Retrorectal hamartoma: A ‘tail’ of two cysts!

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Introduction

Retrorectal hamartomas or tail gut cysts are rare congenital anomalies arising from the vestiges of the embryonic hindgut. They are more common in middle-aged women[1] and usually present as cysts in the retrorectal space.[2] We report the CT scan and MRI appearances, respectively, of two cases with tail gut cysts - one presenting shortly after birth and the other, a 72-year-old man with symptoms for one week. In both, the tail gut cysts were in a right perirectal location. Presentation in such a location in males, at extremes of age, is unusual for tail gut cysts.

Case 1

A 19-month-old male child presented to the pediatric outpatient department for evaluation of a large swelling in the perianal region, which was present since birth and had gradually increased in size. An earlier attempt at aspiration at one year of age revealed that it contained brownish fluid. An initial reduction in size of the mass following the aspiration was followed by gradual increase. On examination, the mass was compressible and nontender. CT scan showed a homogenous, well-defined, cystic lesion in the right perirectal/perianal region, displacing the rectum to the left [Figure 1A]. No solid component, calcification, or postcontrast enhancement was noted in the mass. A small peripheral cyst was also noted [Figure 1B]. No obvious communication with the adjacent bowel was noted.

The mass was excised in toto. Histopathology revealed a cystic lesion lined by cuboidal and columnar epithelium supported by fibroconnective tissue and skeletal muscle fibers, consistent with a cystic hamartoma.

Case 2

A 72-year-old male presented with urinary incontinence for a week. On rectal examination, there was mild tenderness and evidence of extrinsic compression of the rectum and anal canal from the 7 o’clock to 12 o’clock positions, without any mucosal involvement. Pelvic MRI showed a large ovoid, well-defined cystic lesion in close apposition to the right rectal wall, displacing the rectum to the left. Intrallesional hyperintensity on both T1- and T2-weighted images, with layering suggestive of high protein or hemorrhagic contents was noted [Figure 2]. The patient underwent laparotomy and excision of the pelvic cyst. Intraoperatively, a 10 × 15 cm globular cystic mass was seen in the right perirectal region, containing about 800 ml of a dark greenish fluid. The cyst was excised and the patient had an uneventful recovery. Histopathology showed dense acute and chronic inflammatory infiltrate, with extensive hemorrhages and hemosiderin-laden macrophages and occasional epithelioid and foreign body-type giant cells; the findings were suggestive of an infected tail gut cyst.

Discussion

Tail gut cysts are rare developmental cysts which, along with duplication cysts, are categorized under enteric cysts.[1]
The tailgut is the distal-most portion of the hindgut and lies beyond the future anus. The anus develops by around 56 days and following this the tailgut undergoes involution. Failure of regression results in persistent vestiges, which may form into tailgut cysts. The most common location is the presacral retrorectal space, with anterior rectal displacement. The tailgut cyst in our first case appeared to be in a right perirectal location, displacing the rectum and anal canal to the left. In our second case, the cyst was seen to extend from a retrorectal location to the right of the rectum, again with left rectal displacement.

Presentation is usually as an asymptomatic retrorectal mass or with abdominal pain or constipation. Some patients may present with recurrent retrorectal abscesses or following multiple procedures for anal fistulae. On gross appearance they are unilocular or multilocular cystic lesions with mucoid contents. Microscopy shows a variety of epithelia without the presence of villi and crypts. An incomplete muscle layer is often associated with the cyst wall. Infected cysts show cyst wall fibrosis and breakdown of the epithelial lining. Rarely, malignant transformation may occur in the cyst.

Assessment with USG is of limited value, showing only a multilocular cyst with debris and gelatinous material seen as internal echoes. On CT scan, tailgut cysts appear as a discrete, well-margined mass of water or soft-tissue density as seen in our first patient. Calcification of the thin wall may rarely be seen. In case of infection or, rarely, malignant transformation, loss of discrete margins and involvement of adjacent structures may be seen. While histopathology suggested infection of the cyst in our second case, no obvious findings were appreciated on imaging.
A contrast study may have helped clarify the possibility of infection within the cyst. Though typically hypo intense on T1W and hyper intense on T2W images, a heterogeneous appearance on MRI may be seen owing to mucin, proteinaceous material, or hemorrhage within the cyst. T1 and T2 hyperintensity was noted in our second case, which could be attributed to the infection and the extensive hemorrhage found on histopathology. A multilocular appearance with internal septae on T2W images has been described by Kim et al., as being specific for tailgut cysts. However, neither of our cases showed the presence of septae. The presence of a small peripheral cyst accompanying the larger cyst has also been described by Yang et al. A similar finding was identified in the CT scan of our first case (the child). A mass effect in large lesions has been described, with displacement of the adjacent rectum, as was observed in both our patients.

Most previous reports on tailgut cysts have described them as being more common in middle-aged women; their occurrence in infants is rare, with one report of in a two-year-old girl. Both of our cases, however, occurred in males, one at each extreme of age, thus showing that tailgut cysts may occur over a wide age range. Also, unlike the common retrorectal location highlighted in previous reports, a predominantly right perirectal location was observed in both our patients.

We thus highlight the importance of considering tailgut cyst as a possible differential in any case of perirectal cyst, irrespective of age and sex, and the importance of imaging in assessing the characteristics and extent prior to surgery.

References


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