Recurrent Neuroretinitis
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Chief Complaint: Blurred vision.

History of Present Illness: This patient is a 29-year-old white female who noticed that her vision had been getting progressively worse in the right eye (OD) over the past two weeks. She also felt that her central vision was diminished and she had pain with eye movements. The patient went to her local optometrist and was then referred to the University of Iowa for further evaluation.

Past Ocular History: She had a similar episode four years prior to the current episode involving the left eye (OS). At that time she developed gradual onset blurred vision and pain with eye movements. She was seen by her local optometrist who diagnosed her with a “swollen” optic nerve. She was then referred to a neurologist who performed an MRI scan. The scan was reportedly “normal” and she was diagnosed with pseudotumor cerebri. She received no treatment and the vision slowly recovered over the next four months. However, she noted a persistent defect in her inferior nasal visual field OS.

Past Medical History: Three miscarriages at the end of the first trimester, otherwise unremarkable.

Medications: None.

Family History: No eye diseases, and otherwise noncontributory

Social History: Married, lives with her husband. No dietary restrictions. No exposure to cats.

Ocular Exam:
- Visual Acuity (without correction):
  - 20/100 OD
  - 20/20 OS
- Extraocular motility: Full, no nystagmus, no internuclear ophthalmoplegia, normal pursuit, normal saccades
- Pupils: OD 6mm dark, 3 mm light; OS 6 mm dark, 3 mm light; 1.8 log unit relative afferent papillary defect (RAPD) OD
- Intra-ocular pressure: 11 mmHg OD, 12 mmHg OS
- External and anterior segment examination: Normal OU
- Goldmann visual fields (GVF): see below; marked generalized constriction with loss of the smaller isopters OD; inferior nasal constriction OS
- Critical Flicker Fusion: unable to perform test OD; 28.1 Hz (normal>27Hz) OS
- Dilated fundoscopic exam:
  - OD—Normal media, marked optic disc edema, subretinal and intraretinal fluid extending from the optic nerve into the macula, normal vessels, and normal periphery;
  - OS—clear media, pale optic nerve, normal cup-to-disc ratio with peripapillary atrophy, normal macula, normal vessels, small chorioretinal scar in the far inferior periphery.
Figure 1. 30 degree fundus photos of the right (OD) and left (OS) eyes showing optic disc edema and subretinal fluid OD and mild optic nerve pallor OS with peripapillary atrophy.

Figure 2. Image focused on the macula of the right eye showing marked subretinal fluid extending into the fovea from the optic nerve.
**Figure 3.** Optical Coherence Tomography (OCT) scans of the right eye (A) and the left eye (B) showing subretinal and marked intraretinal fluid and resulting thickening of the retina (white triangular area) in the right eye. Note that the fluid is predominately in the outer plexiform layer and there are a few visible highly reflective particles within the fluid. There is some mild thinning of the macula in the left eye with a central macular thickness (CMT) of 265µm.

**Figure 4.** OCT scan of the peripapillary nerve fiber layer OD and OS. Note that there is an increase in the nerve fiber layer thickness OD consistent with disc edema (107 µm; note that the RNFL algorithm failed, underestimating the thickness). There is a decrease in the nerve fiber layer thickness in the superior and inferior arcuate bundles OS consistent with optic atrophy (53 µm).
Figure 5. Goldmann visual fields demonstrate marked constriction of all isopters OD, with loss of the smaller isopters (I1e, I2e). In the left eye there is absolute field loss involving the inferior nasal quadrant with preservation of the central I1e isopter (green).

Clinical Course:

A presumptive diagnosis of neuroretinitis was made based on the patient’s clinical appearance. Specifically, the appearance of the right optic nerve, the profound visual field loss as well as the accumulation of fluid in the outer plexiform layer are very suggestive of this condition. In addition, she had a history of a similar episode in the left eye associated with a "swollen" optic nerve years before. The appearance of optic atrophy and the inferior nasal scotoma in the left eye suggest that the previous episode also affected both the inner and outer retinal layers. However, the patient had very minimal if any exposure to cats. Therefore, a thorough work-up was performed to investigate other etiologies. Blood was drawn for testing of syphilis (RPR), antinuclear antigen (ANA), Bartonella Henselae, complete blood count (CBC), Lyme titers, and angiotensin converting enzyme (ACE). She was treated with a five day course of oral azithromycin and two weeks of oral prednisone and asked to return in one month.

At the next appointment (one month later) she felt that her vision had improved to some extent in the right eye. Examination at that time revealed:

- Visual Acuity (without correction):
  - 20/30 OD
  - 20/20 OS
- Pupils: OD 4 mm dark, 2 mm light; OS 4 mm dark, 2 mm light; No Relative afferent papillary defect (RAPD)
- External and anterior segment examination: Normal OU
- Goldmann visual fields (GVF): see below; mild improvement of the generalized constriction OD
- Dilated fundoscopic exam: OD—Normal media, improved optic disc edema and mild global pallor, resolution of the subretinal and intraretinal fluid in the macula, a prominent macular star, normal vessels, and normal periphery

Laboratory work revealed negative testing for syphilis (RPR), Bartonella Henselae, and Lyme. The CBC, ANA, and ACE levels were all within normal limits. An MRI scan of the brain and orbits was performed on this date due to the atypical nature of the case. There was no mass, enhancing lesion, or demyelinating lesion seen on the scan. One month after this follow up visit, the patient developed symptoms of
decreased vision in the left eye. She was not seen by a physician at this time and the symptoms resolved after about 2 weeks. She returned to the University of Iowa three months following her initial presentation and was felt to have suffered a recurrence of the disease in the left eye. Her visual acuity had decreased to 20/25 OS, there was mild disc edema, and her visual field was worse. She was offered treatment with systemic immunosuppressant medication but declined as she was actively trying to conceive a child.

**Figure 6.** A. Goldmann visual field of the right eye one month after initial presentation demonstrating some improvement, however there is still marked constriction of all isopters and loss of the smaller isopters (I1e, I2e). B. Fundus photo of the right eye showing near complete resolution of the optic disc edema, optic nerve pallor, and the presence of a prominent macular star in the right eye.

**Figure 7.** OCT of the right macula (A) revealing resolution of the outer plexiform layer fluid but now more reflective areas consistent with intraretinal exudates correlating with the macular star figure as well as generalized thinning (CMT 260 µm). OCT measurements of the nerve fiber layer (B) reveal thinning OU (OD 64 µm and OS 56 µm) after 3 months from the acute visual loss.
Discussion:

Neuroretinitis is a clinical diagnosis made based on a constellation of ocular findings. These signs include unilateral painless visual loss, optic disc edema, macular edema, and the later development of a macular star. In the acute phase, vitreal cells are commonly seen. The majority of cases are thought to be due to current or previous infection with Bartonella henselae, a Gram-negative proteobacteria, however, infection with a number of different agents has been postulated to lead to the disease. Most patients with typical neuroretinitis generally suffer a single attack, with slow resolution of symptoms, with or without treatment. The visual prognosis is generally excellent.

In contrast to the above stated clinical course, there exists a subset of patients who suffer repeated attacks of neuroretinitis involving the same or opposite eye. These patients, by definition, have negative serologic studies and thus the disease is referred to as idiopathic recurrent neuroretinitis. While this condition is uncommon, it can be visually devastating. Patients may develop permanent central or cecocentral scotomas, optic disc atrophy, and macular and retinal nerve fiber layer thinning. A number of case reports exists in the literature regarding this disease, its clinical course, and therapeutic options. In the current case, we decided to treat the patient with a short course of oral azithromycin to target Bartonella henselae as well as a short course of steroids to help decrease inflammation. This therapy was initiated upon our first clinical encounter with the patient, when the diagnosis of “typical” neuroretinitis was considered. However, as all of her serologic testing was negative, and she developed recurrent symptoms in the opposite eye, a diagnosis of idiopathic recurrent neuroretinitis was made. At that time we discussed the option with the patient of treatment with systemic immunosuppressive agents, but it was decided not to pursue this treatment, as she was trying to conceive.

To our knowledge, there has only been one case series reported in the literature of idiopathic recurrent neuroretinitis and potential treatment options. In their report, Purvin et. al.describe a retrospective analysis of 30 patients with idiopathic recurrent neuroretinitis treated over a period of 18 years. Of these, seven patients were treated with ongoing immunosuppressive therapy which consisted of low dose oral prednisone and/or oral azathioprine. In each patient, treatment was initiated after several attacks of neuroretinitis which were generally treated acutely with a short course of oral steroids or antibiotics. At some point during their course, the patients began treatment with 50-150 mg/d of azathioprine and/or 10 mg/day of prednisone given on alternate days. Although their study was not designed as a prospective randomized study, the authors did find that treatment as described above decreased the rate of attacks per year by 72% (0.58 attacks per year prior to treatment versus 0.16 attacks per year after treatment).

Diagnosis: Idiopathic Recurrent Neuroretinitis

Differential Diagnosis: Typical Neuroretinitis, Optic Neuritis, Anterior Ischemic Optic Neuropathy (AION)
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<td>• Idiopathic</td>
<td>• Optic disc edema</td>
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<td>• Autoimmune mechanism postulated,</td>
<td>• Macular edema</td>
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<td>possibly related to a</td>
<td>• Fluid in the outer plexiform</td>
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<td>infection (i.e. Bartonella</td>
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<td>Henselae)</td>
<td>• Vitreous cells</td>
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<td>• Late development of a</td>
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<th>SYMPTOMS</th>
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<td>• Decreased vision</td>
<td>• Observation</td>
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<td>• Painless</td>
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<td>• Oral steroids</td>
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REFERENCES


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