Assessment of local sensitivity in incomplete retinal-pigment-epithelium and outer retinal atrophy (iRORA) lesions in intermediate age-related macular degeneration (iAMD)

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Lesions of incomplete retinal-pigment-epithelium and outer retinal atrophy (iRORA) are associated with disease progression in age-related macular degeneration (AMD). However, the corresponding functional impact of these precursor lesions is unknown. We present a cross-sectional study of four patients employing clinical-grade MAIA (stimulus size: 0.43°, ~125 µm) and adaptive optics scanning light ophthalmoscope (AOSLO, stimulus size 0.07°, ~20 µm) based microperimetry (MP) to assess the specific impact of iRORA lesions on retinal sensitivity. AOSLO imaging showed overall reduced photoreceptor reflectivity and patches of hyporeflective regions at drusen with interspersed hyperreflective foci in iRORA regions. MAIA-MP yielded an average retinal sensitivity loss of -7.3 ±3.1 dB at iRORA lesions compared to the in-eye control. With AOSLO-MP, the corresponding sensitivity loss was -20.1 ±4.8 dB. We demonstrated that iRORA lesions are associated with a severe impairment in retinal sensitivity. Larger cohort studies will be necessary to validate our findings.
INTRODUCTION

There is an unmet need for prognostic biomarkers and clinical endpoints, approved by regulatory authorities, in early and intermediate age-related macular degeneration (AMD).[1] In recent years, high-resolution multimodal retinal imaging, including spectral domain optical coherence tomography (SD-OCT), has identified precursor lesions of geographic atrophy (GA) in AMD.[2] These have been further characterized by the International Consensus Atrophy Meeting (CAM) group and termed as incomplete retinal-pigment-epithelium and outer retinal atrophy (iRORA) in SD-OCT imaging, preceding its later stage, complete RPE and outer retinal atrophy, cRORA.[3] While iRORA is highly associated with disease progression,[4] detailed knowledge of its impact on retinal function is very limited, yet essential for its validation as a potential clinical outcome measure. Conventional functional tests such as microperimetry (MP) are inadequate due to the discrepancy between available test stimuli sizes and the tiny iRORA lesion sizes.

Nowadays, adaptive optics scanning light ophthalmoscope (AOSLO) imaging allows cell resolved visualization of photoreceptors by correcting the human eyes natural aberrations.[5] AOSLO systems have been further adapted to allow AOSLO-MP, utilizing the high resolution of the system to facilitate functional sensitivity testing of small retinal areas down to the size of individual photoreceptors.[6–8] AOSLO-MP has helped to investigate the functional relevance of retinal phenotypes in various diseases, including Macular telangiectasia type 2, Choroideremia and Retinitis pigmentosa.[9–11]

The purpose of this cross-sectional study is to juxtapose retinal sensitivity, gauged through clinical-grade MP and AOSLO-MP, at retinal locations in presence and absence of iRORA to determine the lesions’ impact on retinal function.
METHODS

Patients

For this study, patients with large sub-RPE drusen (≥125 µm) associated with iAMD according to the Beckman classification [12] and at least one iRORA [3] lesion (region of choroidal hypertransmission <250 µm, zone of attenuation/disruption of the RPE <250 µm, evidence of overlying photoreceptor degeneration in absence of an RPE tear) were recruited. Exclusion criteria were any history of other retinal diseases, glaucoma, previous history of vitreoretinal surgery, relevant anterior segment disease with media opacity, cRORA or a history of current or previous anti-VEGF treatment. If both eyes of a study patient had iRORA, the eye with a more extrafoveal position of the lesion as well as better BCVA was chosen as study eye. All patients underwent routine ophthalmological examination including best-corrected visual acuity (BCVA), slit lamp and funduscopic examination. Following pupil dilation (0.5% tropicamide, 2.5% phenylephrine), a standardized retinal imaging protocol was performed using SD-OCT raster scanning (241 B-scans, 30°×25° enhanced-depth-imaging [EDI] mode, centered on the fovea, lateral scan distance 30 µm, automatic real-time mode 9 frames; Spectralis, Heidelberg Engineering, Heidelberg, Germany). Throughout this study, a retinal magnification factor of 291 µm/deg of visual angle was assumed.

All patients were recruited at the University Eye Hospital Bonn, Germany, and written informed consent was obtained after explanation of the study’s nature and possible consequences. The study was approved by the institutional Ethics committee (#125/14 & #009/13) and was conducted in accordance with the tenets of the Declaration of Helsinki.

iRORA grading

For precise localization of iRORA, each B-scan of the 241-volume SD-OCT scan was screened
by two medical expert graders (MS, LvDE) with specific expertise in iAMD trials. The largest horizontal diameter of the lesion was annotated within the B-scan. For MP testing, in eye control regions were chosen by eccentricity-matching via mirroring along the foveal vertical meridian. Control regions were allowed to present iAMD-typical structural alterations but required to not contain any iRORA lesions or large blood vessels. In case the requirements were not met, the control region was shifted horizontally or vertically to a valid retinal region, which was found in all cases by shifting less than 100 µm from the initially targeted area. The targeted area was generally smaller than 250 µm edge length, centered on the lesion crossing central B-scan. Per eye, a single iRORA lesion and a single in-eye control region was examined with clinical-grade (MAIA-) and AOSLO-MP. Fundus images showing the MP target locations and AOSLO images of the respective regions are shown in Figure 1.

**MAIA-MP**

For clinical-grade retinal sensitivity testing, the S-MAIA device (CenterVue/iCare, Padova, Italy) was employed (settings: 85 stimuli covering 12° of the central retina, 4-2 dB staircase strategy, stimulus size: 0.43° [Goldmann III]). An initial training session was performed in all patients prior to the main test using achromatic stimuli (400–800 nm). The MAIA test duration was about 8-10 minutes. The MAIA background luminance was 4 apostilb (asb, or 1.3 cd/m²) with a dynamic testing range of 36 dB. The MAIA grid stimulation pattern was aligned according to vessel bifurcations to the en-face IR of the SD-OCT image using Fiji, Image J (U.S. National Institutes of Health). Under detailed consideration of the corresponding OCT B-scans, the MAIA stimuli points at the position of the respective iRORA lesion were identified.

**AOSLO-MP**

A custom dual channel confocal AOSLO was used to simultaneously image the retina
(wavelength 840 nm; field of view: 0.85°) and to deliver visual stimuli at 543 nm (test spot diameter: 0.07°).[7] The custom AOSLO-MP instrument has been described previously in detail.[7] In summary, the setup comprised a broadband laser source (SuperK EXTREME; NKT Photonics, Birkerod, Denmark) that was used to provide multiple light channels. The integrated adaptive optics consisted of a Shack-Hartmann wavefront sensor (SHSCam AR-S-150-GE; Optocraft GmbH, Erlangen, Germany) and a deformable mirror (DM97-08; ALPAO, Montbonnot-Saint-Martin, France) in closed loop. Wavefront sensing was performed using the imaging wavelength. Two acousto-optic modulators (TEM-250-50-10-2FP; Brimrose, Sparks Glencoe, MD, USA) in cascaded configuration allowed generation of high-contrast visual stimuli at a wavelength of 543 nm.[7, 8] The 840 nm AOSLO imaging field created a constant background illumination of 13.2 asb (4.2 cd/m²) against which the 543 nm test spots were shown. Visual stimuli could be presented over a 50 dB dynamic range.[7, 8] The position of the patients’ head was controlled using a dental impression stage (bite bar). Patients were instructed to fixate on a small visual annulus presented via a pellicle beam splitter during imaging and testing. A 4-2 dB descending staircase strategy with three threshold crossings was used. Sensitivity thresholds at both iRORA and control test sites were defined in dB as the average from at least 5 valid repeat runs per location (5 at iRORA, 5 at control region). Runs where catch and lapse trials were notable, or where the patient or the supervisor noted down any issue, were excluded from the analyses. Prior to testing, AOSLO images were recorded at the relevant retinal locations. Patients then performed several practice runs to familiarize themselves with the test procedure. While each single AOSLO-MP test run took less than a minute to complete, the total time for AOSLO imaging and AOSLO-MP took about 1 hour per eye.
Statistical Analysis

A one-sided two-sample t-test was carried out using the \texttt{ttest2} Matlab function to test for statistical significance (p<0.05) of AOSLO-MP sensitivity loss in presence and absence of iRORA.
RESULTS

Four eyes of four iAMD patients (mean age: 73, range: 60 - 85) were included. Detailed patient characteristics are given in Table 1. Multimodal imaging of iRORA and control regions is shown in Figure 1. When averaged across all four test eyes, the loss of sensitivity at the iRORA site relative to the control site was 20.1 ±4.8 dB for AOSLO-MP and 7.3 ±3.1 dB for MAIA-MP (Figure 2).

Table 1:

<table>
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<tr>
<th>Patient</th>
<th>Sex</th>
<th>Eye</th>
<th>Lens status</th>
<th>BCVA [logMAR]</th>
<th>iRORA</th>
<th>control</th>
<th>iRORA</th>
<th>control</th>
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<td>OD</td>
<td>Phakic</td>
<td>-0.1</td>
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<td>9.4 ± 1.2</td>
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<td>m</td>
<td>OD</td>
<td>Phakic</td>
<td>0.1</td>
<td>-</td>
<td>-</td>
<td>12.6 ± 4.1</td>
<td>28.4 ± 2.6</td>
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<td>OS</td>
<td>Phakic</td>
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<td>17</td>
<td>27</td>
<td>14.0 ± 3.0</td>
<td>33.3 ± 2.2</td>
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<tr>
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<td>13</td>
<td>21</td>
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</table>

Table 1: Patient ocular characteristics and retinal sensitivity thresholds. If available, averages ± one standard deviation are reported. Abbreviations: BCVA = Best corrected visual acuity, logMAR = Logarithm of the minimum angle of resolution, MP = Microperimetry, dB = Decibel. AOSLO = Adaptive optics scanning light ophthalmoscope. MAIA-MP was not available in patient #3, AOSLO-MP was not available in patient #4.

Selected cases

Patient #1 presented with hyperreflective material and hyporeflective patches at the photoreceptor level at the iRORA lesion, surrounded by structurally altered photoreceptors. The control region showed an irregular photoreceptor mosaic with large patches of slightly hyporeflective photoreceptors (Figure 1). AOSLO-MP revealed a retinal sensitivity threshold of 9.4 dB at the iRORA lesion and of 34.7 dB (25.3 dB difference, p<0.01 two-sample t-test) at the control region (Figure 2). The corresponding MAIA-MP sensitivity difference was 4 dB.
Patient #2 iRORA lesion and control region displayed morphologies similar to Patient #1. AOSLO-MP was successful, showing a significant retinal sensitivity loss between iRORA lesion and control region (15.8 dB, p<0.01), while MAIA-MP failed due to patient fatigue (Table 1, Figure 2).

Patient #3 displayed patches of healthy-appearing photoreceptors surrounded by less reflective and irregularly arranged photoreceptors, as well as regions of hyporeflective photoreceptors, at the location of the iRORA lesion. The respective control region had a similar appearance (Figure 1). AOSLO-MP revealed a sensitivity loss of 19.3 dB between iRORA and control (p<0.01), the MAIA-MP sensitivity loss was 10 dB (Figure 2).

Patient #4 iRORA lesion appeared as a central black area surrounded by remnant enlarged cone structures. AOSLO-MP failed due to insufficient fixation stability of the patient. MAIA-MP showed a significant retinal sensitivity loss between the iRORA lesion and control region (8 dB, p<0.01) Table 1, Figure 2).
DISCUSSION

We here report the immediate impact of iRORA lesions on retinal function in four patients with iAMD. iRORA is a structural precursor for GA development but can now also be linked to functional impairments detected by AOSLO-MP.

The image signal in confocal AOSLO stems from the reflective properties of the photoreceptors’ outer segments. Patches of less reflective and irregularly arranged photoreceptors, as seen in our patients, are thus highly suspicious of functional and structural impairment. Specifically, the displacement of the cells i.e. by drusen, as well damage to the outer segments are likely causes of the observed changes. The photoreceptor mosaic phenotype at iRORA lesions here reported (see Figure 1), seems to be comparable to previously reported structural changes at retinal locations with iAMD drusen (patient #3) or GA (patient #4).[13, 14] Interestingly, while all iRORA lesions showed abnormalities in photoreceptor arrangement and reflectivity, the impact on function differed across patients. It is hypothesized that with iRORA progression, photoreceptors will further deteriorate. However, photoreceptor loss in areas of iRORA cannot be readily quantified and compared to control regions by confocal AOSLO, given the impact of photoreceptor outer segment health changes on this imaging modality.

In all our patients, a significant loss of retinal sensitivity at the iRORA lesions was detectable. While both MAIA-MP and AOSLO-MP results demonstrate functional loss, its magnitude is about 4 times higher in AOSLO-MP, on average. We suggest that this difference is explained by the smaller and finely targeted stimulus available for AOSLO-MP. The ~6 times larger Goldmann 3 stimulus used for MAIA-MP likely includes areas not affected by the small iRORA lesions and elicits retinal responses by comparably healthy adjacent photoreceptors (compare patient #1 and #3). The steep retinal sensitivity loss detected with AOSLO-MP indicates that iRORA lesions can be considered as a surrogate marker for retinal dysfunction. A Limitation of our study is the fundamentally different setup of the MAIA-MP and AOSLO-MP. The AOSLO-
MP background illumination is about 3.3 times brighter due to light requirements of the adaptive optics components used for wavefront correction and retinal imaging. Furthermore, the MAIA-MP uses white stimuli, while green (543 nm center wavelength) stimuli are used for AOSLO-MP, selected for equal sensitivity in the long- and middle-wavelength sensitive cones. Additionally, we have a small shared patient pool. One patient failed the MAIA-MP (patient fatigue), and one failed the ASLO-MP (insufficient fixation stability), resulting in only 2 patients with shared results for both MAIA-MP and AOSLO-MP out of a total of 4 recruited patients.

**CONCLUSIONS**

This pilot study revealed a statistically significant and severe decrease in retinal sensitivity at iRORA lesions in iAMD patients. Larger cohort studies will be necessary to validate our findings.
ACKNOWLEDGEMENT SECTION

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Role of the funder
The funding organization had no role in the design or conduct of the study.

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Competing interests
J. Ameln: none
M. Saßmannshausen: Heidelberg Engineering (F), CenterVue (F), Carl Zeiss MedicTec (F).
L. Von der Emde: Heidelberg Engineering (F), CenterVue (F), Carl Zeiss MedicTec (F).
T. Ach: Bayer (C), Roche (C), Novartis (C), Novartis (R), Heidelberg Engineering (C), Apellis Pharmaceuticals (C), Nidek (C, R).
F.G. Holz: Acucela (C, F), Allergan (F), Apellis (C, F), Bayer (C, F), Boehringer-Ingelheim (C), Bioeq/Formycon (F, C), CenterVue (F), Ellex (F), Roche/Genentech (C, F), Geuder (C, F), Graybug (C), Gyroscope (C), Heidelberg Engineering (C, F), IvericBio (C, F), Kanghong (C, F), LinBioscience (C), NightStarX (F), Novartis (C, F), Optos (F), Oxurion (C), Pixium Vision (C, F), Oxurion (C), Stealth BioTherapeutics (C), Zeiss (F, C).
W. Harmening: none
**Ethics approval**

The study was approved by the institutional Ethics committee (#125/14 & #009/13) of the University of Bonn.

**Contributorship**

All authors conceived the study. FGH, TA and WMH provided the devices used in the study. JA, MS and LvdE conducted the study. JA, MS and LvdE analyzed the results and wrote the manuscript. JA, LvdE and WMH created the figures. All authors discussed the results and revised the manuscript. WMH is the guarantor and responsible for the overall content of the study.

**Prior presentation**

Part of this work has previously been presented at the Classification of Atrophy Meeting (CAM) 2023.
REFERENCES


FIGURES

Figure 1: Multimodal imaging of four iAMD patients presenting iRORA lesions.

Column 1, Infrared SLO images. OCT B-scan and AOSLO imaging locations are marked (red=iRORA, blue=eccentricity-matched control region). Column 2, OCT-B scan through the center of the iRORA lesion. Column 3, magnification of the iRORA lesions marked by the white rectangle in Column 2. Red arrow heads mark the extent of the lesion as graded from the OCT scan. Columns 4 and 5 show AOSLO images of iRORA lesions and control regions, respectively. Arrow heads indicate the corresponding locations after OCT and AOSLO image registration. In three cases, only a single marker is visible due to the lesion extending beyond the limited field of view of the AOSLO.

Figure 2: Retinal sensitivity at iRORA lesions and control regions in iAMD.

A) Test spot size comparison between MAIA-MP (achromatic stimulus) and AOSLO-MP (543 nm stimulus within a larger 840 nm raster). B) MAIA-MP estimate of retinal sensitivity at the iRORA lesion (17 dB, red) and control region (27 dB, blue) in patient #3. The marker diameter corresponds to MAIA-MP stimulus size on the retina. C) and D) AOSLO-MP test spot size and location at the iRORA lesion (C) and control region (D) in patient #3. Sensitivity thresholds were 14 dB and 33.3 dB, respectively. Both areas show patches of normal hexagonally arranged cone photoreceptors and areas of hyporeflective retina. E) Sensitivity thresholds at the iRORA lesions and control regions for MAIA-MP (patients #1, 3, 4) and AOSLO-MP (patients #1, 2, 3).
Figure 1

[Image of Figure 1 showing OCT and AOSLO images for different patients]

- **Patient 1**: SLO (left), OCT (center left), OCT detail (center right), AOSLO iRORA (right middle), AOSLO Control (right)
- **Patient 2**: Similar setup as Patient 1
- **Patient 3**: Similar setup as Patient 1
- **Patient 4**: Similar setup as Patient 1

Scale bars: 1 mm, 500 μm, 200 μm, and 50 μm.
Figure 2

A Test spot sizes

MAIA-MP

125 µm (Goldmann III)

AOSLO-MP

20 µm

250 µm

E

Threshold [dB]

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<th>Patient ID</th>
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<th>AOSLO</th>
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Control

B MAIA-MP

C AOSLO-MP iRORA

D AOSLO-MP Control

low

Sensitivity

high