Merkel Cell Carcinoma: Case Report and Literature Review

Carcinoma de células de Merkel: Reporte de caso y Revisión de literatura

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Abstract
Merkel Cell Carcinoma is a rare type of aggressive and often fatal neuroendocrine tumor of the skin.

Keywords
► carcinoma
► merkel cells
► hand

Introduction
Merkel cell carcinoma (MCC), primary neuroendocrine carcinoma of the skin,1 is a rare, potentially fatal tumor,2 first described by Toker, in 1972.1,4 Currently, it is the most malignant tumor of the skin and has a high rate of recurrence and mortality.5 The AEIOU mnemonic was used in English to describe the clinical and demographic characteristics of the tumor: asymptomatic, expanding rapidly, immune suppression, older than 50 years, ultraviolet-exposed/fair skin.6 It is a rare tumor, representing less than 1% of malignant cutaneous tumors.7 It occurs mainly on the face and neck (40-60%), followed by the trunk (33%) and rarely on the limbs (10-20%).5,7 It is more frequent in the elderly, mainly Caucasian men. In young people, its prevalence is higher in immunocompromised individuals.8 Several chromosomal changes associated with MCC have been described, the most frequent being located on chromosomes 1, 6, 11 and 16. Currently, MCC recurrence has been linked to the "Merkel cell polyoma virus" (MCPyV).2

Clinically, MCC manifests as a solid, painless, exophytic tumor, skin color or reddish-blue and very aggressive growth, which may increase in weeks to months.2 The initial course of MCC is asymptomatic and the lesion may be...
clinically and histologically similar to basal cell carcinoma, which often delays the diagnosis. The reason for medical consultation is usually the rapidly progressive growth of the tumor\textsuperscript{1,6} and in the initial consultation approximately 26% of patients already have lymphatic metastasis and 8% with distant metastatic disease.\textsuperscript{8} The confirmation of the diagnosis is based on histology and immunohistochemistry, with the expression of epithelial and neuroendocrine markers.\textsuperscript{9} Determining the stage of patients with MCC is of great importance to establish the most appropriate prognosis and treatment.\textsuperscript{1} Tests such as ultrasound, computed tomography (CT), magnetic resonance imaging (MRI) and positron emission tomography (PET)-CT/FDG-PET can be used to complete staging.\textsuperscript{10} The American Joint Committee on Cancer supports the staging system T (tumor) N (lymph node) M (metastases) for MCC. Stage I consists of primary tumors of 2 cm or less, with no evidence of regional ganglion invasion. Stage II includes primary tumors larger than 2 cm (T2 or T3) or a primary tumor with invasion of bone, muscle, fascia or cartilage (T4). Stage III is defined as any primary tumor with regional ganglion disease, and stage IV includes tumors with metastases in addition to the regional lymph nodes.\textsuperscript{5}

The main treatment for MCC is surgery. Treatment consists of radical surgical resection and radiation therapy. Current surgical margin recommendations are based on the size of the primary tumor. In tumors smaller than 2 cm the margins of surgical excision should be 1 cm and 2 cm for tumors larger than 2 cm.\textsuperscript{6} Sentinel lymph node biopsy is important in the stage and prognosis of MCC.\textsuperscript{5} In stages I and II, without local or distant metastases, radical excision should be performed, accompanied by radiotherapy in patients at risk. In stage III, in which there are locoregional metastases, surgical excision of the lesion should be complemented with excision of the lymph nodes and radiotherapy. In stage IV, or the presence of distant metastases, treatment is palliative involving chemotherapy, complemented by radiotherapy.\textsuperscript{2} The role of chemotherapy in the treatment of MCC remains unclear. Currently, chemotherapy is considered as palliative treatment in cases of disseminated disease and not as adjuvant or primary treatment.\textsuperscript{6} Patients should be periodically monitored due to the high rate of recurrence.\textsuperscript{10} MCC is particularly aggressive with a relative mortality of approximately 30% and 50% at 2 and 5 years after diagnosis, respectively.\textsuperscript{5} Although it is a tumor of fast growth and metastasis, there are published data on spontaneous regression.\textsuperscript{2}

We describe a patient with MCC on the back of the wrist. Due to the rarity of the lesion on the upper limb, especially on the dorsum of the wrist, with only two more cases described in the literature, we emphasize the need for early diagnosis and adequate treatment for the good clinical evolution of the patient.

**Case Report**

Female patient, 56 years old, was referred to the Orthopedics consultation for massive swelling of the back of the left wrist, with progressive growth over six months. The objective examination showed a swelling of 3 × 2 cm, solid, adherent to the deep planes, reddish in color (► Fig. 1). Active and passive mobilization of the wrist was painful. Axillary or epitrochlear adenomegalies were not palpable. No neurovascular deficits were detected. NMR of the wrist and hand (► Fig. 2, 3) showed a large tumor, with no apparent involvement of the adjacent major structures, with no apparent bone or tendon invasion. An aspiration biopsy was performed which described a neuroendocrine carcinoma with the possibility of MCC. Chest CT, abdomen and pelvis, and PET-CT, did not show distant lesions. Lymphoscintigraphy marked a sentinel ganglion. Primary surgical treatment was decided. The patient underwent radiographic surgical excision of the sentinel ganglion with a gamma camera and extended excision of the skin lesion (► Fig. 4). The entire extensor tendon compartment of the fingers was resected, and the arthrodesis of the wrist was performed with a plate and screws. Tendon transposition was performed from the radial flexor carpal tendon to the extensor digitorum communis and the abductor pollicis longus (► Fig. 5). The defect is reconstructed using an inverted forearm flap with isolation of the radial pedicle and tunneling to the dorsal area, and coverage of the flap donor area with partial skin graft (► Fig. 6) harvested from the left thigh. The surgery was uneventful. Histological and immunohistochemical examination of the surgical specimen confirmed the diagnosis of MCC and the surgical margins and sentinel ganglion were negative for tumor lesion. The patient underwent
postoperative adjuvant radiotherapy and presented a good clinical and imaging evolution, with no complications or disease recurrence at 12 months after surgery.

Discussion
The benign aspect of MCC, usually in exposed skin areas, contrasts with its extensive microscopic invasion and may delay diagnosis. MCC is a well-known tumor of the skin but is rarely described on the upper limb, especially on the wrist and hand. To date, we are aware of the report of 8 cases of MCC in the upper limb. The differential diagnosis of MCC should include basal cell carcinoma, squamous cell carcinoma, epidermoid cyst, amelanotic melanoma and skin metastases. In cases of MCC of the fingers, resection of the radius involved has been recommended. For tumors of the dorsum of the wrist and hand, a wide local excision is recommended. However, tumor resections at these sites are associated with greater difficulty in reconstruction, due to the limited viability of soft tissues to cover tendons, adjacent neurovascular structures and carpal bones. The case presented elucidates the complexity of this type of surgery, requiring a reverse-flow flap to cover the injured area. Factors of poor MCC prognosis, such as male gender, ganglionic or systemic metastasis and primary tumor location in the head, neck or trunk, have been described. However, there are still only reports and small case series that address the prognostic factors or predictors of survival for the upper limb and hand MCC. This lack of evidence-based data can be a problem for reconstructive wrist and hand surgery, just as malignant hand tumors can be a surgical challenge in balancing tumor eradication and limb function conservation. MCC presents an early locoregional metastasis and recurrence, and both locoregional and distant ganglion metastases are, respectively and independently, associated with worse survival. Patients with upper limb MCC, compared to other locations, have a slightly longer survival, are diagnosed in earlier stages of the disease and have lower rate of regional and systemic metastasis. However, they still have relatively low overall survival for a skin neoplasm. In the present clinical case, treatment with surgical excision and radiotherapy seems, to date, to have been an excellent therapeutic option. However, a longer follow-up, at least five years, will be necessary to confirm the success of this approach.

Due to the rarity of these tumors, mainly located in the wrist, a high index of suspicion is necessary, as well as early
diagnosis and adequate treatment, so that the patient evolves in an appropriate way. Thus, the authors consider that the report of this case may alert and make health professionals aware of this serious and lethal pathology.

**Conclusion**

MCC is an uncommon and aggressive cutaneous malignant neoplasm and rarely described in the upper limb. The primary lesion of the MCC is distinguished by the absence of specific clinical characteristics. Prevention of MCC and early surgical resection plays a crucial role in the patient's prognosis. Currently, the guidelines advocate aggressive clinical behavior and highlight the need for coordination of multidisciplinary care in the approach and treatment of this pathology. Due to the rarity of these tumors and the lack of high quality studies, the MCC of the hand and upper limb has not been adequately described and a high index of suspicion is needed. Thus, the authors consider that the report of this case may alert and raise awareness among health professionals, especially in the pathology of the skin and hand, in the recognition of this neoplasia and assist in the early approach and treatment.

**Conflict of Interests**
The authors declare that have no conflict of interests.

**References**


