Infliximab recovers central cone dysfunction with normal fundus in a patient with ulcerative colitis

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ABSTRACT

Purpose: To report the efficacy of anti-tumor necrosis factor α (anti-TNFα) on autoimmune-mediated macular cone dysfunction in a Japanese woman with ulcerative colitis (UC).

Observations: A 41-year-old woman presented with bilateral visual acuity loss and photophobia. She suffered from UC, and had been treated with prednisolone and 5-aminosalicylate since age 37. Although fundus photographs and optic coherence tomography images were unremarkable, electroretinograms (ERGs) were abnormal. A full-field electroretinogram (full-field ERG) revealed mildly decreased cone responses and oscillatory potential responses bilaterally. Importantly, focal-macular ERG (fmERG) and a multifocal electroretinogram (mfERG) revealed severe macular cone dysfunction in both eyes. Infliximab, a chimeric monoclonal anti-TNFα antibody, was administered to treat recurrent abdominal symptoms and continued at 8-week intervals. Almost 6 months after infliximab therapy, the mfERG response (especially in the central retina), the fmERG response, and visual acuity recovered bilaterally. Abdominal symptoms also improved after infliximab therapy.

Conclusions and importance: Bilateral cone dysfunction with normal fundus were observed in a UC patient, resulting in loss of visual acuity and photophobia. This retinopathy may have been caused by an autoimmune mechanism, such as an autoimmune retinopathy or acute zonal occult outer retinopathy, which is most identifiable by ERG changes. This is the first report demonstrating the efficacy of infliximab in autoimmune retinal dysfunction.

1. Introduction

Common ocular pathologies of inflammatory bowel diseases (IBDs) such as ulcerative colitis (UC), Crohn’s disease, and Behcet’s disease include bilateral uveitis, scleritis, and optic neuropathy. Ocular inflammation is usually controllable with corticosteroids. On the other hand, systemic IBD symptoms are usually treated with oral 5-aminosalicylic acid, prednisolone, azathioprine, and/or intravenous administration of anti-tumor necrosis factor α (anti-TNFα) monoclonal antibodies.

Sometimes, patients with autoimmune retinopathy (AIR) have a history of systemic autoimmune diseases, including IBDs. AIR causes various pathologies, including loss of visual acuity, photophobia, photopsia, and night blindness, despite normal fundus findings in most cases. However, no effective therapeutics have been identified for AIR.

Here, we report a rare case of cone dysfunction in a patient with UC that presented as visual acuity loss and photophobia, despite unremarkable fundus imaging and OCT images, and no evidence of ocular inflammation. Steroid treatment did not control the retinopathy, but infliximab, a chimeric anti-TNFα monoclonal antibody, improved visual acuity and cone function.

2. Case report

A Japanese woman with UC developed bilateral loss of visual acuity and photophobia beginning at 41 years-of-age. The patient began treatment for UC (prednisolone and 5-aminosalicylate) at 37 years-of-age, but continued to suffer from recurrent abdominal symptoms. The patient also had a history of bronchial asthma, depression, and a benign adrenal tumor.

Initially, the patient was assessed at our clinic when she was 42 years-of-age. Upon initial assessment, best-corrected visual acuity (BCVA) was 20/70 (OD) and 20/50 (OS), and intraocular pressure was 19 mmHg (OS) and 14 mmHg (OD). Neither slit-lamp examination nor funduscopy revealed abnormalities associated with inflammation or uveitis in either eye. Funduscopic images (Fig. 1A), fundus

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autofluorescence (Fig. 1B), and fluorescein angiography (Fig. 1C) did not reveal abnormalities in either eye. Optical coherence tomography (OCT) images of both eyes were unremarkable (Fig. 2A). Goldmann perimetry and Humphrey visual field analysis only revealed slightly extended Mariotte blind spots in both eyes (Fig. 1D).

Electrophysiological examinations were performed according to International Society for Clinical Electrophysiology of Vision (ISCEV) standards. A full-field ERG revealed normal responses in dark-adapted 0.01 cd s m\(^{-2}\) (DA 0.01), but slight bilateral reductions in oscillatory potential (OP) responses were observed on bright flash ERG (DA 10.0) (Fig. 3). However, light-adapted single cone-flash response (LA 3.0) with 8Hz stimulus interval revealed a bilateral decrease in both a- and b-wave responses, and LA 3.0 30 Hz flicker ERG revealed a bilateral decrease in amplitude and delayed implicit time (Fig. 3A, B and E). Although cone responses in the full-field ERG were mildly affected, a multifocal electroretinogram (mfERG) revealed widely reduced amplitude in both eyes (Fig. 4A). The foveal response in the right eye was lower than that in the left eye, corresponding with visual acuity.

During the observation period, the patient’s visual acuity and UC symptoms (abdominal pain and diarrhea) gradually worsened. The physician then initiated infliximab therapy (300 mg at week 0, 2, 4, and 6, and then every 8 weeks) to treat recurrent abdominal symptoms. On the day of the first infliximab treatment the patient noticed an immediate change in her subjective vision. The patient’s visual acuity improved gradually as infliximab treatment progressed (Fig. 5B). At 6 months after infliximab treatment, BCVA in the left eye recovered completely (to 20/20) and to 20/22 in the right eye (Fig. 5B). Then, BCVA in the right eye reached 20/20 at 9 months after starting infliximab treatment (Fig. 5B). Good visual acuity in both eyes was preserved during second half of the follow-up period (6–12 months). OCT images revealed no abnormalities and did not differ from images taken before infliximab therapy (Fig. 2B).

Three months after infliximab administration, fmERG still showed continued a reduction of a-wave, b-wave, and OP responses (Fig. 6). Remarkably, fmERG revealed recovery of macular function in both eyes 6 months after infliximab therapy (Fig. 6). Additionally, mfERG responses improved significantly in both eyes after infliximab therapy. The mfERG revealed that central retinal response in the right eye reached the normal range at 9 months after infliximab therapy, and that in the left eye reached the normal range at 6 months post-therapy (Figs. 4B, 5A and 5C). On the other hand, pericentral retinal responses in the mfERG took longer to recover than central retinal responses; indeed, they tended better, but had not recovered completely 1 year after infliximab therapy (Figs. 4B, 5A and 5C–F). On the other hand, one year after initiation of infliximab therapy, the amplitude in the 30 Hz flicker full-field ERG also slightly improved in both eyes, but OP

Fig. 1. Unremarkable fundus ophthalmoscopy images and visual field tests. (A) Fundus ophthalmoscopy, (B) fundus autofluorescence, and (C) fluorescein angiography images demonstrating no abnormalities in either eye. (D) Goldmann perimetry was normal in both eyes (except for extended Mariotte blind spots). R, right eye; L, left eye.
responses (DA 10.0) and cone responses remained modestly decreased (Fig. 3).

Alongside improved visual function, infliximab therapy also alleviated the patient’s abdominal symptoms. Infliximab (300 mg) was continued every 8 weeks for maintaining the visual function and systemic status. During follow-up after visual acuity reaching 20/20 in both eyes (6–12 months), mFERG responses were still unstable (Fig. 5B and C). Interestingly, approximately every 8th week after infliximab administration, the patient complained of worsening abdominal and visual symptoms, suggesting loss of the infliximab effect. Additionally, infliximab re-administration recovered again the subjective visual function, but it took 1–2 weeks after infliximab administration. Indeed, although visual acuity was preserved 1 year after starting infliximab therapy, the area 1 amplitude of the bilateral mFERG response at 12 months after infliximab treatment was lower than previous follow-up points (Fig. 5A and C). Continued infliximab therapy was required to maintain improvements in visual and systemic symptoms.

3. Discussion

Although the patient had UC and loss of visual acuity, there were no clinically detectable inflammatory pathologies, and fundoscopy was normal in both eyes. Electrophysiological examinations revealed cone dysfunction, especially in the macula, which was the likely cause of visual acuity loss and photophobia. These findings were suggestive of an autoimmune retinal disorder such as acute zonal occult outer retinopathy (AZOOR) or AIR. Interestingly, the extent of visual acuity loss was associated with the severity of UC; anti-TNFα therapy alleviated both visual function and systemic symptoms in this case. This suggests that factors involved in the systemic pathology of UC could also drive associated retinal disorders.

Full-field ERG revealed mild cone dysfunction. By contrast, both mFERG and fFERG responses were reduced significantly. Moreover, the improvement after infliximab therapy was clearly observed on mFERG and fFERG. These results suggest that retinal damage was localized to the macula. The time to improved visual acuity and ERG responses suggests that more than 6 months were required to recover full visual function.

AZOOR and AIR cause various pathologies, including visual acuity loss, photophobia, photopsia, and night blindness, despite normal fundus. Often, AIR patients have a history of autoimmune disease. In the present case, the patient had UC and suffered bilateral loss of visual acuity and photophobia, despite the absence of fundus abnormalities. ERGs indicated cone dysfunction with normal rod function. These clinical features suggest AIR, but in the present case immunoblot analysis had not been performed yet.

The findings in this case, including normal fundus appearance, loss of visual acuity, and cone dysfunction are consistent with AZOOR.
However, the patient did not have photopsia, and visual field defects corresponded to the depressed response in the mfERG. Further OCT abnormalities in the photoreceptor layer (outer retinal line), which are characteristic of AZOOR, were not present. These findings did not match the typical phenotype of AZOOR complex, suggesting that AZOOR could be excluded in this case.

Full-field ERG revealed decreased OP responses in both eyes, which is sometimes observed in uveitis. Previously, Takeuchi et al. reported that disappearance or disruption of the outer retinal line in patients with uveitis associated with Behcet’s disease was detectable in OCT, and that this was alleviated by infliximab therapy. That report suggests the efficacy of infliximab for suppressing retinal inflammation. However, our case did not present with clinically detectable inflammation in either eye, and the OCT findings were unremarkable.

Interestingly, the severity of systemic and ocular symptoms was linked. These findings suggest that the pathology of UC could be associated with retinal disorders. IBD increases levels of TNF-α, interleukin (IL)-1β, and IL-6. TNF-α is a classic proinflammatory cytokine produced by antigen-presenting cells and macrophages; the cytokine promotes inflammation (in part) by increasing production of IL-1β and IL-6.11,12
Moreover, Detrick et al. reported significant elevation of IL-6 levels in sera from patients with AIR; indeed, IL-6 is currently being investigated as a novel therapeutic target for AIR. These findings suggest that the therapeutic effect of anti-TNFα therapy in AIR may be mediated via suppression of IL-6. In the present case, UC may work indirectly to develop AIR as a background factor, including the presence of TNFα, IL-6, and other cytokines and chemokines.

Although the patient recovered bilateral visual acuity and central cone responses, full-field cone and OP responses and pericentral retinal responses on mERG remained slightly dysfunctional after treatment. The AIR in the present case could have complicated the pathology, not only by increasing TNFα levels but also through various additional factors such as anti-retinal antibodies and other proinflammatory mediators.

Fig. 4. Improvement of multifocal ERG during infliximab therapy. A multifocal electroretinogram (mERG) was recorded using a commercial mERG system (LE4100; TOMEY, Nagoya, Japan). mERG responses are shown (A) before infliximab treatment, and (B) 9 months after infliximab therapy in the right eye, and 6 months after infliximab therapy in the left eye.
At 1 year after infliximab therapy, visual acuity was good, but mfERG responses had deteriorated, despite ongoing maintenance infliximab treatment. This phenomenon could be caused by the timing of mfERG recording. During the follow-up after visual acuity improvement (6–12 months), subjective visual symptoms worsened before and after infliximab re-administration. It strongly suggests infliximab effects on this retinopathy, while this retinopathy have not been completely improved. On the other hand, mfERG responses of each areas tended to improve gradually during one-year infliximab treatment (Fig. 5C–F). Although at 9 months, P1 amplitude in area 1 of mfERG in the left eye was seemingly decreased, but it might be caused by fixation shift (Fig. 5A and C).

Continuation of infliximab treatment can have a better effect on retinal function. The mfERG results seemed to correspond with the patient’s visual symptoms. MfERG or fmERG could be more sensitive for evaluating retinal function. These findings suggest that in this case of AIR associated with UC, regular infliximab therapy is needed to maintain improvements in visual function. ERGs are key examinations for diagnosis of the autoimmune retinal disorder, as fundus images are typically normal in this type of retinopathy; in most cases, no clinically visible signs of ocular inflammation are present.

Fig. 5. Improved multifocal ERG response and visual acuity during infliximab therapy. (A) Averaged multifocal electroretinogram (mfERG) responses in areas 1–4 during follow-up. (B) One-year follow-up of the minimum angle of resolution logarithm (LogMAR) visual acuity. The dotted line reveals 0 in LogMAR visual acuity (decimal acuity 1.0). One-year follow-up of averaged mfERG P1 amplitude in (C) area 1, (D) area 2, (E) area 3, and (F) area 4. (C–F) The gray area surrounded by a dotted line in each graph denotes a normative range of the mean of normal control (44.0 ± 2.6 years old). Age matched (40–49 years old) six female was recruited for normal control and recorded by mfERG with the same protocol in our institute. The gray area showed the each are of the mean of P1 normative range with standard deviation: area 1, 54.3 ± 10.9 nV/deg²; area 2, 24.4 ± 4.6 nV/deg²; area 3, 19.7 ± 4.5 nV/deg²; area 4, 15.5 ± 3.9 nV/deg².
4. Conclusion

The present study documents a case of UC with macular cone dysfunction with normal fundus and absence of uveitis, which is suggestive of autoimmune retinal diseases including AZOOR or AIR. This is the first report to identify the efficacy of the chimeric monoclonal anti-TNFα antibody infliximab for the autoimmune retinal dysfunction. Thus, infliximab may be a good treatment for autoimmune retinal dysfunction.

Fig. 6. Improved focal-macular ERG responses during infliximab therapy. A focal-macular ERG (fmERG) was recorded using an ER-80 apparatus (Kowa, Tokyo, Japan) at 3 months and 6 months after infliximab therapy. The luminance of white stimulus light and background illumination were 30 and 1.5 cd/m², respectively. A background field with a 45° visual angle was projected onto the eye from the fundus camera. Focal ERGs were elicited by 5 Hz rectangular stimuli (100 ms light on and 100 ms light off). The 15° circular stimulus was carefully and continuously centered on the fovea, as observed on an infrared camera monitor. Three-hundred ERGs were averaged by a signal processor (PuREC, Mayo, Aichi, Japan). Focal ERGs were digitized at 10 kHz, with a band-pass filter of 5–500 Hz for the a-wave and b-wave. The fmERG responses at 6 months after infliximab therapy were better than those at 3 months after infliximab therapy.
diseases.

Patient consent

Informed consent was obtained from the patient. This report does not contain any personal information that could lead to the patient’s identification. This study was approved by the Ethics Committee of Juntendo University (JHS-19-017), and was conducted in accordance with the Tenets of the Declaration of Helsinki.

Author contributions

T.H. designed all of the experiments. M.C.-M., T.H., and M.Y. recruited and followed up the patient. M.C.-M., T.H., M.Y., and A.M. wrote the manuscript.

Declaration of competing interest

The authors declare no conflicts of interest.

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