Ultrasound biomicroscopy in the management of melanocytoma of the ciliary body with extrascleral extension

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Aim: To demonstrate the ultrasound biomicroscopic features of a ciliary body melanocytoma with extrascleral extension, and a conservative approach in its management.

Method: Observational case reports. Two cases of ciliary body melanocytoma were suspected at presentation, confirmed histologically by biopsy, and subsequently monitored for change by serial ultrasound biomicroscopic imaging. The main outcome measures were anatomical and functional preservation of the eye, with avoidance of formal surgical excision.

Results: Ultrasound biomicroscopy allows clear visualisation of the tumours, and the ultrasound characteristic is of low homogeneous internal reflectivity. 5 year follow up with observation only demonstrates success with this conservative management approach. Histopathological evaluation confirmed melanocytoma.

Conclusions: Melanocytoma is a rare tumour. However if considered in the differential diagnosis at presentation and confirmed histologically, further management with use of the ultrasound biomicroscope as an accurate mode of imaging is an acceptable technique for preservation of the eye and avoids surgical excision.

Melanocytoma1 (magnocellular naevus2) is an uncommon tumour3 of the uveal tract which is generally regarded as benign, having the ability for invasion of local tissues,4 but has never been shown to metastasise.1 Classically it is described at the optic nerve head,2 but it is encountered less frequently at the ciliary body6 and there are rare reports of occurrence in the iris,7 sclera,8 conjunctiva,9 and choroid.10

Despite diagnostic accuracy being high in ocular melanotic lesions,11 in previous reports the diagnosis of melanocytoma has been made subsequent to surgical intervention where the tumour has been thought to be malignant melanoma.11 12 Here we report two cases where the diagnosis of melanocytoma of the ciliary body with extrascleral extension was suspected at presentation and confirmed histologically with a biopsy of the extrascleral tumour extension.13

Ultrasound biomicroscopy is a particularly valuable technique for high resolution evaluation of anterior segment tumours in vivo.14 The aim of this report is to exhibit ultrasound biomicroscopic images and features of this tumour, and demonstrate a novel conservative approach to its management by serial ultrasound biomicroscopic imaging.

Patients and Methods

Case 1

A 57 year old white man was referred to the ocular oncology service for evaluation of an asymptomatic pigmented lesion on the superior aspect of his left sclera, present for 10 years, and associated with a ciliary body mass.

On examination his visual acuity was 20/15 bilaterally, and an area of jet black pigmentation was present at the 1 o’clock position on his sclera around an emissary vein. Gonioscopy showed pigmentation in the adjacent drainage angle, but no pigment dispersion, and a normal intraocular pressure (16 mm Hg). A small pigmented tumour was visible arising from the ciliary body.

Ultrasound biomicroscopy was performed, which showed a solid ciliary body tumour with low internal reflectivity and a small anterior cystic element (fig 1A). The remainder of the ocular examination was unremarkable.

In view of the long history of the intensely pigmented episcleral lesion, melanocytoma was suspected and a biopsy was performed.

Pathological findings

Macroscopically a piece of deeply pigmented tissue 3.0 mm×2.0 mm was examined. Microscopically this consisted of plump polyhedral or round heavily pigmented cells with abundant cytoplasm in which nuclear detail was mostly obscured by granular melanosomes (fig 1C). Bleached preparations show round and ovoid nuclei with no nucleoli in the majority of cells (fig 1D). The features are consistent with the diagnosis of melanocytoma; this was confirmed by two independent oculic pathologists.

Follow up

At 5 year follow up, this patient continues to maintain normal visual acuity, and serial ultrasound biomicroscopic imaging has failed to show extension of the lesion (fig 1B).

Case 2

A 77 year old white man was referred with a pigmented lesion on the superonasal aspect of his right sclera.

On examination his visual acuity was 20/30 bilaterally, and an area of deep pigmentation was noted at the 1 o’clock position close to the limbus. Gonioscopy showed increased local drainage angle pigmentation, no pigment dispersion, and normal intraocular pressure. No obvious ciliary body mass was noted. The remainder of the ocular examination was normal.

Ultrasound biomicroscopy revealed a ciliary body mass underlying the episcleral pigmentation, and it also had a homogeneous appearance with low internal reflectivity, measuring 4.0 mm×1.7 mm.

Pathological findings

Macroscopic examination shows a piece of darkly pigmented tissue, measuring 2.0 mm×1.0 mm, which microscopically is composed of highly pigmented plump polyhedral cells. The nuclei are mostly obscured by granular melanosomes, which
on bleached preparations show small round and ovoid nuclei with no nucleoli. Some non-pigmented cells with elongated nuclei were seen in adjacent tissue. No neoplastic features were seen. The appearances are consistent with melanocytoma; this was also confirmed by two independent ocular pathologists.

Follow up
Similarly, at 5 year follow up, there is no deterioration in visual acuity, or change in appearance of the lesion on successive ultrasound biomicroscopy.

DISCUSSION
This report demonstrates ultrasound biomicroscopic images of a ciliary body melanocytoma and the ultrasound image characteristics as observed by this relatively new imaging technique. Also, we have shown that careful monitoring with this instrument allows a novel conservative management approach.

In both of these cases, histopathological study confirmed melanocytoma. The appearances are consistent with “type 1” cells accounting for the majority of the cellular population, and it is likely that the infrequently encountered elongated cells are “type 2” cells. The cytological features of type 1 cells are consistent with a metabolically inactive and benign cell, which correlates well with the pigmentation and lack of growth seen in these cases. Type 2 cells are the metabolically active subtype whose cytological features are believed to account for the propensity towards growth and invasion, and their presence in the extrasceral portion, the advancing edge of this tumour, is compatible with this theory. We appreciate that false negative errors can be associated with biopsy, by sampling a benign area within a tumour and avoiding the area of aggressive growth. Further reliability and confidence in the biopsy could have been afforded by additional fine needle aspiration of the main tumour within the ciliary body, which is regarded as a safe and accurate method for sampling intraocular lesions.

The differential diagnosis of a ciliary body pigmented mass is not too exhaustive. The major diagnostic challenge presented is distinguishing between malignant melanoma, carcinoma, and adenocarcinoma of the pigmented ciliary epithelium. Medulloepithelioma, a congenital tumour, and hyperplasia of the pigmented ciliary epithelium, usually secondary to trauma or surgery, can usually be eliminated by the history.

Ultrasound biomicroscopy is an invaluable tool in the management of such lesions. Precise visualisation and measurement allows accurate objective definition of tumour extent, its interface with surrounding anatomy, and internal features. Correlation of tumour characteristics with histological features is good, and ultrasound biomicroscopic images can have resolving power up to 50 μm, which is analogous to low power light microscopy. A malignant melanoma of the ciliary body also has low internal reflectivity but is more vascular than melanocytoma. Also from features of internal reflectivity, inferences can be drawn about the cellular structure. The low internal reflectivity seen here implies a cohesive internal cytoarchitecture, with lack of vascularity.

There are two characteristics of clinical importance, which were not encountered here, with whose development we would be particularly concerned. Necrosis, which was seen in 10 of 23 cases in one series, and can result in pigment dispersion and secondary glaucoma, and malignant transformation, which is fortunately an infrequently encountered phenomenon.

In view of the essentially benign nature of this lesion, the conservative management approach described above was tested once the initial diagnosis was made based on biopsy. From our observations we conclude this approach is safe with regular follow up, in order to detect any alteration in the character of the tumour, or secondary complications. If such progression is noted, we would advocate formal en bloc surgical excision as a potentially curative management option.

Figure 1 (A) Preoperative ultrasound biomicroscopy of the lesion showing a ciliary body tumour with low internal reflectivity and anterior cystic element. (B) Ultrasound biomicroscopy 3 years after biopsy showing no change in the appearance of the melanocytoma. (C) Histology (haematoxylin and eosin, ×800). Melanocytes packed with melanosomes mostly obstructing nuclei. (D) Melanin bleach preparation (×800) showing tumour cells have round or oval nuclei without nucleoli.
REFERENCES