Medulloepithelioma : Unsuspected intraocular tumor in children.

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Two cases of medulloepithelioma of the eye are described. Both patients presented with leukocoria, endophthalmitis, and intraocular mass. The lesions were mistaken as retinoblastoma on clinical grounds. Clinicians should be aware of this unusual tumor when children, particularly those in the first decade of life, present with such clinical symptoms.

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รายงานผู้ป่วยเด็ก 2 ราย ซึ่งมาจากอินดิเณอร์เรื่อง leukocoria, endophthalmitis, และ intraocular
mass ผู้ป่วยเด็กกล่าวให้รับการผ่าตัดที่ต้นประวัติและเด็กหนุ่มกว้างควรพบต่างๆ ที่เป็น retinoblastoma ซึ่งเป็น
เนื้อเยื่อในพื้นที่พบอยู่ในเด็ก ผ่านการตรวจทางตาสัณห์ระหว่างการผ่าตัด ผู้ป่วยที่ต้องการผ่าตัดที่ต้นประวัติ
ของเด็ก (enucleation) พบว่าเป็น medulloepithelioma ซึ่งเป็นเนื้องอกที่หนาที่พบในเด็กซึ่งอาการ
ทั้งความรู้สึกและเนื้องอกที่มีต่อเนื่อง 3 รายในประเทศไทย ผู้ป่วยมักเป็นเด็กเล็ก หรือเด็กเล็กที่มีอายุน้อยประมาณ
5 ปี อักขรรค์ที่เนื้องอกนี้ดีเสียดีกว่า/do not use ในผู้ป่วยที่มีอาการปัญหาทางเดินกล้าว


Medulloepithelioma or diktyoma of the eye is an uncommon neoplasm usually occurring in young children.\(^1\)\(^2\) To our knowledge only one example has been recorded previously in Thailand.\(^2\) In this communication we have encountered two additional cases. The related medical literature is reviewed.

**CASE REPORT**

**CASE 1**

A 15-month-old girl was hospitalized in August 1984 because of endophthalmitis and leukocoria in the right eye. Her mother had observed that the baby developed a red eye and a white spot in the pupil for 2 months. She received antibiotic and steroid drugs from the ophthalmologists without clinical improvement. The patient was then referred to Chulalongkorn Hospital for further management.

Examination revealed conjunctivitis, corneal edema, flat anterior chamber, rubeosis iridis, seclusio pupil, yellow and opaque lens of the right eye. Tactile tension was normal. The left eye was unremarkable.

Routine laboratory data were within normal limits. Roentgenograms of the chest, skull, and long bones showed no evidence of metastases. Ultrasonogram of the right eye disclosed a retro lental high density mass of 5 mm. in axial length, not connected to the retina or choroid. The vitreous was clear an the thickening choroid was 2 mm. The clinical diagnosis was retinoblastoma. The child underwent an enucleation of the right eye. The postoperative course was uncomplicated. The patient had been followed up for 6 months and remained clinically silent.

The specimen of the tissue in this case as well as in the other (case 2) were fixed in 10% formalin, embedded in paraffin and stained with hematoxylin and eosin. A small 0.7 cm. gray-white nodule was located in the retro lental space of the enucleated eyeball (Fig.1A). Microscopically, the lesion consisted of columnar cells with oval nuclei. The tumor cells were arranged in tubular and papillary pattern with distinct limiting membrane resembling the medullary epithelium of embryonic non-pigmented ciliary epithelium (Fig.1B). Mitotic activity was absent.

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**Figure 1A.** Macrophotomicrograph of retro lental medulloepithelioma (case 1).

**Figure 1B.** A complex convolution of columnar epithelium with distinct limiting membrane. (H&E × 100)
The pathologic diagnosis was benign non-teratoid medulloepithelioma of the right eye.

CASE 2

A 30-month-old girl presented with redness in the right eye, lid edema, and fever for 2 months following a history of trauma to the right eye. She received medications with some clinical improvement. Two weeks later, a small nodule was noted in the pupil of the right eye. The lesion was slowly enlarging. A computed tomographic (CT) scan disclosed increased density and enhancement of the ciliary muscle, anterior chamber and lens of the right globe. The patient was referred to Chulalongkorn Hospital in April 1989 for further management of endophthalmitis and intraocular mass of her right eye.

Examination revealed posterior synerchia of the iris, leukocoria and a mass of 10 disc in diameter which was seen on the optic disc of the right eye. The left eye was normal.

Routine laboratory data were unremarkable. Ultrasonogram demonstrated thickened retina and choroid with calcification. The clinical diagnosis was retinoblastoma. An enucleation of the right eye was performed. The postoperative course was uneventful.

The enucleated eyeball showed a 1.5 cm. grayish white irregular nodule in the retro-lental space (Fig 2). A 0.5 cm. optic nerve was found without gross abnormalities. Microscopically, the tumor displayed a complex convolution of columnar epithelium similar to the lesion in case 1(Fig.3). The inner free surface of the tubules was bounded by a limiting membrane while on the opposite surface, the cells rested on the supporting stroma. Rosette formation and mitoses were absent.

Figure 2. Gross appearance of diktyoma (case 2). The growth appears as an irregular grayish white mass in the retrolental space. The optic nerve is unremarkable.
Figure 3. Layer of columnar cells with oval nuclei presenting a tubular arrangement. (H&E × 100)

The pathologic diagnosis was benign non-teratoid medulloepithelioma of the right eye.

DISCUSSION

Medulloepithelioma has been recognized as a distinctive entity. Diagnosis rests on the characteristic morphological features including the pattern of papillary and tubular arrangement of the primitive non-pigmented ciliary epithelium with a distinct limiting membrane. In the cases presented, the histopathological findings conformed in every detail with this unusual neoplasm.

Medulloepithelioma more commonly has been referred to as “diktyoma” which means net-like. The latter term was introduced by Fuchs in 1908. Grinker proposed the term “medulloepithelioma” because the author considered that this tumor arises from non-pigmented ciliary epithelium which persisted in an undifferentiated form resembling medullary epithelium throughout adult life. According to Zimmerman, medulloepithelioma was a congenital tumor which developed from the immature ciliary epithelium, either during embryonic development in utero or from a persistent anlage of this undifferentiated epithelium during the early postnatal period. The symptoms and signs may not appear until long after birth. Diktyoma may arise from various parts of the eye including the iris, optic nerve, and retina. The tumor has been classified into teratoid and nonteratoid types and each further divided into benign or malignant varieties. Teratoid group contains one or more heteropic tissues that are not normally present in the eyes such as hyaline cartilage, rhabdomyoblasts and cerebral tissues. Nonteratoid group has tissue that resemble the medullary epithelium as in our cases but may contain tissue derived from secondary optic vesicle such as retinal pigmented epithelium, ciliary epithelium, vitreous, and neuroglia. Malignant transformation should be considered when local invasion, frequent
mitoses, and distant metastases are identified. To date more than 100 cases of diktyoma have been published, the majority being in the infants and children with an average age of 5 years. In Thailand there are only 3 cases, including our 2 examples, in which medulloepithelioma has been described. Both cases in our studies were well within the age range reported by others. There is no sex predilection although our patients were both girls.

The tumor is often congenital and occurs spontaneously without any predisposing factors. On reviewing the literature, only 2 cases including our (case 2) had been found after trauma. The relation between medulloepithelioma and trauma is unclear and could be a coincidental finding.

Clinically, most patients presented with leukocoria, decrease visual acuity, signs of severe inflammation and intraocular mass as noted in our examples. Clinicians should keep this unusual tumor in mind when children present with such clinical manifestations.

The ultrasonographic findings and CT features were nondiagnostic in our cases. However Orellana et al have reported the correct pre-enucleation diagnosis of medulloepithelioma in a case of an 8-year-old boy, using ultrasonography and cytologic examination of the vitreous aspirate. It should be noted that the diagnosis is usually established histologically after the eye is enucleated and our cases are unexceptional. Only infrequently has this neoplasm been diagnosed before surgery. Tissue examination thus is essential for proper diagnosis and also assists the surgeons and the family to predict the prognosis. Enucleation is curative for this benign non-teratoid tumor.

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