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Multimodal imaging in *CABP4*-related retinopathy.

**Running Head:** Retinal structure in *CABP4*-related retinopathy.

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ABSTRACT

Purpose: Multimodal imaging has not been documented for \textit{CABP4}-related retinopathy. In this study we describe optical coherence tomography (OCT) and fundus autofluorescence findings for 5 genetically-confirmed cases.

Methods: Retrospective case series.

Results: Four patients with the previously-described homozygous Saudi \textit{CABP4} founder mutation c.81_82insA (p.Pro28ThrfsX44) and 1 patient with the homozygous mutation c.1A>G (p.Met1?) in \textit{CABP4} were examined. The ages ranged between 9-16 years at last follow up, and the duration of follow up ranged from 2-12 years. Foveal thickness was reduced ranging between 175-212 micrometers. Wide field imaging including fundus autofluorescence was unremarkable. All patients presented with a negative electroretinogram, with a variable amount of cone and rod dysfunction. Over follow-up, there was no electroretinographic indication of any progressive retinal dysfunction.

Conclusions: Foveal thinning is a feature of \textit{CABP4} retinopathy. Normal autofluorescence is consistent with inner retinal dysfunction and suggests the condition could be amenable to gene therapy. Retinal dysfunction was stable throughout follow-up.

Keywords: \textit{CABP4}, Congenital stationary night blindness, optical coherence tomography.
Introduction

*CABP4*-related retinopathy, also termed congenital non-progressive cone-rod synaptic disorder (CRSD; MIM #610427), is caused by mutations in the gene encoding calcium-binding protein-4 (CABP4; MIM #608965) on chromosome 11q13. It is characterized by stable low vision, nystagmus, photophobia, a normal or near-normal fundus appearance, and no night blindness. Electretinography shows an electronegative waveform response to scotopic bright flash, near-normal to subnormal rod function, and delayed and/or decreased to nonrecordable cone responses.[1-3] The underlying cause is a dysfunction of the photoreceptor-bipolar synapse, where CABP4 is specifically located on the side of the photoreceptor synaptic terminal. Together with CACNA1F and CACNA2D4, CABP4 plays an important role in the photoreceptor-bipolar cell synapse by allowing the continuous release of glutamate into the synapse. Mutations in the genes encoding these proteins may cause disturbed glutamate release resulting in synaptic dysfunction which manifests clinically as an electronegative electroretinogram, where the a-wave which emanates from the phototransduction is greater than the b-wave which results from activity in the bipolar and Muller cells. Phenotypically such mutations have been described to cause cone dystrophy, cone-rod dystrophy or incomplete congenital stationary night blindness.[1, 4-6]

Albeit rare with only about 10 cases described in the literature (summarized by Khan et al.[1]), *CABP4*-related retinopathy is unique in that it does not cause night blindness, in spite of the protein's function being closely related to other proteins implicated in incomplete congenital stationary night blindness, and that there is a negative electroretinogram, which is hallmark of the latter condition.

In this paper we for the first time present the results of multimodal imaging for *CABP4*-related retinopathy as well as longer-term follow-up of retinal function.
Methods

This study was approved by the institutional research ethics committee and adhered to the tenets of the Declaration of Helsinki. We reviewed the records of patients homozygous for CABP4 mutations known to one of the authors (AOK). All had the Saudi founder mutation (c.81-82insA; p.Pro28ThrfsX44) except one [Patient 2 in Table 1, who was homozygous for the mutation c.1A>G (p.Met1?)], and all had been previously noted in the literature.[1,2]

Retinal structure was analyzed qualitatively and quantitatively with transfoveal horizontal spectral domain optical coherence tomography scans (Spectralis OCT, Heidelberg Engineering, Inc., Heidelberg, Germany). Measurement of foveal retinal thickness, which was defined as the shortest distance from the top of the retinal pigment epithelium line to the internal limiting membrane in the right eyes of patients, was obtained with calipers using the Heidelberg software. In addition, in 2 of the patients (Patients 2 and 5), the Spectralis macular raster consisting of 19 horizontal 6 millimeter line scans, was performed, in spite of nystagmus. This enabled automated software algorithm to display with numeric averages of the macular thickness measurements for each of the 9 map sectors as defined by the Early Treatment Diabetic Retinopathy Study (ETDRS).[7] Retinal thickness in patients was compared to that of controls from a previously published paper by us, [8] using identical protocols (caliper measurement and ETDRS map) in patients and controls. Wide field imaging including fundus autofluorescence was performed in all patients (Optos PLC, Dunfermline, UK). Four of the patients (all except Patient 2) had macular autofluorescence imaging with a fundus camera (Topcon TRC-50DX, Topcon Medical Systems, Inc., NJ, US).

Retinal function was evaluated with full-field electroretinography (ffERG, Nicolet Biomedical Instruments, Madison, Wisconsin, USA), in dark adapted and light adapted state according to ISCEV standards,[9] with a few modifications as follows. Retinal function was evaluated with full-field electroretinography (ffERG, Nicolet Biomedical Instruments, Madison,
Wisconsin, USA), in dark adapted and light adapted state according to ISCEV standards, with a few modifications as follows.[8] Full-field electroretinograms were recorded in a Nicolet analysis system (Nicolet Biomedical Instruments, Madison, Wisconsin, USA), after dark adaptation of subjects for 40 min, dilatation of the pupils with topical cyclopentolate 1% and metaxexedrine 2.5% and topical anaesthesia, with a Burian Allen bipolar contact lens and a ground electrode applied to the forehead. Responses were obtained stimulating with single full-field flash (30 ms) with blue light light (0.81 cd-s/m2: rod response) and with white light (10.02 cd-s/m2: combined rod-cone response). In addition, the dark adapted cone response was measured after stimulation with dim red light (3.93 cd-s/m2). Photopic responses were obtained with a background illumination of 3.4 cd-s/m2 in order to saturate the rods.

**Results**

OCT foveal thickness using caliper measurement at most recent follow-up showed foveal thinning (Table 1, Fig. 1). Foveal thickness ranged between 175-212 micrometers (range of normal: 215-245).[8] In addition, patients 2 and 5 displayed reduced averages of macular thickness measurements for each of the 9 ETDRS map sectors, including the central 1 millimeter sector, compared to controls (data not shown), except for a single value for the outer superior sector in Patient 2 [285 microns for patient 2, compared to median (300) and range (281-332) for controls for that sector]. Due to nystagmus in all patients, the quality of the scans did not allow thickness analysis of the various retinal sublayers. There seemed to be a thinning of the outer plexiform layer in for example Patient 3 with the homozygous Saudi founder mutation c.81_82insA (p.Pro28ThrfsX44), whereas it seemed to be more prominent in Patient 2 with the homozygous mutation c.1A>G (p.Met1?), in CABP4 (Fig. 1). However this may have been due to variable reflectivity and thus visualization of Henle’s fiber layer due to its birefringent optical properties and
oblique orientation, in the context of horizontal displacement of the OCT entry beam through the pupil, due to nystagmus. [10] Thus visualization of the Henle’s fiber layer, which consists of the axons of the photoreceptors, could account for the apparent prominence of the outer plexiform layer. Fundus wide field and macular imaging demonstrated unremarkable fundi including normal distribution of autofluorescence (Fig. 1).

Retinal function was measured in all patients using full field electroretinography, and in 3 out of 5 patients, long-term follow-up of retinal function was possible, ranging over 5-12 years (Table 1). Dark adapted rod responses were present in 3 patients and were normal in 1 of these (Table 1). Dark adapted cone responses (“b1”) were present in 2 of 5 patients, albeit reduced compared to normal (Table 1, Figs. 2-4). Light adapted cone responses and 30 Hz flicker responses were severely reduced in all patients, however the 30 Hz flicker implicit time was not delayed as seen in progressive retinal dystrophies (Table 1, Figs. 2-4). Furthermore, the cone and rod-cone a-waves, emanating from phototransduction activity proximal to the photoreceptor-bipolar synapse, were normal or near-normal. Retinal function was stable over time in all patients (Table 1, Figs. 2-4).

Discussion

In an animal mouse model of \textit{CABP4}-related retinopathy, the outer plexiform layer was thinner in Cabp4 +/- mice, with a reduction in the number of synaptic ribbons and photoreceptor terminals and there were ectopic synapses originating from rod bipolar and horizontal cells extending into the outer nuclear layer.[11] Here, we found thinning of foveal retinal thickness by optical coherence tomography, and in addition, our results strongly suggest macular retinal thinning also outside the fovea in all 9 ETDRS sectors. It is to be noted that the control group used for this study consisted of healthy myopic subjects (range 6–9, median 7.5 diopters of myopia) designed to
match myopic patients from a previous study, [8] and would thus be expected to have thinner-than normal retinas of emmetropic subjects. This further supports a generalized macular retinal thinning in CABP4-related retinopathy. However due to nystagmus the quality of the scans did not permit any quantitative analysis of retinal sublayer thickness.

Retinal function as measured with full field ERG was stable over time and there was no sign of progressive retinopathy. Consistently, all patients had severe reduction of light and dark adapted cone b waves, whereas the cone and rod-cone a-waves, emanating from phototransduction activity proximal to the photoreceptor-bipolar synapse, were normal or near-normal. Rod function was variably reduced and normal in 1 of the patients. One can speculate if CABP4 may have a more prominent role in cone synapses than rod synapses in humans. Consistently, dark adapted Cabp4 -/- mice showed consistently lower ERG a- and b-waves compared to controls, whereas light adapted Cabp4-/- mice showed smaller differences in the a-wave amplitudes than in the b-wave amplitudes, compared to controls.

Fundus imaging was unremarkable in all patients and there was a normal distribution of fundus autofluorescence. This is consistent with the condition being primarily a problem of inner retinal dysfunction and suggests it could be amenable to gene therapy.

In summary, retinal dysfunction was stable over long term follow-up in CABP4-related retinopathy. Foveal retinal thickness was reduced compared to normal, which may be due to primarily reduced thickness of the outer plexiform layer, as seen in Cabp4-/- mice, and could result from a lack of the protein’s function in the photoreceptor-bipolar synapse in this layer. Autofluorescence was unremarkable. These findings further our understanding of the condition.
ACKNOWLEDGEMENTS

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b. Conflict of interest disclosures: None for each of the authors.

c. Contributions to Authors:

Design of the study (PS, AE, AK).

Conduct of the study (PS, AE, AK).

Collection, management, analysis, and interpretation of the data (PS, AE, AK).

Preparation, review and final approval of the manuscript (PS, AE, AK)
References


Figure legends

Figure 1: Wide field imaging and optical coherence tomography findings in CABP4-related retinopathy.

Top panel: Right eye. Wide field imaging and fundus autofluorescence in Patient 3 with the homozygous Saudi founder mutation c.81_82insA (p.Pro28ThrfsX44) in CABP4. There is a normal distribution of autofluorescence.

Lower panel: Right eyes. Optical coherence tomography in Patient 3 (left column) with the homozygous Saudi founder mutation c.81_82insA (p.Pro28ThrfsX44) and in Patient 2 (right column) with the homozygous c.1A>G (p.Met1?) mutation in CABP4. The outer plexiform layer (between arrows) seems more prominent in Patient 2 than in Patient 3. However this may have been due to variable reflectivity and thus visualization of Henle’s fiber layer due to its birefringent optical properties and oblique orientation, in the context of horizontal displacement of the OCT entry beam through the pupil, due to nystagmus, as described by Lujan et al. 2011. [10] Thus visualization of the Henle’s fiber layer, which consists of the axons of the photoreceptors, could account for the apparent prominence of the outer plexiform layer.

Figure 2: Right eye. Long term follow up of retinal dysfunction by full-field electroretinography in Patient 1 with the homozygous Saudi founder mutation c.81_82insA (p.Pro28ThrfsX44) in CABP4. No rod function can be measured. Note electronegative combined rod-cone response. Light
adapted cone responses are reduced. There is no sign of progressive retinal dysfunction over follow-up. A normal full-field electroretinogram is shown for comparison in the right column. ms=milliseconds. µV=microvolts.

Figure 3: Right eye. Long term follow up of retinal dysfunction by full-field electroretinography in Patient 2 with the homozygous homozygous c.1A>G (p.Met1?) mutation in \textit{CABP4}. There is a reduced but measurable rod response. Note electronegative combined rod-cone response. Light adapted cone responses are reduced. There is no sign of progressive retinal dysfunction over follow-up. ms=milliseconds. µV=microvolts.

Figure 4: Right eye. Long term follow up of retinal dysfunction by full-field electroretinography in Patient 4 with the homozygous Saudi founder mutation c.81_82insA (p.Pro28ThrfsX44) in \textit{CABP4}. There is a prominent rod response. Note electronegative combined rod-cone response. Light adapted cone responses are reduced. There is no sign of progressive retinal dysfunction over follow-up. ms=milliseconds. µV=microvolts.
Table 1. Long term retinal function and foveal thickness in 5 patients with \textit{CABP4}-related retinopathy. All patients were male.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age</th>
<th>Gender</th>
<th>Binocular VA at first visit (age 1st visit), binocular VA at follow-up</th>
<th>Refraction at follow-up (OD, OS)</th>
<th>OCT FT OD</th>
<th>Full-field electroretinography (ffERG) at presentation, ffERG at follow up (OD)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
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<td></td>
<td></td>
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<td>Rod a wave</td>
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<td></td>
<td></td>
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<td></td>
<td></td>
<td>A</td>
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<tr>
<td>1</td>
<td>16</td>
<td>M</td>
<td>20/200 (4), 20/200</td>
<td>187</td>
<td>0</td>
<td>0</td>
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<tr>
<td>2</td>
<td>9</td>
<td>M</td>
<td>F&amp;F (2), 20/80</td>
<td>212</td>
<td>34, 138</td>
<td>87</td>
</tr>
<tr>
<td>3</td>
<td>11</td>
<td>M</td>
<td>20/100 (9), 20/200</td>
<td>175</td>
<td>106</td>
<td>117</td>
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<tr>
<td>4</td>
<td>10</td>
<td>M</td>
<td>20/300 (6), 20/300</td>
<td>197</td>
<td>148, 202</td>
<td>192, 302</td>
</tr>
<tr>
<td>5</td>
<td>9</td>
<td>M</td>
<td>20/80 (7), 20/70</td>
<td>204</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>
VA=Visual acuity, OCT FT=optical coherence tomography foveal thickness, OD=right eye, OS=left eye, OU=both eyes, A=Amplitude, It=implicit time. F&F=Fixates and follows.

* Reference material for optical coherence tomography from Oreany et al.[8]