Clinical characteristics and surgical treatment of idiopathic uveal effusion syndrome

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Abstract

● AIM: To investigate the clinical characteristics of idiopathic uveal effusion syndrome (IUES) and to identify effective surgical modalities for its treatment.

● METHODS: This retrospective analysis included clinical data of 33 eyes from 26 patients with IUES at Beijing Tongren Hospital. Records of eye examinations, ocular ultrasound, ocular ultrasound biomicroscopy (UBM), and follow-up surgical treatment were reviewed and analyzed.

● RESULTS: Of 26 patients, 17 (65.4%) were male and 9 (34.6%) were female. The average age of disease onset was 46.8y (range: 22-64y). Seven patients (26.9%) showed retinal detachment in both eyes at presentation. B-ultrasound showed the presence of retinal detachment in one eye or both eyes. All patients had binocular ciliary leakage and detachment. Eyes with retinal detachment underwent four-quadrant partial-thickness sclerectomy and sclerostomy. Subretinal fluid resolution was achieved within 6mo. Recurrence was observed in three eyes and was resolved with re-operation.

● CONCLUSION: Ophthalmic ultrasound and UBM, among others, can be helpful in the diagnosis of IUES. Sclerectomy and sclerostomy are surgical modalities that can successfully treat the disease. Some patients may experience recurrence after surgery; reoperation remains safe and effective for them. Long-term follow-up is essential in such settings.

● KEYWORDS: idiopathic uveal effusion syndrome; sclerectomy; sclerostomy; recurrence; retinal detachment

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INTRODUCTION

Uveal effusion refers to the leakage of fluid from the choriocapillaris, and may cause exudative detachment of the choroid and retina without retinal breaks[1-4]. Uveal effusion may be idiopathic or secondary to choroidal tumors, extensive chorioretinitis, and surgery, among others[2,5-7]. Macular involvement in exudative retinal detachment leads to significant visual acuity impairment[3].

The term “uveal effusion syndrome” (UES) was first introduced by Schepens and Brockhurst[2] in 1963; the disease is diagnosed by exclusion when the other causes of uveal effusion have been ruled out[8]. UES is generally divided into nanophthalmic UES and idiopathic UES (IUES)[8]. Nanophthalmic UES is easily diagnosed because of the characteristic clinical manifestation, short eyeball axis[4,9]; IUES is rarely reported in clinical practice and may be misdiagnosed as another exudative choroidal or retinal detachment disease[10-11]. The causes of IUES remain unknown, and the disease predominantly occurs in otherwise healthy middle-aged people[3,10]. Gass and Jallow[1,3] hypothesized that UES is primarily caused by congenital scleral abnormalities that act as barriers to the transscleral outflow of protein from the suprachoroidal space, and secondarily predisposes one to vortex vein obstruction. According to this theory, Gass[1] proposed a surgical management method for the treatment of UES using scleral sclerectomy and sclerostomy; subsequent studies further confirmed the effectiveness of these surgical modalities[4,12-13].

In this study, we reviewed the records of patients with an IUES diagnosis treated using surgery, in the past ten years, and analyzed the clinical features and surgical outcomes associated with the disease.

SUBJECTS AND METHODS

Ethical Approval This study was approved by Beijing Tongren Hospital Ethical Committee and informed consent was signed by all participants before surgery. The records of 26 patients (33 eyes) diagnosed with IUES, who had undergone surgery in our hospital in the past ten
years, were reviewed. All eyes underwent quadrant partial-thickness sclerectomy combined with partial full-thickness sclerostomy, and all patients were followed up for more than 6mo after surgery.

All patients underwent routine eye examinations, including binocular indirect ophthalmoscopy, ultrasound examination, optical coherence tomography (OCT), and ultrasound biomicroscopy (UBM), and some underwent fluorescein angiography and indocyanine green angiography. A small number of patients did not undergo angiography examinations due to allergies or poor general conditions. Patients were tested for syphilis, human immunodeficiency virus infection and hepatitis B and C infection and underwent routine blood and urine examinations. The exclusion criteria were as follows: the presence of a short eye axis (eye axis <20.5 mm); uveitis; panretinal photocoagulation; intraocular surgery; trauma, and a known cause of ciliochoroidal effusion. Patients with rhegmatogenous retinal detachment and choroidal detachment due to allergies or poor general conditions. Patients were tested for syphilis, human immunodeficiency virus infection and hepatitis B and C infection and underwent routine blood and urine examinations. The exclusion criteria were as follows: the presence of a short eye axis (eye axis <20.5 mm); uveitis; panretinal photocoagulation; intraocular surgery; trauma, and a known cause of ciliochoroidal effusion. Patients with rhegmatogenous retinal detachment and choroidal detachment as a complication were also excluded.

All patients underwent four-quadrant lamellar sclerectomy combined with partial full-thickness sclerostomy. The extent of each quadrant slice was 5×7 mm², about half to two-thirds of the scleral thickness. A 1×2 mm² sclerostomy was made in the center of each sclerectomy site. All the excised scleral pieces underwent pathological examination. The pieces obtained from one patient underwent electron microscopy.

RESULTS

The baseline characteristics of the 26 patients included in this study are summarized in Table 1. Seventeen (65.4%) of the 26 patients were men, and 9 (34.6%) were women. None of the patients had a family history of the disease. The average age of symptom onset was 46.8y (22-64y). Of the 26 patients, 7 (26.9%) had binocular retinal detachment at presentation. The mean duration from symptom onset to surgery was 8.5mo. Three eyes underwent reoperation for recurrent retinal detachment. The time to recurrence in these three eyes was 1, 5, and 9y, after surgery, respectively.

Ultrasound examination of the eyes showed that 24 patients had monocular or binocular retinal detachment (Figure 1A). OCT examination showed only macular exudative retinal detachment in two eyes of two patients (Figure 2A). UBM showed effusion of the ciliary body in the binocular eyes among all patients (Figures 3A-3D). Surgery was performed on the eyes showing retinal detachment, including two in which the subretinal fluid was only confined to the macular region, simulating idiopathic central serous chorioretinopathy. The axial length of the surgical eyeballs ranged from 20.9 to 23.7 mm (average 22.5 mm). The thickness of the sclera near the temporal scleral process was examined by UBM and was found to be an average of 0.76 mm. Fluorescence angiography showed mottled high and low fluorescence in the fundus, which was a leopard-like change.

The average preoperative intraocular pressure (IOP) was 14.8 (9-27) mm Hg, and the postoperative IOP (the first day after surgery) averaged 13.9 (7-24) mm Hg. Surgery was performed on all 33 eyes with retinal detachment. Retinal detachment was characterized by exudative retinal detachment and subretinal fluid shifts with changes in the eye or head position; no break or tear was seen on fundus examination, and no viable proliferations were observed in the vitreous (Figure 4A). Macular retinal detachment was predominantly characterized by neuroepithelial detachment, and more often accompanied by retinal retinoschisis.

During the surgery, we observed the spontaneous release of a pale-yellow suprachoroidal effusion fluid as the choroid was exposed. However, the choroidal effusion fluid was intentionally not drained out. The sclera was abnormally rigid and thick, and scleral texture disorder was noted; therefore, the razor blade knife had to be changed frequently. No complications occurred during the surgery. Pathological examination of the surgically excised scleral slice confirmed the presence of scleral collagen fiber arrangement disorder. Electron microscopy of the scleral slice also showed an irregular scleral fiber arrangement and marked deposits of matrix between the fiber bundles (Figure 5).

All patients were followed-up for at least 6mo after surgery. Retinal detachment was gradually resolved from 1wk to
5mo after surgery (Figures 2B and 4B). Pigmentation was observed in the retinal detachment zone. Choroidal detachment resolution was achieved earlier than retinal detachment resolution (Figure 1B), and ciliary body effusion was gradually resolved (Figures 3E-3H). The visual acuity of 30 eyes increased to varying degrees, and there was no significant

**Figure 2** OCT examination  A: Subretinal fluid confined to the macular region; B: Macular subretinal fluid was absorbed.

**Figure 3** UBM examination  UBM examination showing effusion of the ciliary body (A-D) and resolution of effusion (E-H). A, E: 3 o’clock; B, F: 6 o’clock; C, G: 9 o’clock; D, H: 12 o’clock.

**Figure 4** Retinal image  A: Visible retinal detachment before surgery; B: Retina was attached 3mo after surgery.

**Figure 5** Electron microscopy  A: Electron microscopy showing irregular scleral fiber arrangement; B: Marked deposits of matrix which may be proteoglycans in sclera.
change in the visual acuity in three eyes.

Three patients with recurrent retinal detachment underwent a second surgery—four-quadrant lamellar sclerectomy—combined with partial full-thickness sclerostomy. The scar tissue was removed during surgery and abnormal scleral thickness was observed again. Retinal detachment was resolved within 4mo after the second surgery.

**DISCUSSION**

In this study, we only reviewed the clinical data of hospitalized patients who underwent surgery in the past ten years. Some outpatients who were relieved by medical treatment were not included in the analysis. We found that ophthalmic ultrasound and UBM, among others, aided the diagnosis of IUES. Sclerectomy and sclerostomy were associated with treatment success. However, some patients experienced recurrence after surgery; they underwent successful reoperation.

UES was first reported in 1963 by Schepens and Brockhurst\[1\]; however, only a few cases of the disease have been reported to date\[10,14-15\]. The incidence of IUES is even lower in the existing literature. To the best of our knowledge, the current study, which included 33 eyes from 26 patients, presents the largest number of IUES cases to date. We reviewed the records of IUES patients with an axial surgical eyeball length of ≥20.5 mm, consistent with the criteria employed by Johnson and Gass\[14\].

Uyama et al\[11\] classified 19 eyes from 16 patients with IUES into three groups: type 1 included nanophthalmic eyes with an axial eyeball length shorter than 19.0 mm; type 2 included non-nanophthalmic eyes with a rigid and thick sclera and an axial length averaging 21.0 mm; type 3 included non-nanophthalmic eyes with a normal size and scleral thickness and an axial eyeball length ranging from 25.0 to 22.9 mm. They concluded that sclerectomy was effective in types 1 and 2 only; in type 3 eyes, this treatment modality was ineffective. However, in the type 3 group, only two eyes from two patients were treated.

Johnson and Gass\[14\] investigated 23 eyes of 20 patients with IUES who underwent quadrant partial-thickness sclerectomy with an axial length of ≥20.5 mm (average 23.1 mm), and found that subretinal and/or supraciliochoroidal fluid resolution occurred within 6mo in 19 eyes (83%) after one procedure and in 22 eyes (96%) after one or two procedures.

The causes of UES remain unknown. In 1983, Gass\[11\] hypothesized that the initial cause of UES is congenital scleral dysplasia, which acts as a barrier leading to the obstruction of the transscleral flow of the suprachoroidal protein and secondary venous occlusion. Retinal detachment and choroidal detachment are resolved by sclerectomy, indirectly confirming that scleral dysplasia causes UES\[12-14\].

In 2011, a study that used enhanced depth imaging spectral domain OCT found that the thickness of the choroid was significantly increased in patients with IUES\[15\], adding to the theory of Gass pertaining to the scleral barrier to transscleral flow in the disease. In our study, we observed abnormal scleral thickness values on surgery, as well as irregular collagen fiber arrangement and abnormal deposition, indicating the presence of proteoglycans-like in the scleral matrix, as observed on pathological and electro-microscopic examination, further confirming Gass’ hypothesis. Previous studies have shown the presence of a complex relationship between collagen fibrils and proteoglycans in the extracellular matrix, leading to transscleral fluid outflow obstruction through these scleral abnormalities\[16-17\]. The obstruction of transscleral outflow caused by abnormal sclera first leads to ciliochoroidal detachment; during this period, patients have no clinical symptoms\[10\]. Our study confirmed, using UBM, that all the participants had binocular leakage and ciliary body detachment.

At one time, decompression of the vortex vein was performed for UES treatment\[18-19\]. However, the surgery was associated with a high rate of complications, including bleeding\[18\]. Gass\[1\] proposed the use of sclerectomy for UES. Accordingly, in 1986, Vine\[12\] successfully treated a patient with UES using sclerectomy. Johnson and Gass\[14\] achieved success in the use of quadrantic lamellar sclerectomy and sclerostomy for IUES in 23 eyes of 20 patients. In 2013, a modified ultrasound-guided surgical technique for sclerostomy using a scleral punch for the localization of the site of maximal choroidal swelling was reported and was shown to be effective in UES management\[20\]. In 2017, a new technique for treating patients with uveal effusion syndrome by the fiberoptic-guided CO₂ laser was reported to reduce the risk of bleeding and providing better depth control\[21\].

In some patients, the degree of UES may show spontaneous improvement; however, the disease course is long and may result in serious visual acuity damage. In previous reports, including 20 patients with a natural disease course and a follow-up period of up to 8y, 9 patients showed spontaneous choroidal reattachment within 4mo to 6y, while the remaining 11 did not show improvements at the end of the observation period\[2-14\]. Therefore, timely surgery is required for IUES patients with retinal detachment.

Inconsistent with the findings of previous research\[10\], our study found that the sclera of IUES patients with an axis greater than 20.5 mm is still thickened, and the blade needs to be replaced repeatedly during surgery. Ultrasound examination could not accurately measure scleral thickness. While we attempted to measure the thickness of the sclera near the scleral process by UBM, we did not measure the normal scleral thickness using the same method. We found that the mean thickness of the sclera near the scleral process was 0.76 mm by UBM. A
Characteristics and Treatment of IUES

Previous study showed that UBM may be a more accurate and precise method for the measurement of scleral thickness than magnetic resonance imaging; the study found mean thickness values of 0.65±0.08 and 0.55±0.05 mm for eyes with UES and control eyes, respectively, by UBM[22].

Other clinical features observed in our patients were: a predominance of young and middle-aged patients; male predominance; chronic disease course; asymptomatic initial stage; normal intraocular pressure, slightly higher intraocular pressure in some cases; and absence of inflammatory reaction in the anterior segment and vitreous cavity, consistent with previous studies[14].

This study has some limitations. First, there was no control group. Some patients with exudative retinal detachment experience spontaneous recovery, but this may take a long time. Previous studies have shown that prolonged retinal detachment durations lead to gradual vision impairments[2]. Therefore, surgery is required in cases with exudative retinal detachment, especially when the macula is involved. Second, most of the patients lived far away from the hospital and were spread across the country; therefore, data on the patients’ final vision values could not be obtained because patients tended not to attend the follow-up if retinal reattachment was achieved. Our study did not statistically detect the relationship between patients’ visual acuity and preoperative visual acuity as well as the preoperative lesion time. Previous studies have shown that there is no statistical correlation between postoperative visual acuity and preoperative lesion time[14].

In conclusion, we found that four-quadrant scleral lamellar sclerectomy combined with partial full-thickness sclerotomy was safe and effective for patients with IUES. In patients with relapse, reoperation is still safe and effective. Among IUES patients, the axial axis is normal, but the thickness and composition of the sclera are abnormal. Further research is needed to determine the causes of scleral thickening and component changes; genetics or metabolism may play a role in the disease course.

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REFERENCES


