Guttate endothelial changes with anterior eye inflammation

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Summary

This report details the rapid appearance and gradual disappearance of guttate endothelial changes in a patient who experienced an acute 'red eye' associated with the continuous wear of contact lenses. This episode of acute inflammation was accompanied by a diffuse corneal infiltrate, endothelial bedewing, and the guttate changes.

Guttate endothelial changes in the central cornea are characteristic of primary eye disorders such as cornea guttata and Fuchs's endothelial dystrophy. Similar guttate appearances are also commonly observed in the periphery of the cornea in the aging eye. Guttata represent excrescences of Descemet's membrane, over which endothelial cells may be attenuated or absent. In the early stages guttata appear with the slit-lamp as small shallow depressions in the endothelial mosaic. In later stages they appear as distinct round black holes.

Cornea guttata secondary to active ocular disease is a less well known phenomenon. Wolter and Larson described the occurrence of confluent guttata in the form of lines and geographic patterns in cases of long-standing luetic interstitial keratopathy. Histological studies of such cases have revealed varying degrees of correspondence between the pattern of guttata and the pattern of deep stromal vessels or posterior corneal folds. Endothelial changes have also been reported with a variety of other conditions involving the anterior eye. In these reports, however, the nature of these changes and the time scale for their appearance are not clear. Berliner described circular crevices between cells 'similar to the picture of endothelial dystrophy' occurring in the early stages of interstitial keratitis and guttata changes occurring after the acute inflammation had subsided.

Grayson and Brandreth have both observed the endothelium to take on an appearance resembling cornea guttata during ocular inflammation, this appearance subsiding when the inflammation has passed. They have termed these episodes endothelial oedema. Inomata and Smelser, however, have demonstrated the rapid appearance of excrescences in Descemet's membrane in the rabbit cornea during experimentally induced uveitis.

The purpose of this paper is to report the appearance and disappearance of guttate endothelial changes in a patient during an episode of anterior eye inflammation related to continuous contact lens wear.

Case History

The patient was a 42-year-old woman of good general health. She had congenital high myopia with nystagmus, and light perception only in the left eye. She had 15 years previous experience with a daywear contact lenses in the right eye. As the patient was about to be involved in a continuous-wear contact lens programme, endothelial photographs were taken by a high-magnification slit-lamp technique. The endothelial appearance was normal. A continuous-wear programme was begun. The patient had two episodes of nonulcerative keratitis from which the eye rapidly recovered. A third attempt at continuous-wear was started. At a routine follow-up examination 3 weeks later minor symptoms were reported. There was slight punctate fluorescein staining of the central cornea and slight injection of limbal and bulbar conjunctival vessels. The contact lens fitting was tight (little movement), and the lens front surface was relatively dry. The patient was given a hydrophilic contact lens lubricating solution. The patient slept comfortably that night but awoke in the morning with marked pain in and around the right eye, the sinuses, and teeth. The eyelids were swollen, the eye red and lacri-
mating. The patient removed the lens and took mediation for the facial pain.

The symptoms had very slightly abated when she was examined the following day. The left eye was normal. The right eye presented a typical 'red-eye' reaction associated with continuous contact lens wear. There was considerable conjunctival and ciliary vessel engorgement. There was no fluorescein staining of the epithelium but the cornea was oedematous, and there was a dense infiltrate in the anterior half of the stroma across the entire cornea. There were inflammatory cells (grade 1) in the anterior chamber, but the intraocular pressure was normal. There were fine dust-like particles (measured from the photographs to be approximately 10 μm in diameter) scattered over the posterior surface of the endothelium, giving a widespread bedewing appearance (Fig. 1). There were no large keratic precipitates. The corneal endothelium was irregular in contour, showing numerous round, black areas (approximately 50 μm in diameter) similar in appearance to corneal guttae (Figs. 2, 3). When viewed with a narrow slit-lamp beam these black holes appeared as depressions in the endothelial contour with a raised rim (Fig. 2). These guttate areas were not present before this episode.

On the fourth day after onset of the 'red eye' minor symptoms persisted. The cornea was less oedematous and the stromal infiltrate considerably reduced. The corneal endothelium was less irregular in contour and the guttate appearances much less numerous. There were very few inflammatory cells in the anterior chamber, and a small area of endothelial bedewing was present on the inferior cornea.

One week later the corneal infiltrate had almost disappeared, the guttate endothelial changes were absent, and the eye was quiet.

Two days later the patient resumed continuous wear with the original contact lens. There was no sign of any endothelial changes during the subsequent year of successful lens wear.

Discussion

Acute anterior eye inflammation has been previously reported with therapeutic contact lenses and with
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in this patient had the same composition is unknown. What is clear is that the guttate changes observed are similar in appearance, size, and distribution to those observed in primary endothelial dystrophy, and therefore almost certainly represent accumulations anterior to the endothelium. Whether disturbance of the endothelium was the stimulus for the preendothelial accumulations is speculation, but it is reasonable to assume that both the bedewing and the guttate changes were consequences of the anterior eye inflammation.

References

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