A Case of Multiple Cutaneous Metastases from Hepatocellular Carcinoma Mimicking Pyogenic Granuloma

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Abstract
Hepatocellular carcinoma (HCC) metastasizing to the skin is uncommon and carries a poor prognosis. The most frequent sites of skin metastasis are the face and scalp. Cutaneous metastatic lesions usually appear as nodules, but sometimes show a pyogenic granuloma-like morphology. We report a case of multiple skin metastases from HCC that resembled pyogenic granulomas, and showing HepPar1 as a specific marker of HCC. A key feature in this case was metastasis to the mandibular gingiva and digits. This case emphasizes the need to recognize that gingival and digital masses like benign or inflammatory lesions might represent an initial sign of HCC.

Keywords: Hepatocellular carcinoma; Pyogenic granuloma; Gingival; Acrometastasis; Hepatocyte paraffin-1

Introduction
The most frequent sites of skin metastasis from hepatocellular carcinoma (HCC) are the face and scalp, with lesions appearing as single or multiple, firm, painless, nonulcerative, reddish nodules, measuring 1-2.5 cm in diameter. These metastases represent a sign of poor prognosis, indicating the strong possibility of metastases in other regions of the body, and point to a median survival time of less than 5 months [1]. We report a case of multiple skin metastases from HCC that resembled pyogenic granuloma, including the mandibular gingiva and digits.

Case Report
An 86-year-old Japanese man presented with easy bleeding from a cutaneous nodule on the left elbow that had been present for 1 month (Figure 1). The nodule resembled pyogenic granuloma, measured 10 mm in diameter, and was solitary, soft, berry-shaped and reddish. Medical history included a 6-year history of HCC, originating from liver cirrhosis associated with chronic hepatitis C. He had undergone partial hepatectomy and radiofrequency ablation as initial treatment. The patient remained free of clinical disease for slightly more than 4 years, from which point recurrent HCC in the liver was diagnosed. Treatment for recurrences included transcatheter arterial embolization twice and radiofrequency ablation 3 times. Laboratory investigations revealed slightly elevated serum levels of protein induced by vitamin K absence or antagonist II (PIVKA-II) (93 mAU/ml; normal range, 0–40 mAU/ml). No local recurrence or distant metastasis was evident on computed tomography. We initially suspected pyogenic granuloma based on the clinical appearance. The nodule was removed and sent for histopathological examination.

Histologically, eosinophilic neoplastic cells proliferated with a trabecular pattern within the dermis (Figures 2a-2d). The mass consisted of irregular columns of large, polygonal cells with nuclear atypia and eosinophilic cytoplasm. Tumor cells resembled hepatocytes with nuclear atypia showed positive results for hepatocyte paraffin-1 (HepPar1) and CD10 on immunohistochemical examination. Negative results were obtained for alpha-fetoprotein antibodies and carcinoembryonic antigen.

From these clinical and histopathological findings, we diagnosed cutaneous metastasis from HCC. After 6 months of follow-up, the patient developed local recurrence of HCC and multiple metastases to the face, mandibular gingiva and digits (Figures 3a-3c). PIVKA-II

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Cutaneous metastases from HCC comprise only 0.1% of all metastases [10]. The most common primary site was the lung (44%), followed by kidney (12%), breast (10%) and colon cancers (6%) [11]. Afshar et al. observed only 8 (4%) of 221 cases of HCC metastasis to the hand and wrist [12]. The median survival of patients in the reported cases was 6 months [11]. The mechanisms underlying acrometastasis remain poorly understood. The phalangetal absence of bone marrow has led to alternative theories regarding hematologic spread, including increased blood flow and trauma [13]. One suggestion is that the prostaglandins released following a traumatic experience may be responsible for cell migration and adhesion to bone [14].

Conclusion

Skin metastasis from internal neoplasms should be considered among the differential diagnoses in the evaluation of cutaneous tumors. In particular, this case emphasizes the need to recognize that gingival and digital masses resembling benign or inflammatory lesions such as pyogenic granuloma might represent an initial sign of HCC.

References