Ciliary body adenoma of non-pigmented epithelium
S Mansoor, A Qureshi

CASE REPORT

A 27 year old man presented with blurred vision of two years duration. There was no history of trauma, diabetes, or hypertension. Visual activity was reduced on the right and normal on the left. Intraocular pressure was normal. The right eye had a cataract. Gonioscopy showed a greyish white growth arising from the ciliary body, measuring approximately 1.5 × 0.5 cm, encroaching upon the lens and displacing it temporally. The angles were open and there was no hyperpigmentation. B scan revealed a growth arising from behind the iris from the ciliary body. The retina was intact. The patient underwent surgery, and sectoral cyclectomy with removal of the growth was performed.

PATHOLOGICAL FINDINGS

The ciliary body mass removed consisted of irregular, tan grey, non-pigmented tissue measuring 1.5 × 0.5 × 0.3 cm. Microscopically, the tumour was arranged predominantly in a tubular and papillary configuration. The glandular lumina and intervening areas were composed of mucoid fibrillary material (fig 1). The cells lining these structures were round to polygonal, with round, bland nuclei and abundant eosinophilic cytoplasm. Mitoses were rare. Rare nuclei were enlarged. No significant inflammatory cell component was seen. Areas of calcification were present. The results of immunohistochemical studies were consistent with an origin from the non-pigmented ciliary epithelium (NPCE). The HMB45 antibody was negative, and the tumour stained positive for S100, vimentin, and cytokeratin (fig 2).

DISCUSSION

Primary tumours of the non-pigmented ciliary epithelium are rare. Zimmerman1 proposed dividing these tumours into two classes—congenital and acquired. Congenital tumours arise in the embryonic or early postnatal period, and include medulloepithelioma and glioneuromas. The acquired tumours are seen in adulthood and include Fuch’s adenoma (hyperplasia of the NPCE, pseudoadenomatous hyperplasia, and coronal adenoma),1–3 adenoma, and adenocarcinoma of the non-pigmented ciliary epithelium.

Acquired tumours of the ciliary body have a wide spectrum of clinical and histopathological features and biological behaviour. Many are rare, and most studies have consisted of single case reports.4–6 Classification is based on the presence or absence of pigmentation, the cellular pattern, and malignant behaviour. Acquired neoplasms of the NPCE include Fuchs adenoma, adenoma, and carcinoma of the NPCE.1–5

“Although benign, adenoma of the non-pigmented ciliary epithelium can behave aggressively locally, causing cataract and vitreous haemorrhage”

Our patient developed a true adenoma of the NPCE, a benign tumour with variable histopathological features. The immunohistochemical results were consistent with an origin from the NPCE: S-100 and vimentin were positive. Immunohistochemistry for cytokeratin was also positive—variable staining for cytokeratin has been described in the literature.7 Although benign, adenoma of the NPCE can behave aggressively locally, causing cataract and vitreous haemorrhage.
Adenomas are differentiated from adenocarcinoma of the NPCE by the absence of local infiltrative behaviour and a moderate degree of cellular pleomorphism with nuclear atypia. Rare mitoses are found. This tumour is seen in adulthood, with an age range of 24–70 years (mean, 45), and no sex predilection. The most common presenting symptom is visual loss. A ciliary body adenoma with smooth muscle differentiation has also been reported.

Histopathological differential diagnosis of ciliary body adenoma includes ciliary body adenocarcinoma, adenoma of pigment epithelium, medulloepithelioma, in addition to lesions that occur in the deeper uveal tissue, such as melanoma, melanocytoma, metastatic carcinoma, leiomyoma, and neurilemoma.

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