Long-term Management of Panuveitis and Iris Heterochromia in an Ebola Survivor

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Introduction

The 2013-2016 West African Ebola virus disease (EVD) outbreak has been the largest in history with 28,616 cases and 11,310 deaths in the highest transmission countries of Sierra Leone, Guinea, and Liberia.1 Reports of uveitis have emerged in EVD survivors.2-3 Herein
we discuss clinical features, multimodality imaging, and long-term management of aggressive, sight-threatening panuveitis, in an EVD survivor, providing insight into the pathogenesis of this condition.

Case Report

Acute Ebola virus disease and early post-Ebola syndrome

A 43-year-old physician developed EVD in Kenema, Sierra Leone. After 40 days of hospitalization at Emory University, he was discharged with serum and urine testing negative by quantitative reverse transcription-polymerase chain reaction (qRT-PCR) assay for EBOV RNA. One month after discharge, he experienced blurred vision, sacroilitis, enthesitis, and word-finding difficulties. Ophthalmic evaluation showed visual acuities of 20/15 in both eyes and multiple peripheral chorioretinal scars with hypopigmented haloes bilaterally, consistent with inactive chorioretinitis requiring no intervention (Supplemental Figure 1).

Acute hypertensive anterior uveitis

Fourteen weeks after EVD diagnosis, the patient presented with acute, left hypertensive anterior uveitis with visual acuity of 20/20 and intraocular pressure (IOP) of 44 mmHg in the left eye. Treatment was initiated with topical prednisolone acetate 1% (Pred Forte 1%) 4 times daily, brimonidine 0.2% and dorzolamide 2%/timolol 0.5%, and acetazolamide. Serologies for syphilis, cytomegalovirus, herpes simplex virus-1 and -2, and *Toxoplasma gondii*. HLA-B27 was negative.

Left anterior chamber paracentesis yielded aqueous humor positive for EBOV by qRT-PCR with a cycling threshold of 18.7 and a positive EBOV culture. Pre- and 24-hours post-procedure, conjunctival swab and tear film tested negative for EBOV RNA.

Anterior scleritis, and intermediate uveitis

On day 5 of the acute ocular illness, left visual acuity declined to 20/60 and the IOP was 15 mmHg. Diffuse anterior scleritis and intermediate uveitis prompted addition of oral prednisolone 80 mg, with the topical Pred Forte 1% every 2 hours, timolol 0.5%, and atropine 1% (Supplemental Figure 2).

Left visual acuity declined to 20/150 at day 6 of illness, and examination revealed grade 2+ AC cell, a 0.5 mm hypopyon, and grade 1-2+ vitreous haze. OCT scan showed mild left retinal thickening (Supplemental Figure 2).

Despite symptomatic improvement, left visual acuity worsened to 20/250 at day 9, and IOP declined to 6 mmHg. Corneal edema followed hypotony, with grade 2+ AC cell and grade 3+ vitreous haze (Supplemental Figure 2). Difluprednate 0.05% every 2 hours was initiated.
Optic neuropathy, and panuveitis

Left visual acuity deteriorated to 20/500 and IOP decreased to 2 mmHg at day 10. A left relative afferent pupillary defect (RAPD) developed, indicating optic neuropathy. Iris heterochromia developed at day 11, with a change from blue to green (Figure 1). Left anterior segment OCT (AS-OCT) revealed iris stromal thickening (502 μm) when compared to the right (376 μm). Ultrasound biomicroscopy (UBM) showed ciliary body edema. A dense grade 3+ vitreous haze persisted, prompting B-scan ultrasound, which revealed shallow peripheral, choroidal effusions and optic nerve head edema (Figure 1).

Antiviral therapy with favipiravir

Due to clinical worsening, a 21-day course of oral favipiravir (T-705, Toyoma Chemical, Japan) was initiated. After two loading doses of 2000 mg, favipiravir was administered 1200 mg twice daily. Three days after starting favipiravir (day 18 of illness), visual acuity had declined to finger counting at 2 feet and IOP was 3 mmHg. A periocular triaminolone acetonide injection (40 mg/1 ml) was administered in the Emory University Hospital Serious Communicable Diseases Unit (SCDU). A post-procedure conjunctival swab tested negative for EBOV RNA.

One day after the injection, left visual acuity was hand motions, but IOP increased to 9 mmHg. The patient was discharged on a course of favipiravir, and oral prednisone was tapered, decreasing by 10 mg/day every two weeks. Topical difluprednate 0.05% and atropine 1% were continued.

Long-term follow-up

Forty-five days after initial onset of acute ocular illness, the patient completed favipiravir and remained on oral prednisone 15 mg daily and topical difluprednate 0.05%. Visual acuity had improved to 20/15 with resolution of RAPD and IOP of 10 mmHg. Slit lamp examination showed resolved corneal edema, endothelial pigment without KP, and trace AC pigment. Posterior segment examination showed grade 0.5+ vitreous haze. The iris thickness had decreased to 452 μm by day 32, with resolution of heterochromia (Figure 1).

At one-year follow-up, left visual acuity returned to 20/20 with IOP of 11 mmHg. After 18 months, visual acuity had declined to 20/60 with development of diffuse posterior capsular cataract. Anterior uveitis was observed with diffuse stellate KP, 1+ AC cell and stable posterior segment. Repeat AC paracentesis tested negative for Ebola virus by qRT-PCR. PCR testing was also negative for CMV, HSV, and VZV DNA. Topical prednisolone acetate taper was given over four weeks with resolution of anterior uveitis.

Discussion

The clinical and multimodal imaging features of this aggressive spectrum of ophthalmic pathology highlight the mechanisms of inflammation and infection, which improved following the administration of corticosteroids and antiviral medication.

Iris heterochromia coincided with iris and ciliary body edema by AS-OCT and UBM; this was suggestive of heavy leukocyte infiltration and/or massive protein exudation related to an
extreme inflammatory response associated with active EBOV replication. Immediate improvement of IOP and recovery of ciliary body anatomy following the corticosteroid injection supported the key role of appropriately timed anti-inflammatory therapy.

Although the precise role of favipiravir – a pyrazinecarboxamide derivative that inhibits viral RNA replicase\(^5\) – was unclear, our patient’s disease process worsened initially on topical and systemic corticosteroid. Following the initiation of favipiravir, a periocular corticosteroid injection was administered because of concerns for recalcitrant hypotony, ciliary body shutdown, and irreversible vision loss. The patient’s clinical improvement paralleled the anatomic reduction in iris thickening by AS-OCT.

In summary, diagnostic ophthalmic imaging highlighted the anatomic and pathologic changes that occurred during our patient’s sight-threatening panuveitis. The imaging findings suggested severe reactive inflammation in the presence of EBOV viral replication, emphasizing the need for consideration of management strategies that target infectious and inflammatory processes in post-Ebola uveitis. Potential for uveitis recurrence mandates long-term monitoring.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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**References**

Figure 1. Anterior segment photographs and digital imaging
(A) Slit lamp photograph of right eye with corresponding anterior segment OCT at baseline shows iris thickness of 377 μm. (B) Slit lamp photograph and anterior segment OCT at day 11 show a green iris color with iris thickening at 502 μm. (C) By day 32, slit lamp photograph shows reversal of iris color to original blue color and corresponding decrease in iris edema to 452 μm. (D) Ultrasound biomicroscopy (UBM) show normal ciliary body anatomy of the right eye. (E) UBM of left eye shows ciliary body swelling (green triangles) and supraciliary/choroidal effusion (yellow areas) consistent with progressive panuveitis, choroiditis and evolving hypotony at day 12. (F) Repeat UBM shows decreased ciliary body swelling and resolution of supraciliary/choroidal effusion (yellow arrows) by day 32. (G) B-scan ultrasound shows choroidal thickening at day 11 and repeat at day 32. (H) shows resolution of choroidal thickening.