New Frontiers in the Management of NONINFECTIOUS UVEITIS INVOLVING THE POSTERIOR SEGMENT

Highlights from a roundtable discussion

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Learning Method and Medium
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Content Source
This continuing medical education (CME) activity captures content from a roundtable discussion held on January 23, 2015, in Oxford, England.

Activity Description
Noninfectious posterior uveitis is a worldwide challenge for clinicians. Suboptimal treatment can cause irreversible visual impairment, which is particularly concerning because uveitis often strikes people of working age. Local and systemic corticosteroid therapy is the current mainstay of treatment. Significant developments for improving uveitis management are under way that hope to address the drawbacks of current therapy side effects, improve tolerance to therapy, and improve patient outcomes. The purpose of this activity is to provide clinicians with practical insights on current therapy and new data on emerging treatments that can be applied to daily practice.

Target Audience
This activity intends to educate US and European retina specialists and other ophthalmologists caring for patients with noninfectious uveitis.

Learning Objectives
Upon completion of this activity, participants will be better able to:
• Review elements of the diagnostic evaluation to differentiate noninfectious from infectious uveitis
• Describe the guidelines pertaining to the treatment of noninfectious uveitis
• Review the indications for local and systemic therapies in the treatment of noninfectious uveitis
• Describe the mechanisms and clinical trial data for emerging local therapy for noninfectious uveitis

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Introduction

Uveitis represents a large and diverse group of intraocular inflammatory diseases involving the uveal tract. Although it is an uncommon diagnosis, its incidence and prevalence may be higher than previously thought. Furthermore, uveitis is an important public health issue because of its potential to cause loss of vision; and, its effect on vision and quality of life also may be greater than often realized. Approximately one-third of patients with uveitis may develop vision impairment during the clinical course of disease, approximately 20% may meet criteria for legal blindness, and uveitis has been reported to account for up to 10% of cases of blindness in the United States and for 3% to 7% of cases in Europe.

Permanent vision loss from uveitis is potentially preventable with effective control of the inflammation. The approach to management, however, varies depending on the specific condition. Therefore, accurate diagnosis is the first step for determining appropriate care.

General ophthalmologists often refer patients with uveitis involving the posterior segment to a retina specialist. This review aims to assist retina specialists with their clinical decision-making for such cases. It provides guidance on the initial diagnostic evaluation, offers suggestions on when to refer, and discusses treatment with a focus on the use of local corticosteroids for noninfectious uveitis involving the posterior segment. In addition, it reviews some emerging local therapies that may become part of the therapeutic armamentarium used by retina specialists in the future.

Overview of Uveitis

Uveitis is classified anatomically as anterior, intermediate, posterior, or panuveitis, according to the primary site of inflammation (Table 1). Each of those categories, however, encompasses a number of conditions that can be characterized further along other dimensions. Uveitis can be described by various temporal features (Table 2) and, according to its etiology, in terms of being infectious or noninfectious, with the latter subgroup comprising entities that are either autoimmune, traumatic, idiopathic, or masquerade conditions (ie, neoplastic or drug-induced disorders mimicking uveitis).

TABLE 1. The SUNa Working Group Anatomic Classification of Uveitis

<table>
<thead>
<tr>
<th>Type</th>
<th>Primary Site of Inflammation</th>
<th>Includes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anterior Uveitis</td>
<td>Anterior chamber</td>
<td>Iritis, iridocyclitis, anterior cyclitis</td>
</tr>
<tr>
<td>Intermediate Uveitis</td>
<td>Vitreous</td>
<td>Pars planitidis, posterior cyclitis, hyalitis</td>
</tr>
<tr>
<td>Posterior Uveitis</td>
<td>Retina or choroid</td>
<td>Focal, multifocal, diffuse choroiditis</td>
</tr>
<tr>
<td>Panuveitis</td>
<td>Anterior chamber, vitreous, and retina or choroid</td>
<td></td>
</tr>
</tbody>
</table>

aSUN=Standardization of Uveitis Nomenclature.

*As determined clinically

Anterior uveitis is the most common form of the disease, but intermediate, posterior, and panuveitis are more likely to be sight-threatening. In Western countries, uveitis is more often noninfectious than infectious. Between 25% and 50% of cases overall are associated with a systemic disease, whereas 35% to 50% lack an associated systemic disease or named syndrome. The distribution of etiologies varies among the different forms of uveitis. For example, toxoplasmosis is the most common cause of posterior uveitis whereas in intermediate uveitis, the majority of disease is idiopathic and limited to the eye. Among the minority of intermediate uveitis cases with a systemic disease association, sarcoidosis and multiple sclerosis (MS) are the main contributing conditions.

Diagnosis

A proper diagnostic evaluation of patients with posterior uveitis will allow retina specialists to exclude an infectious cause and then to distinguish between those patients they may manage on their own with local therapy and patients whom they would be inclined to refer or comanage because of the complexity of the disease.

The diagnostic assessment of patients with uveitis begins with a rigorous history and comprehensive ocular examination. The findings from these investigations alone may lead to a diagnosis or else can guide further targeted investigations or referral to another specialist.

A systematic approach to the diagnostic evaluation of patients with uveitis has been described by Jabs and Busingye. Briefly, the ocular history should review onset, duration, and clinical course of the uveitis as well as identify any symptoms the patient is experiencing. Other factors to consider from the history include patient age, sex, race/ethnicity, immune status, and possibility of endemic exposures based on geographic or other characteristics. In addition, a review of systems should be performed to identify any associated systemic disease (Table 3), recognizing, however, that uveitis may be the initial manifestation.

TABLE 2. The SUNa Working Group Descriptors of Uveitis

<table>
<thead>
<tr>
<th>Category</th>
<th>Descriptor</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Onset</td>
<td>Sudden</td>
<td>Insidious</td>
</tr>
<tr>
<td>Duration</td>
<td>Limited</td>
<td>Persistent</td>
</tr>
<tr>
<td></td>
<td>Persistent</td>
<td>&gt;3 months’ duration</td>
</tr>
<tr>
<td>Course</td>
<td>Acute</td>
<td>Episode characterized by sudden onset and limited duration</td>
</tr>
<tr>
<td>Recurrent</td>
<td>Repeated episodes separated by periods of inactivity without treatment ≥3 months in duration</td>
<td></td>
</tr>
<tr>
<td>Chronic</td>
<td>Persistent uveitis with relapse in &lt;3 months after discontinuing treatment</td>
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</table>

aSUN=Standardization of Uveitis Nomenclature.

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Similarly, other testing as part of the workup of a patient with intermediate or posterior uveitis is ordered on the basis of the pretest probability. For example, Western blot for Lyme disease might be ordered when there is reason to suspect the diagnosis according to the patient's history and/or extraocular signs and symptoms. Chest x-ray, serum angiotensin-converting enzyme, and liver function tests may be useful if sarcoidosis is suspected. Skin (Mantoux) or blood (interferon-release gamma assay) tests can be ordered to rule out TB. Testing for Bartonella infection (catscratch disease) might be considered in persons with exposure to cats and a clinical presentation suggestive of this infection. Focal necrotizing retinitis should raise suspicion of toxoplasmosis. Serologic testing is useful in uncertain cases. A negative result can rule out toxoplasmosis; a positive result, however, is not typically helpful given the high prevalence of seropositivity in most populations.

Establishing a specific diagnosis in patients with uveitis involving the posterior segment may necessitate referral or consultation with a colleague with expertise in uveitis care. It is important to note that in some cases, the clinical presentation and the guided investigations may not lead to a conclusive diagnosis and, especially in cases of disease progression, a more invasive approach to the diagnosis may have to be used, including anterior chamber taps, vitreous biopsies, or chorioretinal biopsies. The samples obtained will be analyzed by different techniques, including polymerase chain reaction, cytology, and histopathology, to exclude an infectious process or a malignancy. This is a situation which will be more appropriately guided by the specialist.

Patients in whom early referral should be considered include those with intermediate uveitis presenting with extensive snowbanking, posterior synechiae, neovascularization, or dense vitritis preventing visualization of the fundus because these findings are indicative of a more serious/complex condition and not simply of a pars planitis (Figures 1 and 2), as well as those patients with focal or multifocal retinal or choroidal lesions or occlusive vasculitis. Early referral is also advised if there is any uncertainty about the diagnosis. In addition, any patient whose uveitis is suspected to be part of a multisystem disorder should be referred to the appropriate specialist for further evaluation and management of the extraocular disease.

Currently, in uveitis centers in Germany and France, brain magnetic resonance imaging (MRI) is performed in younger patients with intermediate uveitis according to the request of neurologists who believe that it may allow early diagnosis of MS and thus initiation of disease-modifying treatment that might limit neuronal damage. Further studies are needed, however, to examine the cost-effectiveness of this practice as well as to prove that early intervention can modify MS. It is noteworthy that MRI does not represent standard of care elsewhere. Without question, a patient with intermediate uveitis and neurologic symptoms should be referred to a neurologist for further evaluation. While the role of MRI as a screening tool for MS is questionable, it is important for ruling out MS when planning therapy with an anti-tumor necrosis factor-α (anti-TNFα) agent that can exacerbate demyelinating disease.

Similarly, other testing as part of the workup of a patient with intermediate or posterior uveitis is ordered on the...
Treatment for Uveitis

The goals in treating uveitis are to address any underlying stimulus (ie, infectious cause), to eliminate ocular inflammation, to prevent or treat complications associated with inflammation (eg, cystoid macular edema [CME]), and when needed for chronic and relapsing conditions, to maintain remission using a regimen associated with the least ocular and systemic morbidity. When appropriately applied, these principles will limit or avoid structural damage to ocular tissues.

Whereas infectious uveitis is treated with a pathogen-specific antimicrobial agent with or without a corticosteroid, corticosteroids are the mainstay of treatment for noninfectious disease, especially for the “induction” phase of suppressing active ocular inflammation.

However, immunomodulatory therapy (IMT) may be indicated if a corticosteroid alone does not control the inflammation or if it cannot be tapered to a safe dose. In addition, IMT represents first-line therapy for certain diagnoses whose natural history is known to involve relapses with the potential for permanent structural damage, or those known to be chronic or needing prolonged treatment. Classic examples include Behçet disease, birdshot retinochoroidopathy, Vogt-Koyanagi-Harada disease, sympathetic ophthalmia, multifocal choroiditis with panuveitis, and serpiginous choroiditis when TB is excluded. IMT may also be indicated for management of an underlying systemic disorder.

In 2000, an expert panel developed guidelines on the management of ocular inflammatory disorders using oral corticosteroids and immunosuppressive drugs (antimetabolites, T-cell inhibitors, alkylating agents). The guidelines recommend initiating systemic therapy with a high-dose corticosteroid regimen and adding IMT if the corticosteroid dose cannot be tapered within 3 months to a prednisone equivalent of ≤10 mg/d. Since that report was published, there has been accumulating experience using biologic therapies, and in 2014, guidelines were published on the use of anti-TNFα agents in patients with ocular inflammatory disorders.

Any clinician treating uveitis should be aware of the guidelines, to help inform systemic corticosteroid use and to recognize when IMT and biologic agents are needed so that timely referral can be made, if needed.

Results of a survey examining patterns of treatment for noninfectious posterior uveitis revealed that only 25% of the participating physicians (60 ophthalmologists and 3 rheumatologists) were aware of or used the treatment guidelines. These results also suggested that oral corticosteroids were being overused. Oral corticosteroid treatment was being administered at a mean prednisone equivalent dose of 38 to 46 mg/d for a mean duration of 21 months. Attempts at steroid dose reduction were made in only 10% to 31% of cases. Rates of steroid-related complications were high, and only 12% of patients had been prescribed IMT as a corticosteroid-sparing strategy.

Use of an insufficient initial corticosteroid dose to achieve induction goals of suppressing active inflammation, or failure to initiate IMT when indicated in situations in which suppressive, maintenance therapy is indicated, increases the risk for vision loss from inadequately controlled ocular inflammatory disease. Failure to taper the corticosteroid appropriately exposes patients to an increased risk for corticosteroid-related toxicity.

Corticosteroid treatment

General principles. Treatment with a corticosteroid should be initiated after an infectious etiology or masquerade condition has been excluded. It is often appropriate to begin treatment while the workup is in progress because severe damage may occur while waiting for a definitive diagnosis. However, considering the potential for harm if an infectious uveitis or masquerade syndrome is treated empirically with corticosteroids or immunotherapy, clinicians should never proceed with treatment in the absence of an ongoing diagnostic evaluation.

Not all noninfectious uveitides involving the posterior segment necessitate corticosteroid treatment (Table 4). Considering that a substantial minority of patients with intermediate uveitis have a mild condition and maintain normal visual acuity without any treatment, corticosteroid treatment for noninfectious intermediate uveitis primarily should be considered only if there is macular edema or vitreous haze causing a clinically important loss of vision or if there is evidence of structural complications from the uveitis (eg, synechiae, cataract, glaucoma). Cystoid macular edema detected by OCT represents an indication for treatment, even when vision is good, because chronic CME is a leading cause of vision loss in patients with uveitis.

Topically administered corticosteroids reliably penetrate only into the anterior segment. Therefore, in most cases, corticosteroid treatment for uveitis involving the posterior segment necessitates a local injection or systemic therapy. Systemic corticosteroid treatment may be preferred if the uveitis is bilateral and/or associated with extraocular disease. Patients with these findings may warrant referral to a colleague with appropriate expertise in managing uveitis. Oral corticosteroid therapy may be safer in terms of ocular complication risk (cataract, intraocular pressure [IOP] elevation, or glaucoma) and can be more readily withdrawn than depot injections. However, treatment with an oral corticosteroid is commonly associated with systemic side effects, which many patients are unable to tolerate.

Local therapy with periocular or intravitreal corticosteroid injections can be used to manage intermediate uveitis associated with decreased vision or to manage macular edema in eyes with panuveitis or posterior uveitis.

TABLE 4. Noninfectious Uveitides Involving the Posterior Segment That May Not Necessitate Corticosteroid Treatment

<table>
<thead>
<tr>
<th>Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mild vitritis with no macular edema</td>
</tr>
<tr>
<td>Vascular sheathing without significant vitritis</td>
</tr>
<tr>
<td>Pars planitis with excellent vision and without macular edema or significant vitritis</td>
</tr>
<tr>
<td>Acute posterior multifocal placoid pigment epitheliopathy without macular involvement</td>
</tr>
</tbody>
</table>

Segment That May Not Necessitate Corticosteroid Treatment
Patient selection for local injectable therapy. Patients with uveitis who are most appropriately treated with local corticosteroid injectable therapy alone are those without a systemic disorder who have unilateral or asymmetric intermediate uveitis or posterior uveitis with decreased vision due to macular edema or vitreous haze in the absence of retinitis, chorioretinitis, or oscillusive retinal vasculitis. Patients with bilateral disease with limited vision impairment could also be good candidates. Conditions fitting these descriptions (Table 5) tend to be limited to the eye, relatively unaggressive, and potentially remitting or not needing chronic suppressive therapy, and are therefore suited to local therapy with periocular or intravitreal corticosteroid injections.

In addition, some posterior uveitides associated with systemic disease can be managed with local therapy by the retina specialist in collaboration with the appropriate internal medicine specialist. Such cases would include those in which the ocular inflammation is similar to that in the cases described above and in which the associated systemic disease is relatively mild or because the treatment needed for the systemic disease allows management of the ocular inflammation with local therapy. A good example is sarcoidosis in which systemic disease may be mild and even asymptomatic.

Whether or not local treatment for the uveitis is needed for the latter patients, the retina specialist will continue to have a role as a comanager, with responsibilities for monitoring the ocular response to therapy, uveitis activity over time, and the development of disease- and treatment-related complications.

### TABLE 5. Uveitides Generally Suitable for Local Injectable Therapy

<table>
<thead>
<tr>
<th>Uveitis Condition</th>
<th>Characteristics</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pars planitis or other forms of intermediate uveitis in which there is no need for continuous suppressive treatment</td>
<td>Viritis, snowballs, nonconstant CME</td>
<td>Refer if CME is recurrent or if continuously suppressive therapy is needed</td>
</tr>
<tr>
<td>PIC or MFC without panuveitis but with CNV (needs combined use of steroid and anti-VEGF treatment)</td>
<td>May be bilateral, but flare-up and CNV do not occur in both eyes at the same time</td>
<td>Refer if response is suboptimal or if there is early recurrence</td>
</tr>
<tr>
<td>Acute anterior uveitis (eg, HLA-B27 associated) complicated by macular edema not responding to standard topical therapy</td>
<td>Predominantly anterior or intermediate uveitis that is not itself necessitating chronic immunosuppressive therapy</td>
<td>Refer if the condition advances to need chronic suppressive therapy or if the patient is doing poorly</td>
</tr>
<tr>
<td>Uveitides associated with systemic diseases that either are mild or are predominantly controlled by treatment indicated and administered for the systemic disease (eg, sarcoidosis, MS)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*The content of this table represents the authors’ recommendations. CME=cystoid macular edema; CNV=choroidal neovascularization; HLA-B27=human leukocyte antigen B27; MFC=multifocal choroiditis; MS=multiple sclerosis; PIC=punctate inner choroidopathy; VEGF=vascular endothelial growth factor.

### Periocular corticosteroids

Injection into the sub-Tenon space or into the orbital floor using a transcutaneous or transconjunctival approach are the most common techniques for periocular corticosteroid injection.

A recent retrospective cohort study highlighted the efficacy of periocular corticosteroid injections for controlling vitritis and improving vision when impairment is related to macular edema. The analysis included 1192 eyes of 914 patients who received 1 or more periocular injections for anterior, intermediate, posterior, or panuveitis. By 6 months, inflammation was controlled completely in 73% of eyes and best corrected visual acuity (BCVA) improved to 20/40 or better in 50% of eyes that initially presented with BCVA ≤20/40. Additional injections did not necessarily provide much additional benefit, but did increase the risk for cataract and cataract surgery.

Studies including findings from earlier follow-up of patients treated with a sub-Tenon corticosteroid injection show that inflammation improves within days to weeks and that visual acuity and macular edema improve within weeks to months. However, compared with administration directly into the vitreous, periocular corticosteroid injection may have a slower onset of efficacy and shorter duration of action (2 months vs 3-4 months). Therefore, repeat injections may be needed.

Periocular corticosteroid treatment may be less likely than direct intravitreal administration to cause IOP elevation and cataract, and certainly costs less than intravitreal treatment with a long-acting implant. Therefore, when initiating treatment for uveitis and/or uveitic macular edema with a corticosteroid injection, it would be reasonable to choose periocular administration first.

### Intravitreal corticosteroids

Intravitreal treatment might be reserved for local therapy if the uveitis does not respond sufficiently to a periocular injection or when ongoing therapy is needed for controlling chronic inflammation and uveitic edema. One product, an implant containing dexamethasone 0.7 mg, is approved as intravitreal treatment for uveitis in Europe, although a nonophthalmic preparation of triamcinolone acetonide is commonly used off-label. Four products for intravitreal use are approved for treatment of uveitis in the United States—the dexamethasone implant, 2 triamcinolone acetonide injectable suspensions, and a long-acting implant releasing fluocinolone acetonide (FA) 0.59 mg. The implants are specifically approved for treatment of chronic noninfectious uveitis of the posterior segment whereas the triamcinolone suspensions have a more general indication for uveitis.

The dexamethasone 0.7-mg implant uses a biodegradable delivery system, can be placed in an outpatient procedure, and releases corticosteroid over a period of approximately 6 months in a 2-phase manner with a low concentration of drug release beginning after 3 months. The efficacy of the dexamethasone implant for improving intraocular inflammation and visual acuity in patients with noninfectious intermediate or posterior uveitis was established in a pivotal trial that followed participants to 26 weeks after a single implantation. A multicenter retrospective cohort study evaluated postmarketing experience with the...
dexamethasone implant for treatment of noninfectious uveitis. It included 82 eyes of 63 patients who received a total of 142 injections over a period of up to 35 months. The results showed benefit for improving macular edema as well as for vitreous haze and vision. Reinjection was usually needed at approximately 6 months. IOP elevation (>21 mm Hg) occurred in 33 (40.2%) eyes; all but 1 of those eyes required medical treatment and 2 (2.4%) eyes underwent glaucoma surgery. Four (10%) of 40 phakic patients underwent cataract surgery.

The FA 0.59-mg implant, a nonbiodegradable device impregnated with the drug, is placed through a pars plana incision and sutured to the sclera in a surgical procedure, and releases corticosteroid for approximately 2.5 years. In the pivotal trial that included patients with posterior uveitis, the FA implant significantly reduced uveitis recurrence compared with sham control and maintained or improved vision during 3 years of follow-up. Nearly all phakic patients (93%) underwent cataract surgery over the 3-year study period and the treatment also increased the risk for IOP elevation. During the 3-year study, 78% of eyes receiving an FA implant (0.59 or 2.1 mg) required treatment with an IOP-lowering medication, and 40% required surgery for IOP control.

In the primary end point analysis at 2 years in a study comparing the FA implant with systemic therapy, visual outcomes were similar in the 2 treatment groups, but the implant arm had better control of macular edema and more often had a 2-step improvement in vitreous haze. An analysis of data from 3 years of follow-up determined the FA implant was reasonably cost-effective compared with systemic therapy for individuals with noninfectious unilateral uveitis involving the posterior segment.

Either the sustained-release dexamethasone implant or the intravitreal triamcinolone can be considered a reasonable choice for treating patients with idiopathic intermediate uveitis that is accompanied by macular edema resistant to periocular triamcinolone acetonide injections. The dexamethasone implant is more expensive, and there is no evidence to show it is superior to intravitreal triamcinolone. In Europe, however, intravitreal triamcinolone acetonide is off-label.

Given its longer duration compared with the dexamethasone implant, the FA implant may be a reasonable choice for patients with significant retinal vascular leakage and chronic CME when prolonged therapy is anticipated. Nevertheless, while the benefits can be significant in the right patient, given the substantial risks associated with long-term corticosteroid exposure, it seems prudent that use of the FA implant should be undertaken only after the retina specialist consults with a colleague who is more experienced in management of uveitis to ensure the patient is an appropriate candidate for this long-lasting corticosteroid treatment.

**Corticosteroid injections for anterior uveitis-related macular edema.** A periorcular corticosteroid injection, the dexamethasone implant, or a systemic corticosteroid may be considered for treatment of persistent macular edema associated with anterior uveitis, but should be used only to treat edema that is truly refractory to topical medication.

Unilateral anterior uveitis that is expected to be self-limiting disease is usually treated initially with intensive topical corticosteroid therapy. Prednisolone acetate, 1%, has been used traditionally and given as often as hourly. Topical difluprednate, 0.05%, available in the United States, is noninferior to prednisolone and can be used on a less frequent dosing schedule as an alternative or as rescue therapy. Topical corticosteroid treatment also can be combined with a topical nonsteroidal anti-inflammatory drug (NSAID), although some clinicians believe that topical NSAIDs have little effect in the setting of uveitic macular edema over and beyond that of topical corticosteroids. Anecdotally, topical nepafenac, 0.1%, has been particularly useful, perhaps because of its particularly high ocular penetration, although little evidence exists to suggest that any one NSAID is more effective than another, or effective at all in this setting.

When treating anterior uveitis, physicians should note there may be a lag in the onset of macular edema and in its resolution after starting effective anti-inflammatory treatment with a topical corticosteroid. Therefore, in eyes being treated for anterior uveitis, topical anti-inflammatory treatment should be continued for at least 2 to 4 weeks before macular edema is judged refractory.

**Acetazolamide**

Acetazolamide has been reported to be an effective treatment for uveitic macular edema when given in addition to anti-inflammatory drugs and particularly in eyes with acute rather than chronic macular edema. A meta-analysis of randomized clinical trials investigating acetazolamide found it was ineffective in improving vision. The poor response may be due to late use after chronic edema has caused permanent damage.

**Emerging Local Therapies for Uveitis**

Drawbacks of existing therapies for uveitis, including toxicity and limited efficacy, have created interest in developing new treatments, and particularly for identifying locally administered agents that would avoid systemic adverse effects and have a better safety profile than corticosteroids.

**Anti-TNFα.** A few investigators have evaluated off-label intravitreal administration of anti-TNFα agents in a small series of eyes with noninfectious uveitis involving the posterior segment. Infliximab was reported to improve vitreous haze, macular edema, visual acuity, retinal vasculitis, and retinitis. The agents, however, were poorly tolerated, probably retinotoxic, and even induced panuveitis and vitritis when investigated as treatment for diabetic macular edema (DME) and age-related macular degeneration. No safety concerns were identified with the use of intravitreal adalimumab, but efficacy data showed mixed outcomes.
Methotrexate. A growing number of reports are describing impressive results with intravitreal methotrexate (off-label) for the treatment of uveitis not associated with lymphoma. A multicenter retrospective study reviewing outcomes of 38 eyes (including 16 with intermediate uveitis or pars planitis and 15 with posterior uveitis or panuveitis) treated with methotrexate 400 mcg found improved visual acuity and/or control of intraocular inflammation and/or CME in 79% of eyes.³⁴ Eight eyes relapsed after 1 to 17 months, but the duration of remission in 22 other eyes extended up to 18 months. According to the available evidence, intravitreal methotrexate is a treatment of potential interest as local therapy for a patient with unilateral disease and a contraindication to corticosteroid use.

Anti-vascular endothelial growth factor (VEGF). Intravitreal anti-VEGF therapy also has been used (off-label) in eyes with macular edema associated with uveitis, particularly in the setting of macular edema with neovascularization. Published reports are limited in number, mostly retrospective, and show improvement in macular edema but not in visual acuity.¹⁵ Studies comparing anti-VEGF treatment with intravitreal triamcinolone found better outcomes with the corticosteroid.¹⁴ In addition, the duration of benefit with anti-VEGF treatment is short-lived, so that frequent repeat injections would be needed. Anecdotally, anti-VEGF treatment may also lead to scarring in inflammatory neovascular membranes. In summary, anti-VEGF therapy is not sufficiently anti-inflammatory to be used as a primary treatment for uveitis. It can, however, be used to manage vascular complications (e.g., choroidal neovascularization or retinal neovessels) once inflammation is controlled, or concomitantly with appropriate anti-inflammatory therapy.

Sirolimus. At present, 1 local therapy, intravitreal sirolimus, has completed a phase 3 study for the treatment of noninfectious uveitis involving the posterior segment. Also known as rapamycin, sirolimus acts as an inhibitor of the mammalian target of rapamycin (mTOR) to block T-cell activation, T-cell proliferation, and the production of an array of pro-inflammatory cytokines.³⁵ Sirolimus also has antiangiogenic properties.³⁶ The intravitreal formulation of sirolimus is a clear solution injected with a 30-gauge needle. In situ, it rapidly forms into a drug depot that gradually dissolves, releasing drug for up to 60 days.³⁵

The phase 3 study SAKURA randomized 347 patients to treatment with sirolimus 440 mcg, 880 mcg, or 44 mcg (control).³⁷ In this 2-year study, injections were given every 2 months for the first year and then as-needed.

Percentage of patients with a vitreous haze score of 0 at month 5 was analyzed as the primary efficacy end point. The rates were significantly higher in the sirolimus 440-mcg group compared with the 880-mcg group and the controls (22.5% vs 16.4% and 10.3%, respectively, P=0.025). Differences favoring sirolimus 440 mcg over the control group were also achieved in secondary end point analyses of percentage of eyes with inactive disease (0 or 0.5+ vitreous haze) (52.6% vs 35%, P=.008) and success tapering systemic corticosteroids (76.9% vs 63.6%, P=not significant).

There were no unexpected safety signals associated with injection of sirolimus. Among the 3 study arms there were no cases of culture-positive endophthalmitis and 1 case of culture-negative endophthalmitis. Rates of noninfectious endophthalmitis in the 44-mcg, 440-mcg, and 880-mcg arms were 0, 0.9%, and 3.4%, respectively. Rates of glaucoma and progression of cataract were also low, each occurring in 0.9% of patients in the 44-mcg arm. In the 440-mcg arm, 0.9% of patients developed glaucoma and 1.8% of patients had progression of cataract.

Should sirolimus become licensed for treatment of uveitis, more details on anatomic outcomes and long-term efficacy would be needed to determine its role in clinical practice. As with any new drug, better understanding of efficacy and limitations will depend on broader experience with its use in clinical practice and additional studies conducted post marketing.

Fluocinolone acetonide intravitreal implant. A third long-acting intravitreal implant containing FA 0.19 mg is currently being investigated as treatment for noninfectious uveitis involving the posterior segment in a multinational phase 3 trial. No outcomes data are available yet.³⁸ This product recently became available in both the United States and Europe with an approval for the treatment of DME. It uses a nonbiodegradable delivery system, as does the FA 0.59-mg implant, and also releases drug over 36 months.³⁹

Summary

Appropriate management of uveitis involving the posterior segment is critical for preserving vision as well as for limiting morbidity associated with treatment and extraocular disease, and retina specialists have an important role to play in delivering this care. Corticosteroids remain the most frequently used agent, and while options were initially limited to periocular injections, they have now expanded to the use of intravitreal injections and intraocular devices. Currently, other drugs are emerging and will undoubtedly become therapeutic options, especially considering their potential to reduce the development of local complications, namely cataract and IOP elevation.

As in any indication, however, the safety and efficacy of intervention depends on careful patient selection. Therefore, a proper diagnostic evaluation along with understanding of existing therapeutic guidelines and situations warranting referral will help ensure patients receive the care they require.
Causes and frequency of blindness in patients with intraocular uveitis. Study. Northern California; the Northern California Epidemiology of Uveitis Study.


1. Intermediate uveitis:
   A. Accompanied by neurologic symptoms might prompt brain MRI
   B. Always necessitates anti-inflammatory treatment because of its sight-threatening potential
   C. Is most often associated with sarcoidosis
   D. Refers to disease in which the choroid is the primary site of inflammation
2. A diagnosis of toxoplasmosis:
   A. Is the most common finding in patients with intermediate uveitis in Western countries
   B. Should always be confirmed by serologic testing
   C. Should be suspected in a patient with focal necrotizing retinitis
   D. Should be suspected in a patient with intermediate uveitis and extensive snowbanking
3. The only diagnostic test recommended as routine in the initial workup of adolescent and adult patients with intermediate uveitis is:
   A. HLA-B27 typing
   B. Serology for syphilis
   C. Brain MRI
   D. Chest computed tomography scan
4. Uveitides for which immunomodulatory therapy represents first-line therapy include all the following, except:
   A. Behçet disease
   B. Birdshot retinochoroidopathy
   C. Sarcoidosis
   D. Sympathetic ophthalmia
5. According to expert panel guidelines, IMT is indicated for treating ocular inflammatory disorders when a corticosteroid cannot be tapered to:
   A. Prednisone equivalent ≤5 mg/d within 3 months
   B. Prednisone equivalent ≤7.5 mg/d within 6 months
   C. Prednisone equivalent ≤10 mg/d within 3 months
   D. Prednisone equivalent ≤10 mg/d within 6 months
6. Treatment of uveitis using a sub-Tenon corticosteroid injection:
   A. Is appropriate for anterior or intermediate uveitis, but not for posterior or panuveitis
   B. Is the preferred route for initiating corticosteroid treatment until an infectious etiology is excluded
   C. Might improve inflammation within days
   D. Provides cumulative benefits with repeat injections
7. The FA implant that is approved for treatment of chronic noninfectious uveitis:
   A. Delivers corticosteroid for approximately 6 months
   B. Has been shown less cost-effective than systemic therapy for treatment of noninfectious unilateral uveitis involving the posterior segment
   C. Requires suturing to the sclera
   D. Was associated with a low rate of cataract surgery in the 3-year pivotal trial leading to its approval
8. At month 5 in the phase 3 SAKURA study investigating sirolimus for treatment of noninfectious uveitis involving the posterior segment, differences favoring sirolimus 440 mcg vs control were seen for all the following end points, except:
   A. Rate of glaucoma formation
   B. Percentage of eyes with inactive disease
   C. Percentage of eyes with a vitreous haze score of 0
   D. Success in tapering systemic corticosteroids
9. An FA 0.19-mg implant:
   A. Demonstrated statistically significant efficacy for reducing uveitis recurrence vs sham control in a phase 3 trial
   B. Has a shorter duration of drug release than the FA 0.59-mg implant
   C. Is available in the United States and Europe for the treatment of DME
   D. Uses a biodegradable delivery system and can be delivered in an outpatient procedure
10. Which of the following statements is true about intravitreal infliximab as treatment for noninfectious uveitis involving the posterior segment?
    A. Infliximab has been investigated as an intravenous infusion, but not as an intravitreal injection for the treatment of uveitis
    B. Infliximab can be considered as first-line treatment for ocular manifestations of Behçet disease
    C. Infliximab has been reported to improve vitreous haze, macular edema, and retinal vasculitis
    D. Infliximab exacerbated vitritis when investigated as treatment for noninfectious uveitis involving the posterior segment
NEW FRONTIERS IN THE MANAGEMENT OF NONINFECTIOUS UVEITIS INVOLVING THE POSTERIOR SEGMENT

Activity Evaluation/Credit Request

To receive AMA PRA Category 1 Credit™, you must complete this Evaluation form and the Post Test. Record your answers to the Post Test in the Answer Box located below. Mail or Fax this completed page to New York Eye and Ear Infirmary of Mount Sinai—ICME, 310 East 14th Street, New York, NY 10003 (Fax: 212-353-5703). Your comments help us to determine the extent to which this educational activity has met its stated objectives, assess future educational needs, and create timely and pertinent future activities. Please provide all the requested information below. This ensures that your certificate is filled out correctly and is mailed to the proper address. It also enables us to contact you about future CME activities. Please print clearly or type. Illegible submissions cannot be processed.

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☐ Yes ☐ No I and/or my family member have a financial relationship with New York Eye and Ear Infirmary of Mount Sinai and/or refer Medicare/Medicaid patients to it.

☐ Yes ☐ No I certify that I have participated in the entire activity and claim 1.5 AMA PRA Category 1 Credits™.

Signature Required ___________________________ Date Completed ___________________________

OUTCOMES MEASUREMENT

☐ Yes ☐ No Did you perceive any commercial bias in any part of this activity? IMPORTANT! If you answered “Yes,” we urge you to be specific about where the bias occurred so we can address the perceived bias with the contributor and/or in the subject matter in future activities.

Circle the number that best reflects your opinion on the degree to which the following learning objectives were met:

5 = Strongly Agree  4 = Agree  3 = Neutral  2 = Disagree  1 = Strongly Disagree

Upon completion of this activity, I am better able to:

• Review elements of the diagnostic evaluation to differentiate noninfectious from infectious uveitis                              5  4  3  2  1
• Describe the guidelines pertaining to the treatment of noninfectious uveitis                                            5  4  3  2  1
• Review the indications for local and systemic therapies in the treatment of noninfectious uveitis                        5  4  3  2  1
• Describe the mechanisms and clinical trial data for emerging local therapy for noninfectious uveitis                    5  4  3  2  1

1. Please list one or more things, if any, you learned from participating in this educational activity that you did not already know.

_____________________________________________________________________________________________________________

_____________________________________________________________________________________________________________

2. As a result of the knowledge gained in this educational activity, how likely are you to implement changes in your practice?

4 = definitely will implement changes  3 = likely will implement changes  2 = likely will not implement any changes

1 = definitely will not make any changes

4  3  2  1

Please describe the change(s) you plan to make:

_____________________________________________________________________________________________________________

_____________________________________________________________________________________________________________

3. Related to what you learned in this activity, what barriers to implementing these changes or achieving better patient outcomes do you face?

_____________________________________________________________________________________________________________

_____________________________________________________________________________________________________________

4. Please check the Core Competencies (as defined by the Accreditation Council for Graduate Medical Education) that were enhanced for you through participation in this activity.

☐ Patient Care ☐ Practice-Based Learning and Improvement ☐ Professionalism

☐ Medical Knowledge ☐ Interpersonal and Communication Skills ☐ Systems-Based Practice

5. What other topics would you like to see covered in future CME programs?

_____________________________________________________________________________________________________________

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ADDITIONAL COMMENTS

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