Retinopathy Associated with Biallelic Mutations in PYGM (McArdle Disease)

Identification of ocular associations with systemic diseases can aid diagnosis and phenotyping and can yield pathophysiologic insights. McArdle disease (glycogen storage disease type V) is a rare metabolic myopathy (estimated prevalence, 1/100,000) resulting from biallelic mutations in the PYGM gene, encoding muscle glycogen phosphorylase (reviewed by Lucia et al²). Patients experience exercise intolerance and risk acute rhabdomyolysis, although life expectancy is rarely affected. A case report from 1988 described pattern dystrophy of the retinal pigment epithelium (RPE) in an affected patient. Two further cases have been reported. With only 3 cases, chance association is possible; the 2 entities may be unrelated.

Pattern dystrophies can be associated with mutations in a number of genes, most frequently PRPH2. We describe here a similar macular appearance in 4 further, unrelated patients with McArdle disease, including findings from multimodal retinal imaging and electrophysiologic examination. Genetic screening was performed, both to confirm PYGM mutations and to check for mutations in a number of known genes implicated in macular or pattern dystrophies. The latter screening showed negative results, suggesting that the 2 entities (retinopathy and McArdle disease) are indeed related. This study had research ethics committee approval.

Patient 1, a 64-year-old white man, was referred because of longstanding reduction in right eye vision and an unusual macular appearance. He reported exercise intolerance since childhood and had been diagnosed with McArdle disease after muscle biopsy. Unaided Snellen visual acuity was 20/80 in the right eye (20/60 with pinhole) and 20/20 in the left eye. Over the next 10 years, visual acuity deteriorated in the right eye to counting fingers but remained stable in the left eye. Patient 2, a 63-year-old South Asian man, was referred for reduction in left eye vision over the previous few years. McArdle disease had been diagnosed in late adulthood after muscle biopsy. Visual acuity was 20/17 in each eye; 4 years later, visual acuity was 20/20 and 20/80, respectively.

Patient 3, a 56-year-old white man, was noted previously to have an abnormal retinal appearance by his optometrist. He had been diagnosed with McArdle disease (molecularly confirmed) 2 years before review in our service. He was visually asymptomatic; visual acuity was 20/17 in each eye. Patient 4, a 68-year-old white man, was referred to his local hospital eye service because of visual symptoms; patients 3 and 4 were visually asymptomatic, but an optometrist detected an abnormal fundal appearance. Genetic testing for macular or pattern dystrophy genes was initiated by our service, and genetic confirmation of McArdle disease also was obtained after review in our service (except in patient 3).

This article reports multimodal imaging findings of a distinctive retinopathy in 4 unrelated patients with McArdle disease, similar to the previous case reports. We also report results of retinal electrophysiologic analysis: despite far peripheral abnormalities on ultrawide-field imaging, there was no definite electrophysiologic evidence of generalized retinal, or generalized RPE, dysfunction (full-field electroretinography and electro-oculography light rise results were normal).

Patients 1, 2, and 4 underwent electroretinography recording according to international standards. All demonstrated normal full-field electroretinography results; pattern electroretinograms were undetectable in patient 1 and within normal limits in patients 2 and 4. Patients 1 and 2 also underwent electro-oculography, which showed a normal light rise.

Patients 1 and 2 underwent Sanger sequencing of all coding exons and exon-intron boundaries of PRPH2, thought to be the most likely genetic cause of such a fundus appearance (Manchester Centre for Genomic Medicine, Manchester, United Kingdom). No pathogenic variants were found. All patients underwent screening for disease-causing mutations in a number of additional genes implicated in macular or pattern dystrophies: ABCA4, BEST1, CDH3, EFEMP1, ELOVL4, IMPG1, IMGP2, PROM1, PRPH2, TIMP3; Molecular Vision Laboratory, Hillsboro, Oregon). For patients 2, 3, and 4, the panel also included DRAM2, RPIL1, and TTL5. Results were negative.

Screening of PYGM (Sheffield Diagnostic Genetics Service, Sheffield, United Kingdom) molecularly confirmed McArdle disease in all patients. Patient 1 was homozygous for the frequently reported nonsense mutation p.(Arg50*). Patient 2 harbored the novel homozygous stop mutation p.(Gln176*). Patients 3 and 4 were compound heterozygote: patients 3, p.(Arg50*) and p.(Gly205Ser); patient 4, p.(Arg94Trp) and p.(Gly695Arg).

In all patients, diagnosis of McArdle disease preceded detection of retinal abnormalities (apart from patient 3). Patients 1 and 2 were referred to the ophthalmology service because of visual symptoms; patients 3 and 4 were visually asymptomatic, but an optometrist detected an abnormal fundal appearance. Genetic testing for macular or pattern dystrophy genes was initiated by our service, and genetic confirmation of McArdle disease also was obtained after review in our service (except in patient 3).

Figure 1 depicts the appearance of the fundus on color photography and autofluorescence, the latter showing distinctive reticular areas of stippled hyperautofluorescence. Spectral-domain OCT (Fig S1, available at www.aaojournal.org) confirmed abnormalities at the level of the RPE, outer retina, or both. Ultra-widefield autofluorescence imaging showed abnormalities in the far peripheral retina also, particularly nasally (Fig S2, available at www.aaojournal.org).

Patients 1, 2, and 4 underwent electroretinography recording according to international standards. All demonstrated normal full-field electroretinography results; pattern electroretinograms were undetectable in patient 1 and within normal limits in patients 2 and 4. Patients 1 and 2 also underwent electro-oculography, which showed a normal light rise.

Patients 1 and 2 underwent Sanger sequencing of all coding exons and exon-intron boundaries of PRPH2, thought to be the most likely genetic cause of such a fundus appearance (Manchester Centre for Genomic Medicine, Manchester, United Kingdom). No pathogenic variants were found. All patients underwent screening for disease-causing mutations in a number of additional genes implicated in macular or pattern dystrophies: ABCA4, BEST1, CDH3, EFEMP1, ELOVL4, IMPG1, IMGP2, PROM1, PRPH2, TIMP3; Molecular Vision Laboratory, Hillsboro, Oregon). For patients 2, 3, and 4, the panel also included DRAM2, RPIL1, and TTL5. Results were negative.

Screening of PYGM (Sheffield Diagnostic Genetics Service, Sheffield, United Kingdom) molecularly confirmed McArdle disease in all patients. Patient 1 was homozygous for the frequently reported nonsense mutation p.(Arg50*). Patient 2 harbored the novel homozygous stop mutation p.(Gln176*). Patients 3 and 4 were compound heterozygote: patients 3, p.(Arg50*) and p.(Gly205Ser); patient 4, p.(Arg94Trp) and p.(Gly695Arg).

In all patients, diagnosis of McArdle disease preceded detection of retinal abnormalities (apart from patient 3). Patients 1 and 2 were referred to the ophthalmology service because of visual symptoms; patients 3 and 4 were visually asymptomatic, but an optometrist detected an abnormal fundal appearance. Genetic testing for macular or pattern dystrophy genes was initiated by our service, and genetic confirmation of McArdle disease also was obtained after review in our service (except in patient 3).

This article reports multimodal imaging findings of a distinctive retinopathy in 4 unrelated patients with McArdle disease, similar to the previous case reports. We also report results of retinal electrophysiologic analysis: despite far peripheral abnormalities on ultrawide-field imaging, there was no definite electrophysiologic evidence of generalized retinal, or generalized RPE, dysfunction (full-field electroretinography and electro-oculography light rise results were normal).

Screening results for mutations in a number of macular dystrophy genes were negative. In particular, the findings could not be attributed to mutations in PRPH2, which would have been most likely to result in a similar retinal phenotype. Patient 4 had been diagnosed previously with Stargardt disease, but genetic testing yielded no mutations in ABCA4. The results of this study are of clinical significance because they support the association of this retinopathy with McArdle disease and can reduce the likelihood of misdiagnosis; this is increasingly important because particular genetic causes of retinopathy (including ABCA4) are subject to a number of novel treatment trials.
Why might muscle glycogen phosphorylase deficiency affect the retina? The enzyme catalyses conversion of glycogen to glucose-1-phosphate in skeletal muscle. Liver, muscle, and brain have different isoforms of the enzyme. In human RPE, glycogen is present, and both muscle and brain isoenzymes have been demonstrated. Intracellular glycogen is likely to act as a buffer in glucose delivery from choroid to photoreceptors via the RPE. It is possible that deficiency of a glycogen phosphorylase isoenzyme may disturb this process, leading to degeneration. This is one hypothesis; other mechanisms are possible, and further work is needed.

Acknowledgments

The authors thank Priya Francome-Wood, Moorfields Eye Hospital, and Andrew Wakelin, Association for Glycogen Storage Disease UK.

**Figure 1.** Retinal appearance on color imaging (left panels) and fundus autofluorescence (FAF) imaging (right panels). A, Color fundus images from patient 1. B, Short-wavelength (488-nm) posterior pole (55°) FAF images from patient 1. C, D, Corresponding images from patient 2. E, F, Corresponding images from patient 3. G, H, Pseudocolor image and 488-nm FAF image from patient 4. In all cases, yellow reticular lesions are visible on color imaging, with atrophy in patient 1 and small areas of pigmentary change in patients 2, 3, and 4. Fundus autofluorescence (FAF) shows reticular hyperautofluorescence (likely to result from loss of photoreceptor outer segments, accumulation of fluorophore in the retinal pigment epithelium [RPE], or both) in all cases, with additional dark areas indicating RPE atrophy in patient 1.

**Omar A. Mahroo, PhD, FRCOphth**
**Kamron N. Khan, PhD, FRCOphth**
**Genevieve Wright, MSc**
**Zoe Ockrim, FRCOphth**
**Renata S. Scalco, MD**
**Anthony G. Robson, PhD**
**Adnan Tufail, MD(Res), FRCOphth**
**Michel Michaelides, MD(Res), FRCOphth**
**Ros Quinlivan, MD, FRCPath**
**Andrew R. Webster, MD(Res), FRCOphth**

1. UCL Institute of Ophthalmology, University College London, London, UK; 2. Genetics and Medical Retina Services, Moorfields Eye Hospital, London, UK; 3. Department of Ophthalmology, King’s College London, St Thomas’ Hospital Campus, London, UK; 4. Section of Ophthalmology...
and Neuroscience, Leeds Institute of Biomedical and Clinical Sciences, University of Leeds, Leeds, UK; MRC Centre for Neuromuscular Disease, National Hospital for Neurology and Neurosurgery, London, UK.

Financial Disclosure(s): The author(s) have no proprietary or commercial interest in any materials discussed in this article.

Supported by the Wellcome Trust (grant no.: 206619/Z/17/Z); Fight for Sight UK; Thomas Pocklington Trust; NIHR Biomedical Research Centre at Moorfields Eye Hospital and the UCL Institute of Ophthalmology; Retinitis Pigmentosa Fighting Blindness; The Macular Society UK; and Foundation Fighting Blindness USA. The funding organizations had no role in the design or conduct of this research.

HUMAN SUBJECTS: Human subjects were included in this study. This study had research ethics committee approval. All research adhered to the tenets of the Declaration of Helsinki. All participants provided informed consent.

No animal subjects were used in this study.

Author Contributions:
Conception and design: Mahroo, Khan, Webster
Analysis and interpretation: Mahroo, Khan, Robson, Tufail, Michae-
ides, Quinlivan, Webster
Data collection: Mahroo, Khan, Wright, Oekrim, Scallo, Tufail, Michae-
ides, Quinlivan, Webster
Obtained funding: N/A
Overall responsibility: Mahroo, Khan, Webster

Correspondence:
Omar A. Mahroo, PhD, FRCOphth, Moorfields Eye Hospital, 162 City
Road, London EC1V 2PD, United Kingdom. E-mail: o.mahroo@ucl.ac.uk.

References

1. Lucia A, Nogales-Gadea G, Pérez M, et al. McArdle disease: what do neurologists need to know? Nat Clin Pract Neurol. 2008;4(10):568–577.
2. Leonardy NJ, Harbin RL, Sternberg Jr P. Pattern dystrophy of the retinal pigment epithelium in a patient with McArdle’s disease. Am J Ophthalmol. 1988;106(6):741–742.
3. Casalino G, Chan W, McAvoY C, et al. Multimodal imaging of posterior ocular involvement in McArdle’s disease. Clin Exp Ophthal. 2018;101(3):412–415.
4. Alsberge JB, Chen JJ, Zaidi AA, Fu AD. Retinal dystrophy in a patient with McArdle disease. Retin Cases Brief Rep. 2018 Aug 1. https://doi.org/10.1097/ICB.0000000000000790 [Epub ahead of print].
5. Senanayake P, Calabro A, Hu JG, et al. Glucose utilization by the retinal pigment epithelium: evidence for rapid uptake and storage in glycogen, followed by glycogen utilization. Exp Eye Res. 2006;83:235–246.

Telemetric Intraocular Pressure Monitoring after Boston Keratoprosthesis Surgery

The use of the Boston keratoprosthesis Type I (BI-KPro) implantation to restore vision in corneal blindness has considerably increased worldwide. Patients with severe corneal opacification without realistic prognosis for success of a corneal transplant can benefit from this treatment. Secondary glaucoma has been identified as one of the 2 most common complications after BI-KPro surgery occurring on average in almost every third patient and being the most frequent reason for long-term vision loss.

Because of the physical properties of the BI-KPro, measurement of intraocular pressure (IOP) by established methods is not feasible after surgery. Therefore, IOP can be estimated only by finger palpation. This technique is prone to high variability, resulting in an unmet clinical need for new concepts to reliably monitor IOP after keratoprosthesis surgery.

Continuous IOP measurement with an implantable sensor was described decades ago. Despite different approaches, no system has reached clinical applicability until now, mainly because of technical difficulties.

The telemetric ARGOS-IO IOP sensor (Implandata Ophthalmic Products GmbH, Hanover, Germany) combines an implantable IOP sensor ring consisting of a microelectromechanical system application—specific integrated circuit with a handheld reading device to measure IOP (Fig 1). An electromagnetic inductive connection between the coil of the sensor and the activated reader powers the circuit, initiating a pressure reading and enabling telemetric data transfer.

We report the 1-year results of a prospective, open-label, multicenter, single-arm clinical trial aimed to assess the safety, tolerability, and performance of the ARGOS-IO telemetric IOP sensor implanted in eyes undergoing BI-KPro surgery.

This clinical trial was conducted in full accordance with the Declaration of Helsinki (ICH-GCP, ISO14155:2011). All patients provided written informed consent before enrollment. (ClinicalTrials.gov identifier: NCT02945176).

A total of 13 patients with an indication for BI-KPro implantation were successfully screened and initially enrolled, 12 of whom successfully received the implant (Table S1, available at www.aaojournal.org). The surgical approach consisted of the typical open sky approach for BI-KPro implantation. Dependent on adequate capsular support, the sensor was placed in the ciliary sulcus with or without additional suture fixation to the sclera. The pressure sensor was calibrated using direct intracameral manometry in the anterior chamber to measure current IOP as reference.

To assess safety of the implant, the study analyzed all serious adverse events during the first 12 months after implantation, as well as adverse events and severe adverse device events occurring in enrolled patients.

The primary objective to evaluate the performance of the study device was the comparison of telemetric IOP measurement to invasive IOP measurement using intracameral manometry.

Eight of 12 enrolled patients completed the study with the sensor implanted in the eye. One patient voluntarily withdrew from the study with the sensor left in place. In 3 patients, the sensor was explanted, either after dislocation in a patient with aniridia (n = 1) or after necessity for multiple additional ocular surgeries (n = 2). In these 2 cases, the sensor was removed to avoid potential additional complications, not because the sensor device caused or was thought to cause the surgical revision.

Of 168 adverse events, 16 adverse events in 4 patients were rated as possibly related to the medical device by the investigators. Those potential (severe) adverse device events were anterior chamber cells, cystoid macular edema, hypotony, iris adhesion, pigment dispersion, vitritis, increased IOP, and formation of a retroprosthetic membrane (Table S2, available at www.aaojournal.org).

Surgical intracameral manometry was performed in a total of 24 visits in 9 patients. The study protocol entailed manometry in 4 visits in