Adaptive optics imaging in the most common inherited retinal degenerations

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ABSTRACT
Adaptive optics is the imaging technique of the retina. It is applied in visualizing microscopic changes in the retinal photoreceptors and the retinal blood vessels. Introducing adaptive optics as a diagnostic tool facilitated diagnosis of inherited retinal diseases, such as Stargardt disease, cone-rod dystrophy, and retinitis pigmentosa. These diseases are often inherited in heterogenous patterns. Thus, genetic testing often fails to give a certain diagnosis. Optical coherence tomography, fundus autofluorescence, fluorescein angiography, visual field examination and electrophysiological testing are vital for diagnosing patients with inherited retinal diseases. Adaptive optics imaging complements diagnosis and monitoring of conditions such as Stargardt disease and cone-rod dystrophy. In this article we sum up the main characteristics of inherited retinal diseases in adaptive optics imaging.

KEY WORDS: adaptive optics, Stargardt disease, cone-rod dystrophy, retinitis pigmentosa.

INTRODUCTION AND BASICS OF ADAPTIVE OPTICS
Adaptive optics (AO) is a powerful tool in in vivo imaging of the microstructures of the retina. This imaging technique was primarily developed in astronomical telescopes in order to reduce atmospheric irregularities. The aberration-correcting system provided a high-quality image of distant objects [1].

Attempts to eliminate the optical aberrations of the eye were started by Dreher et al. [2]. The first use of AO in imaging the microstructures of the retina was reported in 1997 [3]. There are two main technologies of adaptive optics used in visualizing the retina’s photoreceptors: split-detector (split-detector adaptive optics scanning light ophthalmoscopy – SD-AOSLO) and confocal (confocal scanning laser ophthalmoscopy – cSLO) – both confocal and spectral images may be taken simultaneously [4, 5].

The classical retinal imaging adaptive optics camera consists of three main pieces: a wavefront sensor, a wavefront corrector and a control system. The aberrations are measured by the wavefront sensor and the corrector, whereas the controller interprets the sensor-collected data and controls the interaction between the sensor and corrector. There are two sources of light: one of them used to illuminate the retina and the second to measure and correct the wavefront aberrations [1, 4]. Lately, improved sensorless adaptive optics technology with enhanced quality parameters has been introduced [6]. The imaging is non-invasive and safe. It provides the resolution of 2 µm. It may be performed with or without the patient’s pupil dilation; however, the pupil dilation is usually performed before the examination. The combination of AO and optical coherence tomography provides even greater resolution of the image and enables 3D visualization [6, 7].

PRINCIPLES OF ADAPTIVE OPTICS IN OPHTHALMOLOGY
One of the main goals of AO in ophthalmology is visualization of the photoreceptors: rods and cones. The photoreceptors’ parameters examined in AOSLO are: cone density, cone spacing, Voronoi analysis, reflectivity, regularity, metrics for the preferred orientation of cones and local spatial anisotropy [5, 7-12]. The regions of the photoreceptor loss allow visualization of the lying underneath retinal pigment epithelium (RPE) cells [13, 14]. Photoreceptor pathologies are found in e.g. diabetic retinopathy (DM) and age-related macular degeneration (AMD) [15, 16]. Figure 1 shows an adaptive optics image of photoreceptors in a normal retina.

Adaptive optics non-invasively provide a precise image of the retinal microvasculature. Early changes in the microcirculation, e.g. in the course of DM and AMD, are detected in AO imaging [4, 15]. The imaging of the retinal nerve fiber...
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CHARACTERISTICS OF THE INHERITED RETINAL DISEASES: STARGARDT DISEASE, CONE-ROD DYSTROPHY, RETINITIS PIGMENTOSA

Stargardt disease

Stargardt disease is one of the most common inherited retinal diseases, with a prevalence of around 1 in 10,000 [5, 17]. It is a bilateral dystrophy which affects mainly the macula. It leads to loss of central vision and dyschromatopsia. The dynamics of vision deterioration differs individually depending on the location of the foveal lesion [18]. The presence of central scotoma is seen in visual field examination, microperimetry being used for monitoring the progression of the disease [19]. The macular abnormalities in Stargardt disease are shown in Figure 2.

Stargardt maculopathy is a genetic condition in most cases associated with mutations in the \( \text{ABCA4} \) gene (MIM601691) coding a protein located in outer segments of the photoreceptors. More than 200 pathogenic variants of \( \text{ABCA4} \) mutations leading to Stargardt disease have been identified while the three most common mutations are responsible for only 27% of the cases [20]. SD-OCT (spectral-domain optical coherence tomography) shows changes in the central macula such as loss of outer retina structure. The dynamics of Stargardt disease progression can be defined using SD-OCT through ellipsoid zone loss evaluation [21].

Fundus autofluorescence (FAF) shows loss of central autofluorescence in the central macula and an increased signal in surrounding regions resembling bulls-eye maculopathy. With the use of FAF testing we can monitor progression of the morphological changes of the macula. The mean progression of definitely decreased autofluorescence lesions is determined as 0.51-0.76 mm\(^2\)/year, and of total decreased FAF as 0.35 mm\(^2\)/year [22, 23]. According to the FAF image characteristics of the fovea (including foveal sparing) and homogeneity of the background, Stargardt disease eyes may be divided into three subtypes. The RAE (rate of atrophy enlargement) differs between these three subtypes [24].

Fluorescein angiography (FFA), used more commonly in the past, showed a “dark choroid” sign in the central macula due to blocking of the choroid signal by lipofuscin deposits in the central macula. The image from electrophysiological tests, such as multifocal electroretinography (mfERG), shows decreased macular function with a preserved peripheral signal [5, 17]. An FFA image of Stargardt disease is shown in Figure 3.

**Characteristics of adaptive optics imaging in Stargardt disease**

Spectral as well as confocal AO imaging is a potent tool in imaging pathologies in photoreceptor morphology of eyes with Stargardt disease. The cone spacing is increased and cone density is decreased in Stargardt subjects’ retina compared to the normal retina [5, 17, 25], as shown in Figure 4. The confocal AO images show “dark spaces” within photoreceptors’...
structure which represent non-wave guiding cones visualized in SD-AOSLO. The AOSLO changes may be found before the OCT and FAF detectable changes [25].

The introduction of SD-AOSLO technique revealed that the cone density in Stargardt disease is in fact higher than was expected based on confocal AOSLO imaging and improves the quality of the imaging, making it more accurate in IRDs [5, 17].

There is a correlation between increased autofluorescence and loss of photoreceptor cells. It has been found however that AOSLO image reflectivity does not correlate with lipofuscin accumulation. AOSLO images show a hyperreflective perifoveal ring and its margin marks the margin of the scotoma [26].

Adaptive optics encounters some difficulties in Stargardt disease evaluation. Central scotoma and poor fixation make it unable to examine some eyes thoroughly. The accumulation of lipofuscin prevents AOSLO photoreceptor visualization in certain areas [27]. The cone spacing and density differ between the examined areas within the same eye and there has not been established a universal method to evaluate the degree of cone atrophy [26]. A possible solution may be creating cone density deviation maps (CDD) that present variation in foveal structure and may be compared with OCT and microperimetry results [28].

Adaptive optics provides visualization of the photoreceptors’ morphology in Stargardt disease, which complements FAF visualization of lipofuscin accumulation and OCT cross-section of the retinal layers [27].

Cone-rod dystrophy
Cone-rod dystrophy (CRD) is a retinal dystrophy where the dysfunction is higher in cones than in rods. The prevalence ranges from 1/30,000-1/40,000 [29]. It is genetically polymorphic, with multiple variants of inheritance [30, 31]. The eye fundus examination shows perifoveal atrophy and “bull’s eye” appearance (Figure 5). The changes include loss of RPE, photoreceptor loss and lipofuscin accumulation. The symptoms include photoaversion, progressive visual acuity deterioration and poor color vision due to cone dysfunction. In advanced stages, CRD leads to bilateral blindness [29, 31]. The diagnostic process of CRD includes genetic testing, OCT, ERG and FAF. The scotopic and photopic ERG cone and rod responses are extinguished in CRD [31].

Characteristics of adaptive optics imaging in cone-rod dystrophy

The photoreceptors’ AO imaging in CRD seen in AOSLO show disruption of its mosaic and increased cone spacing and rod spacing, as seen in Figure 6. The level of the changes increases with the age of the subject. The photoreceptors in CRD show higher spacing, lower density, and lack of waveguiding cones within atrophic regions. In the lowered-density regions, cone spacing measurements correlate with mfERG, microperimetry results and visual acuity changes [7, 13, 30, 32, 33].

The AOSLO imaging of the nine confirmed autosomal-dominant cone-rod dystrophy (AD-CRD) patients from the

Figure 4. Stargardt disease: adaptive optics image of the perifoveal photoreceptors. Note the dark spaces and decreased cone density

Figure 5. Cone-rod dystrophy: photograph of the posterior pole

Figure 6. Cone-rod dystrophy: adaptive optics image of the perifoveal photoreceptors. Note the decreased cone density
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same family showed that the pattern of photoreceptor damage varies between the subjects with the same type of mutation, regardless of age [34].

**Retinitis pigmentosa**

Retinitis pigmentosa (RP) is an inherited progressive bilateral degeneration of the photoreceptors. Like other IRDs mentioned above, the disease may be caused by mutations in multiple genes, with a heterogeneous inheritance pattern [35]. The loss of rods precedes the loss of cones; the changes result in reduced visual acuity, constricted visual field and, eventually, blindness [11, 32, 36, 37]. Retinitis pigmentosa may be isolated or a part of another syndrome, e.g. Usher syndrome, where sensorineural hearing loss is also present. The characteristics of RP in the ophthalmoscopic examination include presence of bone cells, which starts in the outer regions of the retina and proceeds to the macular region in more advanced stages, blood vessel narrowing, optic nerve atrophy and macular changes: signs of dystrophy or tapetal macular reflex. An eye fundus image of RP is shown in Figure 7. The phenotype differs even among subjects with the same type of mutation [38].

Visual field examination shows progressive narrowing of the visual field. SD-OCT and time-domain OCT are used to visualize the photoreceptors layers. It shows thinning or loss of outer retinal layers throughout the macula and hyper-reflective profiles in the subretinal space at the fovea. However, it lacks resolution for singular photoreceptor visualization and evaluation [36, 38]. The ERG responses are distinguished, in more advanced stages below the level of test reliability [38].

Changes in the macula in retinitis pigmentosa eyes are not uncommon. Diseases such as epiretinal membrane and macular hole may be found due to ischemia of macular RPE as well as changes in the vitreous body structure. Such conditions may be treated surgically [37, 39].

**Characteristics of adaptive optics imaging in retinitis pigmentosa**

Cone loss is observed in RP as well as Usher syndrome. The cone spacing in RP eyes is higher than in healthy eyes [14, 32, 35, 38]. Regions of photoreceptor atrophy and hyper-reflective structures in outer retinal layers are visualized [38], as seen in Figure 8.

The interpretation of the images of the photoreceptors in the retinitis pigmentosa shows a high level of variability between the interpreters due to the cone loss areas in the retina. The image interpretation quality seems to be higher in SD-AOSLO compared to confocal AO [5, 40].

Foveal visual acuity loss and foveal sensitivity changes in retinitis pigmentosa are noted when morphological changes in the retina are already in an advanced stage [41].

The efforts to determine the rate of disease progression with AOSLO seem to be successful – it has been shown to be consistent with the progression in SD-OCT and microperimetry findings [36] and to be a successful tool in monitoring the disease progression during the experimental treatment [11].

**USE OF ADAPTIVE OPTICS IN OTHER INHERITED RETINAL DEGENERATIONS**

The AOSLO images of patients with Bornholm eye disease (BED), an X-linked cone dysfunction, proves the disruption of the cone mosaic with a high level of intersubject variability between the patients with different disease-causing mutations [42].

Achromatopsia (ACHM) is also a subject of AO retinal studies. Split-detector optics demonstrates a decreased number of photoreceptors as well as variations in cone density maps between the regions within the retina and between the eyes of ACHM subjects [9, 43].

The AO studies of patients with occult macular dystrophy (OMD) show regions of loss of photoreceptors, RPE visualization and lower cone density than in healthy eyes and a correlation of the morphological changes with the microperimetry [44-46].

![Figure 7. Retinitis pigmentosa: photograph of the posterior pole](image1)

![Figure 8. Retinitis pigmentosa: adaptive optics image of the perifoveal photoreceptors. Note the dark spaces, decreased cone density and hyper-reflective structures](image2)
LIMITATIONS

The challenges of adaptive optics include obtaining good-quality images in poor fixation, as well as in nystagmus – the eye tracking devices overcome those difficulties [47–49].

The interpretation of the images is also prone to mistakes made by non-experienced examiners. The machine learning of AO interpretation is vital to establish objective interpretation and forming right conclusions.

The need for larger study cohorts in IRDs is emphasized [50].

SUMMARY

Numerous ophthalmic researchers have emphasized the role of longitudinal studies of patients with inherited retinal diseases. The research is limited by the low prevalence and difficult diagnostic process in some cases. Classical and well-known techniques, such as OCT, fundus autofluorescence, fluorescein angiography, and electroretinography used in diagnosing and monitoring the IRD patients play a vital role.

With the course of scientific development, the role of novel ophthalmic diagnostic solutions is becoming more significant. Adaptive optics imaging gives hopes for better understanding of the morphological basics and natural course of retinal diseases and with the introduction of experimental therapies also for monitoring the effectiveness of introduced treatment. Being able to monitor the rare retinal conditions is crucial for the further discovery and development of medical solutions.

DISCLOSURE

The authors declare no conflict of interest.

References