Age related macular degeneration (AMD) is the most common cause of vision loss in people over 50 years of age. In non-exudative AMD, loss of vision is caused by atrophy of the retinal pigment epithelium (RPE), outer retina, and choriocapillaris. In the exudative type of AMD, ingrowth of new vessels from the choroid into the subretina and/or sub-RPE space results in vision loss.

The survival of the photoreceptors cells is critically dependent on the normal function of the RPE cells. The accumulation of extracellular deposits (drusen) between the RPE and the choroid is known to occur in AMD. The location, composition, and structure of drusen associated with ageing and AMD have been studied. Moreover, the RPE overlying drusen have altered lipofuscin autofluorescence, suggestive of additional changes in RPE metabolism due to AMD. Furthermore, clinicopathological studies have demonstrated thinning of the retina related to atrophy of retinal photoreceptors. Therefore, documentation of topographic changes of the choriotreital surface and quantitative assessment of retinal thickness may be helpful in identifying and monitoring the presence of abnormalities of the RPE.

Scanning laser ophthalmoscopy (SLO) has been used to map the topography of the retinal surface. The technique of optical coherence tomography (OCT) has been used to provide high resolution cross sectional images of the retina and permit visualisation of abnormalities in the retinal structure. The retinal thickness analyser (RTA) has been used to map the retinal thickness in the posterior pole. A prototype version of the RTA has been applied to map the topography of the retinal surfaces, in addition to mapping the thickness of the retina.

In the present study, a prototype version of the RTA was applied to patients with atrophic AMD to assess pathological alterations in the vitreoretinal and chorioretinal surface heights and the retinal thickness and determine their relation to visual acuity and RPE hypopigmentation.

METHODS

The system for retinal topography and thickness mapping is based on a prototype version of the commercially available RTA and has been described previously. The system generates an optical section image of the retina by projecting a laser onto the retina at an angle and viewing the reflection and/or scattering of the laser light from the vitreoretinal and chorioretinal interfaces. The vitreoretinal and chorioretinal interfaces correspond to layers close to the inner limiting membrane and retinal pigment epithelium, respectively. By scanning the laser beam, a series of 20 optical section images encompassing a 2 mm x 2 mm retinal area was acquired in 330 ms. The digitised images were analysed to determine retinal elevations and thickness by a dedicated software algorithm, as described in detail previously. Two two dimensional arrays containing the topographic data for the vitreoretinal and chorioretinal surfaces were created. Since retinal thickness is defined as the depth separation between the vitreoretinal and chorioretinal surfaces, the system for retinal topography and thickness mapping is high resolution cross sectional images of the retina and permits visualisation of abnormalities in the retinal structure. The technique of optical coherence tomography (OCT) has been used to provide high resolution cross sectional images of the retina and permit visualisation of abnormalities in the retinal structure. The retinal thickness analyser (RTA) has been used to map the retinal thickness in the posterior pole. A prototype version of the RTA has been applied to map the topography of the retinal surfaces, in addition to mapping the thickness of the retina.

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Abbreviations: AMD, age related macular degeneration; OCT, optical coherence tomography; RPE, retinal pigment epithelium; RTA, retinal thickness analyser; SLO, scanning laser ophthalmoscopy
Pearson correlation was performed to determine the correlation coefficient and significance level of the association between the topographic variations, retinal thickness, and visual acuity of the retinal area that was used for fixation.

The presence of RPE hypopigmentation, excluding drusen, was established by inspection of colour fundus photographs of the eyes. In eyes with RPE hypopigmentation, the relative chorioretinal and vitreoretinal surface height was determined by averaging surface heights in the area of RPE hypopigmentation and subtracting the 10th percentile height measurements in the surrounding 2 × 2 mm retinal area. Similarly, in eyes without RPE hypopigmentation, relative surface heights were also determined in areas comparable in size to hypopigmented areas. The 10th percentile height measurement was used as baseline rather than minimum height in order to eliminate outliers that may occur due to single point noise related artefacts. Statistical t test (unpaired) was applied to determine the significance level of the difference in relative surface height and thickness in eyes with and without RPE hypopigmentation.

Subjects
Patients diagnosed with non-exudative AMD were referred from the vitreoretinal service at the University of Illinois at Chicago (UIC). Approval for use of human subjects was obtained from the institutional review board of the UIC. Patients with clear media, pupil dilatation of 5 mm or more, and visual acuities of 20/200 or better were included in the study. The study was explained to the patients, and signed consent forms were obtained. Imaging was performed in 22 white patients, 12 females and 10 males, 10 left and 12 right eyes. The patients’ mean age was 74 (SD 8) years and the best corrected visual acuity was 0.5 (0.3) in logarithm of minimum angle of resolution (logMAR) units (20/90 (20/68) Snellen).

The eye was dilated with 2.5% phenylephrine hydrochloride and 1% tropicamide. Fundus examination was performed using direct or indirect ophthalmoscopy and slit lamp biomicroscopy with a 78 dioptre precorneal, 90 dioptre precorneal, or Goldmann contact lens. The patient was seated in front of the modified slit lamp biomicroscope, and an internal fixation target was presented for viewing during optical section imaging. Imaging was performed at nine contiguous retinal areas.
covering a 20 degree field centred on fixation. After optical section imaging, a colour fundus photograph of the eye was obtained using a clinical fundus camera. The patients were classified in two groups based on slit lamp examination of the fundus and evaluation of colour fundus photographs of a 30 degree field centred on the fovea and according to a previously published study.\(^2\) Group 1 (n = 11) included patients with multiple drusen, areas of increased pigmentation or hyperpigmentation associated with drusen, areas of depigmentation or hypopigmentation of the RPE, and presence of multiple foci of atrophy less than 175 \(\mu\)m in diameter. Group 2 (n = 11) included patients with areas of hypopigmentation or depigmentation or apparent absence of RPE that were more than 175 \(\mu\)m in diameter, in which choroidal vessels were more visible than in surrounding areas.

**RESULTS**

An example of imaging performed in the left eye of a 75 year old patient with atrophic AMD is presented (Fig 1). A box overlaid on the fundus photograph indicates the area of imaging. Optical section images were analysed to map the topography of the vitreoretinal and chorioretinal surfaces and the thickness of the retina. The vitreoretinal surface displayed a depression corresponding to the area of reduced thickness. The chorioretinal surface showed topographic variations.

The relation between the topographic variations, retinal thickness, and visual acuity is shown (Fig 2). Variation in the vitreoretinal surface height was moderately correlated with visual acuity \((r = -0.4; p = 0.03; n = 22)\). Increase in variation of chorioretinal surface height was correlated with decrease in visual acuity \((r = -0.5; p = 0.01; n = 22)\). The retinal thickness was not associated with visual acuity \((r = 0.2; p = 0.2; n = 22)\).

The average relative height of the vitreoretinal and chorioretinal surfaces and thickness of the retina in eyes with and without RPE hypopigmentation is shown (Fig 3). Relative height of the vitreoretinal surface in eyes with RPE hypopigmentation was significantly less than eyes without RPE hypopigmentation \((p = 0.005)\). Relative height of the chorioretinal surface was similar in eyes with and without RPE hypopigmentation \((p = 0.4)\). Retinal thickness in eyes with RPE hypopigmentation was less than eyes without RPE hypopigmentation \((p = 0.04)\).

**DISCUSSION**

Since loss of vision in AMD is attributed in part to anatomical changes in retinal and subretinal tissues, the ability to quantitatively document topographic and thickness variations in these tissues may improve the diagnosis, monitoring, and management of this disease. The clinical method of slit lamp biomicroscopy is subjective and has limited resolution for evaluation of fine details in retinal structures. Slit lamp biomicroscopy provides information on retinal structural abnormalities primarily by comparison with adjacent locations and does not allow quantification or documentation of retinal thickness. Additionally, elevations of the chorioretinal surface are often difficult to appreciate on slit lamp biomicroscopy. While fluorescein angiography is useful for delineating areas of geographic atrophy due to the hypopigmentation of the RPE, it does not provide quantitative evaluation of the associated changes in the overlying retinal tissue.

Scanning laser ophthalmoscopy is inadequate for mapping of the chorioretinal surface and retinal thickness in absolute units, because of its limited depth resolution.\(^2\) Optical coherence tomography allows high resolution cross sectional retinal imaging and visualisation of intraretinal structures.\(^3\) In contrast, our system provides in only 330 ms, 20 serial optical section images encompassing a 2 mm \(\times\) 2 mm retinal area, which are analysed to generate maps of the vitreoretinal and chorioretinal surface height and retinal thickness with spatial resolution of 100 \(\mu\)m. Since the analysis algorithm detects the peak locations corresponding to the retinal interfaces, changes in the reflection and scatter of light due to media opacities that result in broadening of the lines, do not significantly affect the measurements. In the presence of severe media opacities, the peak corresponding to the vitreoretinal interface may not be identified. Likewise, in the presence of subretinal fluid or blood, the imaging of the chorioretinal interface may be limited. However, these factors did not limit the interpretation of the data that was obtained in the current study. Since a significant reduction in RPE pigmentation will result in an indistinguishable chorioretinal interface, the image analysis algorithm identified indistinct peaks as artefacts and excluded such data from the thickness maps.

Progressive vision loss due to geographic atrophy, the advanced form of atrophic AMD, occurs in 3.5% of people aged 75 years and over in the United States.\(^6\) Enlargement of areas of atrophy and visual acuity loss have been associated in the geographic atrophy form of AMD.\(^6\) Additionally, increased fundus autofluorescence has been shown to be present in areas of geographic atrophy of RPE associated with AMD.\(^7\) We expected decreased thickness at the fovea to be correlated with reduced visual acuity, based on the premise that abnormalities in the subfoveal pigment epithelium would result in roughly proportionate changes in the foveal thickness and visual acuity. However, we found a lack of association between thickness and visual acuity in the present study. One explanation for this finding is that since patients with subfoveal atrophy might prefer to fixate with extrafoveal sites, we would be correlating thickness with acuity of extrafoveal location. Some of these extrafoveal sites tend to be less involved in the atrophic process and have better visual acuity than the diseased fovea, thus weakening the correlation.
between thickness and visual acuity. Additionally, decreased visual acuity in these patients is attributed to the combined effects of pathological loss of photoreceptor function and the use of extrafoveal fixation. Furthermore, the reduction in the retinal thickness due to partial photoreceptor cell layer atrophy, undetectable by our technique, may have contributed to the lack of correlation between thickness and visual acuity. While, in the current study, we did not determine the precise fixation location of the patients, methods such as SLO perimetry are available that can be used in future studies. The findings further imply that in atrophic AMD, changes in the height of the chorioretinal surface that may be attributed to changes in the retinal pigment epithelium and Bruch’s membrane are related to the visual acuity loss. Future prospective studies will be required to establish the value of assessment of chorioretinal topographic variations in prediction of visual acuity loss.

It is hypothesised that RPE dysfunction leads to accumulation of intracellular material in the RPE and extracellular material in the form of basal laminar and linear deposit, thus changing the composition of Bruch’s membrane and its permeability to nutrients. In the areas with excessive accumulated extracellular debris, one response to the metabolic distress may be RPE atrophy, followed by choriocapillaris ischemia and photoreceptor atrophy.9 In the present study, we found that eyes with areas of RPE hypopigmentation had significantly less elevated vitreoretinal surface and lower retinal thickness when compared to eyes without RPE hypopigmentation. The depression on the vitreoretinal surface was found to be larger than the chorioretinal surface, suggesting atrophy of the photoreceptor layer and associated intraretinal layers as the result of RPE loss. This result is in accord with findings on histopathology of geographic atrophy of the RPE that demonstrated a much greater alteration of the contour of the vitreoretinal interface, due to loss of rods and cones, outer nuclear layer, and overall retinal thickness, than that of the chorioretinal interface.10 Overall, mapping of retinal topography and thickness provides objective and quantitative measurements of retinal structural abnormalities and shows promise as an adjunct for the evaluation of retinal structural changes caused by AMD.

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