Autofluorescent hyperreflective foci on infrared autofluorescence adaptive optics ophthalmoscopy in central serous chorioretinopathy

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ABSTRACT

Purpose: To test the hypothesis that hyperreflective foci in central serous chorioretinopathy (CSCR) are auto-fluorescent and may represent macrophages that have engulfed outer retinal fluorophores from the retinal pigment epithelium (RPE) and photoreceptors.

Methods: Enrolled subjects underwent spectral domain and swept-source optical coherence tomography, adaptive optics flood-illumination, and adaptive optics scanning laser ophthalmoscopy (AOSLO), including near-infrared autofluorescence (AO-IRAF). For the AO-IRAF imaging, retinal fluorophores were excited using 795 nm light and imaged using the red-shifted emission spectra. FAF has been quantified as normal (absence of signal) or hyperfluorescent (strong signal). The RPE contains lipofuscin, and melanin that can be excited using different wavelengths (OCT) and visible in simultaneously acquired confocal AOSLO images in active stage. The hyperautofluorescent foci in the patient with active CSCR disappeared coincident with clinical resolution.

Conclusions: We show here the first AO-IRAF images from patients with CSCR, demonstrating hyper-autofluorescent punctate foci, colocalized with hyper-reflective foci on confocal AOSLO images and in OCT. The autofluorescence of these foci may be driven by the accumulation of photoreceptor and RPE fluorophores within macrophages during the active stage of the disease.

1. Introduction

Central serous chorioretinopathy (CSCR) typically affects males between 20 and 50 years old. Neurosensory retinal detachment involving the macular region occurs due to leakage from the choroid through the retinal pigment epithelium (RPE). CSCR pathogenesis is not fully understood, however, it is believed that choroidal abnormalities are the primary underlying pathophysiology. Subretinal fluid (SRF) in CSCR can sometimes resolve spontaneously if diagnosed cases, however, this can lead to atrophy of the retina or RPE and potentially choroidal neovascularization. The most utilized treatment for CSCR is currently photodynamic therapy (PDT) with about 15–30% of patients having recurrence. The diagnosis and workup of CSCR typically involves multi-modal imaging. Structural imaging with optical coherence tomography (OCT) and scanning laser ophthalmoscopy (SLO) can quickly reveal and quantify features such as SRF. More subtle fluid-leakage can be detected with fluorescein angiography (FA). Fundus autofluorescence (FAF) can reveal additional information about the health of the RPE and the regions of interest are often described as hypofluorescent (minimal or absence of signal) or hyperfluorescent (strong signal). The RPE contains lipofuscin, and melanin that can be excited using different wavelengths of light and imaged using the red-shifted emission spectra. FAF has been used extensively to evaluate RPE status in CSCR. However, FAF

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performed on conventional clinical devices lacks the resolution needed to image CSCR at a cellular level in vivo.\(^1\)\(^2\)

Cellular level imaging of the living retina can be achieved with adaptive optics ophthalmoscopy (AOO).\(^3\) AOO utilizes an adaptive optics sub-system to correct for ocular aberrations and can achieve near diffraction-limited images of the living retina.\(^4\) Adaptive optics scanning laser ophthalmoscopy (AOSLO) produces en face images of the retina over a small field-of-view (FOV), typically around 1–2\(^\circ\) with a resolution of few microns that can be stitched together to form larger montages of the retina. AOSLO can implement all the same modalities as SD-OCT including confocal reflectance and autofluorescence but with microscopic details, including individual photoreceptor and RPE cells due to the improved lateral resolution.\(^5\)\(^6\)\(^7\)

OCT demonstrates hyper-reflective foci in CSCR that are considered to be accumulated shed photoreceptor foot plates, secondary to RPE dysfunction.\(^8\)\(^9\) Also, there are hyper-reflective structures in the outer plexiform and outer nuclear layers and around the elongated outer segments of the photoreceptors in CSCR.\(^10\) Recently, Iacono et al. showed that the hyper-reflective foci seen in spectral-domain OCT (SD-OCT) are colocalized with the hyperautofluorescent areas seen in the granular AF phase of the disease using Spectralis.\(^11\) However, these images obtained using commercial devices lacked the resolution to evaluate the fluorescence signal from individual cells.\(^12\)

Hyperreflective clusters on cross-sectional OCT in the outer retina of central serous chorioretinopathy have been imaged using AOSLO reflectance with both confocal and split-detection imaging techniques. These investigations identified three distinct patterns of these clusters and showed that they have a regular granular morphology\(^13\) and that they colocalize with the hyper-reflective foci previously described in SD-OCT. Those OCT hyperreflective dots are associated with longer persistence of the subretinal fluid,\(^14\) poorer best-corrected visual acuity (BCVA), and the need for treatment in CSCR.\(^15\) These OCT hyperreflective clusters have been hypothesized to be either proteins, lipids, macrophages, aggregates, or retinal pigmented epithelial cells.\(^16\)

The aim of this study was to use clinical multi-modal imaging in combination with AOSLO reflectance and autofluorescence to investigate the cellular origin of the hyperreflective foci seen in clinical structural images. We evaluated the microscopic autofluorescence of the hyper-reflective foci by using AO-IRAF. We hypothesized that these hyper-reflective foci could be hyper-autofluorescent on AO-IRAF due to the presence of fluorophores such as lipofuscin and melanin that can constrain the hypotheses regarding their cellular origin.

### Table 1

**OCT imaging parameters.**

| Parameter                  | Patient 1 | Patient 2 | Patient 3 | Active | Resolved |
|----------------------------|-----------|-----------|-----------|--------|----------|
| Size X [pixel]             | 384       | 1024      | 384       |        |          |
| [mm]                       | 5.9       | 5.6       | 4.4       |        |          |
| Size Z [pixel]             | 496       | 496       | 496       |        |          |
| [mm]                       | 1.9       | 1.9       | 1.9       |        |          |
| Scaling X [μm/pixel]       | 5.78      | 5.49      | 11.40     |        |          |
| Scaling Z [μm/pixel]       | 3.87      |           |           |        |          |
| ART Mode (averaged)        | ON (10)   | ON (16)   | ON (10)   | ON (16)|          |
| Quality [dB]               | 27        | 20        | 27        | 18     |          |
| EDI Mode                   | OFF       | ON        | OFF       |        |          |
| # of B-Scans               | 25        | 37        | 25        | 37     |          |
| Pattern Size [\(\mu m\)]  | 20 × 20   | 20 × 15   | 20 × 20   | 15 × 15|          |
| [μm]                       | 5.9 × 5.9 | 5.6 × 4.2 | 5.9 × 5.9 | 4.4 × 4.4|          |
| B-scan sampling [μm]       | 247       | 117       | 122       |        |          |

### Table 2

**Patient characteristics at the time of imaging.**

| Patient | Eye | VA  | CMT (μm) | Description                                |
|---------|-----|-----|----------|-------------------------------------------|
| 1       | OD  | 20/ | 253      | Chronic SRF in superonasal macula          |
|         | M   | 25  |          |                                            |
| 2       | OS  | 20/ | 227      | Resolved SRF after period of chronicity     |
|         | M   | 20  |          |                                            |
| 3       | OS  | 20/ | 232      | Resolved SRF after ICGA-guided half-fluence PDT 2 months prior |
|         | M   | 25  |          |                                            |

M: male/female, VA: visual acuity, CMT: central macular thickness, SRF: subretinal fluid, ICGA: indocyanine green angiography, PDT: photodynamic therapy, OD/OS: right/left eye.

was obtained from each volunteer after the risks and benefits of participation were explained both verbally and in writing. To ensure safe imaging, all light levels were kept below the American National Standard Institute (ANSI) laser safety limits and were calculated in accordance with best practices for multi-wavelength ophthalmic imaging.\(^21\)

Three patients with chronic CSCR were included. Only one eye per patient was studied. Each patient underwent a complete ophthalmic examination that included the measurement of BCVA, a slit-lamp examination, and a fundus examination. Chronic CSCR was defined by persistence of subretinal fluid for more than 4 months, in association with RPE alterations as seen on FAF.

Patients underwent clinical multi-modal imaging with SLO and SD-OCT (Spectralis HRA + OCT, Heidelberg Engineering GmbH, Heidelberg, Germany); relevant scan parameters are presented in Table 1. SLO reflectance images as well as infrared autofluorescence (IRAF) and blue autofluorescence (BAF) images were acquired.

AOO was carried out using both a flood-illumination adaptive optics (FIAO) fundus camera (rtx1-e, Imagine Eyes, Orsay, France) and a research-grade custom-built fluorescence AOSLO that has been described in detail previously.\(^22\) For AOSLO, the confocal reflectance and AO-IRAF channels were used to simultaneously acquire several image sequences of 90-s each across a 1.5\(\circ\) × 1.5\(\circ\) field of view at a frame rate of 30 Hz. A 795 nm centered (16 nm FWHM) super luminescent diode (MS-795-G15, Superlum, Dublin, Ireland) provided confocal illumination and fluorescence excitation while AO-IRAF emission was collected between 814 and 851 nm using a bandpass filter (FF01-832/37, Semrock, USA) in the AF detection channel. Wavefront sensing was carried out using a 909 nm laser diode.

Each 90-s confocal AOSLO video (consisting of 2700 individual images) was desinusoized and registered using strip-wise registration software developed in-house.\(^23\) AO-IRAF images were co-registered to the confocal image and averaged to increase the signal-to-noise ratio (SNR). The averaged AO images were montaged together using Adobe Photoshop (2019, Adobe Systems, Inc., San Jose, CA, USA). The brightness and contrast of individual AO images were manually adjusted to approximately balance these values across the montage.

As a final step, AOSLO and FIAO montages were manually co-registered with clinical images, using vessels and other prominent structural features as landmarks. All modalities were re-scaled to match the scale of the AOSLO images, allowing structures to be compared across modalities. Co-registered Spectralis SLO images were used to determine the precise location of SD-OCT cross sections with respect to the en face modalities.

To quantify the number and size of the hyperautofluorescent foci in the AO-IRAF images, ImageJ was used. A signal intensity profile was plotted (both horizontal and vertical) to which a standard Gaussian function was fitted to obtain the full-width half maximum (FWHM) of the foci.
3. Results

Patient characteristics at the time of imaging are summarized in Table 2.

Fig. 1 shows the imaging findings in patient 1. Near-infrared reflectance showed a granular alteration in reflectivity in the area of chronic SRF. The OCT cross-section showed a small pocket of SRF with disruption of the architecture of the outer retinal bands, including loss of external limiting membrane, ellipsoid and interdigitation zones. A small focus of RPE hypertransmission was also detected.

In FIAO and confocal AOSLO images (Fig. 1c and d), the photoreceptor mosaic was seen in the periphery of the field of view, but was not visible in the area of SRF, giving way to patchy signal with ill-defined boundaries. On AO-IRAF (Fig. 1e), the autofluorescent signal was attenuated in the area of SRF but, within this area, multiple hyper-AF foci were seen (white arrowheads). Approximately 13 foci emitted
The mean FWHM was 11 ± 4 μm.

Fig. 2 illustrates findings in patient 2. A small, shallow, RPE elevation remained without any SRF. The slightly hyper-reflective zone in the FIAO image corresponds to this RPE elevation. However, it is less prominent in the reflectance AOSLO image. The AO-IRAF shows no clear signs of hyper- or hypo-autofluorescence.

Fig. 3 shows the multimodal imaging of patient 3 with recurrent CSCR. Near-infrared fundus autofluorescence performed on the commercial SLO showed multiple central heterogeneous hyper-autofluorescent foci (not shown in Fig. 3). AO-IRAF revealed approximately 45 hyper-autofluorescent granular foci localized in the outer retina, regularly spaced from one another. The FWHM of these foci was measured to be 17 ± 4 μm, with an average hyperautofluorescent area of approximately 350 μm². These AF foci co-localized with the hyperreflective foci in the AOSLO reflectance image. The area without fluid in the superonasal part of the macula did not show any specific pattern. The FIAO image showed multiple hyper-reflective foci in the posterior pole but no more granular pattern either in the fundus or AO-IRAF.

4. Discussion

In this study, we used AOO including AO-IRAF to study the OCT-hyperreflective foci of CSCR, previously studied by AOSLO reflectance, to test the hypotheses that these foci are hyperautofluorescent. We reproduced previously reported findings that these foci appear as granular clusters on AOSLO reflectance, and we noted their presence only in eyes with SRF, and only in areas of SRF.

In eyes with active CSCR, the hyperreflective granular clusters were also hyperautofluorescent on AO-IRAF. They were not detectable by AO-IRAF, however, in eyes or regions without SRF, specifically patient 2 with inactive CSCR and patient 3 with resolved SRF after PDT. This could be attributed either to the absence of the autofluorescence sources due to structural changes accompanying the resolution of SRF or to the absence in these regions of a focal plane between the strong signal from the RPE and the discrete one from the outer retina. It is also possible that due to significant SRF, and the fact that the excitation beam is focused using photoreceptors as a guide, there is more uncertainty where the focal plane is situated, which alters our ability to acquire the AF signal. In patient 2 with inactive CSCR, the visualization of the photoreceptor
mosaic in the AO-IRAF image may have been due to the waveguiding properties of photoreceptors imposing the photoreceptor mosaic structure on the RPE AF signal. We believe this to be the case because i) photoreceptors are not known to be autofluorescent in healthy subjects, ii) we have observed this phenomenon in previous experiments, and iii) photoreceptors are not known to be autofluorescent in healthy subjects, suggesting that the IRAF signal is an indicator of cellular origin. We believe this to be the case because i) the IRAF signal is an indicator of cellular origin, ii) other groups have described similar phenomena in previous experiments, and iii) the IRAF signal is an indicator of cellular origin. Other limitations of the present study include the small sample size, lack of previous treatment history, and lack of follow-up on the SRF case (shown in Fig. 1). These shortcomings will be addressed in future study of a larger sample size using the “healthy” fellow eye as control.

Macrophages can originate from the activation of retinal microglia or by infiltration of monocytes into the retina from the retinal vessels, or choroid, and migrate into the subretinal space in several retinal disorders such as age-related macular degeneration (AMD), retinal detachment, and diabetic retinopathy. Microglial activation induces local proliferation, migration, secretion of cytokines, chemokines and neurotoxins, and enhanced phagocytosis. In a study of photoreceptor clearance by microglia, Hisatomi et al. defined two types of infiltrating cells in the subretinal space: a cell of c. 10 μm diameter with lysosomes and finger-like processes, and a cell of c. 15 μm diameter containing melanin granules. It is therefore possible that the IRAF structures observed in chronic CSCR represent activated microglia on infiltrating monocytes involved in the phagocytosis of photoreceptor outer segments.

Retinal pigmented epithelial cells are also the primary source of autofluorescence in the retina, including melanin that resides mainly in the apical processes of the cell. When injected into the vitreous under experimental conditions, RPE cells seem to undergo phenotypic and behavioral transformation and adopt macrophage-like characteristics. Phenotypic transformation is also observed in degenerative disease of the macula and neurosensory detachments. Given that melanin is a likely source for the OSLO IARAF findings in this study, it is also possible that the OCT-hyperreflective foci under study might be transformed migratory RPE cells playing a role in phagocytosing photoreceptor outer segments shed by detached retina.

5. Conclusion

This paper presents, to our knowledge, the first AO-IRAF images of the hyperreflective foci commonly seen by OCT in CSCR. These hyperreflective foci were hyper-autofluorescent only in eyes with SRF and only in the areas of SRF. While we cannot yet conclude the cellular basis for this IRAF signal, candidate cells could include resident macrophages or RPE cells with a modified phenotype, providing the basis for subsequent experiments and further investigation. Additional studies involving donated eyes are necessary to confirm the presence of photoreceptor fragments in macrophages.

Patient consent

Written consent was obtained from all subjects to publish AO and clinical images. The report does not contain any personal information that could lead to the identification of the patient.

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Authorship

All authors attest that they meet the current ICMJI criteria for Authorship.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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