ADRENAL GANGLIONEUROMA IN AN ADULT - CASE REPORT

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INTRODUCTION

Neuroblastic tumors arise from neural crest cells and encompass a spectrum from neuroblastoma – an undifferentiated, malignant tumor – to ganglioneuroma – a well – differentiated, benign neoplasm (Leavitt et al. 2000). Neuroblastomas and ganglioneuroblastomas represent the second most common group of solid extracranial neoplasms of infancy and childhood and account for approximately 8% of malignancies in these age groups. Most of these tumors are diagnosed by the age of 4, although rare examples have been reported in adults. Ganglioneuroblastoma is an intermediate type of tumor in the spectrum of ganglion cell tumors, containing both primitive neuroblastomatus and mature ganglioneuromatous elements of sympathetic cell origin (Shimada et al. 1999). Ganglioneuromas, ganglioneuroblastomas, and neuroblastomas are histologically differentiated by their stage of neuroblast maturation (Sovak et al. 2008). Ganglioneuromas are composed of mature ganglion cells and are considered benign tumors. Ganglioneuroblastomas and neuroblastomas are less mature and are considered more aggressive and dangerous (Lonergan et al. 2002).

CASE REPORT

A 48-year old man examined for progressive weakens in lower limbs and back pain on Department of Neurology. Computerized tomography (CT) observed a large, irregular mass retroperitoneal left. Mass was 25 – 20 – 15 cm diameters. Right kidney and adrenal was normal. Multiple lymphadenopathies in paraaortic and aorta-caval lymph regions were observed. Patient had no clinical sign of endocrinal tumor and laboratory findings were negative for hormone-active tumor. Surgical exploration was made and adrenal tumor with ipsilateral kidney was removed. The lymph nodes in aorta-caval and paraaortic regions could not be removed. In pathologic examination; the left nefrectomy material was macroscopically 27 – 20 – 16 cm and inside it was seen compressed kidney measuring 11 – 7 – 3 cm and all around it is big tumoral mass measuring 20 – 19 cm (Figure 1). Tumor mass was well circumscribed and white in colour (like fish meat) and have areas of hemorrhage, necrosis and calcification. The histologic examination showed that the lesion was composed of large cells with abundant cytoplasm with large vesicular nuclei and prominent nucleolus, representing mature ganglion cells with no or minimal residual neuroblasts and background “schwannian stroma” comprised of organized fascicles of neurotic processes, mature Schwann cells, and fibroblasts (Figure 2).
DISCUSSION

Ganglioneuromas are rare, benign, fully differentiated tumors that contain mature Schwann cells, ganglion cells, fibrous tissue, and nerve fibers. These tumors have no immature elements (such as neuroblasts), atypia, mitotic figures, intermediate cells, or necrosis. The presence of any of these tissue characteristics excludes the diagnosis of ganglioneuroma (Shin et al. 2002). Ganglioneuromas can grow almost anywhere along the paravertebral sympathetic ganglia, and they can sometimes grow in the adrenal medulla (Maweja et al. 2007). These tumors can arise de novo and result from the maturation of a ganglioneuroblastoma or neuroblastoma into a ganglioneuroma. They may also develop within a neuroblastoma treated with chemotherapy. Metastases from ganglioneuromas are exceedingly rare. Metastasis is thought to be the end result of matured ganglioneuroblastomas or neuroblastoma metastases rather than true ganglioneuroma metastases. Ganglioneuromas secrete catecholamines in as many as 37% of cases (Geoerger et al. 2001). Overall, patients with ganglioneuroma have a favorable prognosis. Ganglioneuromas are usually asymptomatic, regardless of their size, and they are typically discovered on a routine radiograph; however, abdominal pain, dyspnea, cough, and palpation of an abdominal mass may be clinical indicators of a ganglioneuroma. These tumors may be hormonally active, and hypertension, diarrhea, flushing, and virilization may occur as a result of catecholamine, vasoactive intestinal polypeptide, or androgenic hormone. Nonetheless, emergency situations caused by catecholamine secretion are rare (Patterson et al. 2009).

Treatment depends on surgical resection whenever possible, chemotherapy for tumors unlikely to be completely resected and radiotherapy for cases with regional nodal disease. Rousseau et al. 1999, reported a ganglioneuroblastoma of the left adrenal gland with liver metastasis. They had complete remission from combination therapy with chemotherapy and radiotherapy after surgical resection. Hiroshige et al. 1995 presented a case of well-differentiated ganglioneuroblastoma treated with surgical resection.

CONCLUSION

Surgical resection is sufficient for low risk and non-metastatic tumors. Combination therapy with chemotherapy and radiotherapy is necessary for disseminated cases.

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References