Introduction

Medulloepithelioma presents as an intraocular congenital tumor that typically presents during the first decade, is a tumor of the non-pigmented ciliary epithelium. WHO International histological classification of tumors of the eye states two separate classifying entities of medulloepithelioma both teratoid or nonteratoid which can present either as a benign or a malignant variant.1

Small medulloepitheliomas often go undetected due to anterior placement of iris. Even after symptoms develop, clinical suspicion of tumor is often overlooked due to wide variation in presentation. Medulloepitheliomas being slow-growing tumors are not visible until they enlarge enough to protrude into the pupil, distort the iris, or invade adjacent tissues.1 The decreased vision in medulloepithelioma are due to subluxation and dislocation of the lens with cataract formation. With subsequent ciliary body epithelium involvement, the aqueous production is hampered, leading to signs of glaucoma as well as hypotony.

Acquired tumors of the nonpigmented ciliary epithelium are classified as adenomatous hyperplasia (Fuchs adenoma), adenoma, and carcinoma.2 Surgical enucleation is considered standard therapy for intraocular medulloepithelioma after a diagnosis is established. Extrascleal extension occurs in 10-19% of cases3 and carries a poor prognosis often leading to metastasis at various sites and death.

We present a peculiar case of a patient presenting only with loss of vision following blunt trauma with underlying non-teratoid regionally malignant medulloepithelioma which metastasised to the cervical and sub mandibular lymph nodes that presented only with trauma and was completely unaware about the diseased condition. The diagnosis of an underlying tumor was only made subsequently. Due to the peculiarity of the tumor presentation, diagnosis is often delayed.

Case report

A healthy 4-year old girl presented with blunt trauma injury to her left eye by falling off the staircase following which she reported to the OPD on November 2014 (Figure 1). On examination, the patient had Perception of light, lens substance was dislocated and pushed posteriorly, iris rupture with neovascularization was noted while only the temporal pupil was reacting to light. The cornea was clear bilaterally. Due to significant media opacity, the posterior segment examination was deferred clinically by bio microscopy and a USG B-scan was advised. Examination of his right eye was clinically sound. He presented with no systemic ailments or any significant previously treated medical or ophthalmic history.

On USG B scan, Posterior capsule integrity seemed lost, with lens matter in anterior part of vitreous cavity and retina was attached (Figure 2). On November 24, the child was undertaken for...
a Pars Plana Lensectomy and vitrectomy. On table a mass was seen extending from the ciliary body circumferentially and a biopsy was taken. Histopathology reports revealed cluster of some round and epithelial cells. Cells were basophilic and pleomorphic and showed some pigmented cells also (Figure 3) (Figure 4).

Figure 2 On USG B scan, lens matter was seen displaced into the anterior part vitreous cavity, with loss of posterior capsule integrity.

Figure 3 Biopsy of iris tissue showing cluster of round and epithelial cells.

Figure 4 Giemsa stain revealed basophilic cells showing pleomorphism.

Post operatively, MRI was advised of brain and orbit and it revealed a residual mass extending into anterior and lateral aspect of the globe (Figure 5). Regional metastasis to the cervical and sub-mandibular lymph nodes were noted and it nearly confirmed the diagnosis.

Due to histologically malignant features and a radiological confirmation, an enucleation of the left eye was planned under GA (Figure 6).

There occurred no recurrence in the subsequent follow up visits and patient was doing well and thus, this trauma prevented the further dire consequences of a Medulloepithelioma.

**Discussion**

Due to the unavailability in singular characteristics of medulloepithelioma, there is limited information on long-term survival. Metastatic intraocular medulloepithelioma is a very rare disease, so there have been no performed clinical human and animal trials to reinforce evidence-based therapy. This case highlights medulloepithelioma of the ciliary body in a 4-year-old child which masquerades as a dislocated lens matter along with neovascularization. The presenting complain of an ipsilateral blunt ocular trauma was initially assumed to be responsible for the diagnosis of dislocated lens with subsequent secondary inflammatory pain. Ciliary body medulloepithelioma being the commonest ciliary body neoplasm in childhood, the term “medulloepithelioma” was coined in 1930s by Grinker. It is thought to arise from the undifferentiated medullary tissue of the embryonic retinal pigment epithelium destined to form the nonpigmented ciliary body epithelium subsequently after birth. It is most commonly found at the ciliary body. It can have a presentation of a benign or malignant like condition or may also exhibit teratoid features. Several clinical features suspicious of ciliary body medulloepitheliomas, were present in our case. These tumors occur mostly during the first decade of life as ipsilateral,
solitary flesh-coloured solid lobulated mass arising from the ciliary body. Other differential diagnoses simulating a ciliary body mass in the first decade, such as retinoblastoma, malignant melanoma, juvenile xanthogranuloma, or ocular metastatic lesion, should also be considered. Due to the close juxtapostioning of the tumor to the intraocular contents complications such as cataract, ectopia lentis, secondary rubeotic iris may present former to ciliary body mass presentation. Most commonly involved structure is the lens and can present with a manifold of lens-related pathological entities such as a lens opacification, lens notching (referred to as a “lens coloboma”) and ectopia lentis. Thus, when noted a coloboma of lens without associated coloboma of uveal tissue and optic nerve head should prompt an underlying ciliary body medulloepithelioma. Given the presenting history of an ipsilateral blunt ocular injury in our case, the initial diagnosis of a traumatic cataract was very rational. The pre operative observations of iris tissue neovascularization with cysts on table might possibly have suggested of an underlying neoplastic process. With an earlier clinical diagnosis of an underlying neoplastic condition, the pars plana lensectomy would not have been the probable treatment of choice.

Surgical removal of the tumor thus provided a definitive histological diagnosis with a cluster of basophilic cells showing pleomorphism. The characteristic rosettes and fleurettes of retinoblastoma maybe seen occasionally but retinoblastoma have an eosinophilic content on contrary to medulloepithelioma. The final call for enucleation of the diseased eye is usually based on a painful eye with poor visual potential, and a strong clinical suspicion of malignancy showing voluminous tumor growth. The ciliary body medulloepitheliomas being locally invasive, complete excision is curative and associated with a good disease free survival. In our case, due to an early diagnosis the local scleral and extra scleral invasion of the tumor was hampered, thereby not necessitating the need for adjuvant chemotherapy after enucleation.

Conclusion

On conclusion, the differential diagnosis of an intraocular neoplasm should be considered when a child presents with a masquerade of severe chronic uveitis associated with cyclitic and fibrotic membranes, with other features of IOP alteration. An ultrasound biomicroscopy should be carried out to exclude a ciliary body neoplasm due to a posterior location of the ciliary body which is manner times missed on a conventional USG B-scan.

Acknowledgements

None.

Conflict of interest

The author declares no conflict of interest.

References