Salzmann's Nodular Corneal Degeneration

62-year-old woman presents with 3 years of progressive decrease in vision

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Chief Complaint
A 62-year-old woman presents with a 3-year history of painless, progressive loss of vision in both eyes.

History of Present Illness
Over the last 3 years, there has been gradual, painless loss of vision in both eyes (right greater than left) at both distance and near. She has had 3 updates of her spectacle prescription that have not provided satisfactory vision. There is no diurnal variation in vision.

Past Ocular History
- Myopia, presbyopia, astigmatism
- Dry eye syndrome, with relief of symptoms with artificial tears
- No ocular surgeries

Medical History
- Hypertension

Medications
- No ocular medications. Systemic medications: metoprolol 50 mg twice daily

Family History
- No family history of glaucoma, macular degeneration, blindness or known ocular diseases.

Social History
- Does not use tobacco or drink alcohol.

Ocular Exam
Visual acuity (VA) with correction:
- Right eye (OD): 20/200, pinhole to 20/50-1
- Left eye (OS): 20/25+2, pinhole to 20/20

Manifest Refraction
- OD -2.00 + 5.00 x 20 VA 20/60
- OS -0.75 + 7.50 x 165 VA 20/25

Intraocular pressure (applanation): OD 16 mmHg, OS 18 mmHg

Pupils: Equal, round, 4 mm in dark, 2 mm in light, no relative afferent pupillary defect

Visual fields: Full to confrontation both eyes (OU)

Motility: Full OU

Slit Lamp Exam
- External: Normal OU
- Lids & lashes: Inspissated meibomian glands with thick waxy secretions and telangiectasia, normal lashes OU
• Conjunctiva & sclera: Clear and quiet
• Cornea:
  o OD: Greyish nodules between the corneal epithelium and Bowman’s layer in the paracentral superonasal region extending centrally into the visual axis; iron line at inferior border of the lesion
  o OS: Greyish nodules between the corneal epithelium and Bowman’s layer in the paracentral superior and superotemporal regions extending centrally into the visual axis; iron line at inferior border of the lesion. See Figure 1.
• Anterior chamber: Deep and quiet OU
• Iris: Normal architecture OU
• Lens:
  o OD: 2-3+ nuclear sclerosis
  o OS: 1-2+ nuclear sclerosis
• Vitreous: Normal OU

**Dilated Fundus Exam**

Disc: Normal, cup-to-disc ratio 0.3 OU
- Vessels: Normal OU
- Macula: Normal OU
- Periphery: Normal OU

**Figure 1a-b:** Slit lamp photos of the left eye before treatment

**Other Tests**

Topography was ordered to characterize the corneal nodules seen on slit lamp exam in the setting of significant astigmatism. Figure 2 shows the crab-claw like pattern of irregular astigmatism superiorly on each eye due to the superior Salzmann’s nodules. Placido images show irregular mires most noticeable superiorly on each eye corresponding to the Salzmann’s nodules.

**Diagnosis**

A diagnosis of Salzmann’s nodular degeneration was made on the basis of the characteristic clinical appearance and the clinical history of progressive increasing irregular astigmatism associated with

**Figure 2:** Topography before treatment.
worsening of best corrected visual acuity. She was also diagnosed with nuclear sclerotic cataracts on the right greater than left eye.

**Treatment and Clinical Course**

To improve visual acuity and to decrease the irregular astigmatism a superficial keratectomy was recommended to remove the Salzmann’s nodules.

**Course**

The patient underwent superficial keratectomy on her right eye. Postoperatively, there was complete clearing of the subepithelial opacities with excellent clarity of the underlying stroma. One month later, reliable keratometry and intraocular lens (IOL) calculations were obtained and an uneventful phacoemulsification and implantation of a posterior chamber IOL was performed (see Figure 3). Several months later, she underwent superficial keratectomy on her left eye with similar results, followed by cataract surgery on her left eye one month later. Postoperatively both eyes achieved best-corrected visual acuity of 20/20, with substantial reduction in the astigmatic error (Table 1; Figure 4). At the time of this writing, the vision and examination remains stable for 47 months with no recurrence of Salzmann’s nodular corneal degeneration. The mild dry eye continues to be well controlled with artificial tears.

**Table 1**

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<tr>
<th>Pre-operative Manifest Refraction</th>
<th>Post-operative Manifest Refraction</th>
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<tr>
<td>-2.00 + 5.00 x 20 VA 20/60 OD</td>
<td>-0.25 + 0.75 x 045 VA 20/20 OD</td>
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<tr>
<td>-0.75 + 7.50 x 165 VA 20/25 OS</td>
<td>-1.00 + 1.00 x 150 VA 20/20 OS</td>
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**Pathology**

Salzmann’s nodules are characterized by hyaline deposited anterior to a disrupted or even displaced Bowman’s layer, posterior to an atrophic corneal epithelium. Figure 5 for is an example of the typical histopathology of Salzmann’s nodules.
**Figure 5:** Salzmann’s nodule histopathology. A. Atrophic corneal epithelium overlies a nodular, hyalinized fibrous plaque between the epithelium and Bowman’s layer. A portion of Bowman’s layer is absent at the right side of the lesion. H&E, 50x. B and C. Periodic acid Schiff stain highlights the thickened, irregular epithelial basement membrane. 50x, 100x. (click any image for enlarged view)

**Discussion**

Salzmann’s nodular corneal degeneration was described by Katz in 1930 following a 1925 case series on the condition published by Maximilian Salzmann, although Salzmann concedes that a case published by Ernst Fuchs in 1901 appears to be the first published case of this corneal degeneration [1].

Salzmann’s nodular corneal degeneration is characterized by bilateral gray-white elevated nodules anterior to Bowman’s layer, which may be visually significant, cause foreign body sensation, or be asymptomatic. It is most common in middle-aged women. The cause of this degeneration is unknown; however, it has been associated with chronic ocular surface inflammation [1-6]. A case series of 152 eyes concluded that the most common associations were, in descending order, meibomian gland dysfunction (MGD), contact lens wear (especially hard contact lenses), peripheral vascularization, pterygium, keratoconjunctivitis sicca, and exposure keratitis [1]. Another case series of 180 eyes also identified MGD as the most common comorbidity [2].

Conservative treatment consists of management of the underlying etiology, such as MGD. Eyelid hygiene and doxycycline may be considered to treat the MGD prior to surgical treatment. Contact lens cessation or re-fitting may also be beneficial. Medical management is successful in preventing the need for surgery in most cases; the indications for surgery are (1) discomfort; (2) reduced vision due to progressive increase in astigmatic error. Surgery is uniformly successful in removal of the nodule and almost invariably successful in reducing the astigmatic error in allowing for improvement in best corrected visual acuity, usually with a much weaker spectacle prescription. Stabilization of the refractive error usually takes 3 to 6 weeks. In rare cases of neglected nodules within the visual axis, anterior stromal haze may compromise the final visual results.

Whereas subtle recurrences are common in most cases over a 5 to 15 year period, visually significant recurrences are uncommon (5 to 20%) and can be minimized with meticulous control of the etiology responsible for the condition [1, 2]. Those cases can be treated with repeat superficial keratectomy. If anterior stromal haze persists in the visual axis after treatment of primary or recurrent nodules, phototherapeutic keratectomy can be offered [1].

Superficial keratectomy of astigmatism-inducing Salzmann’s nodules should always be performed prior to cataract surgery. Stable keratometry readings and IOL calculations cannot be reliably obtained until 3 to 6 weeks after surgery. Toric IOLs should be avoided because of the risk of recurrent nodules and induction of post-operative astigmatism.

Since the diagnosis of Salzmann’s nodular corneal degeneration is clinical, several more examples appear in figures 6-10.
| Figure 6a: | Slit lamp photo of the right eye of another patient with a superior Salzmann’s nodule. |
| Figure 6b: | Slit lamp photo of the left eye of another patient with a superior Salzmann’s nodule. |
| Figure 6c: | Corneal topography. The right eye has an against the rule irregular astigmatism with superior steepening. The placido image shows irregular mires superiorly. The left eye has an asymmetric against the rule irregular astigmatism with superonasal steepening and corresponding irregular mires on placido image. |

<p>| Figure 7a: | Slit lamp photo of the left eye of another patient with Salzmann's nodular corneal degeneration. Nodules are peripheral and most evident superonasally and superotemporally in this photo. |
| Figure 7b: | Slit lamp photo of the right eye. Salzmann's nodules are annular in the superior periphery. |</p>
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<th>Figure 8a: Slit lamp photo of the right eye of another patient with Salzmann's nodular corneal degeneration. Nodules are annular in the mid-periphery.</th>
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<td>Figure 8b: Slit lamp photo of the left eye of another patient with Salzmann's nodular corneal degeneration. Nodules are annular in the interior mid-periphery.</td>
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<td>Figure 8c: Corneal topography. The right eye shows a very irregular astigmatism with superior and nasal steepening. Placido image shows irregular mires in the mid-periphery in all directions. The left eye has a very irregular astigmatism with marked superior and temporal steepening. The placido image has irregular mires with some nasal steepening.</td>
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<td>Figure 9: Slit lamp photo of the right eye of another patient with Salzmann’s Nodular Corneal Degeneration. Nodules are nasal and inferonasal.</td>
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<td>Figure 10: Slit lamp photo of another patient with Salzmann’s Nodular Corneal Degeneration. Several small nodules, superior and inferior approach the visual axis.</td>
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For a summary of the key facts regarding Salzmann’s nodular corneal degeneration, see Table 2.

### Table 2

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<th>Epidemiology</th>
<th>Signs</th>
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<td>• Onset often around 50 years of age, but has been reported in ages 13-92 [2]</td>
<td>• Bilateral gray-white elevated corneal subepithelial nodules in the paracentral or central cornea or near the limbus with associated pannus, most commonly in an annular distribution [4].</td>
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<td>• Female patients account for 75-90% of all cases [1,2].</td>
<td>• There is a subset of Salzmann’s nodules with associated vascularization [5].</td>
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<td>• Often bilateral [1,2].</td>
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<td>• Deposits are progressive</td>
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| • Associated chronic ocular surface inflammation  
  o Meibomian gland dysfunction  
  o Contact lens wear  
  o Keratoconjunctivitis sicca  
  o Exposure keratitis  
  o Trachoma | |

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<th>Diagnosis</th>
<th>Treatment</th>
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| • Clinical: based on slit lamp exam  
  • Topography: may be helpful for making the diagnosis. Nodules cause irregular astigmatism.  
  • Histopathology: Thin epithelium, thick basement membrane with displaced Bowman layer, and non-specific stromal scarring | • Superficial keratectomy (SK) is standard to reduce irregular astigmatism. In a large case series SK was successful in 90.2% of eyes [2].  
  • Phototherapeutic keratectomy has been used in some cases following SK, and a case review has not identified a significant difference in outcomes compared to repeat SK [1].  
  • Recurrence after SK is common, in one series recurrence occurred in 21.9% after a mean follow-up time of 61 months [2]. Visually significant recurrences occur in only 5-20% of patients [1]. |

### Conclusion

Salzmann’s nodular corneal degeneration is a non-inflammatory, slowly progressive nodular corneal degeneration that may induce ocular surface discomfort and progressive astigmatism with decreased best corrected visual acuity. The diagnosis is based on clinical examination. Topography is helpful in evaluating the contribution of the nodule to visual impairment. Superficial keratectomy is a successful treatment option, and visually significant recurrence afterward is relatively rare.

### Differential diagnosis [4]

- Corneal scarring
- Spheroidal degeneration

### References


**Suggested Citation format**