Case Report

A rare case of cytomegalovirus papillitis in patient with immunodeficiency

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A 26-year-old male diagnosed with AIDS came with sudden blurred vision and central scotoma in left eye since 2 weeks before admission. His visual acuity was counting finger at 5 meters with normal IOP and anterior segment. The posterior segment revealed edematous optic nerve covered by exudates and hemorrhages. Due to low CD4+ count and serological test result, we considered a HIV-related opportunistic ocular infection, specifically HSV infection. As visual acuity worsened during treatment with acyclovir, we performed PCR examination from aqueous tap which revealed positive CMV DNA. Unfortunately, the visual acuity had worsened to no light perception before he received any specific anti-CMV agent. CMV papillitis is an unusual presentation of CMV retinitis. PCR examination from aqueous or vitreous tap should be performed while waiting for serological test result, especially in doubtful cases. Therefore, appropriate diagnosis and management can be established early to prevent irreversible visual loss.

Keywords: AIDS, cytomegalovirus, immunodeficiency, papillitis

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Cytomegalovirus (CMV) papillitis is defined as greater than 270 degrees of disc edema/blurring of the disc margins as seen on direct examination and on color fundus photographs caused by CMV.\(^1\) CMV ocular infection will develop in 12% to 46% of patients with acquired immune deficiency syndrome (AIDS) in their lifetime. Of those patients with AIDS in whom CMV retinitis develops, CMV papillitis reportedly develops in up to 4% as well. Lestari\(^2\) reported the prevalence of CMV retinitis in Cipto Mangunkusumo Hospital (CMH) is 5.8%, there is no papil involvement reported in those cases. The majority of these patients have CD4 counts of less than 50 cells/mm\(^3\)\(^1-3\).

Diagnosis of CMV papillitis in the setting of human immunodeficiency virus (HIV)/AIDS is essentially clinical, based on the features just described. In CMV papillitis, the optic nerve head is edematous with focal hemorrhages, and an afferent pupillary defect usually is present. Polymerase chain reaction (PCR)-based analysis of the aqueous or vitreous samples may provide critical diagnostic information of high sensitivity and specificity that allow the clinician to differentiate CMV from other herpetic causes of necrotizing retinitis and from toxoplasmic retinochoroiditis in immunocompromised patients with atypical lesions.\(^1,3\) In this case report, we demonstrated a rare case of ocular CMV infection that manifests as CMV papillitis in HIV/AIDS.

### CASE REPORT

A male, 26 years old, came to Cipto Mangunkusumo Hospital, Jakarta on January 2014 complaining sudden blurred vision in left eye and central scotoma since two weeks before he was admitted. It was not red, it did not swell, and it was painless. Patient did not get any treatment for his eye before. There was no history of spectacles usage, hypertension, and diabetes mellitus. In the past medical history, this patient had been diagnosed with pulmonary tuberculosis, cryptococcal meningitis, and positive HIV test result. From social history, patient was a transmigrant from Sumatra Island. There was history of promiscuity with men and and intravenous drug user (IVDU).

Ophthalmological examination showed left eye’s visual acuity was 5/60, normal intraocular pressure (IOP), clear anterior segment, good right reflex with positive relative afferent pupillary defect (RAPD), and clear lens. In the posterior segment we found few cells in vitreous, and papil was hyperemic covered by exudates and peripapillary hemorrhages. There were flame-shaped hemorrhages, with turtous vein, and macular reflex hard to be evaluated. Right eye was in normal condition. We assessed this patient with specific opportunistic infection with several differential diagnosis including CMV, herpes simplex virus (HSV), cryptoccocus, toxoplasmosis, and syphilis infection (Figure 1 and Figure 2).

At the second and the third week after initial visit, the eye condition was getting worse, eventually leading to no light perception. Ancillary test revealed some abnormal result as shown in the Table 1. Based on clinical and laboratory results, we considered an active HSV infection, so we consulted the patient to Department of Internal Medicine and the patient got 800 mg acyclovir four times daily (Figure 3).

Due to worsening condition and confusing diagnosis, we performed PCR examination from aqueous sample, as shown in Table 1. Polymerase chain reaction (PCR)-based analysis of the aqueous or vitreous samples may provide critical diagnostic information of high sensitivity and specificity that allow the clinician to differentiate CMV from other herpetic causes of necrotizing retinitis and from toxoplasmic retinochoroiditis in immunocompromised patients with atypical lesions.\(^1,3\)

**Figure 1.** Normal funduscopy (right); hyperemic papil covered by exudates and peripapillary hemorrhages, flame-shaped hemorrhages (left). At this initial visit, the visual acuity of the left eye is 5/60

**Figure 2.** Normal funduscopy (right); the condition of the left eye is getting worse with hyperemic papil with extensive exudates and peripapillary hemorrhages, flame-shaped hemorrhages. At the third week follow-up, the visual acuity of the left eye is no light perception.
ous tap to confirm the diagnosis. It was revealed that the patient had a positive CMV deoxyribonucleic acid (DNA). Therefore, we consulted to Department of Internal Medicine for a change to specific anti-CMV medication. The patient then received 900 mg valganciclovir twice a day for three weeks.

During the follow-up, five weeks after the first visit, the patient developed immune recovery uveitis with raising intraocular pressure up to 40 mmHg. He complained of pain of the left eye, accompanied with conjunctival and ciliary injection, the corneal edema and large keratitic precipitates were also noted. In anterior chamber, we found massive cells (+4) and flare (+2). The lenses were clear and the posterior segment was still the same. The patient received anti-glaucoma medication and topical steroid (Figure 3).

In seventh week after the initial visit, patient came with subsided peripapillary hemorrhages and exudates after administration of oral ganciclovir 900 mg twice daily in fourth week. This condition showed a good response to valganciclovir treatment. Based on the clinical manifestation, we diagnosed this patient with CMV papillitis (Table 1).

Patient has been treated with highly-active anti-retroviral therapy (HAART) medication and also the anti-tuberculosis agent. HAART which consists of evafirenz, lamivudine, tenofovir disoproxil fumarate.

**DISCUSSION**

Opportunistic infections (OIs), which have been defined as infections that are more frequent or more severe because of immunosuppression in HIV-infected persons. It is important to recognize

![Figure 3](image1)

**Figure 3.** Immune recovery uveitis. Left eye shows conjunctival and ciliary injection, deep anterior chamber. Yellow arrow shows massive cells (+4) and flare (+2) in the anterior chamber. Red arrow shows keratitic precipitate

![Figure 4](image2)

**Figure 4.** Left eye was showing subsided peripapillary hemorrhages and exudates after administration of oral valganciclovir 2 x 900 mg in third week

Table 1. Laboratory examination of clinical importance

<table>
<thead>
<tr>
<th>Test</th>
<th>Value</th>
<th>Interpretation</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Allergy-Immunology</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CD4+ percentage</td>
<td>1%</td>
<td>Decreased</td>
<td>31–60</td>
</tr>
<tr>
<td>CD4+ absolute</td>
<td>4 cells/mL</td>
<td>Decreased</td>
<td>410–1590</td>
</tr>
<tr>
<td><strong>TORCH test</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anti-rubella IgG</td>
<td>2732.0 IU/mL</td>
<td>Reactive</td>
<td>&lt;5.0 (non-reactive)</td>
</tr>
<tr>
<td>Anti-rubella IgM</td>
<td>0.2 COI</td>
<td>Non-reactive</td>
<td>&lt;0.8 (non-reactive)</td>
</tr>
<tr>
<td>Anti-CMV IgG</td>
<td>24.7 Au/mL</td>
<td>Reactive</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>Anti-CMV IgM</td>
<td>0.4 COI</td>
<td>Non-reactive</td>
<td>0.7</td>
</tr>
<tr>
<td>Anti-HSV II IgG</td>
<td>0.24</td>
<td>Non-reactive</td>
<td>&lt;0.90 (non-reactive)</td>
</tr>
<tr>
<td>Anti-HSV II IgM</td>
<td>0.91</td>
<td>Reactive</td>
<td>&lt;0.90 (non-reactive)</td>
</tr>
</tbody>
</table>

The laboratory results showed an active HSV type II infection so the patient got treated by acyclovir 5 x 800 mg. By the time, we observed that the ocular manifestation was not improved with the treatment, so we considered another etiology. Based on the literature, the most common opportunistic infection on AIDS with very low CD4+ count (<200 cells/mL) is CMV infection. To confirm the diagnosis, we did the aqueous tap and PCR examination and revealed a positive result CMV infection. TORCH: Toxoplasma, rubella, cytomegalovirus, herpes simplex; IgG: immunoglobulin G; IgM: immunoglobulin M; anti CMV: anti cytomegalovirus; anti HSV: anti- herpes simplex virus

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that the relationship between OIs and HIV infection is bidirectional.

Human immunodeficiency virus can cause systemic or organ diseases. Several systemic OIs include Mycobacterium tuberculosis infection, Cryptococcus infection, cytomegalovirus disease, toxoplasmosis infection. At the initial visit, our patient was already diagnosed with pulmonary tuberculosis and cryptococcal meningitis, still ongoing treatment. With the presenting systemic OIs, it is likely that this patient might have a specific organ disease.\textsuperscript{4,5}

Iqbal\textsuperscript{6} stated an association between CD4+ cells count with ocular complication of HIV with the commonest infection is CMV. Cullen et al\textsuperscript{7} stated that neuro-ophthalmic manifestations of HIV tend to present at an advanced stage of the disease when CD4 cell counts are depleted below 200 cells/\textmu L. In HIV-infected patients, opportunistic infections such as cytomegalovirus, toxoplasmosis, syphilis, and tuberculosis are by far the most common causes of optic nerve disorders.

In our cases, we found that patient was complaining sudden blurred vision of his left eye, diagnosed as AIDS since three months before came to this hospital (October 2013) with initial symptoms were loss of consciousness due to cryptococcal meningitis, starting HAART medication two months later (November 2013) and initial CD4+ count at first visit was 4 cells/mL (n: 410-1,590 cells/mL). Initial presentation showed visual acuity 5/60, normal IOP, quiet eye with normal anterior segment, positive RAPD, and posterior segment manifestation that is edematous papil, peripapillary hemorrhages and exudates, and hemorrhages in retina. From this condition, our primary consideration was opportunistic infection with differential diagnosis caused by CMV, HSV, Cryptococcus, Toxoplasma, and syphilis. The laboratory examination and PCR testing show positive tendency to CMV infection. Based on this result we gave specific anti-CMV infection, valganciclovir 900 mg twice daily.

Cytomegalovirus papillitis has been described in immunocompromised patients and is usually a rapidly blinding disease.\textsuperscript{8} Many studies reported the incidence of CMV papillitis found less than 10% in CMV retinitis cases. CMV retinitis accounts for 75–85% of CMV disease and over 90% of blindness in AIDS patients. Freeman et al\textsuperscript{9} report papillitis is present in up to 4% of patients with CMV retinitis. Gross et al\textsuperscript{10} noted 4.1% patients with CMV retinitis also had CMV papillitis at initial presentation. Similarly, Rosecan et al\textsuperscript{11} noted optic nerve involvement in 8.3% eyes with CMV retinitis. Roarty et al\textsuperscript{12} also reported a patient with severe CMV papillitis and visual acuity of no light perception vision, and Palestine et al\textsuperscript{13} noted CMV papillitis in 7% eyes with CMV retinitis with no light perception vision upon initial presentation. Until now, there is no report of isolated CMV papillitis cases in without retinal manifestation in AIDS patient.

Successful management of active CMV retinitis reported by Figueiredo\textsuperscript{14} in an active CMV retinitis infection. In his report the patient had good visual outcome, from hand movement with good projection to 5/12 after induction doses of oral valganciclovir.

The Centers for Disease Control and Prevention (CDC), the National Institutes of Health, and the HIV Medicine Association of the Infectious Diseases Society of America\textsuperscript{15} give recommendation for treating CMV retinitis. For sight threatening lesions which is adjacent to the optic nerve or fovea, the recommended initial therapy comprised intravitreal injections of ganciclovir (2 mg/injection) or foscarnet (2.4 mg/injection) for 1–4 doses over a period of 7–10 days to provide higher intraocular levels of drug and faster control of the infection until steady state intracellular ganciclovir concentrations are achieved. This initial therapy might be accompanied by valganciclovir 900 mg twice daily for two until three weeks then followed by once daily doses.

In our case, he given a systemic valganciclovir 900 mg twice daily in first three weeks, to control the presumably active CMV infection and to prevent the CMV involvement to the contralateral eye and also to improve patient’s systemic condition. Although the guideline of CMV papillitis’ management has not been established yet, this CDC recommendation is applicable in this case.

Based on the case explained above, we concluded the screening recommendations for patients with AIDS and low CD4+ T-cell counts should be performed routinely. PCR examination from aqueous
or vitreous tap should be performed earlier while waiting for the serological testing result, especially in doubtful cases which severity progress rapidly. An appropriate diagnosis and management can be established and performed earlier to prevent the irreversible visual loss.

Conflicts of interest
The authors affirm there is no conflict of interest in this study.

REFERENCES


