Giant Mature Sacrococcygeal Teratoma in an Adult - a Case Report
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Citation

Abstract
Teratoma is a congenital germ cell tumor, composed of a wide diversity of tissue, containing all three germ cell layers. The sacrococcygeal area is the commonest site of extragonadal teratomas in infants, but presents as a rare diagnostic dilemma in adults. We present a rare case of giant mature sacrococcygeal teratoma in an adult female patient.

CASE REPORT
A 17-year-old female belonging to rural area presented with complaints of a mass in the sacrococcygeal area since birth and gradually increasing in size. There were no neurological complaints of lower limb weakness or bowel and bladder dysfunction. On examination, a huge swelling approximately 25 x 25cm in size was present in the sacrococcygeal area (Figure I).

Figure 1
Figure I Clinical photograph of sacrococcygeal teratoma

It was of variable consistency and not fixed with skin. On per rectal examination, the rectal mucosa was free from masses. The patient was anemic (hemoglobin: 8g%) with normal sugar and urea level. His chest X-ray was normal. Alpha fetoprotein was 1 IU/ml (N: 0.55-2.64), LDH was 307 IU/l (N: 114-240 IU). CT scan showed a huge (25 x 25cm) fatty mass with some soft tissue and few calcifications with disfigurement of the sacrum (Figures II and III).

Figure 2
Figure II CT scan image of sacrococcygeal teratoma
The mass was completely excised from the sacral approach (Figure IV).

**DISCUSSION**

When the current focus on sacrococcygeal teratoma is on prenatal assessment, fetal intervention [1] and outcome after surgery in the pediatric age group [2], sporadic cases of sacrococcygeal teratoma in the adult age group come as a surprising rarity. A survey of literature reveals few case reports and even fewer series of cases. [3-8] In fact, a recent comprehensive review could find only 92 published cases in the literature, confirming its rare incidence. [9]

Its incidence has been reported to be 1:87,000 with a female to male ratio of 10:1. [10] Its rarity is further compounded by varied symptoms, which are due to its size, compression of pelvic viscera or complications like infections, bowel obstruction, pelvic abscess, pilo-nidal sinus, degeneration/fistulas, or malignancy. A 20-30% risk of secondary infection and a 1-12.5% risk of malignant degeneration has been calculated. [11] Most adult teratomas described are small, found in female patients and rarely present externally. [12]

The differential diagnosis includes anterior sacral meningocele, anal duct or gland cyst, necrotic rectal leiomyosarcoma, extraperitoneal adenomucinosis, cystic lymphangioma, pyogenic abscess, neurogenic cyst, and necrotic sacral chordoma. [13] Preoperative biopsy should be avoided as it can cause tumor spread, or abscess formation and should not be performed if the tumor is potentially resectable. [3] Good appropriate imaging in the form of CT or MR scan remains the cornerstone of decision making in its management.

The treatment of sacrococcygeal teratomas is mainly surgical. Increasing experience with retro-rectal/presacral tumours and better imaging has resulted in evolving classifications and algorithms for their management. [14-17] The approach depends on the peculiarities of each case (size, location and invasion of the nearby viscera), with most teratomas amenable to resection from posterior sacral approach. Resection of the coccyx is considered necessary to prevent local recurrence [12]; however, it is logical in case of its involvement by the neoplastic mass or suspected malignant transformation. Anterior or combined approaches are rarely required for tumours invading the viscera. Prognosis is directly related primarily to local control, which may be difficult to achieve for malignant lesions.

As late occurrence of malignancy following resection of a histologically mature sacrococcygeal teratoma has been reported [18], our patient has been kept on 3-monthly follow-up.

**References**

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