Spindle cell carcinoma of the conjunctiva: A rare entity

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An 85-year-old male presented with painless bulging lesion over the cornea. Clinical history, diagnostic imaging studies, and histopathologic sections were evaluated. The patient clinically displayed an vascularized conjunctival lesion located at the superior bulbar conjunctiva with extension onto cornea covering 2/3 of his pupillary aperture superiorly. His visual acuity was counting fingers at 4 m. The patient underwent a total excision of the lesion including conjunctival and corneal parts. Histopathologic evaluation revealed spindle cell carcinoma which involves the whole conjunctival squamous epithelium with significant polarity loss, nuclear enlargement with hyperchromasia and pleomorphism, and mitotic activity. Diagnosis of spindle cell carcinoma is challenging because of overlapping histopathological features with other spindle cell tumors. The detailed pathologic examination is very important for the decision of proper treatment.

Key words: Conjunctiva, spindle cell carcinoma, squamous epithelium

Spindle cell carcinoma is a rare and unusual biphasic malignant tumor, which involves sarcomatoid proliferation of pleomorphic spindle cells and squamous cell carcinoma (SCC). SCC, with the spindle cell component, is an uncommon phenomenon and a rare type of malignant tumor.[1,2] Spindle cell carcinoma is a poorly differentiated variant of SCC that rarely occurs in the conjunctiva.[3,7]

We aimed to present a case with conjunctival spindle cell carcinoma to emphasize the importance of detailed pathologic examination to differentiate the cell type for the decision of proper treatment. Informed consent was obtained from the patient.

Case Report

An 85-year-old male referred to our clinic with decreased vision in the right eye. He did not have any history of trauma and did not complain about pain. He described a pedunculated lesion without any ulceration, which grew slowly over 3 months. In his ophthalmic examination, best-corrected visual acuity was counting fingers at 4 m in his right eye and 0.1 in his left eye. His intraocular pressure was 18 mmHg in both eyes. Anterior segment examination revealed a large vascularized lesion located in the superior bulbar conjunctiva with extension onto cornea closing 2/3 of the pupillary area [Fig. 1]. The left eye revealed no pathology in the anterior segment of the eye. Fundoscopic examination of the right eye could not be performed; on the left eye, optic nerve was pale. The patient underwent an excisional biopsy of the lesion removing the whole tumor in the conjunctiva as well as on the cornea and cryotherapy to the conjunctival margins.

Histopathologic evaluation revealed in situ carcinoma which holds the whole conjunctival squamous epithelium with significant polarity loss, nuclear enlargement with hyperchromasia and pleomorphism, and mitotic activity. The stroma was rich in atypical cells forming herds and bundles of spindle or epithelioid cells, with hyperchromatic nuclei and pleomorphism mixed with inflammatory cells [Fig. 2]. In immunohistochemical staining, atypical stromal cells were stained positive with vimentin, pancytokeratin (cytokeratin AE1/AE3), epithelial membrane antigen (EMA), smooth muscle actin (SMA), CD99, p63, and calponin and were stained negative with caldesmon and MyoD1. Positivity of EMA and pancytokeratin

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revealed the epithelial origin of the tumor [Fig. 3]. In addition, p63 positivity is indicated the squamous differentiation. Vimentin, SMA, and calponin positivity proved the sarcomatous component with myoepithelial differentiation [Fig. 4].

Because of the malignant potential of the lesion, an orbital magnetic resonance imaging was taken, which showed no posterior and orbital extension.

In his medical history, our patient had a nonmetastatic carcinomatous gingival lesion for which he had received radiotherapy 4 years ago. However, his conjunctival lesion was accepted as primary because the new lesion was far from the radiotherapy treatment area and in situ carcinoma was detected in the superficial epithelium.

After surgical excision, his vision improved to 0.2. He was referred to the oncology department for further evaluation. Unfortunately, the patient lost to follow-up. Contact with a relative revealed that he was taking home care because of severe senile health problems.

Discussion

SCC of the conjunctiva is a rare malignancy; however, it is reported to be the most common malignant tumor of the ocular surface.[8] SCC has the potential to penetrate the corneoscleral lamella into the anterior chamber and can breach the orbital septum to invade the soft tissues of the orbit, sinuses, and brain. These tumors may metastasize via lymphatics or blood during the disease.[9] Surgical excision with or without cryotherapy and radiotherapy remains the widely accepted treatment for SCC of the conjunctiva.[9,10]

Spindle cell carcinoma of the conjunctiva is relatively rare, with only a few cases reported in literature.[3-7] Cervantes et al. reported a total of 287 cases of SCC of the conjunctiva, in which only two cases were documented as spindle cell carcinoma.[11] Spindle cell carcinoma is a poorly differentiated variant of SCC which is considered to be more aggressive and can also affect the progress and outcome of the disease.
**Conclusion**

Because of their possible aggressive behavior, conjunctival malignancies are known to be sight- and life-threatening.\[9\] The detailed pathologic examination is very important to differentiate the cell type for the decision of proper treatment.

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**Conflicts of interest**

There are no conflicts of interest.

**References**