A 50-year-old man was admitted to our neurosurgery department with the chief complaint of difficulty in standing and walking due to leg weakness for nearly 2 days. He also described back pain while sitting and standing. Neurological examination demonstrated paraparesis with impaired anal sphincter tonus. He had no history of trauma or any systemic illness.

Thoracic magnetic resonance imaging (MRI) and computerized tomography (CT) revealed expansile mass lesion that was compressing the spinal cord at T9 level. Homogenously enhancing mass lesion was found to originate from the anterior portion of the spinous process and both laminas of T9 vertebra (Fig. 1). Routine blood tests were uneventful except a calcium value of 14.3 mg/dL (8.4–10.2). As blood parathormone (PTH) test also revealed a very high value of 547.45 pg/mL (15–68.3), endocrinology consultation was ordered to rule out primary hyperparathyroidism. Neck sonography demonstrated hyperplasia and multiple nodules on the thyroid gland. However, any cystic or solid pathologic lesion compatible with adenoma were not seen on the parathyroid regions. Parathyroid scintigraphy with Tc-99m

INTRODUCTION

Brown tumors also called as osteoclastomas, are rare nonneoplastic lesions that arise in the setting of primary or secondary hyperparathyroidism. Parathyroid adenomas or hyperplasia constitute the major Brown tumor source in primary hyperparathyroidism while chronic renal failure is the leading cause in secondary hyperparathyroidism. Most of the patients with the diagnosis of primary hyperparathyroidism present with kidney stones or isolated hypercalcemia. However, nearly one third of patients are asymptomatic and hypercalcemia is found incidentally. Skeletal involvement such as generalized osteopenia, bone resorption, bone cysts and Brown tumors are seen on the late phase of hyperparathyroidism. The symptoms include axial pain, radiculopathy, myelopathy and myeloradiculopathy according to their locations. Plasmocytoma, lymphoma, giant cell tumors and metastases should be ruled out in the differential diagnosis of Brown tumors. Treatment of Brown tumors involve both the management of hyperparathyroidism and neural decompression. The authors report a very rare spinal Brown tumor case, arisen as the initial manifestation of primary hyperparathyroidism that leads to acute paraparesis.

Case Report

50-year-old man was admitted to our neurosurgery department with the chief complaint of difficulty in standing and walking due to leg weakness for nearly 2 days. He also described back pain while sitting and standing. Neurological examination demonstrated paraparesis with impaired anal sphincter tonus. He had no history of trauma or any systemic illness.

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MIBI revealed focal activity retention on the inferior portion of the right thyroid lobe. Abdomen and thorax CT were uneventful. Endocrinology department insisted for the urgent parathyroidectomy in order to minimize hypercalcemia-related complications. So, patient with the diagnosis of primary hyperparathyroidism underwent surgery for parathyroidectomy first. Pathological parathyroid tissue was found and excised. Total thyroidectomy was also performed since frozen examination of a suspected thyroid nodule was reported as micropapillary carcinoma. The day after parathyroid and thyroid surgery blood calcium and PTH levels decreased to 10.2 mg/dL and 10.42 pg/mL, respectively. Patient underwent spine surgery. Tumor was excised in piecemeal fashion. Wide posterior decompression (T8 partial/T9 total laminectomy-bilateral T9 transvers process and pedicle excision) was followed by T8–10 posterior instrumentation and fusion (Fig. 2). Paraparesis resolved postoperatively. Patient was mobilised with a brace on postoperative day 1. Absolute pathology reports were consistent with the parathyroid adenoma and spinal Brown tumor (Fig. 3).

DISCUSSION

Brown tumors occur due to increased osteoclastic activity in the setting of either primary or secondary hyperparathyroidism. Parathyroid adenomas or hyperplasia constitute the major BT source in primary hyperparathyroidism while chronic renal failure is the leading cause in secondary hyperparathyroidism. BTs are localized form of osteitis fibrosa cystica (OFC) that is commonly associated with hyperparathyroidism. OFC is a process that is characterized by hyperstimulation of osteoclastic proliferation via longstanding excessive PTH production causing bone resorption and bone marrow fibrosis.

Histopathologically, BTs are composed of osteoclast-like giant cells and hemosiderin with a fibrovascular stroma. The name Brown tumor comes from the accumulation of hemosiderin that gives the surrounding stroma a brown color. BTs look like similar histologically to other giant cell lesions such as giant cell tumor, aneurysmal bone cyst and reparative giant cell granuloma. Only the clinical manifestation and endocrinologic status help to differentiate BTs from other giant cell lesions.

BTs usually develop in the third or fourth decades of life. In the beginning, BTs that develop in primary hyperparathyroidism were seen much more common. However, they are seen more often recently as a result of secondary hyperparathyroidism that is closely related with increased life expectancy of chronic renal disease patients who require hemodialysis. Most of the patients with the diagnosis of primary hyperparathyroidism present with kidney stones or isolated hypercalcemia. However, nearly one third of patients are asymptomatic and hypercalcemia is
found incidentally. Skeletal involvement such as bone resorption, bone cysts, BTs and osteopenia is seen on the late phase of hyperparathyroidism.9,10,24,27

The symptoms of spinal BTs include axial pain, radiculopathy, myelopathy and myeloradiculopathy according to their locations.9,11,12,13,26,27 Radiographically, BTs do not have a pathognomonic appearance. It can mimic multiple myeloma, metastases, sarcomas and other giant cell lesions9,11,12,13,26,27. X-Rays demonstrate solitary or multiple, lytic, expansile lesions. Bone cortex might be thinned and expanded but it is not penetrated by the tumor frequently. On CT scan, relatively well demarcated soft tissue mass with local bone erosion and expansion is observed. MRI is the diagnosis modality of choice in evaluating the location, extent of spinal tumor and neural compression. On MRI, BTs are commonly seen hypointense on T1-weighted images, hypointense or hyperintense on T2-weighted images and homogenous enhancement is noted after contrast injection. Intratumoral hemorrhages lead to fluid-fluid level appearance on MRI.9,14

Fig. 3. A: There is marked resorption of bone trabeculae with marrow fibrosis (left sided) and clustering of osteoclasts that create a brown tumor (right sided) (×40, hematoxylin and eosin staining). B: Clusters of giant cells and hemosiderin-laden macrophages in the fibrous cellular stroma are seen on higher magnification (×200, hematoxylin and eosin staining).

Table 1. Literature review of spinal Brown tumors seen in primary hyperparathyroidism

| Authors (year) | Sex | Age | Affected spinal level | Symptoms | Treatment |
|---------------|-----|-----|-----------------------|----------|-----------|
| Shaw and Davies (1968) | M | 35 | T10 pedicle | Paraparesis, urinary retention | Resection of lesion and parathyroidectomy |
| Shuanggshi et al. (1972) | F | 40 | L4 posterior element | Paraparesis, radicular pain | Resection of lesion and parathyroidectomy |
| Sundaram and Scholz (1977) | F | 63 | T10 body and pedicle | Paraparesis, urinary retention | Resection of lesion and parathyroidectomy |
| Siu et al. (1977) | F | 64 | T10 | Paraplegia, urinary retention | Resection of lesion and parathyroidectomy |
| Ganesh et al. (1981) | M | 40 | T2 body and pedicle | Paraparesis, radicular pain | Parathyroid adenoma excision only |
| Yokota et al. (1989) | F | 58 | T5 pedicle | Paraparesis, numbness | Resection of lesion and parathyroidectomy |
| Doras et al. (1990) | F | 54 | T9 pedicle | Paraparesis | Resection of lesion |
| Kashkari et al. (1990) | F | 51 | T6/T7 body | Paraparesis | Resection of lesion, instrumentation, fusion and parathyroidectomy |
| Sarda et al. (1993) | F | 23 | T3/T4 | Paraparesis, radicular pain | Resection of lesion and parathyroidectomy |
| Motateau et al. (1994) | M | 57 | L4–5 facet joint | Radiculopathy | Resection of lesion, instrumentation and fusion |
| Mustonen et al. (2004) | M | 28 | L2 posterior elements | Radiculopathy, numbness | Resection of parathyroid adenoma only |
| Haddad et al. (2007) | F | 26 | T2/T3 posterior elements, T4 body | Paraparesis, numbness | Resection of lesion and parathyroidectomy |
| Khalil et al. (2007) | M | 69 | T2 body and pedicle | Radiculopathy | Resection of lesion |
| Altan et al. (2007) | F | 44 | Left lateral side of S2 | Radiculopathy, low back pain | Resection of lesion and parathyroidectomy |
| Hoshi et al. (2008) | F | 23 | Sacrum | Radicular pain | Resection of parathyroid adenoma only |
| Lee et al. (2013) | M | 65 | L2 body and L1 posterior elements | Low back pain, radicular pain | Resection of lesion, instrumentation, fusion and parathyroidectomy |
| Khalafbari and Moharamzad (2014) | M | 16 | L2 pedicle and posterior elements | Paraparesis, sphincter dysfunction | Resection of lesion, instrumentation, fusion and parathyroidectomy |
| Khalafbari and Moharamzad (2014) | F | 46 | L3 posterior elements | Paraparesis, low back pain | Resection of lesion and parathyroid adenoma |
| Khalafbari and Moharamzad (2014) | F | 52 | C6 lamina and pedicle | C6 radiculopathy, neck pain | Resection of lesion, instrumentation, fusion and parathyroidectomy |
| Khalafbari and Moharamzad (2014) | M | 38 | T7 body, pedicle and lamina | Paraparesis, sphincter dysfunction | Resection of lesion, instrumentation, fusion and parathyroidectomy |
| Sonmez et al. (present study) | M | 50 | T9 pedicle and posterior elements | Paraparesis, sphincter dysfunction | Resection of lesion, instrumentation, fusion and parathyroidectomy |
Plasmocytoma, lymphoma, giant cell tumors and metastases should be ruled out in the differential diagnosis of BTs[1-3,5,6,19]. When a solitary lesion that correspond to BT is found on the spine, differential diagnosis might include true giant cell tumor, aneurysmal bone cyst and giant cell reparative granuloma. Also, hyperparathyroidism associated bone changes such as loss of lamina dura of the teeth roots and generalized demineralization of the medullary bone of the jaw could help the surgeon to differentiate BT from other giant cell lesions[41].

Treatment of BTs involve both the management of hyperparathyroidism and neural decompression. Total or subtotal parathyroidectomy is the gold standard for the treatment of primary hyperparathyroidism[3,5,12,19]. Parathyroid surgery rapidly decreases the excessive amount of PTH thus achieving complete regression of the lesions with remineralization[3,6,40]. Parathyroidectomy and medical treatment may also be used together. Neurological status is the main determinant in case of neural compression in the treatment of BTs[3,7]. Immediate surgical decompression could be necessary for achieving good outcome. Unless the decompression procedure does lead to instability, patients can be mobilized in braces without spinal fixation. However, spinal instrumentation and fusion is needed if the tumor is too large, affects multiple levels or pathological spinal fractures exist[3,10,12,19].

Our search in English literature demonstrated 20 spinal BT patients with primary hyperparathyroidism (Table 1). 12 of 20 patients (60%) were women. The patients’ age ranged from 16 to 69 with a mean of 45.3 years. Thoracic spine was the most affected part of the spine (55%) followed by lumbar, sacral and cervical regions. Almost all the patients were presented with major neurologic deficit. 15 patients (75%) underwent double surgeries for both removing the mass and to treat the primary hyperparathyroidism. 6 patients (35%) underwent instrumentation and fusion surgery whilst 11 patients (65%) underwent only decompression surgery. After surgery, improvement of symptoms were observed in all patients.

CONCLUSION

BTs should be kept in mind in the differential diagnosis of lytic, expansile spinal tumors, especially with the histopathological diagnosis of giant cell tumors. Treatment consists of both treating the underlying primary pathology and decompression of the neural structures, if necessary. Urgent surgery can be needed to preserve neural function and stabilize the spine.

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