Retinitis Pigmentosa Sine Pigmento
Masquerading as Normal Tension Glaucoma

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INTRODUCTION

Retinitis pigmentosa (RP) is a heterogeneous group of diseases with progressive photoreceptor degeneration. In some RP patients, the typical bone spicule formation is lacking and the retina may appear normal in the early stages of the disease. This poses a diagnostic challenge to the ophthalmologist. We herein report a case of unilateral progressive visual field loss and optic atrophy due to RP sine pigmento which was initially masqueraded as normal tension glaucoma (NTG).

CASE REPORT

A 42 years-old gentleman was referred to our unit with progressive blurring of vision in his right eye for 6 months, especially affecting peripheral vision and in dim conditions. He had a history of hypertension, was an exsmoker, with no family history of glaucoma/Retinitis pigmentosa (RP). Initial examination revealed a visual acuity of 6/6 and normal color vision OU, with no relative afferent pupillary defect. Autorefraction yielded – 6.50 DS (OD) and –7.50 DS (OS). The intraocular pressure (IOP) by applanation tonometry was 17 mm Hg (OD) and 18 mm Hg (OS). Fundal exam showed myopic fundi OU with pink disks and vertical cup-to-disk ratios of 0.6 (OD) and 0.5 (OS). There was inferior rim thinning and peripapillary atrophy of the right optic disk (Figs 1A and B). Examination was otherwise unremarkable. Central corneal thickness was 554 µm (OD) and 567 µm (OS). Automated white-on-white threshold perimetry (program 24- 2, SITA fast, model 750, Humphrey Instruments, Dublin, CA) showed right constricted field (superior > inferior) extending in an arcuate fashion from the blind spot (Fig. 2A). Optical coherence tomography showed a borderline- thinned peripapillary retinal nerve fiber layer thickness of 88.75 µm OD and 90.89 µm OS. Magnetic resonance imaging of the brain, orbit and pituitary gland with contrast was unremarkable. In view of right inferior optic disk rim thinning and normal IOP together with corresponding superior constricted visual field, normal tension glaucoma (NTG) was suspected. However, as the severity of the visual field loss was out of proportion to the structural damage (mean deviation = –15.66 dB) and a repeat visual field examination one year after presentation revealed a rapid mean deviation decline of –7.99 dB (Fig. 2B), a review by medical retinal specialist was initiated. At this point, his visual acuity and color vision remained normal OU but there was a right-sided relative afferent pupillary defect, with progressive changes in the retina (Fig. 1B). The left eye remained normal. Full field electroretinogram (ERG) of his right eye showed non-recordable scotopic rod responses. For the photopic cone
responses, the a-wave was nonrecordable and the b-wave was delayed with a smaller amplitude than normal. The 30 Hz response also had a reduced amplitude. The left eye responses were normal. The final diagnosis was thereby confirmed by the ERG finding.

DISCUSSION

We initially considered NTG as our provisional diagnosis owing to an enlarged vertical cup-to-disk ratio, inferior disk rim thinning and corresponding visual field loss, although the highly asymmetrical involvement was not typical and the structural-functional relation was out of proportion. With time, there was a rapid decline of visual field without significant changes in media opacity (mean deviation deteriorated –7.99 dB in a year). These made the diagnosis of NTG less likely. Another diagnostic clue in this case is the progressive constriction of visual field despite adequate IOP reduction. RP is usually a bilateral disease, but unilateral cases have been reported.1,2

We have also documented, from our patient, that progression from "normal" fundal appearance to a phase with vessel attenuation, disk pallor and rapid visual field progression can occur within a year’s time for unilateral RP sine pigmento.

In summary, RP sine pigmento should be considered as a differential diagnosis in young NTG suspects who had out-of-proportion structure-function correlation, even without a family history or obvious retinal pigment epithelium abnormalities such as bone spicule formation. An ERG is helpful to confirm the diagnosis in such cases.

REFERENCES