Uveal Melanoma Mimicking Advanced Coats’ Disease in a Young Patient

Naina Gupta\textsuperscript{a} William Terrell\textsuperscript{a} Lynn Schoenfield\textsuperscript{a, b} Claudia Kirsch\textsuperscript{c} Colleen M. Cebulla\textsuperscript{a}

\textsuperscript{a}Havener Eye Institute, Department of Ophthalmology and Visual Science, and \textsuperscript{b}Department of Pathology, The Ohio State University Wexner Medical Center, Columbus, Ohio, and \textsuperscript{c}Department of Radiology, North Shore Long Island Jewish Health System, Hofstra Medical School, Manhasset, N.Y., USA

Key Words
Uveal melanoma · Necrotic choroidal melanoma · Coats’ disease

Abstract
Background/Aims: To report a case and the unique histopathology of a necrotic uveal melanoma mimicking advanced Coats’ disease in a young adult. Method: A 26-year-old male presented with a blind, painful eye, total exudative retinal detachment, and bulbous aneurysms consistent with Coats’ disease. No masses were visualized on ultrasound or CT scan, and the patient underwent enucleation of the eye. Results: Histopathology of the involved eye confirmed a necrotic uveal melanoma with persistent spindle cells forming a collar around residual tumor vessels. Conclusion: Careful consideration is needed in approaching any patient with a blind, painful eye and opaque media, even in younger populations.

Case Report
This single case report was considered exempt from our Institutional Review Board. A 26-year-old incarcerated male, with a past medical history of hepatitis C, presented to the emergency department complaining of 1 week of headache and a red, painful right eye (OD), associated with nausea and vomiting. He noted a 1-year history of atraumatic, painless blindness OD. Visual acuity was ‘no light perception’ OD and 20/20 in the left eye (OS), with a relative afferent pupillary defect OD and intraocular pressures of 42 mm Hg OD and 15 mm Hg OS. Slit lamp examination OD showed extensive anterior chamber flare and florid iris neovascularization. A total exudative retinal detachment was visible near the lens (fig. 1a), with turbid, yellow subretinal fluid, small subretinal hemorrhage, and numerous bulbous aneurysms within the retinal vasculature (fig. 1b). Gonioscopy revealed neovascularization of the iris and angle, with angle closure OD. OS examination was normal. B scan, though limited due to the patient’s pain, revealed a mobile retinal detachment with shifting subretinal opacities and no solid mass (fig. 1c). CT with contrast showed diffuse increased attenuation through the entire right globe, suggestive of hemorrhagic and/or proteinaceous products (fig. 1d); no mass was visualized. The findings were consistent with stage 5 Coats’ disease [9].

Introduction
Occult ocular neoplasms have been discovered after enucleation [1, 2] or evisceration [3] of blind, painful eyes and may present in unusual ways, particularly when necrotic [4–8]. Here, we present a case of necrotic, uveal melanoma with atypical presentation, mimicking advanced Coats’ disease.
Despite medical therapy, the patient’s pain was uncontrolled. He underwent an uncomplicated enucleation OD with no gross external abnormalities of the globe.

Histopathology (fig. 2) showed an extensively necrotic (75%) choroidal melanoma in the temporal posterior globe, which was obscured by the extensive intraocular proteinaceous fluid. It spared the ciliary body and iris, and the anterior edge was 10 mm from the limbus and 2 mm from the optic nerve. Its largest basal diameter was 14 mm, and its height was 7 mm (pT3a). The melanoma was predominantly spindle cell type (with <10% epithelioid cells). There was no extension into the sclera, nor was there vascular invasion. Necrotizing scleritis was present. Immunohistochemical stains for both S-100 and melan A were positive, consistent with the diagnosis of melanoma. Similar to necrotic retinoblastomas, the melanoma consisted of large dilated vessels surrounded by a collar of tumor cells, 20–30 cells thick, with intervening necrosis.

The patient underwent a metastatic workup including an analysis of the complete blood count and lactate dehydrogenase, a comprehensive metabolic panel, a CT of the chest/abdomen/pelvis, and brain MRI. Chest CT revealed right hilar lymphadenopathy, with transbronchial biopsy showing lymphocyte aggregates without metastasis. All other testing was normal.

**Discussion**

Choroidal melanomas are rare in young adults [6], and spontaneous necrosis is similarly rare [7]. This patient’s presentation in particular is unique, with its extensive necrosis, similarity to advanced Coats’ disease, lack of enhancing mass on imaging, and unusual histopathology. The average presenting age of uveal melanoma is 55, with previous studies showing them to be uncommon in younger individuals [6]. Histologically, the globe was filled with proteinaceous fluid, obscuring a necrotic melanoma in the posterior globe. The predominant spindle cell morphology is not typical of prior reported necrotic melanomas [4]. Frasier and Font [4] found that only 23% of uveal melanomas associated with inflammation were predominantly spindle cell type; most tumors were listed as having an epithelioid, mixed, or ‘necrotic’ cell type. The spontaneous necrosis of uveal melanomas is thought to develop due to the melano-
ma outgrowing its blood supply centrally in a watershed area [10]. In this case, the residual viable tumor persisted along prominent tumor vessels. Like many retinoblastomas, it consisted of large dilated vessels surrounded by a collar of melanoma cells, which were 20–30 cells thick, with intervening necrosis.

An enucleation was performed rather than evisceration due to the inability to fully evaluate the posterior pole. While computed tomography (CT) is often the imaging modality of choice in the acute setting, increased attenuation from blood products, melanin, or proteinaceous material associated with a tumor can mask the iodinated contrast enhancement of the lesion, as has been reported with brain tumors; newer CT techniques such as dual-energy CT scanning are being developed to try to delineate tumoral enhancement that can be masked by hemorrhage [11]. MRI may also be limited by paramagnetic effects that can cause variable T1 and T2 signals [12]. Previous reports have been published describing the evisceration of occult melanomas in blind painful eyes [1–3]. The incidence of evisceration of occult melanomas is likely underreported.

In conclusion, an occult necrotic uveal melanoma must be considered in inflamed eyes with exudative retinal detachments, even in young patients. This case highlights the importance of enucleation rather than evisceration in blind painful eyes with opaque media and the potential limitations of modern imaging to detect these lesions.

Acknowledgments

Supported by the National Eye Institute of the National Institutes of Health under Award Number K08EY022672. Additional funds were provided by the Ohio Lions Eye Research Foundation and the Patti Blow Fund.

Statement of Ethics

This case report adheres to the tenants of the Declaration of Helsinki.

Disclosure Statement

The authors of this article have no conflicts of interest to disclose.
References