Introduction

Circumscribed pigmented lesions of the ocular fundus are not uncommon. They can be congenital, developmental, neoplastic or the result of infectious or inflammatory processes. Pigmented neoplastic lesions of ocular fundus, both benign and malignant, can arise from the retinal pigment epithelium (RPE) or from choroidal melanocytes. It is important for the ophthalmologist to properly diagnose these lesions to avoid erroneous diagnosis and misdirected treatment.

Discussion

Lesions Arising from the Retinal Pigment Epithelium

Congenital hypertrophy of the Retinal Pigment Epithelium (CHRPE) is an uncommon benign fundus lesion. The typical unifocal, unilateral lesion occurs in about 0.5% of individuals [1]. CHRPE lesions are usually asymptomatic and detected on routine ophthalmic examination. The typical unifocal CHRPE is a gray to black, well-defined, minimally elevated fundus lesion having a diameter in the range of 2–5 mm (Figure 1). The lesion frequently undergoes at least partial depigmentation with aging, developing discrete intraretinal atrophic foci (lacunae that tend to enlarge and coalesce over the years). No diagnostic studies are generally indicated for characterization of these retinal lesions.

Fluorescein angiography (FA) and indocyanine green angiography (ICGA) show a well-defined blocking defect at the retinal pigment epithelial level. Transmission of choroidal fluorescence takes place through lacunae in the lesion [2].
region of the fundus. These lesions are hypofluorescent on FA and hypoautofluorescent on fundus autofluorescence (FAF) [3]. Atypical multifocal bilateral CHRPE presents oval or irregularly shaped lesions, variable in size (usually between 0.1 and 2.0 mm in maximal diameter), widely separated rather than clustered; it is associated with familial colonic adenomatous polyposis syndrome [1].

Reactive Hyperplasia of the Retinal Pigment Epithelium is a result of an ocular insult, such as inflammation or trauma. It can appear as a diffuse or sessile lesion or as a nodular mass. Its dark black color and associated signs of prior inflammation or trauma can help to differentiate the lesion from choroidal melanoma. Major complications or progression are quite unusual [4].

Adenoma and adenocarcinoma of the RPE have respectively benign and malignant histopathologic features and they are generally diagnosed in adulthood, with a mean age of 53 years [5]. Unlike uveal melanoma, these tumors seem to have no predilection for race. They may have a slight predilection for females [6]. Clinically, these lesions are usually solitary and unilateral and begin as a small, deep retinal tumor that is dark brown to black in color. They usually grow very slowly and invade the overlying sensory retina. Some instances, they have arisen from solitary congenital hypertrophy of the RPE [7]. FA can document the retinal feeder vessels and show early hypofluorescence and late minimal hyperfluorescence of the tumor, without visibility of choroidal vessels. Ultrasonography (US) typically demonstrates the tumor to be elevated and to have medium to high internal reflectivity. Enhanced depth imaging optical coherence tomography (EDI-OCT) shows the tumor surface as irregular and dense posterior optical shadowing (Figure 2) [6].

Combined hamartoma of the retina and RPE is a congenital, nonhereditary lesion with an uncertain etiology. This lesion is composed of an admixture of pigment epithelial cells, proliferating blood vessels and glial tissue. It appears as an ill-defined gray retinal mass with tortuous or straightened retinal blood vessels and it is characteristically located on or adjacent to the optic disc, but it can be seen in extrapapillary areas of the fundus. FA shows abnormal retinal blood vessels in the mass and gradual late staining of the lesion. Optical coherence tomography (OCT) shows an irregular lesion with vitreoretinal traction [8].

Lesions Arising from Choroidal Melanocytes

Choroidal nevus is the most common primary intraocular tumor. In Caucasians, it has been estimated that approximately 2 to 10% of persons older than 50 years have at least one choroidal nevus [9]. It is much more common in persons of lightly pigmented races and it appears to be equally common in men and women. The typical choroidal nevus appears as a small gray to brown tumor. Many choroidal nevi exhibit characteristic surface alterations such as drusen and RPE pigment clumping. Most choroidal nevi are 5 mm or less in basal diameter and 1 mm or less in thickness. Blurred or distorted vision attributable to a choroidal nevus may be the result of an accumulation of serous subretinal fluid [10], cystic degeneration of the retina overlying a macular choroidal nevus [11] or choroidal neovascularization [12]. Diagnosis of a typical choroidal nevus is based almost exclusively on clinical features; fundus photograph can be useful to compare subsequent follow-up evaluations.

US is commonly used for the measurement of tumor thickness; most choroidal nevi are nearly flat and can be difficult to detect. Choroidal nevus is characterized by high internal reflectivity on US. On FA nevus remains typically hypofluorescent. EDI-OCT can image choroidal nevi, even those undetectable by US: choriocapillaris thinning over the nevus, partial or complete choroidal shadowing deep to the nevus, RPE atrophy, photoreceptor loss and inner segment–outer segment junction irregularity are some of the EDI-OCT features of choroidal nevus (Figure 3). On FAF imaging pigmented nevus generally appears more hypoautofluorescent from the intrinsic melanin that absorbs light compared with less pigmented nevi that appears more hyper-autofluorescent. The estimated annual rate of malignant transformation of a pre-existing nevus into melanoma reported in several studies varied from 0.2 to 1% [13].

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cells). Clinically, the melanocytoma is a dark brown tumor that involves the substance of the optic disc. The surface of the tumor commonly appears striated because of insinuation of the darkly pigmented cells of the tumor between axons in the nerve fiber layer. If the melanocytoma compresses the optic disc, it can produce prominent visual field defects. Lesion enlargement can occur but is usually slow. The diagnosis can be made by opthalmoscopic recognition of its characteristic clinical features. In most cases, FA of a melanocytoma of the optic disk demonstrates hypofluorescence throughout the angiogram, because the cells are deeply pigmented and closely compact with relatively little vascularity. In cases with optic disk edema, there is hyperfluorescence of the edematous portion of the optic disk. ICGA also shows the lesion to be generally hypofluorescent. US shows high reflectivity of the lesion (Figure 4). Malignant change is estimated to occur in about 1-2% of cases [14].

Choroidal melanoma is the most common primary intraocular malignant tumor, with an incidence of 5.1 cases per million per year [15] Choroidal melanoma can originate from a pre-existing choroidal nevus. In 1995, Shields et al. identified several risk factors predictive of nevus growth into melanoma: thickness more than 2 mm, subretinal fluid, symptoms of flashes/floaters and blurred vision, orange pigment, margin less than or equal to 3 mm from optic disk, ultrasonographic hollowness and absence of halo. Tumors with three or more risk factors likely represent small melanoma, demonstrate malignant transformation in 50% of cases at 5 years and treatment should be considered early [16]. The classic appearance of choroidal melanoma is pigmented domed-shaped or mushroom-shaped tumor with an associated exudative retinal detachment.

The mushroom-shaped configuration is due to the breaking of the overlying Bruch’s membrane and RPE, forming a nodular eruption beneath the retina that progressively enlarge. Although some choroidal melanomas are detected on ophthalmic evaluations prompted by visual symptoms (e.g., blurred vision, visual field defect, flashes, floaters), many patients who have such a tumor are asymptomatic at the time of detection. Small choroidal melanomas can be detected by EDI-OCT and it is also a useful tool for differentiating choroidal nevus from small choroidal melanoma. It has limited use with heavily pigmented tumors, which cast a significant posterior shadow. US is an important diagnostic tool used to define tumor extent and shape, and to measure tumor dimensions of choroidal melanoma. Typical features of a posterior uveal melanoma on US include acoustic hollowing, choroidal excavation and orbital shadowing (Figure 5). Spontaneous vascular movement may be noted in a highly vascularized tumor.

On A-scan the four cardinal acoustic hallmarks of malignant melanoma are a regular internal structure with similar height of the inner tumor spikes or regular decrease in height, low to medium reflectivity, solid consistency with no after movement of tumor spikes, and echographic sign of vascularization with a fast, spontaneous, continuous, flickering vertical motion of single tumor spikes. The characteristic features on FA include gradually increasing fluorescence that starts at or before the arterial phase, increases in intensity well into the recirculation phase, and persists for at least 45 min. The tumors that have broken through Bruch’s membrane characteristically reveal “double circulation” pattern because of superimposition of the intravascular fluorescence of the intact retinal vasculature over the fluorescence of large caliber vessels within choroidal tumor. ICGA allows better visualization of intrinsic vasculature in choroidal melanoma with maximal fluorescence at an average of 18 min from injection. Choroidal melanoma generally shows on FAF slight intrinsic hyperautofluorescence and the brightness increases with pigmented tumours, larger tumours, and those associated with disrupted RPE [17, 18].

**Pseudomelanoma**

There are conditions that can clinically simulate melanoma, leading to diagnostic ambiguity. The most frequent pseudomelanomas included choroidal nevus, peripheral exudative hemorrhagic chorioretinopathy (PEHCR), congenital hypertrophy of the retinal pigment epithelium and idiopathic hemorrhagic detachment of the retina or RPE [18] This section covers selected nonneoplastic-simulating conditions that are not discussed elsewhere in this article. Peripheral exudative hemorrhagic chorioretinopathy (PEHCR) is a peripheral retinal degeneration of the elderly manifesting with exudative and hemorrhagic changes.
This condition can closely simulate melanoma with a homogenous deep brown mass in the periphery (Figure 6). However, PEHCR shows blockage on FA, dark hypautofluorescence on FAF, echodensity with retraction cleft on US and spontaneous resolution [19].

Figure 6: Fundus photography (Left) and US imaging (Right) of PEHCR.

Idiopathic hemorrhagic detachment of the retina occurs after bleeding into the subretinal space and in some cases large amount of blood may accumulate, creating a hemorrhagic detachment of the retina, whereas idiopathic hemorrhagic detachment of the retinal pigment epithelium occurs in a variety of conditions such as senile disciform macular degeneration and angiod streaks, and it may complicate other conditions characterized by breaks in Bruch’s membrane. In many of these cases there is a clinically detectable source of the blood, such as an intraretinal microaneurysm [20].

Conclusion

A variety of pigmented lesions can simulate choroidal melanoma. Suspicious choroidal nevus is still the lesion most difficult to differentiate from choroidal melanoma. Most other pseudomelanomas can lead to diagnostic ambiguity. Some elements can still generate confusion, but careful assessment of the clinical and instrumental characteristics can help to avoid diagnostic errors.

References


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