Abstract. — Melanoma has long been known to be a malignancy that can present in a plethora of forms. This often makes diagnosing it a challenge, but lends support to the concept of frequently keeping it in the clinician’s differential. We present the unique case of a rapidly enlarging axillary mass in an elderly man that initially did not appear concerning for malignancy. Pathologic diagnosis was particularly difficult due to the cystic nature of the mass, and the uncommon histochempl staining pattern of this malignant melanoma. We discuss the variety of therapies directed at symptom relief and control when discovered in advanced stages. This was a rare presentation of a malignant melanoma that highlights the need to maintain a suspicion for the disease in uncertain clinical situations.

Key Words: Melanoma, Axillary mass.

Introduction

Pathologies arising in the axilla are vast and related to a number of different organ systems. This complex anatomical region contains major nervous, vascular, lymphatic, and musculoskeletal structures, covered by a complex dermis. Masses found in this region may be secondary to derangements that are congenital1,2, iatrogenic3, infectious4,5 or proliferative6-9. Often times the clinician will need to employ not only an exhaustive history and physical exam but also call for imaging and often biopsy.

Melanoma has long been known as a malignancy that can present in many different forms and in many different locations1,10-12. As a result, a long held dogma is that the clinician should include it in their differential diagnosis frequently.

Case Report

An 85 year-old man with advanced Parkinson’s disease was transferred to our Hospital from skilled nursing facility with a recent history of a rapidly enlarging, erythematous and painful right axillary mass (Figure 1). He had become aware of the mass 3 months prior to presentation. He was on Coumadin for atrial fibrillation. He had an MRI, which showed an 11.7 cm complex fluid collection with differential diagnosis of hematoma or abscess. Incidentally, imaging also noted a remote right humeral neck fracture. He was started on a 10 days course of trimethoprim/sulfamethoxazole, did not respond and was transferred to the hospital.

On exam he had a 15 cm, tense, warm and mobile mass in the right axilla, with skin ecchymosis and signs of peripheral venous congestion (Figure 1). There was no appreciable locallymphadenopathy. The remainder of the right upper extremity was non-edematous and the neurovascular exam was normal. His only abnormal blood test was an INR of 2.2. Repeat MRI showed a 15 cm mass with central cystic component, encompassed by a thick lobulated rind of soft tissue. Computed tomography of the chest revealed locallymphadenopathy and a 1cm, non-calcified nodule in the right middle lobe of the lung. Under ultrasound guidance approximately 0.5 liters of turbid, yellow fluid was aspirated and sent for cytology. This showed extensive inflammation, coagulative-type necrosis and scant cells that labeled positive for microphthalmia-associated transcription factor (MITF) (Figure 2).

Given the concern for the viability of the overlying skin, an attempt at fine needle aspiration was preferred over incision. Although infection was a possibility, there was a definite concern for malignancy. No organisms were identified and histochemical stains for acid-fast bacilli and fungi were negative. The few cells that were obtained were identified as malignant. As part of staging, a PET scan was obtained displaying avid uptake in several discrete areas of soft tissue, lung, and the axial skeleton. A repeat attempt,
surgery. As seen in this case, this malignancy is capable of assuming many different macroscopic and histologic guises, including that of an abscess. Numerous phenotypes exist and range from desmoplastic to clear cell and melanoma notoriously can mimic other neoplasms. Helpful clues to diagnosis include, intracytoplasmic melanin pigment as well as intranuclear cherry-red nucleoli. Although the melanin was scanty in this case, there were focal findings of characteristic nucleoli. This case exemplifies the fact that the diagnosis of melanoma usually involves selective immunohistochemistry (IHC) testing. 

MITF is purportedly a very sensitive nuclear stain for melanoma but in some studies it had poor specificity. In our case, the suspicion for melanoma was confirmed when the specimen stained strongly for the more specific cytoplasmic antibody HMB45. Typically, malignant melanomas also do not label for cytokeratins or pan-lymphocyte common antigen, findings that were also seen in this case.

Melanoma is most commonly diagnosed as a primary cutaneous lesion, and the recommended diagnostic maneuver is full thickness excisional biopsy. The mainstay for metastatic disease is medical therapy. However, there is a role for surgical resection if disease is limited to one organ. Further, neoadjuvant therapy should be evaluated in this setting. In particular, if metastatic disease resection can improve quality of life it should be considered. If metastatic foci are not amendable to excision, radiation therapy has been shown to be effective for local control and symptom improvement. 

At the time of presentation, our patient’s functional status was poor, (Eastern Cooperative Oncology Group functional status of 4), and he had evidence of extensive metastatic burden. Discussions at our Tumor Board, yielded the recommendation for radiation therapy, as we did not believe that the patient would tolerate either medical therapy or surgical resection.

**Discussion**

Melanoma continues to be a diagnostic challenge in medicine, pathology, radiology and with core biopsy directed at the rind of the mass, yielded ample tissue that was sent for immunohistochemical staining, confirming the diagnosis of malignant melanoma.

**Figure 1.** Erythematous and painful right axillary mass.

**Figure 2.** Aspirated cells stained for microphthalmia-associated transcription factor (MITF).

**Conflict of Interest**

None. There was no funding associated with this study.

**References**

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