Cleroderma or systemic sclerosis (SSc) is a chronic multi-system disorder predominantly affecting the skin, musculoskeletal, gastrointestinal, pulmonary and renal systems. Corneal involvement in SSc is rare. Filamentous keratitis, exposure keratitis (secondary to lid changes), peripheral ulcerative keratitis, pellucid marginal degeneration and increase in the central corneal thickness (CCT) have all been reported to be associated with SSc.

Keratoconus is a progressive non-inflammatory corneal ectasia. It is a bilateral and asymmetric corneal disease characterized by localized corneal thinning that leads to protrusion of the thinned cornea.
In this paper we describe a patient with SSc and keratoconus. To our knowledge, in the literature there is no other case report that describes presence of keratoconus in patients with SSc.

CASE REPORT

A 50 year-old female admitted to the hospital with the complaint of decreased vision, burning and grittiness in her eyes. The patient had been followed for seven years by her rheumatologist with the diagnosis of SSc. On presentation, her best corrected visual acuity was 8/10 in the right eye and 5/10 in the left eye. Manifest refractions were +0.50 (+1.50α98) diopter (D) in the right eye and -0.50 (-2.50α170) D in the left eye. Keratometry values were 49.47α93° D, 47.18α3° D in the right eye; 53.59α86° D, 50.15α176° D in the left eye. White to white cornea diameter was 11.79 mm in the right eye, 11.75 mm in the left eye. “Scissors” reflex was present in both eyes on retinoscopy. Slit lamp examination revealed normal conjunctival fornices, clear corneas with no evidence of inflammation, vascularization or ulceration. There wasn’t Vogt’s stria, Fleischer ring and apical scar. There was no history of eye surgery or inflammatory eye disease. Corneal sensation was intact bilaterally. Intraocular pressure measured with Goldmann applanation tonometry was 17 mmHg bilaterally. Minimal nuclear sclerosis of the lens was present in both eyes. Fundus examinations of both eyes did not reveal any pathology. CCT measured by an ultrasonic pachymeter was 455 micrometer (μm) in the right eye and 423 μm in the left eye. Schirmer tear test (after topical local anesthesia) was 2 mm / 5 minute bilaterally. Break-up time was 1 second bilaterally.

Her physical features of scleroderma included tightness of the skin, per-oral puckering and facial telangiectasia of her face as well as flexion contracture on her fingers (Figures 2a-b). The computerized visual field test and fundus angiography were within normal limits.

For her dry eye symptoms, patient was prescribed non-preserved topical lubricants. The patient was offered hard contact lenses but she did not accept to use lenses, so that glasses were prescribed.

DISCUSSION

Scleroderma or systemic sclerosis (SSc) is a chronic multi-system disorder predominantly affecting the skin, musculoskeletal, gastrointestinal, pulmonary and renal systems. Although the exact etiology is unknown, recent evidence suggest that immune activation plays a pivotal role in the pathogenesis. The most apparent clinical features of SSc are Raynaud’s phenomenon and dermal thickening with attachment of the skin to underlying tissues, which is manifested by taut skin, pursed lip, mask-like face and tapering of the digits.

The most common ocular findings of SSc include keratoconjunctivitis sicca, shallow conjunctival fornices, thickening and tightness of the eyelids and occlusive choroidal vascular disease. The most frequent ocular manifestation of SSc is keratoconjunctivitis sicca, which has been reported to be present in 37-79% of patients. Depending on its severity, keratoconjunctivitis sicca can be present with mucous strands in the precorneal tear film, superficial keratopathy, punctate keratopathy or filamentary keratitis. In the presented case, the tests revealed dry eye, but mucous filaments, superficial and filamentous keratitis were not present in the biomicroscopy.

Type I collagen is the major component in the cornea, comprising about 68% of the dry weight of the cornea. In addition, collagen types III, V, VI, XII, and XIV have all been detected in the corneal stroma. The cornea is particularly vulnerable to collagen vascular diseases due to its collagen composition and relation to the rich vascular supply of the conjunctiva and episclera. Peripheral corneal thinning is the result of obliterator microangiitis from deposition of immune complexes in the limbal vasculature, especially in the setting of scleral inflammation. Collagenases and proteases, released by infiltrating leukocytes and activated stromal keratocytes, degrade stromal collagen.
Corneal involvement in scleroderma is rare. Peripheral corneal thinning associated with scleroderma is more often inflammatory and ulcerative. Filamentous keratitis, exposure keratitis (secondary to lid changes) and peripheral ulcerative keratitis have all been reported to be associated with SSc.2–5,11 Extracellular matrix overproduction by fibroblasts is the hallmark of SSc. Activated fibroblasts overproduce extracellular matrix proteins, such as collagen type I, III, V, VI and VII, tenasin, proteoglycans, fibronectin, laminin and fibrillin-1. Ultrastructural studies showed collagen fibrils were increased in histopathology of SSc.12 Two studies found statistically significant increased CCT in patients with SSc compared to matched controls.5,11 However two studies showed no difference in CCT.
of patients with SSc compared to matched controls. They concluded that this phenomenon may be due to medical treatment of the scleroderma patients. So this issue remains unclear if the CCT of SSc patients are affected in the disease process.

Corneal ectasias, such as pellucid marginal degeneration (PMD) in a SSc case have been reported in the literature. PMD is a non-inflammatory peripheral corneal thinning and it is not associated with signs of inflammation, vascularization or infiltrates. Corneal topography typically shows flattening in the vertical meridian with marked steepening inferiorly below the site of thinning that extends into the infero oblique meridian. In contrast, the thinning in keratoconus occurs at the apex of conical shape, corneal protrusion and the steepening is the greatest at the apex of the cone and reduces concentrically towards the periphery, as with this case (Figure 1). It has been suggested that PMD and keratoconus may be different manifestations of the same etiological factor.

In 1953, Agatston reported a case describing retinopathy in a SSc patient. He claimed that scleroderma may also be associated with other eye disorders including keratoconus. However no association of scleroderma with keratoconus was reported in the literature yet. To the best of our knowledge, our case is the first report in the literature presenting keratoconus in a scleroderma patient.

In conclusion, keratoconus can be associated with scleroderma. Since scleroderma is usually associated with increased central corneal thickness, in this first report in the literature, we presented the rare association of scleroderma with keratoconus.

### References