A Promising Therapeutic Approach for Treatment of Posterior Uveitis: Recombinant T Cell Receptor Ligand Protects Lewis Rats from Acute and Recurrent Experimental Autoimmune Uveitis

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Abstract

Introduction: Chronic autoimmune uveitis is a major cause of vision loss from intraocular inflammation in humans. In this study we report that a recombinant TCR ligand (RTL220) composed of the α1 and β1 domains of MHC class II molecules linked to the uveitogenic interphotoreceptor retinoid-binding protein (IRBP) 1177–1191 peptide is effective in the suppression of acute and recurrent experimental autoimmune uveitis (EAU). Material and Methods: EAU was induced with IRBP1177–1191 peptide or by adoptive transfer of specific T cells in Lewis rats. The rats received 5 doses of RTL220 subcutaneously every other day starting at the onset of clinical signs of EAU. Results: The administration of RTL220 resulted in a delayed onset and a significant amelioration of the disease severity at clinical levels and showed protection of the retina from inflammatory damage at histological levels. In treatment of recurrent EAU, RTL220 administrated at the first or second onset of clinical disease significantly inhibited EAU, modulated immune responses and provided protection from relapses of uveitis. The systemic and local proinflammatory cytokines were significantly reduced, including IL-17. There was local and systemic increase in IL-10 and reduction in the expression of the proinflammatory chemokines CCL2, CCL3 and CCL5. Conclusions: Our studies demonstrate a successful treatment of acute and recurrent EAU with RTL220, which effectively suppressed the recurrence of inflammation and reversed clinical and histological EAU by altering cytokine and chemokine expression. These findings strongly support a possible clinical application of this novel class of peptide/MHC class II drugs for patients with autoimmune uveitis.

Key Words

Experimental autoimmune uveitis, rat · RTL, treatment · Cytokine · Chemokine

Introduction

Uveitis, a sight-threatening inflammatory disease of the eye, can result from immune-mediated responses against ocular antigens. Recurrent uveal inflammation, which occurs in ~50% of the uveitis cases, is the major morbidity because following a resolution of an acute episode, the patient’s vision is frequently decreased by chronic subclinical inflammatory reactions that may eventually lead to blindness [1]. The conventional treatment includes corticosteroids and immunosuppressive agents, which are limited by their side effects and only efficient in approximately half of the recurrent ocular inflamma-
tion patients [2, 3]. Animal models of experimental autoimmune uveitis (EAU) have been useful in testing new therapeutic approaches to treat intraocular inflammation [4].

For uveitis to develop, autoreactive T cells must be activated outside the eye and then pass the blood-ocular barrier, enter the eye and crossreact with ocular autoantigens. In autoimmune uveitis, Th1 and Th17 cells play an important role in the pathogenicity of the disease [5]. IRBP is the major uveitogenic retinal antigen eliciting EAU [4, 6]. In immunologically normal mice or rats, EAU is a T-cell-mediated disease that targets the photoreceptor cells where the target antigens are located, which often leads to an irreversible destruction of photoreceptor cells and subsequently to the loss of vision [4, 7]. Mice and rats genetically predisposed to a predominantly Th1 or Th17 response differ in the severity of EAU that they develop [5].

MHC class I and class II are molecules that influence the immunological susceptibility of many types of noninfectious uveitis [8]. Vogt-Koyanagi-Harada (VKH) disease, sympathetic ophthalmia and birdshot retinochoroidopathy are conditions that show a significant MHC association with both MHC class II molecules and MHC class I molecules, indicating an autoimmune pathogenesis for uveitis [9]. Over the years, a number of different approaches to treatments have been investigated [2, 10, 11]. Blocking the antigen presentation function of MHC class II molecules by competitor peptides has been proposed as a possible therapeutic approach for MHC-associated autoimmune diseases [12–14]. Recombinant TCR ligands (RTLs) consisting of covalently linked β1 and α1 domains of MHC class II molecules that are tethered to specific antigenic peptides represent minimal TCR ligands (RTLs) consisting of covalently linked peptide-major histocompatibility complex (MHC) class II molecules by competitor peptides have been proposed as a possible therapeutic approach for MHC-associated autoimmune diseases [12–14]. Recombinant TCR ligands (RTLs) consisting of covalently linked β1 and α1 domains of MHC class II molecules that are tethered to specific antigenic peptides represent minimal TCR ligands and are a new class of drugs treating autoimmune inflammation [15, 16]. These compounds are developed to bind selectively to the specific TCR of interest and were shown to effectively prevent and reverse clinical and histological signs of experimental autoimmune encephalomyelitis (EAE) in an antigen-specific manner [17, 18].

We have previously demonstrated that RTL201 bearing the nonretinal antigen, myelin basic protein peptide MBP72–85, completely suppressed autoimmune anterior uveitis associated with EAE in Lewis rats [19]. In the present study, our goal was to design a new RTL bearing the retina-specific antigenic peptide IRBP1177–1191 peptide (RTL220) and determine its efficacy in the treatment of acute and, in particular, recurrent EAU (R-EAU). T cells responsive to IRBP and arrestin (S-antigen) have been found in the blood of patients with uveitis, which suggests that these proteins may be the major autoantigens involved in human uveitis [4, 20, 21]. Our findings show a RTL220 can successfully treat R-EAU by altering cytokine and chemokine expression. These results support a possible clinical application in the therapy of autoimmune uveitis.

### Material and Methods

#### Animals

Female Lewis rats (Harlan Sprague-Dawley, Inc., Indianapolis, Ind., USA), 6–8 weeks old, were used in these studies. The rats were housed at the Oregon Health and Science University Animal Care Facility according to institutional and federal guidelines. All animal experimentation procedures adhered to the ARVO Resolution on the Use of Animals in Research and have been approved by the institutional animal committee. The experimental groups consisted of 3 or 6 rats and the experiments were repeated at least twice.

#### Generation and Production of RTL220

The major epitope of IRBP is composed of amino acid residues 1177–1191 (IRBP1177–1191; also called R16 [22, 14]) and was used to design and express our new RTL220 gene. The RTL220 construct is comprised of a single gene encoding a polypeptide of 220 amino acids containing the β1 and α1 domains of the rat MHC class II (RT1.B) and the covalently tethered peptide IRBP1177–1191 (ADGSSWEGVGVVPDV) (fig. 1). The plasmid sequences were confirmed as described previously [15]. The construct was expressed in *Escherichia coli*. RTL220 was purified by chromatography following the same methodology that was used to make the ‘empty’ (RTL101; β1o1 chain only) and antigen-coupled RTL constructs described previously [15]. RTL203 bearing the covalently tethered cardiac myosin peptide CM-2 was used as an irrelevant control [19].

#### Induction and Assessment of EAU

**Acute EAU.** Lewis rats were immunized with 100 μl of 20 μg IRBP1177–1191 in complete Freund’s adjuvant supplemented with 2 mg/ml of *Mycobacterium tuberculosis*. The rats were evaluated for clinical signs of EAU daily by biomicroscopy. The intensity of inflammation was scored blind on an arbitrary scale of 0–4 as follows: 0 – no disease, 1 – engorged blood vessels in the iris and abnormal pupil configuration, 2 – hazy anterior chamber, 3 – moderate opaque anterior chamber with the pupil visible and 4 – opaque anterior chamber, obscure pupil and proptosis.

**Recurrent EAU.** R-EAU was induced by adoptive transfer of IRBP1177–1191-specific T cells as described recently [23]. Donor Lewis rats were initially immunized with 20 μg of IRBP1177–1191 in complete Freund’s adjuvant and *M. tuberculosis*. Ten days later, their spleens were removed, and the suspension of splenocytes was stimulated with 20 μg/ml IRBP1177–1191 peptide for 48 h. Blasts were collected and injected intraperitoneally at 5 × 10⁶ dose per recipient rat. The rats were examined daily for clinical signs of inflammation by biomicroscopy for about 39 days following the T cell transfer.
**Treatment with RTLs**

RTLs at 300 μg per dose were injected subcutaneously in the back at the time of IRBP immunization or at the onset of clinical EAU signs. The rats received 5 doses of RTL220 every other day. To treat R-EAU, 5 dosages of 300 μg of RTL220 were administered starting at the first or second onset of clinical EAU followed by weekly injections of 300 μg of RTL220 subcutaneously for the duration of the experiment.

**Histology**

The eyes were removed and fixed in formalin and then processed for paraffin embedding. Tissue sections (5 μm) were stained with hematoxylin and eosin for histopathological scoring. EAU was scored on a scale of 0 (no disease) to 4 (maximum disease), based on the presence of inflammatory cell infiltration of the iris, ciliary body, anterior chamber and retina, as follows: 0 = normal anterior and retinal architecture, no inflammatory cells in these structures; 1 = mild inflammatory cell infiltration of anterior segment and retina; 2 = moderate inflammatory cell infiltration of anterior segment and retina; 3 = massive inflammatory cell infiltration of anterior segment and retina, disorganized anterior segment and retina, and 4 = massive inflammatory cell infiltration of anterior segment and retina, disorganized anterior segment and retina, and photoreceptor cell damage [24].

**Cytokine/Chemokine Analysis**

The eyes were collected at the end of the experiment. Cytokines and chemokines were extracted from each eye with PBS by homogenization [25]. The soluble fraction was collected by centrifugation from 2 individual eyes of different rats from 2 experiments (IL-13 in 3 eyes) and then used as a cytokine source for determination. The aliquots from each eye were tested in duplicate. Splenocytes obtained from individual spleens were cultured at 5 x 10^5 cells/well in a 96-well flat-bottom culture plate in RPMI stimulation medium with 20 μg/ml IRBP peptide for 48 h. Supernatants were harvested and stored at −80°C until testing for cytokines in triplicate. A customized rat Beadlyte multiplex cytokine detection kit (Millipore) was used to detect the following cytokines and chemokines: IFN-γ, IL-1β, IL-4, IL-6, IL-17, TNF-α, IL-10, IL-2, IL-13, MCP-1, MIP-1α and RANTES simultaneously as described by the manufacturer’s protocol. The signals were analyzed on a Luminex 200 (Bio-Rad), and the data were analyzed using Bio-Plex Management software (Bio-Rad).

**Statistical Analyses**

Statistical analysis was performed with Prism Graph Pad software using a 2-tailed Student t test or the significance between the controls and treatment groups was determined by 1-way analysis of variance (ANOVA). Differences with a p value <0.05 were considered significant and are denoted by an asterisk.

**Results**

**RTL220 Inhibits IRBP1177–1191-Induced EAU**

The studies were designed to determine the efficacy of a new RTL220 bearing the retina-specific antigenic IRBP1177–1191 peptide in the treatment of acute and, in particular, recurrent EAU. The therapeutic effects of RTL220 were evaluated in Lewis rats with EAU induced with IRBP1177–1191. Inflammatory infiltrates are found in both anterior and posterior parts of the eye. Therefore, we were able to identify clinical signs using biomicroscopy. Inflammatory changes in the posterior part and retinal damage were assessed by histology. The rats uniformly developed eye inflammation, which occurred on days 9–10 following the antigen injection. Typically, the inflammation diminished after 25 days.

Two treatment regimens consisting of 5 doses of 300 μg RTL220 administered every 2 days were tested in the following groups: (1) treatment started on day 1, concurrent with immunization of the IRBP antigen, and (2) treatment started at the onset of clinical inflammatory signs as assessed by biomicroscopy. The controls included: untreated rats, treated with empty RTL101 without peptide and RTL203 carrying the irrelevant cardiac myosin peptide CM-2. Figure 2 shows collective results from 1 typical experiment. RTL220 effectively reduced the clinical signs of EAU (p < 0.0004 for all groups, n = 6; 1-way ANOVA). In the treatment regimens, we observed delayed onset by 3–4 days and a considerable ame-
Inhibition of IRBP1177–1191-induced EAU with RTL220 in Lewis rats. Time course of EAU in RTL-treated rats with 5 doses of 300 µg of RTL220, RTL101 or RTL203 subcutaneously starting on the day of IRBP immunization (a) or at the onset of clinical signs (b). The data present an average of 10 eyes for 5 rats and SD from a representative experiment. The severity of inflammation was determined as described in the methods. The significance between the controls and treatment groups was determined by 1-way ANOVA, overall p = 0.0004. c Overall severity of EAU. Bars indicate CDIs calculated for each group (n = 5) in this experiment.

d Level of cytokines in individual eyes from unimmunized rats (control), immunized with IRBP peptide (untreated) and not treated, and RTL220-treated rats (RTL220) from rats treated at the onset of clinical signs (n = 4 eyes). Beadlyte multiplex cytokine detection kit (Millipore) was used to measure cytokines as described in the methods. The significance between the control and treatment group was determined using Student's t test (* p < 0.05) and is denoted by an asterisk.

e Summary of histology scores for each group. Bars represent average histology scores for anterior and posterior segments of all eyes (n = 10) in experimental groups. f Representative histology of the retina from different treatment groups as shown in a and b. Arrows indicate inflammatory cells and retinal waves.
loration of disease severity and duration compared to the control groups (fig. 2a, b). The rats that were treated starting at disease onset showed complete suppression of EAU after 5 days of mild uveitis (fig. 2b). Nonspecific RTL203 and empty RTL101 did not alter the course of EAU, demonstrating the specificity of the RTL treatment. The histological results from acute EAU agreed with clinical assessments. The control eyes (untreated, RTL203 and ‘empty’ RTL101) showed the presence of inflammatory cells in the anterior chamber, vitreous and iris and infiltrating the retina. Positive clinical scores in some treated EAU rats were consistent with the presence of some lingering inflammatory cells in the anterior portion of the eye. The posterior segment was mostly free of infiltrating cells (day 18). RTL220 treatment of active EAU, administered at disease onset, essentially eliminated the infiltration of inflammatory cells into the eye and fully preserved the retina. RTL220 treatment that started on day 1 with the disease induction was less efficient in preventing eye inflammation than treatment starting at onset (fig. 2e, f) and markedly lowered the severity of inflammation in the retina to 30%, compared to untreated control rats. Although diminished mild anterior inflammation was detected in the anterior chamber during the course of disease, the posterior segment was clear of inflammatory cells.

To compare the overall severity of disease in the course of the illness we calculated the cumulative disease index (CDI) and the results are presented for both eyes of each rat in experimental and control groups in figure 2a and b (fig. 2c). The CDIs indicated that RTL220 significantly inhibited EAU in rats treated at the time of immunization (p < 0.05) as well as at the onset of clinical EAU (p < 0.01) compared to untreated controls. Pro- and anti-inflammatory cytokines were examined in the eyes of rats treated with RTL220 at the onset of clinical EAU and IBRP-injected untreated rat eyes at the end of the experiment on day 18 after immunization. Figure 2d shows the significant reduction in the proinflammatory cytokine levels of IL-2, IL-1β and IL-17 (p < 0.05 denoted by an asterisk) in RTL-treated eyes as compared to untreated rats. TNF-α, INF-γ (not shown) and IL-6 change was reduced but did not reach statistical significance. Anti-inflammatory IL-13 was increased, IL-10 was slightly increased, and IL-4 was unchanged in treated and untreated eyes.

**RTL220 Suppresses Recurrence of IRBP1177–1191-Induced EAU**

We wished to examine whether RTL220 can also affect the course of recurrence of EAU. To study these effects, we used the Lewis rat model of recurrent uveitis induced by adoptive transfer of 5 × 10⁶ IRBP1177–1191-specific T cells. R-EAU is characterized by early disease onset on day 3 after T cell transfer in all rats and by 3 relapses over 40 days that varied in individual rats. We compared untreated rats to rats treated with 5 doses of 300 µg injected every 2 days, followed by 300 µg of RTL220 once a week for the duration of the experiment. Regimen 1 started at the onset of the first attack of R-EAU and regimen 2 at the second attack of EAU. Because the clinical course varied, which is consistent with the previously published data [23], we show representative results for each eye of 2 out of 6 rats per group from both treatment regimens and untreated controls in figure 3. The regimen starting at the first onset of clinical signs drastically reduced the severity of EAU and stopped the recurrences of EAU in all treated rats.
Treatment of Uveitis with RTL

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Day 39 retinal histology

Cytokine level (pg/ml)

- IL-1β
- TNF-α
- IL-6
- IL-12
- IFN-γ
- IL-10
- IL-4

Cytokines

- Untreated
- 1st onset
- 2nd onset

Cytokine level (pg/ml)

- Eye

normal retina
untreated control

RTL220 at 1st onset
RTL220 at 2nd onset
(n = 6) – a score of 4 at the peak in control rats (fig.3a, b) compared to a score of 0.5–1 at the peak in RTL220-treated rats (fig. 3c, d). When RTL220 was administrated at the second onset of inflammation, (fig. 3e, f; regimen 2) more than 50% reduction in severity of the second bout of inflammation (score 2–3) compared to control rats (score 1–2) was observed. The initial 5-dose treatment without weekly follow-up treatments had a diminished therapeutic effect with relapses (data not shown).

The CDI shows a significant difference between RTL220-treated (2.625 ± 2.5) and control groups (32.9 ± 2.62) for the treatment started at the first attack of inflammation (p < 0.0001, n = 6). For the second treatment regimen, the results also showed marked suppression of R-EAU, but the CDI did not reach statistical significance. The histology of R-EAU rats performed on eyes collected at the end of the experiments (39 days after immunization) also confirmed the clinical results (fig. 3h). The retinal morphology of RTL220-treated rats showed a fully preserved morphology compared to that of the untreated rats, in which we found retinal weaving and local loss of photoreceptor cells, as well as signs of neovascularization in some rat retinas.

To further analyze the therapeutic effects of RTL220 we examined inflammatory cytokines and chemokines present in R-EAU in the eye and periphery, then compared it to the untreated rats. Splenocytes obtained from individual spleens of the different treatment groups were stimulated with the immunizing peptides for 48 h and supernatants were evaluated for cytokine secretion. Our results showed that both regimes of RTL220 treatments of EAU led to a significant reduction in proinflammatory cytokines secreted by spleen and eye cells (fig. 3g). The reduction in IL-17, IL-2 and IFN-γ was observed for splenic secretion, whereas only IL-2 and IL-1β were decreased in treated eyes. The systemic IL-10 levels were increased compared to the untreated controls, especially in the rats treated at the second onset. Other cytokines in the eyes were similar to the levels of those of the untreated controls, possibly due to collection of the samples at the end of the experiment.

The chemokines CCL2, CCL3 and CCL5 were upregulated at the onset of clinical symptoms, but the RTL220 treatment showed a significant suppression of the secretion of these chemokines in the periphery (fig. 4). The levels of CCL2 and CCL5 were reduced in the periphery and in the eye, which agrees with the lack of inflammatory cells in the eye. The level of CCL2 was suppressed in the spleen of treated animals, but the levels of CCL3 in the eye were measured very low, below 10 pg/ml.

**Discussion**

The ultimate goal of new immunotherapies is to inhibit the ongoing disease process by modulating effector mechanisms. Our aim was to suppress ongoing inflammation with RTL220 and also protect the neuronal retina from retinal damage due to inflammation. Indeed, our studies showed that RTL220 suppressed clinical and histological signs of acute and recurrent IRBP-peptide-induced EAU. The prophylactic outcome was less successful in complete suppression, although the treatment prior to onset showed considerable protection of the retina from EAU. Importantly, treatment with RTL220 effectively abolished clinical and histological signs of relapses not only when delivered with the first onset of clinical disease but with later attacks of inflammations. The overall effect of this immunosuppression was a marked reduction in infiltrating cells in the eyes of the rats treated with RTL220 and decreased production of proinflammatory cytokines, such as IL-17 and IL-2, the major mediators of eye inflammation, and increased in anti-inflammatory IL-10 cytokine. The lack of inflammation in the eye may be due to an altered proinflammatory cytokine and chemokine expression in the periphery, thus preventing cell recruitment to the eye.

Visual loss is more common in posterior than anterior uveitis because of irreversible damage to the retina, which may be a consequence of the influx of the inflammatory cells and secretion of proinflammatory cytokines. Chronic uveitis involves ongoing priming and recruitment of new T cells into the effector pool and thus requires long-term interventional medical therapy. These studies offer a new treatment for recurrent uveitis using RTL immunotherapy targeting pathogenic T cells that in effect change their properties from pro- to anti-inflammatory.

RTL treatment acts in an antigen-specific manner, since RTLs without immunizing peptide (RTL101) or nonspecific peptide (RTL203) had no effect on the suppression of EAU. However, the antigen specificity of RTL raises a question as to whether treatment with single RTLs would be successful in uveitis patients where target antigens may be unknown. This is an ongoing project in our laboratory and future studies of treating animals immunized with the multiantigenic retinal homogenate will examine the effectiveness of such an approach. However, recently published studies on RTL treatment of EAE mice injected with spinal cord homogenate or combinations of 2 different peptides to induce disease showed that treatment with single RTLs could reverse EAE as long as targeted T cells were present in the periphery [26]. The
authors believe that treatment with single RTLs can induce a cytokine switch in cognate T cells that inhibits both, the target and bystander T cells, providing new evidence for the potential applicability of RTL therapy in multiple sclerosis. Such a scenario could be possible in uveitis.

RTL treatment is a platform technology that we hope could be applicable for the treatment of many autoimmune diseases mediated by T cells. Thus far, RTLs have been successfully used in mouse experimental models of multiple sclerosis and rheumatoid arthritis, and these studies have revealed some details of an underlying mechanism of the therapeutic activity of RTLs. RTL engagement with TCR in the absence of CD4 binding was found to result in rapid TCR phosphorylation, calcium mobilization and reduced extracellular, signal-related kinase activity, as well as in a deviation from a Th1 to a Th0 cell phenotype based on cytokine production [27, 28]. It has been suggested that elevated levels of IL10 induced in Th1 cells by RTLs have important regulatory implications for autoimmunity because IL-10 is known for its anti-inflammatory effects on Th1 cell and macrophage activation in EAE [18, 29]. Besides the anti-inflammatory effect of IL-10 in EAE, it has been reported that intracocular expression of IL-10 by intravitreal injection of AAV2/2-tetON-vIL-10 protected from S-Ag-induced EAU in Lewis rats with vIL-10 expressed over a long period of time [30]. Our experiments showed an increased secretion of IL-10 by splenocytes from RTL220-treated rats, suggesting that changes in cytokine expression may suppress the recruitment of inflammatory cells to the eye.

In general, RTL therapies in mice and rats inhibited the systemic production of pathogenic cytokines by the targeted specific T cells but also inhibited downstream, local recruitment and retention of inflammatory cells in the CNS as well as in the eye. In EAE studies using the C57BL/6 model, in which IA^b^-restricted T cells specific for myelin oligodendrocyte glycoprotein peptide (MOG-
35–45) are implicated in disease pathology, RTL551 (carrying covalently tethered MOG35–55 peptide) treatment of mice strongly and selectively reduced the secretion of IL-17 and TNF-α, the latter of which was associated with the downregulation of chemokines and their receptors, and the inhibition of vascular cell adhesion molecule-1 and intercellular adhesion molecule-1 expression on endothelial cells [31]. IL-17 was also found to play a role in the pathogenesis of EAU, showing that systemic and local IL-17 response was correlated with disease severity in EAU mice [32]. Targeting IL-17, even late in the disease process, ameliorated pathology, indicating an effector role for this cytokine in the pathogenesis of EAU [33]. Our results showed a considerably reduced systemic and local secretion of IL-17 after RTL220 treatment of acute and recurrent disease. Moreover, CCL2, CCL3 and CCL5 were suppressed in the eyes with EAU. It is widely accepted that synthesis and secretion of inflammatory chemokines play an important part in the pathogenesis of ocular inflammation [34, 35]. Both CCL2 and CCL5 are associated with infiltrating inflammatory cells and are potent chemoattractants for T lymphocytes and macrophages, which are related to infiltrating cells observed in the posterior segment of eyes with EAU [34–36]. Thus, decreased chemokine levels by RTL220 may reflect a reduced number of recruited T lymphocytes and macrophages invading the retina during uveitis.

In summary, we demonstrated for the first time a successful treatment of acute and recurrent EAU. RTL220 effectively suppressed the recurrence of inflammation and reversed clinical and histological EAU by altering proinflammatory cytokine and chemokine expression, which possibly stopped the recruitment of pathogenic T cells into the eye. Such efficacy of RTLs in the therapy of anterior [19] and posterior uveitis (this study) strongly supports the possibility of clinical application of this novel class of peptide/MHC II drugs for patients with presumed autoimmune uveitis.

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References


Erratum

To the article by Adamus G, Karren LJ, Mooney J and Burrows GG, entitled ‘A promising therapeutic approach for treatment of posterior uveitis: recombinant T cell receptor ligand protects Lewis rats from acute and recurrent experimental autoimmune uveitis’ [Ophthalmic Res 2010;44:24–33, DOI: 10.1159/000281815], the following Disclosure Statement is to be added:

OHSU and Dr. Burrows have a significant financial interest in Artielle Immunotherapeutics Inc., a company that may have a commercial interest in the results of this research and technology. This potential conflict of interest has been reviewed and managed by OHSU.