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Case report

Clinical and pathological correlation of cotton wool spots in secondary angle closure glaucoma

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1. Introduction

This case report extended from the original report, to our knowledge, is the first to provide histopathological co-localization of retinal damage corresponding to a long faded CWS in angle closure glaucoma.\textsuperscript{1} Cotton wool spots (CWS) are a well-described retinal finding occurring in association with many different well understood and idiopathic systemic retinal-vascular diseases, most commonly diabetes and hypertension.\textsuperscript{2} Typical CWS appear as transient gray to white areas with indistinct borders in the retinal nerve fiber layer (RNFL) usually less than 1/3 disc diameter, often adjacent to small retinal vessels. CWS contain clusters of cytoid bodies which form due to focal obstruction of orthograde or retrograde axonal transport in retinal ganglion cell axons.\textsuperscript{3,4} CWS usually fade over a few weeks following initial presentation and have only recently been associated with lasting retinal or disc damage.\textsuperscript{5,7} Most CWS will appear in the posterior fundus without any loss of visual acuity unless the central macula is involved. Traditional assumptions held that following the resolution of a CWS there were no pathological sequelae. Subsequent investigation with scanning laser ophthalmoscopy has shown localized loss of nerve fiber layer thickness corresponding with prior CWS\textsuperscript{3,5}. Therefore, CWS are not as innocuous as previously assumed. The incidence of CWS has rarely been reported in association with glaucoma\textsuperscript{(3)} and subsequent co-localized pathologic examination following resolution of CWS has not previously been reported.

2. Case report

An 18 year old female was diagnosed with unilateral secondary angle closure as a result of ring melanoma of the ciliary body OS, detailed in a previous report\textsuperscript{(1)}. She was initially followed for iris abnormalities that were misdiagnosed initially as iridocorneal endothelial

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Fig. 1. Fundus Photo OS during and after resolution of cotton wool spots and correlating histopathology from superior temporal macula. Relevant findings include a collection of glial cells in the nerve fiber layer and partial collapse of the middle nuclear layer. AGNO3, Epon 1μ section X 27.5 & 125.

Fig. 2. An Octopus VF in Nov 1976 OS revealed an early nasal step clearly progressed by August 1979.
syndrome (ICE) and followed over three years during which she maintained an uncorrected visual acuity of 20/20 OD and between 20/20 and 20/25 OS. During the last three months of follow up prior to her histologic diagnosis of low grade iris melanoma, she developed increasingly frequent IOP spikes (30–60 mmHg) refractive to medical treatment and gonioscopy consistent with progressive angle closure secondary to an expanding iris-ciliary body ring melanoma. A CWS in the superior arcuate RNFL was first documented with fundus photography 5/6/79. The CWS had completely faded by subsequent fundus exam and photo 7/6/1979 (Fig. 1), the last before enucleation OS. The patient did not experience any vascular occlusive phenomenon or undergo radiotherapy prior to the appearance of the CWS on clinical exam. Her last Octopus visual field test (Fig. 2) demonstrated a stable early superior nasal step in the left eye. Disc cupping was minimal and stable in annual stereo disc photos prior to her spiking IOP and increasing angle closure (Fig. 3). Once the melanoma diagnosis was established and the tumor considered inoperable, enucleation was recommended. Low grade iris-ciliary body melanoma was proven histologically after enucleation in August of year 3 of follow up and confirmed partial angle closure by tumor extension in ring fashion in the ciliary body, iris and angle.

This report augments a prior report. On review, 36 years later prompted by a recent complex imaging study, Iμ Epon retinal step sections originally stained with a novel AgNO3 solution demonstrated a retinal nerve fiber layer (RNFL) scar and inner nuclear layer (INL) collapse in the prior location of the CWS. Light microscopy (LM) and transmission electron microscopy (TEM) (Fig. 4) shortly after enucleation had demonstrated temporal quadrant laminar optic nerve (ON) retrograde axonal transport block.

Wolter previously mentioned the occurrence of CWS in angle closure glaucoma and described the morphology of terminal end bulb swelling in nerve fiber layer axons via a silver stain and light microscopy but offered no explanation of the pathophysiology. This case again demonstrates that angle closure glaucoma can lead to a CWS. Prior literature has concluded that the underlying mechanism for CWS is localized retinal arteriolar insufficiency. Glaucoma has generally not been listed among the possible causes. Malignancy has been reported as a possible etiology of CWS as it can result in localized vascular insufficiency due to shunting of blood flow or metastatic blockade of small retinal vessels. In this case, an asymptomatic CWS appeared in the posterior fundus far from the malignant site in the iris and ciliary body and there was no evidence of spread of the spindle melanoma beyond those sites. Since the appearance of the CWS corresponded with a period of unstable highly spiking IOP we speculate that the CWS most likely was triggered by high IOP, documented as elevated intermittently to 60 mmHg.

3. Discussion

To our knowledge, this delayed follow-up case report is the first to describe a CWS appearing during the course of secondary angle closure glaucoma with subsequent co-localization of a retinal nerve fiber layer (RNFL) scar and inner nuclear layer (INL) collapse corresponding to the prior CWS. Other investigators have shown alterations in clinical glaucoma testing with OCT and scanning laser ophthalmoscopy corresponding to prior occurrence of CWS in AIDS and systemic hypertension. Although detection of peripapillary disc hemorrhages has been regarded as a sign of ongoing glaucomatous disease, especially in low tension glaucoma, the finding of a CWS in the setting of angle closure glaucoma warrants caution in attributing loss of RNFL thickness to glaucomatous injury. Although we are reporting a correlation between CWS and angle closure glaucoma, the exact mechanism of occurrence is unclear. We theorize that an acute rise in intraocular pressure may result in focal retinal vascular insufficiency due to abrupt change in ocular perfusion pressure. Our patient’s young age and good general health seem inconsistent with systemic vascular compromise. Although our patient experienced high IOP in the setting of angle closure, it is unlikely that the CWS found on examination is specific to angle closure,
and could also occur in the setting of high IOP due to other mechanisms.

4. Conclusions

In summary, CWS can occur in angle closure glaucoma and ophthalmologists should be aware of this possibility as the subsequent testing of VFs, nerve fiber layer integrity and disc contours can be altered as a result of focal damage associated with the CWS. Eye care providers are encouraged to note and document CWS in glaucoma as they are not generally recognized as an associated sign. The localized loss of the RNFL as a sequelae of CWS can potentially be confused with progression of glaucomatous injury.

5. Patient consent

Written consent to publish case details has been obtained from each patient. Approval was obtained in writing and in compliance with HIPPA and Declaration of Helsinki.

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Conflicts of interest

The following authors have no financial disclosures (AB, CN, SM, DM).

Authorship

All authors attest that they meet the current ICMJE criteria for Authorship.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.ajoc.2018.02.028.

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