Clinical and laboratory characteristics of ocular syphilis, co-infection, and therapy response

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Purpose: To describe the clinical presentation of patients diagnosed with presumed latent ocular syphilis and congenital ocular syphilis at tertiary referral center in Turkey, and to compare the clinical findings with patients described in other studies, specifically focusing on demographics and co-infections.

Methods: This is a retrospective study reviewing the medical records of patients diagnosed with ocular inflammation between January 2012 and June 2014 at a tertiary referral center in Turkey. Ocular syphilis was diagnosed on the basis of non-treponemal and treponemal antibody tests, and cerebrospinal fluid analysis. All the patients diagnosed with ocular syphilis were tested for human immunodeficiency virus (HIV), *Toxoplasma gondii*, rubella, cytomegalovirus, and herpes.

Results: A total of 1,115 patients were evaluated between January 2012 and June 2014, and 12 patients (1.07%) were diagnosed with ocular syphilis based on the inclusion criteria. None of the patients were seropositive for HIV. Two patients were seropositive for *T. gondii*-specific IgG. Clinical presentations include non-necrotizing anterior scleritis, non-necrotizing sclerokeratitis, anterior uveitis, intermediate uveitis, posterior uveitis, panuveitis, and optic neuritis. All of the patients showed clinical improvement in the level of ocular inflammation with intravenous penicillin 24 million U/day for 10 days. Three patients received additional oral methotrexate as an adjunctive therapy. Two cases received low-dose trimethoprim–sulfamethoxazole.

Conclusion: Ocular syphilis is an uncommon cause of ocular inflammation in HIV-negative patients. Central retinochoroiditis is the most common ocular manifestation, and it is the most common cause of visual impairment. Ocular syphilis might present associated with co-infections such as *T. gondii* in developing countries. Oral methotrexate might be beneficial as an adjunctive therapy for ocular syphilis in resolving the residual intraocular inflammation and cystoid macular edema after specific therapy with intravenous penicillin.

Keywords: ocular syphilis, scleritis, uveitis, methotrexate, human immunodeficiency virus, *T. gondii*

Introduction

Syphilis is an infectious disease caused by the spirochete *Treponema pallidum* which is most often transmitted sexually in acquired disease or vertically during pregnancy in congenital syphilis (CS).\(^1,2\) Four stages have been described in acquired disease, and overlap can exist between the stages.\(^3,4\) The primary stage is characterized by a painless ulcerated lesion known as a chancre, which is thought to be the site of entrance into the host.\(^3\) The secondary stage typically occurs 2 weeks to several months after infection and is associated with a skin rash on the palms of the hands and soles of the feet.\(^4\) A period of latency can occur after the secondary stage, and recurrences often present during the 1st year after diagnosis.\(^5\) Patients can remain in this stage indefinitely or develop tertiary syphilis.\(^5\)
During this stage, serious systemic manifestations occur, including cardiovascular and neurologic symptoms as well as gumma formation throughout the body.6–8 Ocular syphilis is considered to occur during any stage of infection, and it is known as “great masquerader”.9 CS has been classified as early CS and late CS.10 Early CS is characterized by organ involvements including liver, kidney, bone, and hematological and mucocutaneous manifestations.10 Ocular manifestations including interstitial keratitis, chorioretinitis, salt–pepper fundus, glaucoma, cataract, and chancres of the eyelid have been mostly described in late CS.11 Every patient with ocular inflammation is recommended to be tested for syphilis.9 Since non-specific treponemal tests, including venereal disease research laboratory test (VDRL) and rapid plasma reagin test (RPR), normalize during the latent and tertiary stages, testing for syphilis should include both non-specific treponemal tests and specific treponemal antibody tests including fluorescent treponemal antibody absorption (FTA-ABS) and the microhemagglutination assay for T. pallidum12 (MHA-TP). Treatment for presumed latent ocular syphilis is the same as for neurosyphilis, specifically 10–14 days of intravenous penicillin (IV PCN) 3–4 million units every 4 hours.13 In this study, we describe the clinical presentation of patients diagnosed with presumed latent ocular syphilis and congenital late syphilis at a tertiary referral center in Turkey. We compare the clinical findings with patients described in other studies, specifically focusing on demographics and co-infections.

Materials and methods
This is a retrospective study reviewing the medical records of patients diagnosed with ocular inflammation between January 2012 and June 2014 at a tertiary referral center in Turkey. This study adheres to the tenets of Declaration of Helsinki. Institutional review board/ethics committee approval was obtained from the Dusaha Goz Hospital, Ankara Turkey. Written informed consent was obtained for all individuals who enrolled in this study. The results of non-treponemal antibody tests VDRL and RPR, and specific FTA-ABS and MHA-TP, cerebrospinal fluid (CSF) FTA-ABS and CSF protein levels were analyzed. The diagnosis of presumed latent ocular syphilis and presumed congenital late syphilis was done according to the criteria described in the literature.2,13 All the patients diagnosed with presumed latent ocular syphilis and presumed congenital late syphilis were tested for human immunodeficiency virus (HIV) and TORCH (Toxoplasma gondii, rubella, cytomegalovirus, herpes). Clinical presentation, demographic information, and response to treatment were collected for each patient. Best corrected visual acuity (BCVA) was tested by using a Snellen chart. Snellen fractions of BCVA were converted to LogMAR. Intraocular pressure was measured by using non-contact tonometer (NT-530, Nidek Co., Ltd., Gamagori, Japan). Slit-lamp biomicroscopy with SL-D7 (Topcon, Tokyo, Japan) and dilated fundus examination using an indirect ophthalmoscope (Keeler, West Berkshire, UK) were performed for all patients. Fundus fluorescein angiography (FFA), indocyanine green angiography (ICGA), and spectral domain optical coherence tomography (SD-OCT) were done by Spectralis Heidelberg Research Architecture plus optical coherence tomography (HRA + OCT; Heidelberg Engineering, Heidelberg, Germany). Visual field analysis was performed by Humphrey HFA II (Carl Zeiss Meditec AG, Jena, Germany). None of the patients had been treated for syphilis before admission to our clinic. In all, 4 (33.3%) of the 12 patients were receiving oral prednisone less than 3 months duration for their intraocular inflammation at presentation. None of the patients were on topical prednisolone acetate on admission. All of the patients received treatment with IV PCN 24 million U/day for 10 days. IV methylprednisolone for 3 days followed by high-dose oral prednisone was added to IV PCN treatment for the patient with papillitis. Oral methotrexate between 15 and 20 mg/week doses for 6 months was added to three patients 4 weeks after IV PCN therapy. Low-dose trimethoprim–sulfamethoxazole (trimethoprim, 160 mg plus sulfamethoxazole, 800 mg; one tablet twice daily, 2 days per week) as primary prophylaxis against T. gondii was given for 1 year for 2 patients who were seropositive for specific IgG for T. gondii.

Results
Medical records of 1,115 patients diagnosed with ocular inflammation were evaluated between January 2012 and June 2014. In all, 12 (1.07%) patients, 17 eyes were diagnosed with ocular syphilis. Briefly, 10 patients, 13 eyes had diagnosis of presumed latent ocular syphilis, and 2 patients, 4 eyes had diagnosis of presumed congenital late syphilis. Mean age at presentation was 43.75 years (range 8 and 67 years), and all of the patients were Caucasian. In all, 7 (58.3%) were male and 5 (41.7%) were female. Five (41.6%) patients had bilateral involvement. The initial BCVA of the patients were between 20/400 and 20/20 (0–1.3 LogMAR). Intraocular inflammation was recommended to be tested for syphilis.9 Since non-specific treponemal tests, including venereal disease research laboratory test (VDRL) and rapid plasma reagin test (RPR), normalize during the latent and tertiary stages, testing for syphilis should include both non-specific treponemal tests and specific treponemal antibody tests including fluorescent treponemal antibody absorption (FTA-ABS) and the microhemagglutination assay for T. pallidum12 (MHA-TP). Treatment for presumed latent ocular syphilis is the same as for neurosyphilis, specifically 10–14 days of intravenous penicillin (IV PCN) 3–4 million units every 4 hours.13 In this study, we describe the clinical presentation of patients diagnosed with presumed latent ocular syphilis and congenital late syphilis at a tertiary referral center in Turkey. We compare the clinical findings with patients described in other studies, specifically focusing on demographics and co-infections.

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Table 1 Demographics and clinical characteristics of the cases

<table>
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<tr>
<th>Cases</th>
<th>Age (years)</th>
<th>Sex</th>
<th>Initial BCLA, LogMAR</th>
<th>Clinical manifestations</th>
<th>Laboratory findings</th>
<th>Imaging</th>
<th>Treatment</th>
<th>Follow-up</th>
<th>Final BCLA, LogMAR</th>
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<tr>
<td>1</td>
<td>35</td>
<td>F</td>
<td>20/20 (OD) 0.0</td>
<td>Anterior scleritis</td>
<td>SLE OD discloses 2+ diffuse scleral injection at the nasal quadrant</td>
<td>IV PNC 24 million U/day for 10 days</td>
<td>12 months</td>
<td>20/20 (OD) 0.0</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>43</td>
<td>M</td>
<td>20/20 (OS) 0.0</td>
<td>Sclerokeratitis</td>
<td>SLE OS discloses 2+ scleral injection at the nasal quadrant associated with limbal corneal thinning</td>
<td>IV PNC 24 million U/day for 10 days</td>
<td>6 months</td>
<td>20/20 (OS) 0.0</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>55</td>
<td>F</td>
<td>20/20 (OD) 0.0</td>
<td>Non-granulomatous iridocyclitis</td>
<td>SLE OD discloses 10 small KP 1+ flare in the AC and ½+ cell in the AV</td>
<td>IV PNC 24 million U/day for 10 days</td>
<td>18 months</td>
<td>20/20 (OD) 0.0</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>49</td>
<td>M</td>
<td>20/32 (OD) 0.0/0.22</td>
<td>Non-granulomatous iridocyclitis</td>
<td>SLE OD discloses 5 small KP ½+ cell ½+ flare in the AC ½+ cell in the AV, and SLE OS discloses 30 small KP 1+ flare 1+ cell in the AC and 1+ cell in the AV</td>
<td>IV PNC 24 million U/day for 10 days</td>
<td>24 months</td>
<td>20/20 (OU) 0.0</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>67</td>
<td>M</td>
<td>20/32 (OS) 0.3/0.22</td>
<td>Intermediate uveitis</td>
<td>SLE OD discloses 10 small KP 1+ cell 2+ flare in the AC and ½+ cell in the AV, and SLE OS discloses 15 small KP 2+ flare 1+ cell in the AC and ½+ cell in the AV</td>
<td>IV PNC 24 million U/day for 10 days</td>
<td>12 months</td>
<td>20/25 (OD) 0.1/0.0</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>62</td>
<td>F</td>
<td>20/32 (OD) 0.22</td>
<td>Intermediate uveitis</td>
<td>SLE OD discloses 1+ V cell 1+ V haze and fundus ex discloses peripheral chorioretinal pigmentary changes</td>
<td>IV PNC 24 million U/day for 10 days</td>
<td>18 months</td>
<td>20/20 (OD) 0.0</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>52</td>
<td>F</td>
<td>20/400 (OS) 1.3</td>
<td>Intermediate uveitis</td>
<td>OCT OS discloses foveal thickness of 693 µm associated with PED and ERM formation</td>
<td>IV PNC 24 million U/day for 10 days</td>
<td>24 months</td>
<td>20/20 (OS) 0.0</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>46</td>
<td>M</td>
<td>20/32 (OD) 0.22</td>
<td>Papillitis</td>
<td>Fundus ex OD discloses 2+ hyperemia and 2+ swelling of the optic disc HVF 24/2 discloses central scotoma and increased blind spot</td>
<td>IV PNC 24 million U/day for 10 days</td>
<td>24 months</td>
<td>20/20 (OD) 0.0</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>50</td>
<td>M</td>
<td>20/200 (OS) 1.0</td>
<td>Posterior uveitis, branch retinal artery occlusion</td>
<td>FFA OS discloses absence of arteriolar filling inferotemporally associated with vascular occlusions and aneurysmal changes, areas of non-perfusion</td>
<td>IV PNC 24 million U/day for 10 days oral MTX 17.5 mg/week for 6 months</td>
<td>15 months</td>
<td>20/40 (OS) 0.3</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>18</td>
<td>M</td>
<td>20/80 (OU) 0.6</td>
<td>Posterior uveitis, focal chorioretinitis</td>
<td>FFA early phase OU discloses staining at the macula and choroid in the posterior pole, increased staining at the intermediate phase and persistence of staining at the late phase ICGA early phase discloses areas of hypofluorescence intermediate phase discloses staining at the choroid associated with areas of hypofluorescence late phase discloses areas of hypo and hyperfluorescence associated with increased staining at the choroid OCT OU discloses CME with a foveal thickness of 307 µm OD, and 266 µm OS</td>
<td>IV PNC 24 million U/day for 10 days low-dose TMP–SMX prophylaxis for 1 year</td>
<td>12 months</td>
<td>20/50 (OU) 0.4</td>
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(Continued)
### Table 1 (Continued)

<table>
<thead>
<tr>
<th>Cases</th>
<th>Age (years)</th>
<th>Sex</th>
<th>Initial BCVA, LogMAR</th>
<th>Follow-up</th>
<th>Final BCVA, LogMAR</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>11</td>
<td>8</td>
<td>M</td>
<td>20/80 (OU) 0.6</td>
<td>12 months</td>
<td>N PNC 24 million</td>
<td>FTa-ABS positive Mha-TP, prophylaxis for 1 year</td>
</tr>
<tr>
<td>12</td>
<td>31</td>
<td>F</td>
<td>20/40 (OU) 0.3</td>
<td>18 months</td>
<td>N PNC 24 million</td>
<td>FTa-ABS positive Mha-TP, prophylaxis for 1 year</td>
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**Clinical findings**

<table>
<thead>
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<th>Imaging</th>
<th>Laboratory findings</th>
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<tr>
<td>FT-A-ABS positive MHA.</td>
<td>Positive for T. pallidum</td>
</tr>
<tr>
<td>TP positive IV, negative</td>
<td>Indocyanine green angiography</td>
</tr>
</tbody>
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**Clinical manifestations**

- Posterior uveitis
- Focal chorioretinitis

**Follow-up findings**

- SD-OCT of right and left eyes of the second patient showed areas of hypo and hyperfluorescence associated with focal chorioretinitis.

**Treatment**

- FFA early phase of right and left eyes with vitritis and branch retinal artery occlusion disclosed venous staining, leakage, and areas of non-perfusion.
- SD-OCT of right and left eyes of patient with focal chorioretinitis disclosed areas of hypo and hyperfluorescence associated with increased staining of choroid.

**Abbreviations:**

- AC, anterior chamber; AV, anterior vitreous; BCVA, best-corrected visual acuity; CSF, cerebrospinal fluid; CME, cystoid macular edema; DC, discontinue; ERM, epiretinal membrane formation; Ex, examination; F, female; FFA, fundus fluorescein angiography; FTA-ABS, fluorescent treponemal antibody absorption; HIV, human immunodeficiency virus; HVF, Humphrey visual field; ICGA, indocyanine green angiography; IG, immunoglobulin; IV-PNC, intravenous penicillin; KP, keratic precipitates; M, Male; Mha-TP, micro-hemagglutination assay for *Treponema pallidum*; MTX, methotrexate; OCT, optical coherence tomography; OD, right eye; Os, left eye; OU, both eyes; PeD, pigment epithelial detachment; SLE, slit-lamp examination; T, toxoplasmosis; TMP–sMX, trimethoprim–sulfamethoxazole; V, vitreous.

*Note: *The eyes with posterior uveitis had clinical manifestations of vitritis, branch retinal artery occlusion inferotemporally associated with vessel sheathing and intraretinal hemorrhages (Figure 4A), and focal chorioretinitis characterized by yellowish lesions with fomed centers at the level of the pigment epithelium in the macula and paramacular region in both eyes of two patients (Figure 5A–D). Focal chorioretinitis was the most common posterior segment finding in our study (29.3%). FFA late phase of the left eye with vitritis and branch retinal artery occlusion disclosed venous staining, leakage, and areas of non-perfusion (Figure 4B). FFA early phase of right and left eyes of the patient with focal chorioretinitis disclosed staining at the macula and choroid in the posterior pole, intermediate phase revealed increased staining, and late phase disclosed persistence of staining at the posterior pole (Figure 6A). Indocyanine green angiography early phase of right and left eyes of the patient with focal chorioretinitis disclosed areas of hypo and hyperfluorescence in the posterior pole, intermediate phase disclosed staining of choroid associated with areas of hypo and hyperfluorescence, and late phase disclosed areas of hypo and hyperfluorescence associated with increased staining of choroid (Figure 7A). The resolution of vitritis and intraretinal hemorrhages associated with persistence of arterial sheathing inferotemporally was noted for the patient with vitritis and branch retinal artery occlusion OS at the 3rd month of the MTX therapy (Figure 4C).
Ocular syphilis

with panuveitis disclosed creamy-yellow superficial retinal accumulations and optic disc congestion (Figure 10A and B). FFA early phase showed increased choroidal fluorescence OU (Figure 10C) and venous phase showed increased staining at the optic disc and bifurcation of the vessels OU (Figure 10D). The resolution of vitritis, the creamy-yellow superficial retinal accumulations and optic disc congestion were noted in right and left eyes of the patient with panuveitis at the 3rd month of the MTX therapy (Figure 10E and F). FFA arterial phase disclosed decreased choroidal fluorescence in right eye, and venous phase indicated the resolution of staining at the optic disc and bifurcations of the vessels in left eye (Figure 10G and H). Fundus examination of the patient with right papillitis disclosed 2+ hyperemia and 2+ swelling of the right optic disc (Figure 11A). Humphrey visual field analyzer 24-2 disclosed central scotoma and increased blind spot OD (Figure 11B). The resolution of optic disc congestion and edema OD with improvement of HVF 24-2 central scotoma in right eye at the 1st month of follow-up visit was noted for the patient with papillitis (Figure 11C and D). None of the patients were seropositive for HIV, and 2 (16.6%) of the 12 patients were seropositive for T. gondii-specific IgG antibody. The CSF FTA-ABS analysis of all cases was negative. Elevated CSF protein was found in 2 cases who were seropositive for T. gondii-specific IgG antibody.

The patients who have received IV PNC only were followed-up 6 months to 2 years without recurrences. The patients who have received both IV PNC and oral MTX were followed-up 15–24 months period without recurrences. The final BCVA of the patients were between 20/50 and 20/20 (0.4–0.0 logMAR) with a mean LogMAR visual acuity of 0.06. Focal chorioretinitis was found as the most common cause of visual impairment.

Discussion

Syphilis is a sexually transmitted infection (STI) with a steady increase in the number of infected cases in the USA since the middle of 1980s.14 A dramatic rise in incidence of primary
Figure 3 Represents the spectral domain optical coherence tomography of the left eye at pre-treatment and post-treatment phases.

Notes: (A) Spectral domain optical coherence tomography of the patient with intermediate uveitis disclosed left cystoid macular edema with a foveal thickness of 693 \( \mu \text{m} \) associated with the retina pigment epithelial detachment and epiretinal membrane formation. (B) Spectral domain optical coherence tomography of the patient with intermediate uveitis disclosed regression of left cystoid macular edema and the retina pigment epithelial detachment with a foveal thickness of 362 \( \mu \text{m} \) associated with persistence of epiretinal membrane formation at the 3rd month of the MTX therapy.

Abbreviations: MTX, methotrexate; OS, left eye.

Figure 4 Represents the color fundus photo and fundus fluorescein angiography of the left eye at pre-treatment and post-treatment phases.

Notes: (A) Color fundus photo of left eye with posterior uveitis discloses vitritis and branch retinal artery occlusion inferotemporally associated with vessel sheathing and intraretinal hemorrhages. (B) Fundus fluorescein angiography late phase discloses venous staining, leakage, and areas of non-perfusion. (C) Color fundus photo discloses resolution of vitritis and intraretinal hemorrhages associated with persistence of arterial sheathing inferotemporally at the 3rd month of the MTX therapy. (D) Fundus fluorescein angiography late venous phase discloses decreased areas of non-perfusion, vascular leakage, and staining associated with the areas of capillary occlusions inferotemporally.

Abbreviations: MTX, methotrexate; OS, left eye.
and secondary syphilis in the USA has been reported from 2.1 per 100,000 people in 2000 to 4.5 per 100,000 in 2011. The Centers for Disease Control and Prevention (CDC) has revealed that 72% of all primary and secondary syphilis cases occurred in men who have sex with men (MSM) in 2011 which has increased from 7% in 2000. In a study of MSM presented to sexually transmitted disease (STD) clinics, a higher proportion of primary and secondary syphilis has been noted in cases co-infected with HIV (10.1%) than HIV-negative cases (2.6%). Since 2001, threefold rise in incidence of syphilis and HIV infection in MSM have been reported in the age group of up to 25 years at the University Hospital of Dresen. The strong association between ocular syphilis co-infection with HIV has been considered to be related to genital ulcerative diseases that facilitate HIV infection. Our study disclosed the incidence of ocular syphilis among patients with ocular inflammation between January 2012 and June 2014 as 1.07%. There was no comparable study related to the incidence of ocular syphilis in Turkey. However, RPR seropositivity among the blood donors at the central blood bank of Turkey has been reported to increase from 0.04% in 1987 to 0.2% in 2002. None of our cases were found seropositive for HIV. A low rate of HIV co-infection has also been reported in the first national seroprevalence survey in Saudi Arabia. This might be related to the sexual behaviors of individuals living in mostly Muslim populations. A steady increase, approximately 15-fold in the incidence of ocular syphilis between 1999 and 2004, has also been reported in UK. The British Ocular Syphilis Study (BOSS) has revealed a total of 41 new cases (63 eyes) of intraocular syphilis between 2009 and 2011 with an annual incidence of 0.3 per million UK adult population. The mean age, sex distribution, and laterality of ocular involvement in BOSS were as 43.8 years, 90.2% male, and 56% bilateral involvement, respectively. Our study disclosed the mean age as 43.8 years, 58.3% male, and 41.6% bilateral involvement that were compatible with the BOSS study. Syphilis is considered to display diverse ophthalmologic manifestations. A case series study of ocular syphilis in 18 eyes of 12 patients in a Singaporean population between 2004 and 2009 disclosed that anterior uveitis and panuveitis were reported two most common findings. The other case series study including 22 patients with syphilitic uveitis followed at Singapore National Eye Center between 1995 and 2006 has revealed that non-granulomatous iridocyclitis was the most common finding (62.1%) at presentation followed by panuveitis (27.5%), posterior uveitis (13.7%), and intermediate uveitis (10.3%). This study also disclosed that vitritis (65.4%) was the most common posterior segment finding. The most common posterior segment finding in our study was found as posterior...
uveitis presenting as chorioretinitis rather than vitritis. In a study involving 35 eyes of 19 Chinese patients with syphilitic uveitis, posterior uveitis has been disclosed as the most common finding. In a recent study from the USA involving 61 eyes of 35 patients, independent of HIV status panuveitis was reported as the most common presentation in ocular syphilis. Our study disclosed that both non-granulomatous iridocyclitis (20.9%) and posterior uveitis (20.9%) were two most common presentations followed by panuveitis (11.7%) and intermediate uveitis (11.7%). Ocular syphilis was also
Figure 7 Represents the indocyanine green angiography of the right and left eye.

Notes: Indocyanine green angiography early phase of right and left eyes with focal chorioretinitis disclosed areas of hypofluorescence (Ai and Aii), intermediate phase disclosed staining of choroid associated with areas of hypofluorescence (Bi and Bii), and late phase disclosed areas of hypo and hyperfluorescence associated with increased staining of choroid in the posterior pole (Ci and Cii).

reported to manifest as scleritis, episcleritis, or optic neuritis. Non-necrotizing anterior scleritis (5.8%), non-necrotizing sclerokeratitis (5.8%), and optic neuritis (5.8%) were the least common findings in our study population. Since *T. pallidum* is a non-culturable bacteria, the diagnosis is considered to depend on mainly serological tests. Three syphilis testing algorithms – traditional algorithm, reverse algorithm, and the European Centre for Disease Prevention and Control (ECDC) algorithm have been reported among the serological diagnostic tests. The traditional algorithm
Figure 8 Represents the spectral domain optical coherence tomography of the right and left eyes at pre-treatment phase.

Notes: Spectral domain optical coherence tomography of right and left eyes of patient with focal chorioretinitis revealed right cystoid macular edema with a foveal thickness of 307 µm (A), and left foveal thickness of 266 µm (B). Spectral domain optical coherence tomography of right and left eyes of second patient with focal chorioretinitis disclosed macular thinning with right foveal thickness of 126 µm (C), and left foveal thickness of 141 µm (D).
Figure 9 represents the spectral domain optical coherence tomography of the right and left eyes at post-treatment phase.

Notes: Spectral domain optical coherence tomography of right and left eyes of two patients with focal chorioretinitis discloses right foveal thickness of 131 µm (A), left foveal thickness of 147 µm (B), for the first patient, and right foveal thickness of 368 µm (C), left foveal thickness of 266 µm (D), for the second patient at the 6th month of follow-up exam.
uses a non-treponemal test, VDRL or RPR for screening, and screen-reactive samples are tested by a treponemal assay such as MHA-TP of FTA-ABS for confirmation. The reverse screening algorithm uses an automated, treponemal test such as enzyme immunoassay (EIA). Samples that are reactive by EIA are then tested by RPR to assess disease and treatment status. This technique allows clinical laboratories a rapid syphilis testing. However, the results of EIA screening and RPR are often reported discordant such as reactive by EIA and non-reactive by RPR. Centers for Disease Control and Prevention (CDC) recommends to perform MHA-TP to further analyze the discordant sera. A non-reactive MHA-TP result suggests that the results of EIA screening are falsely reactive, while a reactive MHA-TP result supports either a past, successfully treated syphilis or late/latent syphilis. The European Centre for Disease Prevention and Control algorithm uses a treponemal immunoassay that is followed by a second, different treponemal assay as a confirmatory test in high-prevalence populations. We used the traditional algorithm for the serological diagnosis of syphilis in our study.

*T. pallidum* disseminates to the central nervous system within days after exposure. Neurological involvement is reported to occur during any stage of the infection, and it is manifested as asymptomatic neurosyphilis, acute meningeal syphilis, meningoarterial syphilis, paretic neurosyphilis, and tabetic neurosyphilis. Syphilis-specific CSF diagnostic tests including CSF VDRL, FTA-ABS, or PCR for *T. pallidum* DNA are recommended whenever any additional signs of neurosyphilis such as headache, hypoacusis, tinnitus, cranial nerve abnormalities, and motor and sensory symptoms occur. The majority of cases of neurosyphilis are reported in HIV-infected patients. This is considered to be associated impaired control of syphilis infection due to compromised host immune activity as a result of HIV co-infection. Especiably, HIV-patients with a CD4 count of <350 cells/µL seem to be under a higher risk of developing CSF and neurological abnormalities most likely meningitis.

Co-infections with other causes of STI such as *Neisseria gonorrhoea*, Herpes simplex virus 2, *Chlamydia trachomatis*, Hepatitis B and Hepatitis C virus and *Trichomonas vaginalis* are reported to be prevalent in developing countries. The incidence of syphilis between 1990 and 1996 in Slovak Republic has been reported to increase nearly five times, and the patients have been most commonly co-infected with *Neisseria gonorrhoea*. Increased number of cases of
Figure 11 (Continued)
Figure 11 Represents the color fundus photo and Humphrey visual field 24-2 of the right eye at pre-treatment and post-treatment phases.

Notes: (A) Color fundus photo of the patient with right papillitis discloses 2+ hyperemia and 2+ swelling of the optic disc. (B) Humphrey visual field 24-2 discloses right central scotoma and increased blind spot. (C) Color fundus photo of the right eye discloses resolution of optic disc congestion and edema at the 1st month of follow-up visit. (D) Humphrey visual field 24-2 discloses improvement of central scotoma at the 1st month of follow-up visit.
Syphilis in the Black Sea region of Turkey has been reported between the years 1992 and 1994 related to unprotected sex, promiscuity, and prostitution. The incidence of syphilis in pregnant women has also noted to increase in Slovak Republic between 1990 and 1996 with the stable number of cases with CS. TORCHES (toxoplasmosis, rubella, cytomegalic inclusion disease, herpes virus including Epstein-Barr, syphilis) was defined as congenital co-infection with more than one pathogens, and it was considered as an infectious etiology of congenital cataract. CS and congenital toxoplasmosis are more frequent infectious causes of congenital blindness or visual impairment. Rare cases of CS co-infected with congenital rubella and congenital toxoplasmosis co-infected with congenital rubella have been described. However, co-infected with T. gondii in congenital ocular syphilis has been scarcely described in the literature. Our study disclosed that two of the cases who were siblings with congenital late syphilis were seropositive for T. gondii-specific IgG antibody. The cause of their visual impairments was related to posterior uveitis presented as central retinochoroiditis.

Chorioretinitis in the macula and delay in diagnosis were related to visual loss in patients with ocular syphilis. The BCVAs of our patients with CS showed slight improvement from 20/80 to 20/50 with IV PCN and low-dose TMP–SMX which was considered as prophylaxis related to the presence of risk factor that was seropositivity for syphilis.

MTX has been considered to have an adjunctive role to suppress intraocular inflammation, reduce uveitic macular edema, and prevent the recurrences of the disease. Cases 1–3 received adjunctive therapy with MTX after specific therapy with IV PCN to resolve the residual intraocular inflammation and CME.

Conclusion
Ocular syphilis is uncommon cause of ocular inflammation in HIV-negative patients. Central retinochoroiditis is the most common ocular manifestation, and it is the most common cause of visual impairment. Ocular syphilis might present associated with co-infections such as T. gondii in developing countries. Oral MTX might be beneficial as an adjunctive therapy for ocular syphilis in resolving the residual intraocular inflammation and CME after specific therapy with IV PCN.

Disclosure
The authors report no conflicts of interest in this work.

References