MALIGNANT PARAGANGLIOMA WITH VERTEBRAL METASTASIS

Case report

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ABSTRACT - A paraganglioma is a rare tumor, composed of chromaffin cells, originated from the neural crest, a group of cells associated to the autonomous nervous system. When the primary location is the adrenal gland, the tumor is called pheochromocitoma. The malignant paraganglioma is a very rare presentation; it is diagnosed by local recurrence after total resection of the primary mass, or findings of distant metastases. We present a case report of a 29-year-old woman with cervico-brachial pain. In 1995 she underwent a carotid body tumor resection. Magnetic resonance imaging (MRI), plain X-rays and computerized tomography scan revealed multiple lesions in C5, T5 and T12. She underwent a surgical procedure to correct the cervical lesion. The histological and immunohistochemical assays revealed a malignant paraganglioma. She received adjuvant radiotherapy, showing clinical improvement after treatment, presenting no symptoms after one year. The therapeutic approach is based on the total resection of the tumor. The treatment of distant metastases can be made with adjuvant measures such as conventional radiotherapy, I¹³¹-MIBG, or chemotherapy, especially in malignant pheochromocitomas.

KEY WORDS: paraganglioma, vertebral metastasis, spinal tumor

Paragangioma maligno com metástase vertebral: relato de caso

RESUMO - O paragangioma é tumor raro, composto de células cromafins, associado ao sistema nervoso autônomo. Quando localizado na glândula supra-renal, o tumor é chamado feocromocitoma. Descreve-se um caso de paciente do sexo feminino, 29 anos, que se apresentou com cervicobraquialgia e que havia sido operada em 1995 para exérese de tumor glômico da carótida cervical. RM, RX e TC revelaram múltiplas lesões acometendo o corpo vertebral de C5, T5 e T12. Foi submetida à ressecção cirúrgica radical da lesão cervical, com substituição do corpo vertebral por prótese de titânio. A histopatologia e o estudo imunohistoquímico da lesão confirmaram o diagnóstico de paragangioma maligno. As outras lesões foram tratadas com radioterapia. Um ano após os procedimentos, a paciente apresenta-se assintomática. O tratamento destas lesões consiste na associação da ressecção cirúrgica radical do tumor e medidas complementares como radioterapia convencional, aplicação de I¹³¹-MIBG, ou quimioterapia, principalmente nos paragangliomas malignos.

PALAVRAS-CHAVE: paragangioma maligno, metástase vertebral, tumor vertebral

A paraganglioma is a rare tumor, composed of chromaffin cells, originated from the neural crest, a group of cells associated to the autonomous nervous system. Paragangliomas can be found in several locations. When the primary location is the adrenal gland, the tumor is called pheochromocitoma; in the head and neck region, the paragangliomas can be most found in the carotid body, jugular and vagal glomus; other locations also include the retroperitoneum, para-aortic region, bladder, filum terminale, skull, larynx, and others. The presentation of the tumor is variable: in most cases it is represented as a non-symptomatic slow-growing mass, compressing the anatomic structures around it. On the other hand, the tumor can secrete catecholamines, inducing a typical clinical history represented by hypertension, paroxysms and headache. Malignant paragangliomas are a very rare presentation, limited to less than ten cases registered between 1985-1996¹; the annual incidence in retrospective studies is 1/10 000 000². It is diagnosed by local recurrence after total resection of the primary mass, or findings of distant metastases. The time intervals for recurrence may be from months to many years.

This study presents a case report of a 29-year-old woman with vertebral metastases 5 years after the
diagnosis and surgical removal of a carotid body tumor. She underwent a surgical procedure to correct the vertebral lesions, and adjuvant radiotherapy.

CASE

In 1995 a 29-year-old woman presented a cervical mass on her left side, which evolved to a progressive cervical pain irradiating to the left ear. The physical examination was normal, except for a decrease on the left carotid and temporal arteries pulse. All laboratorial exams were normal; a MRI of the neck showed a 4.7 x 3.3 x 3.1 cm lobular mass with an intermediate T1-weighted sign and a high heterogeneous T2-weighted sign, enhanced with gadolinium, on the left carotid bifurcation, with a mass effect over the ipsilateral sub-mandibular gland, suggesting a carotid body tumor. A carotid-vertebral angiography showed an encasement of the left carotid by the tumor. A resection of the mass was performed followed by a carotid-carotid internal saphenous vein by-pass. The outcome was positive; a carotid-vertebral duplex scan showed an adequate flow, demonstrating a good functioning of the graft. Histopathology revealed a 4.0 x 3.0 x 2.5 cm grayish-brown mass, with necrotic areas. The microscopy showed a pattern of cords and nest cells, typical of a carotid body tumor. The patient was discharged from the hospital without any symptom.

In October 2001, the patient was readmitted to the hospital presenting a cervico-brachial pain; the patient was well until the last six months. The neurological exam showed a hyposthesia of the left territory from C4 to C5 (Fig 1). The erythrocyte sedimentation rate (ESR) was 37mm. Plain X-rays of the cervico-thoracic spine revealed a pathologic fracture of C5, T5 and T12. CT scan of the cervico-thoracic spine showed a fracture of T5 and T12, with a severe lesion on vertebral body of C5 (Fig 1). Cervico-thoracic MRI examination revealed a destructive lesion of C5, with displacement of the vertebral posterior wall towards the spinal canal; bone destruction of T5 and T12, with invasion of the spinal canal on T5 level suggesting a secondary implant (Fig 1a 4). In November 2001 a surgical procedure was proposed, comprising the C5 corpectomy, identifying an hemorrhagic tumor tissue; the setting of a titanium cage filled with an osteoinductive coral graft, and the osteosynthesis of C4 to C6 using four 16-mm screws and a 16-mm titanium plate (Fig 5).

Histopathologic assessment revealed vertebral bone infiltration with a pattern of spread cells suggestive of epithelial origin, with an early result of adenocarcinoma. A full diagnostic trial was performed, including a thoracic-abdominal-pelvic CT scan, endoscopy, colonoscopy, mam-
mography, pelvic and intravaginal ultrasound scan, measurement of CEA, CA-125, CA-19.9, CA-15.3, β-HCG, α-FP, and regular laboratory trials. An endoscopy of gastrointestinal tract five revealed inactive gastric ulcers at the antrum (with normal serum gastrin, Helicobacter pylori and histopathologic analysis) and a left kidney cyst. A hypothesis of malignant paraganglioma was then suggested. A revised histopathologic assessment was requested; and an immunohistochemical study was performed in both primary and secondary lesions with enolase, chromogranin, synaptophysin, S-100. Both lesions showed a brown aspect on microscopy, although the secondary lesion demonstrated a weaker staining to enolase and S-100 protein; the final revisited diagnosis was the presence of secondary implants of a malignant paraganglioma.

The patient has received adjuvant radiotherapy for cervico-thoracic spine 4 000cGy, with improvement in her symptomatology.

**DISCUSSION**

Paragangliomas can occur in several locations, usually related to mesodermal branchial archs. These tumors arise from neuro-ectodermal cells, forming neuroendocrine tumors divided in four categories: typical and atypical carcinoid, oat cell carcinoma, and giant cell neuroendocrine carcinomas.

Neuroendocrine cells, similar to chromaffin cells, are found spread on extra-adrenal tissues, as nodules and aggregated cells, associated with the adrenal gland, composing the paraganglionic tissue. These extra-adrenal paraganglia are in close association with the autonomous nervous system, and can be divided in three groups, according to their anatomic location: branchiometric, intra-vagal, and aortic-sympathetic. Intra-vagal and branchiometric paraganglia are related to the parasympathetic system; the general location is next to the great arteries and cranial nerves of the head and neck, including the carotid body. The intra-vagal paraganglia are distributed along the vagus nerve. The aortic-sympathetic paraganglia are related to the sympathetic ganglia, distributed along the abdominal aorta.

Some of the branchiometric paraganglia, specially the carotid body, act as chemoreceptors responding to variations in oxygen tension and carbon-dioxide concentrations; therefore their other denomination: chemodectomas.

Paragangliomas, however, are found in other locations; the possible explanation for this fact is the migration of the paraganglionic tissue to unconventional sites such as the skull, parasellar region, ponto-cerebellar angle and cauda equina, acting as a differential diagnosis on backache.

When the tumor is located in the adrenal gland, it is denominated pheochromocitoma. It usually begins as a glandular hyperplasia, then it forms micro nodules, macro nodules, and when larger than 1cm it is denominated pheochromocitoma.
**Pathology** - Paraganglionic tissue is composed of 2 types of cells: type 1 or chief neuroectodermal cells, bearing catecholamines granules; and type 2 or sustentacular cells. The histological aspect of the tumor on hematoxiline-eosine and reticuline stain is the classical “Zellballen” pattern, presented as nests or cords of type 1 cells surrounded by type 2, or sustentacular cell. On immunohistochemical analysis, type 1 cells contain neuron specific enolase, chromogranin and leu-enkephalin; type 2 cells contain S-100 protein. In some cases, particularly those that were recurrent, locally aggressive and malignant, the organoid pattern was less apparent; central necrosis on the Zellballen was found, as nuclear pleomorphism. But the diagnosis of malignant paraganglioma is only made by findings of local recurrence or distant metastasis. Linnoila et al. described the decreased expression of neuropeptides related to malignant paraganglioma, using immunohistochemical assays with leu and met-enkephalin, somatostatin, pancreatic polypeptide and VIP. Possible explanations would be the decreased synthesis or elevated secretion rate or defective intracellular storage. They conclude that no matter the mechanism, the decreased expression of neuropeptides is related to a worse prognosis. Kliewer et al. demonstrated a maximum staining intensity with enolase and chromogranin in benign paragangliomas. Malignant paragangliomas showed minimal staining to S-100 protein on type 2 cells.

**Symptomatology, diagnosis and treatment** - The clinical aspect of a paraganglioma is mainly related to the mass effect of the tumor on adjacent structures. The classical triad of headache, diaphoresis and palpatations, and other symptoms of a hyperadrenergic state, such as hypertension, paroxysms, and cardiac, gastrointestinal and metabolic manifestations are typical of a pheochromocitoma, being less apparent in a paraganglioma and rarely present in a malignant paraganglioma.

The diagnosis of a paraganglioma is based on image methods that can be associated to biochemical assays in cases that the functioning tumors are secreting catecholamines. The “gold-standard” method is the MRI. It shows great accuracy demonstrating the size and density of the tumor and its relation to the adjacent structures. Plain X-rays and CT are valuable methods in showing bone and vertebral metastasis. Ultrasound scan is a limited but important method in diagnosing retroperitoneal and pelvic paragangliomas, showing less sensibility in tumors located in other sites. Angiography is an important tool for the diagnosis of vascular commitment caused by the tumor, as on planning pre-operative embolization.

Scintillography is an essential method of determining metastatic disease. The use of I\textsuperscript{131}- MIBG (meta-iodobenzylguanidine), a structural analog of guanethidine, with uptake in the adrenergic granules has been elucidated in some studies. This method can show the presence of the disseminated disease even with a non-functioning tumor, being useful on the diagnosis of primitive neural crest tumors, such as neuroblastomas and paraganglioma. I\textsuperscript{131}- MIBG has a sensitivity of 87-91% and a specificity of 90-94%. The use of I\textsuperscript{131}- MIBG for the treatment of paragangliomas is discussed below.

Biochemicals assays are employed on the diagnosis of a functioning tumor. The usual methods are the measurement of the urinary catecholamines le-
vels and its metabolites on a 24-hour urine sample. Levels of metanephrine, vanilmandelic acid and free catecholamines can be measured. The number of samples depends upon the degree of suspicion; the presence of abnormal levels in a patient with a typical clinical history is very suggestive of the diagnosis.

The success in treating benign and malignant paragangliomas is based on the early diagnosis, complete resection of the tumor using an adequate catecholamine blockade, and an excellent anesthesia. The strategy must be carried on the adequate study of the tumor and adjacent structures and vessels; a complete resection of the primary mass is the treatment of choice. Usually the tumor presents as a latent slowly growing mass, the resection of isolated secondary masses shows a better prognosis, as demonstrated on pulmonary metastasis of tumors of the head, trunk and extremities. The use of $^{131}$-MIBG radiotherapy (RT) has been implicated on partial remission of individuals with malignant pheochromocitomas. The intensity in the uptake of the radiotracer by the tumor cells is not fundamental, as even the non-secreting tumors show an adequate response to the treatment. Although good results have been described with $^{131}$-MIBG therapy in treating bony metastasis, the main role of $^{131}$-MIBG is diagnostic and palliative instead of curative. Conventional RT has been described for treatment of malignant paragangliomas. A series of 84 patients with chemodectomas of the head and neck demonstrated 73% local control for 25 years with radiotherapy solely, compared to 54% for 15 years with surgery solely. The recommended doses are 4 500-5 000cGy during 4-5 weeks. The combined use of RT and chemotherapy has been proposed in treatment of malignant pheochromocitomas, using a three-drug trial composed of Cyclophosphamide, Dacarbazine and Vinristine; these two methods would act in sequence in different sites, with the final result being better than the isolated use of one method or the other. The previous use of chemotherapy would be implicated in increasing sensibility and uptake of $^{131}$-MIBG by the tumor. The treatment of choice for malignant paragangliomas with vertebral metastasis is decompressive surgery. According to the National cancer database on malignant paragangliomas of the head and neck, the 5-year relative survival rate was 59.5% (76.8% for regionally confined carcinoma and 11.8% for distant metastasis). Among the patients who were followed-up until death, those treated with adjuvant irradiation had a longer median survival (45 months) compared with those patients who were treated with surgery solely (12 months).

In conclusion, we report a rare case of a para-

ganglioma with vertebral body metastasis in 29-year-old patient six years after the complete removal of a carotid body tumor, and its surgical and radiotherapy aspects. A long-term follow-up is mandatory for all patients, as previously demonstrated. The time intervals for local recurrence and distant metastasis can vary from months to many years after the initial diagnosis. The use of scintillography with $^{131}$-MIBG is a powerful tool for the identification of the metastasis, and a palliative method for treating the malignant paraganglioma. The primary management of a malignant paraganglioma should be directed towards the complete surgical resection of the primary tumor, regional lymph nodes and distant metastasis. The therapeutic proposal for our patient was the resection of secondary implants in C5 given the main destructive site at this topography, associated with adjuvant RT. The patient has shown clinical improvement, having no symptoms after 6 months of adjuvant therapy.

REFERENCES