (fig. 1B). A lateral radiograph (fig. 1C) and anteroposterior tomogram (fig. 1D) revealed pressure erosion of the spinous process of L3 and an expanding lesion with a calcified rim that extended from the L4 level inferiorly to the level of the sacrum.

The calcified cystic lesion, not visible on the myelogram, was readily apparent on the tomograms, while the eggshell-thin calcified rim could be seen on the lateral radiograph projecting posteriorly into the soft tissues (fig. 1D). These studies were followed by a CT scan of the lumbar region, which demonstrated that water-soluble contrast material, not previously visible, had actually entered into the cystic, calcified, postoperative meningocele (fig. 1E). This false meningocele extended posteriorly from the operative site into the soft tissues of the back. The calcified rim was smooth and well defined and extended posteriorly to just below the skin of the back. The contrast material readily identified on the CT scan could not be appreciated on the initial radiographic studies, even in retrospect.

The patient was reoperated, and, after the calcified roof of the CSF-filled cyst was removed, a 1 cm opening in the dura was visualized and CSF could be identified leaking through this rent in the dura. The calcified, cystic, false meningocele was totally removed and the dural rent was closed with a dural patch. The patient had an uneventful postoperative course and became pain-free.

Discussion

Our case is an example of a postoperative false meningocele that calcified in its peripheral margin. A remarkably similar case of a calcified postoperative meningocele was reported by Rosenblum and DeRow [1] in 1963. We believe that this is the first time this entity has been identified by CT.

A noncalcified, postoperative, false meningocele was reported in 1946 by Hyndman and Gerber [2], and additional reports followed. However, this postsurgical complication appears to be uncommon. Previous reports [1–13] all emphasized that the etiology of these lesions is an opening in the dura that allows CSF to leak into the soft tissues. It appears that these false lumbar meningoceles occur when the dura is inadequately closed after intentional or inadvertent opening of the dura at surgery.

When CSF accumulates in the soft tissues, only a limited amount can be absorbed. When this point is reached a fibrous reactive membrane forms that walls off the collection of CSF. Eventually this nonabsorbing fibrous wall may become calcified, as happened in our case. The interval between operation and onset of symptoms is variable.

Congenital cysts have also been reported, and they may or may not be lined with an arachnoid membrane. They are

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AJNR 5:326–328, May/June 1984 0195–6108/84/0503–0326 $00.00 © American Roentgen Ray Society
usually found in the thoracic region and are thought to be secondary to impaired venous drainage from the vertebral bodies in the presence of kyphosis [2]. Other acquired cysts may develop secondary to trauma in which there is a laceration in the dura. One case of an acquired arachnoid cyst was thought to be secondary to a rent in the dura after severe paroxysm of coughing with herniation of the arachnoid through the dural tear [11].

The older literature reports that these cysts may or may not fill with Pantopaque and may even require placing the patient supine for demonstration. These cysts are demonstrated to better advantage with water-soluble contrast material, and the use of Pantopaque is strongly discouraged. Myelography and even tomography after myelography may not reveal contrast material within the cyst; however, postmyelography CT readily demonstrates the accumulation of even very low concentrations of contrast material, as seen in our case. It is postulated that, with the added technique of postmyelography CT, contrast material will probably be demonstrated in virtually all cases.

Although postoperative false meningoceles have been reported, they are probably rare. However, if symptoms do recur in an individual patient, an acquired cyst should be considered in the differential diagnosis. With surgical removal
of these acquired abnormalities and repair of the dural laceration, the prognosis is good because the lesion is benign. With metrizamide myelography followed by spinal CT, this diagnosis is readily confirmed and the anatomy can be demonstrated to excellent advantage.

REFERENCES