Primary malignant melanoma of lacrimal sac

Min Ren¹, Ji-Hong Zeng², Qing-Li Luo², Feng Bi¹, Jun Chen²

¹Department of Oncology, West China Hospital, Sichuan University, Chengdu 610041, Sichuan Province, China
²Department of Ophthalmology, West China Hospital, Sichuan University, Chengdu 610041, Sichuan Province, China

Correspondence to: Jun Chen. No.37, Guoxue Alley, Wuhou District, Chengdu 610041, Sichuan Province, China. clfx828@126.com

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Dear Sir,

We write to present a case report of primary malignant melanoma of the lacrimal sac. Tumors of the lacrimal sac are rare, and most of them are benign and of epithelial origin. Malignant melanoma of the lacrimal sac is even rarer accounting for 5% of lacrimal sac tumors and 0.7% of ocular melanoma. To our knowledge, only 27 cases with complete clinical information have ever been reported worldwide in English literature. The disease has an insidious onset and masquerades as a chronic dacryocystitis. As a result, it is difficult to identify at an early stage. However, an early diagnosis is essential, because it is potentially lethal progression if therapy is delayed or inadequate.

CASE HISTORY

A 65-year-old female patient complained of a 1y history of epiphora and a slowly progressive swell in the right medial canthal area (Figure 1A). Two months before seeking medical attention, she experienced occasional bloody discharge from the right eye and epistaxis from the right nasal cavity, without any pain or visual disturbance. The patient had no previous history of regional surgery or trauma, nor history of a cutaneous or mucosal primary melanoma. Physical examination revealed a localized well-defined non-tender palpable mass at the right medial canthus (Figure 1B). The mass had a relatively clear boundary, neither infiltrating the surrounding soft tissue nor extending into the orbital cavity or nasolacrimal duct. There was no associated enlargement of the cervical lymph nodes. A dacryocystectomy was performed. A dark-red soft mass, without surrounding tissue invasion, was removed from the fossa of the right lacrimal sac. Its transverse section was completely dark (Figure 1C). Intraoperative frozen section of the mass revealed a malignant melanoma. Wide local excision was then performed, including surrounding soft tissues within 5 mm and the local peristomeum. Histopathological examination of the surgical specimen showed solid growth of malignant cells characterized by melanin pigmentation (Figure 2A). The depigmented tumor cells showed hyperchromatic nuclei with prominent nucleoli and numerous heteromorphic nuclei (Figure 2B). The features confirmed the diagnosis of malignant melanoma. A complete metastatic workup including CT of the head and neck, chest, abdomen and pelvis failed to reveal lymphadenopathy or other evidence of metastasis. By the ninth post-operative month, the patient had no evidence of local recurrence or metastasis. One-year follow-up revealed local recurrence without metastasis (Figure 1D). Then, resection of the recrudescent melanoma and local adjuvant irradiation therapy were ordered.

DISCUSSION

The skin is most commonly affected by malignant melanoma, accounting for about 4% of all skin cancers and resulting in more than 77% of skin cancer deaths. Compared with cutaneous malignant melanoma, mucosal melanoma is rare and behaves far more aggressively. Patients with mucosal melanoma generally have a poorer prognosis despite aggressive treatment. Primary malignant melanoma of the lacrimal sac is extremely rare, because melanocytes are not normally found in the tissues of the lacrimal duct system. Most researchers believe that these melanocytes originate in the neural crest or migrate from the conjunctival sac during embryologic development.

Primary malignant melanomas of the lacrimal sac have an insidious onset and develop similar symptoms to chronic dacryocystitis, the features of which along with limited early visibility result in delayed diagnosis and poorer prognosis. Gleizal et al. noted that patients with primary malignant melanomas of the lacrimal sac presented with epiphora (65%), bloody discharge (35%) and pain (10%). Most patients develop a palpable or visible mass at the medial canthus, usually six months after epiphora. The average age of onset was 59y (range 38-80y). There was no significant gender difference.

The treatment varies with size and extent of the tumor, which may range from only a dacryocystectomy to wide en bloc resection, post-operative radiotherapy, chemotherapy and immunotherapy in a variety of combinations. Recently,
Shuman et al. retrospectively studied 52 patients with head and neck mucosal melanoma. With a median follow-up of 97 mo, they found that the median overall survival and the disease-free survival are 52 mo and 15 mo respectively and surgery is the primary treatment modality. Positive surgical margins predict poorer overall survival, but the outcomes of the patients requiring re-excision to achieve negative margins are not significantly different from those of the patients initially with negative surgical margins. Early diagnosis and thorough excision are the key factors associated with improved overall survival. They also demonstrated that adjuvant radiotherapy and/or chemotherapy is not linked with an improved outcome. However, some researchers thought otherwise, who believe high-energy radiotherapy to be beneficial, regardless of the fact that melanoma is found radioresistant, as wide surgical resection in the head and neck region is often difficult. Furthermore, Shuman suggested that a systemic treatment is needed because head and neck mucosal melanomas see high rates of locoregional recurrence and distant metastasis.

CONCLUSION

In summary, malignant melanoma of the lacrimal sac is extremely rare and could be difficult to diagnose because of its similar symptoms and signs to chronic dacryocystitis. In dealing with persistent dacryocystitis, especially in older patients, ophthalmologists should consider the possibility of a malignant melanoma. CT scanning is important for diagnosis. While the optimal management of the condition is not established, early diagnosis and wide excision with negative surgical margins seem to be the most effective approach. Further studies with larger patient populations and longer follow-up are needed to determine the outcomes of treatment.

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REFERENCES