Kaposi’s Sarcoma of the Bulbar Conjunctiva in an Immunocompetent Patient

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Key Words
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Abstract
Kaposi’s sarcoma is an uncommon endothelial malignant tumor, first described by Moricz Kaposi in 1872. It is frequently encountered in patients with acquired immunodeficiency syndrome (AIDS), immunosuppression or organ transplantation. Conjunctival Kaposi’s sarcoma is a rare tumor, mostly in patients with AIDS. We present a selected case with a superior bulbar conjunctival lesion in a patient who is HIV negative and immunocompetent, with an unremarkable general medical history. The lesion was surgically excised with no adjunctive treatment, and histological examination confirmed the diagnosis. There was no recurrence or metastasis 5 years after surgery.

Introduction
Kaposi’s sarcoma (KS) is an uncommon spindle cell malignancy of endothelial cell origin, first described by Moricz Kaposi in 1872. It is frequently encountered in patients with acquired immunodeficiency syndrome (AIDS), immunosuppression, organ transplantation or human herpes simplex virus-8 (HHV-8) infection [1–3]. Conjunctival sarcoma is rare and is encountered mostly in patients with AIDS, in some cases as the first clinical presentation [4, 5]. We report a case of an immunocompetent patient with a superior bulbar conjunctival KS, managed successfully with surgical excision alone.
Case Report

A 70-year-old male presented with a 3-month history of a gradually growing mass on his left superior bulbar conjunctiva (fig. 1a). Examination revealed a dark red, firm, painless mass with a thinner stalk attachment on the conjunctiva and no evidence of orbital or scleral invasion (fig. 1b). This was an isolated ocular involvement in a HIV-negative patient, with no clinical evidence of immunosuppression and unremarkable general health.

We applied our standard protocol for a potentially malignant conjunctival lesion. Surgical excision included a 2-mm margin of healthy tissue around the conjunctival stalk and the underlying Tenon’s capsule followed by meticulous hemostasis with a bipolar cautery. Additional intervention like cryotherapy or radiotherapy was reserved for a possible incomplete excision depending on the histological diagnosis. The bare sclera defect was allowed to granulate with secondary intention healing. Conjunctival surface reconstruction was avoided in order to monitor the surgical site for possible recurrence of the tumor. The bulbar conjunctiva healed with a smooth surface, no signs of inflammation and no evidence of recurrence 1 year after surgery (fig. 2a).

Histological examination confirmed the diagnosis of a stage b and c KS, characterized by the presence of spindle cells positive on CD31, CD34 and Ulex Europaeus markers with minimal pleomorphism and mitotic activity. They are forming slits containing extravasated red blood cells, whereas extracellular or intracellular periodic acid-Schiff-positive hyaline bodies can also be found [6] (fig. 2b). Histology revealed that the tumor was excised with a clear margin of healthy conjunctival tissue with no deep invasion to the underlying tissues. The specimen was also tested negative for HHV-8 infection by means of polymerase chain reaction (PCR). In view of the special clinical characteristics of such a malignancy, the patient was referred to the immunology and infectious diseases department for evaluation and subsequent postoperative follow-up of his general health. He was diagnosed immunocompetent with normal immunoglobulin levels and no evidence of HIV or other viral infection. His immune condition remained unremarkable throughout the observation period of 5 years.

Discussion

Ocular KS is rare in HIV-negative patients with less than 30 cases being reported before 1982. Only 20% of AIDS patients suffering from KS have ocular manifestations (lids or conjunctiva) and in 12% of cases ocular involvement is the primary manifestation of systemic disease [7–9].

Ocular lesions have slow growth and can be cured with simple surgical excision whereas cryotherapy is an adjunctive intervention which was not used in our case because of the clear tumor margins and the lack of systemic involvement. Radiotherapy with chemotherapy is reserved for multifocal aggressive disease [10–12].

Simple complete surgical excision was curative in our case with no evidence of recurrence or distant metastasis and no immune-related general health problems in the postoperative observation of 5 years. Differential diagnosis of similar lesions should include arteriovenous shunt, pyogenic granuloma, malignant blue nevus, bacterial angiomatosis and vascular malformations. Isolated KSs of the conjunctiva unrelated to AIDS have been reported previously [13].

This case may strengthen the evidence that such conjunctival malignancies may occur as an isolated incidence and do not necessarily relate to AIDS or immunosuppression [14]. Simple surgical excision of the conjunctival lesion, with no adjunctive measure, can be curative with no evidence of recurrence or distant metastasis in a long-term follow-up of 5 years.
Disclosure Statement

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Fig. 1. a Solid mass on the superior bulbar conjunctiva displacing the upper eyelid upwards. b CT scan imaging of the orbits shows no scleral or orbital extension of the lesion.
Fig. 2. a Appearance of the bulbar conjunctiva 12 months after surgery with little visible scar tissue and no evidence of recurrence. b Histology section shows the characteristic neoplastic spindle cells forming slits filled with extravasated red blood cells. HE, ×200.

References

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