Short Communication

A rare case of oguchi disease exhibiting the classic Mizuo-Nakamura phenomenon

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ABSTRACT

Introduction: Oguchi disease also called congenital stationary night blindness is an autosomal recessive genetic condition first identified in Japan with a few cases reported in other countries like Europe, America, India and Pakistan. We report one such case exhibiting the Mizuo-Nakamura phenomenon with a reduced amplitude on the b-wave of the rod electroretinogram advocating this diagnosis.

Case Report: A 40-year-old man presented with complaints of non-progressive decrease in vision as the day progressed for 6 months with normal vision during the day in both eyes. There was no history of trauma or any other ocular or systemic disease. His parents had a second-degree consanguineous marriage and his father experienced similar symptoms. Visual acuity for distance was 6/6 and N6 for near in both eyes. His anterior segment and posterior segments were normal. When the patient was put to dark adaptation for 3 hours, there was a change in his fundus background colour from grey to red (Mizuo-Nakamura phenomenon). The b-wave amplitude was decreased on the rod electroretinogram. The cone electroretinogram on the other hand was normal. All this was confirmed by his genetic study which favoured this diagnosis.

Conclusion: This disease presents with features similar to retinitis pigmentosa in terms of mutations in the SAG gene. This highlights the importance of the Mizou-Nakamura phenomenon in formulating a confirmatory diagnosis and treatment protocol.

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1. Introduction

Oguchi’s disease is a rare autosomal recessive disorder. It is also known as congenital stationary night blindness and was initially described in the year 1907 by a Japanese clinician by name Chuta Oguchi.1 Although mostly found in Japan, few cases have also been reported in India, Pakistan and the Far West. The patient presents with diminished vision during the night with fairly normal vision during the day. This decrease in vision is non-progressive and is similar in symptomatic presentation to conditions like Retinitis Pigmentosa, Stargardts disease, Juvenile retinoschisis and Progressive cone dystrophy. Although these conditions have unique fundus findings and electroretinogram readings which are confirmatory of their diagnosis, Oguchi’s disease has a normal fundus picture with a decrease in the amplitude of the b-wave on the rod electroretinogram with a normal cone electroretinogram.2 The patient experiences a characteristic phenomenon, named after its founders, Mizuo and Nakamura. This unique morphological and functional abnormality of the retina is a typical golden yellow or sometimes a silver-grey discoloration of the retina and posterior segment in the presence of light which disappears after dark adaptation only to reappear again in the presence of light.3 These features along with genetic testing is confirmatory as the mutations in the SAG gene involved in Oguchi’s disease may also be involved in few cases of familial retinitis pigmentosa.
2. Case Report

A 40-year-old man presented with complaints of non-progressive decrease in vision as the day progressed for 6 months with normal vision during the day in both eyes. There was no history of trauma or any other ocular or systemic disease. His parents as a second-degree consanguineous marriage and his father experienced similar symptoms. Visual acuity for distance was 6/6 and N6 for near in both eyes. His anterior segment and posterior segment were normal with no evidence of bony spicules or atrophy. Visual field examination, using the Humphrey’s visual field analyser and colour vision was normal bilaterally. The fundus showed a golden to grey discolouration in light (Figures 1 and 2). When subjected to dark adaptation for 3 hours, the fundus colour changed to red (Mizuo-Nakamura phenomenon) (Figures 3 and 4). The rod electroretinogram showed reduced amplitude in the b-wave (Figure 5) while the cone electroretinogram was normal suggestive of retinal dysfunction rather than degeneration (Figure 6). Examination of the man’s mother and his two brothers were unremarkable. The father could not be evaluated as he had passed away in a road traffic accident two years ago.

3. Discussion

Oguchi’s disease is mainly caused by a mutation. This gene codes for arrestin which is located on chromosome 2q37 described as Oguchi type 1. On the other hand, mutations in GRK 1 gene coding for rhodopsin kinase is present on chromosome 13q34 called Oguchi type 2. Mutations in the SAG gene can also cause Retinitis Pigmentosa. Therefore, genetic testing must also be correlated clinically in order to establish a definitive diagnosis. The electroretinogram has a negative a-wave followed by a positive b-wave indicating bipolar cell activity. A decrease in amplitude of the b-wave
in the rod electroretinogram is seen in Oguchi’s disease whereas there is a decrease in b-wave amplitude on the cone electroretinogram in Progressive cone dystrophy.

4. Conclusion

It is of primary concern to thoroughly evaluate the patient before establishing a definitive diagnosis and thereby planning a management protocol. A genetic study although confirmatory must be correlated clinically in a case of Oguchi’s disease where it is important to evaluate the fundus in both light as well as in dark.

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6. Conflict of Interest

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References


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