Surgically induced necrotizing scleritis after primary pterygium surgery with conjunctival autograft

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Background: Although pterygium excision with conjunctival autograft is a widely performed surgical procedure, surgically induced necrotizing scleritis (SINS) following such surgery is extremely rare.

Methods: A 68-year-old man underwent nasal pterygium excision with conjunctival autograft uneventfully. On postoperative day 17, the conjunctival graft was avascular, with epithelial defect. Although topical steroid and antibacterial treatments were continued, the graft and sclera melted, with the ischemic sclera showing gradual thinning. The thinning area spread to the adjoining cornea, and active inflammation with epithelial defect was observed adjacent to the site of thinning.

Results: Systemic and microbiological examination was noncontributory. The patient was suspected of having SINS, and administration of oral prednisolone was started. Although the necrotic area was reduced temporarily, medication was discontinued due to nausea, and the area of thinning increased. Conjunctival flap surgery was later performed, and the graft was well accepted.

Conclusions: SINS must be considered in the differential diagnosis of patients with scleritis following pterygium surgery, especially if radiation or mitomycin C has not been used.

Keywords: scleritis, pterygium, pterygium surgery, conjunctival autograft, SINS

Introduction
Surgically induced necrotizing scleritis (SINS), a local autoimmune reaction occurring near previous surgical wounds, has been reported after cataract surgery, trabeculectomy, retinal detachment, and strabismus surgery.1 We report SINS after primary pterygium excision with conjunctival autograft, an extremely rare occurrence, with only two previously reported cases.2,3

Case report
A 68-year-old man with no history of systemic or ocular disease was referred to Tokyo Dental College Ichikawa General Hospital for treatment of primary pterygium. Best-corrected visual acuity (BCVA) was 20/20 in both eyes. Slit-lamp examination revealed nasal pterygium in his right eye, and pterygium excision and conjunctival autograft transplantation without mitomycin C were performed uneventfully.

On postoperative day 1, slit-lamp examination revealed a well-adapted, epithelialized graft, with corneal epithelial defect at the site of pterygium excision. Topical treatment with 0.5% levofloxacin and 0.1% betamethasone five times daily was initiated. On postoperative day 17, the conjunctival graft was avascular, with epithelial defect;
the initial corneal epithelial defect had diminished with no symptoms. Although topical steroid and antibacterial agents were continued, the graft and sclera melted, with the ischemic sclera showing gradual thinning. On postoperative day 29, although the scleral bed was covered by keratinized conjunctival epithelium, active inflammation with epithelial defect was observed adjacent to the site of thinning (Figure 1). By postoperative day 36, the thinning area had spread to the adjoining cornea, and active inflammation persisted. Bacterial and fungal cultures of the lesion were negative. Tests for rheumatoid factor and antinuclear antibody were also negative. A diagnosis of SINS was suspected, and administration of oral prednisolone (20 mg per day) was commenced. Five days later, the conjunctival and scleral inflammation had decreased and vascularization was observed on the inferior scleral side. However, prednisolone was discontinued due to nausea, and the area of thinning increased in size (Figure 2A and B). Resection of necrotic tissue and lamellar keratoplasty were planned but not performed, as the patient ceased visiting the hospital. Conjunctival flap with Tenon’s capsule was later performed at another hospital, and the graft was well accepted.

**Discussion**

In this case, SINS occurred approximately 2 weeks after surgery, with ischemia of the conjunctival graft and underlying sclera, as previously reported. In earlier cases, the extent of the disease has been limited to the sclera. Here, however, scleral thinning and inflammation spread to the adjoining cornea. Scleral necrosis and melting can occur after pterygium surgery due to the use of adjunctive irradiation, mitomycin C, or excessive cauterization of the sclera. These etiologies should be ruled out before diagnosing SINS. In this case, no radiation or mitomycin C was used, and cauterization was kept to a minimum during surgery. In fact, hundreds of pterygium surgeries have been performed at our institute, in the same manner, with no other cases developing necrotizing scleritis. The spread of the disease to the cornea eliminated cauterization as a likely primary cause. In addition, reduced inflammation after systemic immunosuppression supported our diagnosis, indicating the utility of immunosuppressive treatment in identifying SINS.

Treatment of SINS after pterygium surgery has included immunosuppression with systemic steroids, cyclophosphamide, or tacrolimus, and surgical intervention, including resection of necrotic tissue, amniotic membrane transplantation, and scleral or corneal tissue patch grafts. In this case, systemic steroids had to be discontinued, and the administration of tacrolimus and further surgical procedures were considered. Conjunctival flap with Tenon’s capsule is an option, especially if obtaining a scleral or corneal graft is difficult. Resection of necrotic tissue and conjunctival flap with Tenon’s capsule were performed, and the graft was well accepted.
This case suggests that SINS should be considered as a possible diagnosis in patients with scleritis after pterygium surgery, especially if radiation or mitomycin C has not been used. Ischemia and melting in the conjunctival graft and underlying sclera indicate SINS, and a risk exists of it spreading to the adjoining cornea.

Disclosure
The authors report no conflicts of interest in this work.

References