INTRODUCTION

Teratomas are benign germ cell tumors which include embryonic ectoderm, mesoderm, and endoderm. They are usually found out of their anatomical origin [1]. They comprise 3–5% of childhood tumors and about 20% are malignant [2].

Teratomas can appear in all midline sites, but usually are found in sacrococcygeal area, gonads, mediastinum, cervicofacial region and intracranial fossa. Spinal teratomas are rare. In this study we describe a case of conus medullaris teratoma which was diagnosed based on imaging studies. The patient underwent surgery. We did bilateral laminectomy. The mass lesion had an obvious and rigid attachment to the conus medullaris. The wall of the lesion was resected as much as possible, but total resection of the lesion's wall could not be done due to changes in neural monitoring. Previous related studies are reviewed.

Key Words

Conus medullaris; Teratoma; Intramedullary spinal cord neoplasms.

CASE REPORT

A 12-year-old girl was referred because of progressive gait impairment and back pain since 4 months ago. In the physical examination, deep tendon reflexes (DTRs) of lower extremities were normal and had no sphincter disorders. She had gait disorder as bradykinesia and was obviously incapable of fast movements. Lower extremity force was 4/5.

In the imaging studies and dorsolumbar MRI, a heterogeneous well-defined intradural intramedullary mass in conus medullaris was found. A hypersignal area in T1W images consistent with fat was noted, so teratoma was included in the differential diagnosis (Fig. 1). The patient underwent surgery with a possible diagnosis of a conus medullaris teratoma.

In the operating room, the patient underwent general intravenous anesthesia. After connecting the neuromonitoring leads—somatosensory evoked potential (SSEP) and motor evoked potential (MEP), we utilized warming blankets to prevent intraoperative hypothermia. We also used narcotics and propofol to prevent disturbance in patient’s neuromonitoring.

After mapping the site of surgery by using C-arm, we did bilateral laminectomy of T12 and L1 by using micro-drill and kerison ronguer (number 2 and 3). The mass lesion had an obvious and rigid attachment to the conus medullaris and filum terminale, cutting the filum terminal was not possible. After
incising over the lesion, yellowish liquid containing patches of hair and small white solid tissues was found and then drained totally and specimens were sent for pathologic evaluations. The procedure last approximately 120 minutes.

After the surgery no complication and no new motor neurologic deficits were noted and follow up showed no residual tumoral tissue (Fig. 3). Pathologic report confirmed the diagnosis of teratoma (Fig. 4). In the four month follow up after the surgery patient has no pain and her gate became normal.

As the patient was under 18 years old, we obtained the written informed consent from patient's parent.

**DISCUSSION**

Spinal teratoma was first described by Virchow and Gowers in 1863 [6]. Most cases of spinal teratoma are diagnosed in the first two decades of life, Nsir et al. [7] noted that so far, few cases of conus medullaris teratoma are published in the literature and most of them were adults. According to the previous studies about one-third of cases accompanied by total resection [7].
Işık et al. [8] reported a case of intramedullary teratoma of conus medullaris in 2008 and stated that they could resect the tumor totally with no neurological symptoms. Öktay et al. [9] presented a 12-year-old male patient with conus medullaris teratoma with 95% tumor resection. In further follow up examinations, all symptoms were faded away. Moreover, the authors declared that intraoperative electrophysiological monitoring can help maximum resection of the tumor. Hejazi and Wittmann [10] reported total resection of conus medullaris teratoma in two adult patients. The authors mentioned that intramedullary teratoma of conus medullaris is very rare and should be considered as a differential diagnosis of cauda equina tumors. They suggested that surgical resection should be done before onset of neurological deficits. Most cases of spinal teratoma in adults are found in thoracolumbar region and specially conus medullaris area [11].
The pathogenesis of spinal teratoma is not exactly known, but there are two theories. One theory has hypothesized that these tumors are originated from primordial germ cells that has migrated incorrectly during embryogenesis in dysraphism. Another theory suggests although germinomas are originated from germ cells, other non-germinoma germ cell tumors such as teratomas occur because of misfolding and misplacement of embryonic cells into the lateral mesoderm [12].

In addition to congenital malformations in adults with teratoma, evidence of trauma or surgery such as lumbar punctures has been observed in these patients [13]. Iatrogenic cysts may occur because of entering dermal and epidermal tissues during closure of myelomeningoceles or entering epithelium during spinal epidural injections [14].

Diagnosis of teratoma is based on histopathologic findings and the specimen should contain three germinal layers (ectoderm, mesoderm, and endoderm); however, the existence of only two layers does not rule out the diagnosis [13].

The tumor has three pathologic subtypes: mature, immature, and malignant. The mature type is the most common form [15]. Mature and immature types are considered benign specially if treated immediately. Mature teratomas consist of differentiated components such as chondrocytes, squamous epithelial cells, endocrine tissues, mucosal cells, and neural elements. Immature types are more aggressive and contain primitive, undifferentiated, and embryonic tissues. Malignant tumors—as the name implies—contain some components of malignant germ cells and are originated from yolk sac or endodermal sines. The malignant forms are associated with high levels of serum alpha fetoprotein and poor prognosis [6,11,16].

Teratoma’s association with spinal malformations such as spina bifida, partial sacral agenesis, hemivertebrae, myelomeningocele, tethered cord and diastematomyelia has been reported in literature [11]. Since most cases of spinal teratomas are associated with anomalies such as split cord and dysraphism, these lesions may be non-neoplastic and occur due to dysembryogenic mechanisms [16,17].

Spinal teratomas may be extra-dural, intra-dural or intra-medullary. Intramedullary types are rare cases and they tend to occur in thoracolumbar area [8]. Moreover intradural, extramedullary teratoma cases are more common in childhood [11]. Oktay et al. [9] presented a 12-year-old male patient with conus medullaris teratoma with 95% tumor resection. In further follow up examinations, all symptoms were faded away. Moreover, the authors declared that intraoperative electrophysiological monitoring can help maximum resection of the tumor.

In these patients the most common symptoms are weakness of extremities (paresis), sensory alterations and abnormalities in DTRs which depend on the location of the tumor [6]. MRI is the gold-standard diagnostic method for spinal teratoma, however other imaging modalities like plain X-ray and CT scans can be used as well. Abnormalities in these modalities are bone and vertebral anomalies, erosions and increased interpeduncular space [6]. MRI shows teratoma as a heterogeneous mass containing solid and cystic components and usually fat tissue is also found in the lesion like our case which is helpful for preoperative diagnosis [18]. Differential diagnosis of spinal teratomas is ependymoma, astrocytoma, and complicated neurenteric cysts [19].

As the natural history of these tumors is unknown, the treatment method of choice in children with teratoma is complete resection of tumor [20,21]. Total resection of the lesion should be performed to prevent recurrence, but if total resection may accompany neurological damages and deficits, resection must be stopped [6]. Extensive laminectomy in 30–80% of patients leads to spinal cord instability; therefore simultaneous laminoplasty is recommended [22].

The recurrence rate of tumor depends on histopathologic type, as we can see symptomatic recurrence in mature teratomas even in cases with subtotal resection is rare. Recurrence rate is estimated about 10% in previous studies and it was reported mostly in immature and malignant forms [8]. As the tumor is slow growing, the patients need long-lasting follow ups and serial imaging for many years [23,24].

Adjuvant radiotherapy is recommended for teratomas containing malignant components, but there is no evidence of the efficacy of adjuvant chemotherapy in these patients [9,23,25]. In very rare cases of immature types with evidence of malignancy, postoperative adjuvant chemotherapy may alter the aggressive behavior of this tumors and lead to complete and permanent cure [10,26]. Also the age of the patient and surgical approaches may influence the clinical outcomes [27].

Conflicts of Interest

The authors have no potential conflicts of interest.

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ORCID iDs

Behzad Aminzadeh https://orcid.org/0000-0001-6569-4670

Ehsan Keykhosravi https://orcid.org/0000-0002-8762-2042

REFERENCES

4. Nonomura Y, Miyamoto K, Wada E, et al. Intramedullary teratoma of


