A case of aberrant lacrimal gland and fistula

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SUMMARY A case of aberrant lacrimal gland and fistula sited extraorbitally is presented in view of its rare incidence.

An aberrant lacrimal gland is a rare abnormality. We report here a case observed in the Clinic of Ophthalmology, Numune Hospital, Ankara.

Case report

A 9-year-old boy was seen in September 1987 with the complaint of constant watering since birth from the middle of the right temporal region, where the skin was never free of tears. The watering increased on weeping.

On examination, a small orifice could be seen in the skin in the middle of the right temporal region. The opening was round and 1 mm in diameter. Clear transparent fluid could be seen continually flowing from the orifice (Fig. 1). There was no evidence of excoriation, inflammation, or swelling round the opening. Apart from this condition both eyes, including the lacrimal apparatus, were normal in all respects. There was no other congenital abnormality in the eyes or elsewhere.

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Fig. 1 Clear transparent fluid continually discharged from the orifice.

A quantitative biochemical assay of the transparent fluid coming from the orifice, which was consistent with that of tears, was done. A fine catheter could be passed into the fistula for about 0.5 cm in the direction of the external canthus. Radiological examination after injection of radio-opaque dye showed the dye in the fistula. The investigations confirmed that the fluid coming from fistula was tear secreted by an aberrant lacrimal gland.

The patient was operated upon. After skin incision had been made and a probe passed, the fistula and the gland, which was 0.5×0.8×1.6 cm in size, were dissected out from the surrounding tissues. The gland with the fistula was completely removed. The postoperative course was normal.

Microscopic examination of the specimen showed a glandular tissue similar to that of lacrimal gland (Fig. 2).

Discussion

An aberrant lacrimal gland is an unusual congenital and developmental abnormality and differs from an ectopic gland.1 The first case of aberrant lacrimal gland associated with other congenital abnormalities was reported by Gördüren in 1962 in Turkey,2 and in 1963 a second case was also reported by the same author.3 Firat and Emüler4 described two cases of aberrant lacrimal gland in the bulbar conjunctiva. Slem et al.5 reported a case of aberrant lacrimal gland in the orbit.6 Our case is a typical example of an aberrant lacrimal gland, and we believe it to be the sixth case from this country to be published.

Dr Ayşe Ayhan examined the sections of the specimen and we are indebted to her for the histopathological diagnosis.
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Fig. 2 Glandular acini observed in the microscopic sections. (Haematoxylin and eosin, ×240.)

References


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