A rare pediatric case with radiological findings: pelvic cystic schwannoma

**ABSTRACT**

Schwannomas are peripheral nerve sheath tumors usually detected in adults which are extremely rare in pediatric population and when present they are commonly associated with Neurofibromatosis type 2. While frequently observed in the head, neck, and extremities, they could be detected anywhere in the body including abdominal cavity. The most common site for intraabdominal schwannomas is stomach and pelvic schwannomas are extremely rare. The imaging characteristics are quite diverse, and they could seldom be pure cystic. Herein, we describe a case in the pediatric age group diagnosed with pelvic cystic schwannoma.

**Keywords:** Schwannoma, intraabdominal, pelvic, pediatric radiology.

A 14-year-old male who was otherwise healthy admitted with non-specific abdominal pain. Abdominal ultrasound revealed a round mass with the size of 35 x 29 x 21 mm located adjacent to the right lateral wall of the bladder. The mass had smooth borders, was heterogeneous with hyperechoic areas centrally and hypo echogenicity peripherally, and there was no blood flow on color Doppler examination (Figure-1a-b). For further evaluation magnetic resonance imaging (MRI) was acquired; a hyper intense complex cystic lesion with hypo intense areas on T2-weighted (w) image (Figure-1c) and no enhancement with intravenous gadolinium-based contrast medium administration was detected (Figure-1d). There was also suspicious diffusion restriction within the lesion. Patient had no signs of neurofibromatosis (NF). In the robotic surgery, the mass was found on the right side of the bladder and near the internal iliac and obturator artery posteriorly, and total excision of the lesion was performed. The lesion was diagnosed as cystic schwannoma on further pathological examinations.

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Schwannomas are usually detected in adults between the 2nd-5th decades. They are extremely rare in the pediatric population and commonly associated with NF type 2. Schwannoma, the most common neoplasm of the peripheral nervous system, is predominantly a benign lesion. Although it is frequently detected in the head, neck, and extremities, it could be observed anywhere in the body. In the abdomen, the most common site is the stomach, pelvic schwannomas are extremely rare and constitute approximately 0.3-3.2% of all lesions (1). Since schwannomas are benign, they may remain asymptomatic for a long time prior to diagnosis and reach large sizes. Symptoms usually develop because of compression to adjacent organs or nerves (2).

The imaging characteristics of schwannoma is quite diverse. On MRI, it tends to be iso-hypo intense on T1-w image compared to muscle and hyper intense on T2-w image. Marked and homogenous enhancement with contrast medium administration is typical and may be diagnostic. However, related with the cystic degeneration and hemorrhage the signal intensity of the lesion may be heterogeneous. When present, heterogeneity has been shown to correlate histologically with a greater ratio of Antoni B tissue than Antoni A. Seldom, cystic degeneration may reach advanced levels and cause the mass to appear as pure cystic as in our case (3).

Schwannomas, when particularly located within the abdomen and are completely cystic, have wide range of differential diagnosis. Dermoid and epidermoid cysts, tail gut cyst, lymphangioma, a lymph node with cystic degeneration, and bladder diverticulum are included in the differential diagnosis of a smoothly circumscribed complex cystic solitary lesion located in the pelvic region, as in our case (4). Surgical excision and following histopathologic examination would be the gold standard for the diagnosis.

Although the pure cystic schwannoma with pelvic location is extremely rare, it should be kept in mind in the differential diagnosis of pelvic cystic lesions.

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References