Association of Visual Function Measures with Drusen Volume in Early Stages of Age-Related Macular Degeneration

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Received: October 28, 2019
Accepted: February 1, 2020
Published: March 30, 2020


PURPOSE. To assess which visual function measures are most strongly associated with overall retinal drusen volume in age-related macular degeneration (AMD).

METHODS. A total of 100 eyes (16 eyes with early AMD, 62 eyes with intermediate AMD, and 22 eyes from healthy controls) were recruited in this cross-sectional study. All subjects underwent several functional assessments: best-corrected visual acuity (BCVA), low-luminance visual acuity (LLVA), visual acuity (VA) measured with the Moorfields Acuity Chart (MAC-VA), contrast sensitivity with the Pelli–Robson test, reading speed using the International Reading Speed texts, and mesopic and dark-adapted microperimetry. Drusen volume was automatically determined based on optical coherence tomography using an approach based on convolutional neural networks. The relationship between drusen volume and visual function was assessed with linear regressions controlling for confounders.

RESULTS. Mean drusen volume and MAC-VA differed significantly among all AMD stages and controls ($P < 0.001$). In univariate linear regression, LLVA, MAC-VA, contrast sensitivity, and mesopic and dark-adapted microperimetry were significantly negatively associated with the overall drusen volume (all $P < 0.006$). After controlling for AMD stage, age, and the presence of subretinal drusenoid deposits, MAC-VA and mesopic and dark-adapted microperimetry were still significantly associated with drusen volume ($P = 0.008$, $P = 0.023$, and $P = 0.022$, respectively).

CONCLUSIONS. Our results suggest that MAC-VA, as well as mesopic and dark-adapted microperimetry, might indicate structural changes related to drusen volume in early stages of AMD.

Keywords: Automatic segmentation of drusen, drusen volume, age-related macular degeneration, contrast sensitivity

Age-related macular degeneration (AMD) is the leading cause of visual impairment in the elderly in developed countries, with a worldwide prevalence of 8.1% for early AMD and 8.69% for any AMD in people over 45 years of age.1,2 Due to current demographic trends, the burden of AMD is estimated to grow to 288 million by 2040.1 Late stages can lead to a severe loss of visual acuity, whereas early stages of the disease are often not associated with obvious visual symptoms. In conventional visual function tests under high luminance and high contrast, patients with early and intermediate stages of the disease usually achieve good scores; however, they often report difficulties and vision loss under low lighting, low contrast, and changing light conditions.3–5 Therefore, standardized visual function tests under low luminance and low contrast have attracted increasing interest, particularly with regard to early stages of AMD, as they might allow better monitoring of the disease and aid in predicting disease progression.6 Usually the first clinical sign of AMD are drusen located between the basal lamina of the retinal pigment epithelium (RPE) and the inner collagenous layer of Bruch’s membrane (BM), in the sub-RPE-basal laminar space.7,8 Retinal cells overlying drusen exhibit structural and molecular abnormalities indicative of photoreceptor degeneration and Müller glial activation, suggesting that photoreceptor cell function is compromised as a consequence of drusen formation.9 Drusen are among the most important biomarkers for staging AMD.10–15 A recently developed convolutional neural network (CNN)-based approach for a fully automated segmentation of drusen in optical coherence tomography (OCT) images14 allows us to compute the overall drusen volume. Compared to manual segmentation of drusen,
automated segmentation has been shown to be highly reproducible and accurate.\textsuperscript{13,16} The relationship between visual function tests under low luminance and low contrast and measurements of retinal structure associated with AMD progression, such as drusen volume, has not been well described so far. It may be that structural and functional measures provide complementary information about disease status. Thus, we evaluated the relationship between drusen volume and a battery of visual function tests under low luminance and low contrast.

**METHODS**

We conducted a cross-sectional study at the Department of Ophthalmology, University of Bonn, Germany, from January 2017 until January 2019. The study was approved by the Institutional Review Board of the University Bonn (approval ID 013/16). Written informed consent was obtained from all participants following an explanation of all non-invasive tests involved. The protocol followed the tenets of the Declaration of Helsinki.

Sixteen eyes from patients with early AMD, 62 eyes from patients with intermediate AMD (iAMD), and 22 healthy eyes were recruited from patients from the AMD outpatient clinic, the self-help organization Pro Retina, and family members of patients. Participants were categorized as early AMD, iAMD, or healthy controls (no apparent aging changes), based on the classification described so far. It may be that structural and functional characteristics and the lighting conditions in which these tests were administered, we presumed a more cone-mediated function in the BCVA and reading test and a partially rod-mediated function in the LLVA, MAC-VA, contrast sensitivity, and, in particular, mesopic microperimetry.

Prior to microperimetry testing with the S-MAIA device, pupillary dilatation was performed. The S-MAIA performs fundus tracking using a line-scanning laser ophthalmoscope with a super-luminescent diode illumination with a central wave light of 850 nm for mesopic testing and an additional light-emitting diode projecting red (627 nm) stimuli for dark-adapted testing. The dark-adapted testing with red (627 nm) stimuli is more influenced by cone-mediated function, reflecting a mixture of both rod- and cone-mediated responses.\textsuperscript{21} As previously described, a customized stimulus grid was used that consisted of 33 points located at 0°, 1°, 3°, 5°, and 7° from fixation.\textsuperscript{25} First, mesopic testing was performed. Patients were not dark-adapted, but the room light was switched off just before the examination. After mesopic testing, patients underwent 30 minutes of dark adaptation while waiting in the examination room (light was switched off, light level was <0.1 lux), and then dark-adapted testing was performed. The microperimetric outcome measure was the global mean sensitivity (in dB) and mean sensitivity at eccentricities of 0°–1°, 3°, 5°, and 7°. All tests were performed in one eye, with the non-study eye covered with an eye patch.

**Automatic Segmentation of Drusen**

For automated segmentation of drusen, the pipeline reported by Gorgi Zadeh et al.\textsuperscript{14} was used. In the first step, this drusen segmentation pipeline automatically segments RPE and BM bands, using a CNN, which transforms an input B-scan into RPE and BM probability maps. For the final hard segmentation of RPE and BM bands, probability maps are converted into cost maps so that pixels with higher probability have lower costs. Dijkstra’s algorithm is used to find a path with the minimum accumulated costs from the left to the right of each map.\textsuperscript{20} The extracted paths are considered as final RPE and BM band segmentation.

In the second step, an ideal (normal) RPE is estimated through a rectification of RPE and BM bands.\textsuperscript{14} In the rectification step, both RPE and BM are shifted vertically and column-wise until the BM band becomes a straight horizontal line, then a low-degree polynomial is fitted on the shifted RPE band and transformed back into the original image coordinates and is regarded as the drusen-free RPE. Finally, any area between the RPE and drusen-free RPE is classified as drusen. To eliminate falsely detected drusen, those with a height of 2 pixels or less are removed from the final charts and employ a high-contrast, high-pass letter design with a gray background of the same mean luminance as the letters to simulate lower contrast situations.\textsuperscript{19} The letters are also referred to as vanishing optotypes, because, for normal vision, the detection and recognition thresholds are very similar, and the letters seem to disappear soon after the recognition limit has been reached.\textsuperscript{13} Contrast sensitivity was measured using a Pelli–Robson chart presented at a distance of 1 meter.\textsuperscript{20–22} For the IreST, patients wore their best near correction and were asked to read one paragraph aloud while they were timed with a stopwatch.\textsuperscript{23} BCVA and reading speed were performed under photopic conditions, whereas LLVA, MAC-VA, and contrast sensitivity were performed under mesopic conditions. Based on the test characteristcs and the lighting conditions in which these tests were administered, we presumed a more cone-mediated function in the BCVA and reading test and a partially rod-mediated function in the LLVA, MAC-VA, contrast sensitivity, and, in particular, mesopic microperimetry.
segmentation. More details on the automated drusen segmentation pipeline can be found in the Supplementary Material.

**Statistical Analysis**

Descriptive statistics were performed to assess baseline demographic variables for the AMD groups and controls. Due to the sample size, most results were not normally distributed (by the Shapiro–Wilk test), so non-parametric tests were used for analysis. The Kruskal–Wallis test was used for group comparisons. Pairwise differences were calculated using the nonparametric Wilcoxon rank-sum test.

The relationship between drusen volume and demographic variables was assessed with univariate linear regression. In the overall cohort, separate univariate linear regression models against drusen volume were performed for each of the visual functional tests. If the relationship between a functional test and average person measures reached a $P < 0.05$ in univariate analysis, multiple regression was used to ensure that the findings were not confounded by different demographic characteristics across groups. Statistical analyses were performed using the statistical software SPSS Statistics (IBM, Armonk, NY, USA). $P < 0.05$ was considered statistically significant.

**RESULTS**

**Sociodemographic and Clinical Characteristics of the Participants**

The 100 eyes studied included 16 eyes with early AMD (16%), 62 eyes with iAMD (62%), and 22 eyes from healthy controls (22%). The mean age of participants was 67.5 ± 8.1 years, and there were more female participants (66%) than male. The controls were significantly younger than the participants ($P = 0.001$). Women, n (%) 13 (59.1) 11 (68.80) 42 (67.7) Eyes, n 22 16 62 Subretinal drusenoid deposits, n (%) 0 5 (31.3 %) 12 (19.4) Pigment changes, n (%) 0 0 31 (50)

**Table 1. Characteristics of Participants in Each Group**

**AMD Group**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Control</th>
<th>Early</th>
<th>Intermediate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (y), mean ± SD</td>
<td>59.8 ± 6.3</td>
<td>70.0 ± 6.7</td>
<td>69.7 ± 7.3</td>
</tr>
<tr>
<td>Eyes, n</td>
<td>22</td>
<td>16</td>
<td>62</td>
</tr>
<tr>
<td>Women, n (%)</td>
<td>13 (59.1)</td>
<td>11 (68.80)</td>
<td>42 (67.7)</td>
</tr>
<tr>
<td>Subretinal drusenoid deposits, n (%)</td>
<td>0</td>
<td>5 (31.3 %)</td>
<td>12 (19.4)</td>
</tr>
<tr>
<td>Pigment changes, n (%)</td>
<td>0</td>
<td>0</td>
<td>31 (50)</td>
</tr>
</tbody>
</table>

**Figure.** Boxplot showing drusen volume (in mm$^3$) for controls and early and intermediate AMD (excluding outliers). Each boxplot includes the maximum (upper whisker), upper quartile (top of the box), median (horizontal line in box), lower quartile (bottom of the box), and minimum (lower whisker) values.

In univariate linear regression, LLVA, MAC-VA, contrast sensitivity, and mesopic and dark-adapted microperimetry were still significantly decreased in early AMD compared to controls (all $P < 0.05$). BCVA and MAC-VA were also significantly decreased in early AMD compared to controls ($P = 0.016$ and $P = 0.006$, respectively), but there was no significant difference in all other functional tests between the two groups (all $P > 0.05$). When comparing early AMD to iAMD, BCVA and reading speed did not differ significantly ($P = 0.31$ and $P = 0.07$, respectively), but there was a significant decrease in LLVA, MAC-VA, contrast sensitivity, and mesopic and dark-adapted microperimetry in the iAMD group compared with the early AMD group (all $P < 0.05$) (Table 2). Univariate linear regression revealed no significant association between drusen volume and age ($\beta$ coefficient $= 0.001; P = 0.642$).

**Relationship Between Drusen Volume and Visual Function Tests**

In univariate linear regression, LLVA, MAC-VA, contrast sensitivity, and mesopic and dark-adapted microperimetry were significantly negatively associated with the overall drusen volume (all $P < 0.006$) (Table 3). After controlling for AMD stage, age, and the presence of SDD, MAC-VA and global mesopic and dark-adapted microperimetry were still significantly associated with drusen volume ($P = 0.008$, $P = 0.023$, and $P = 0.022$, respectively). For mesopic and dark-adapted microperimetry, mean sensitivity at 0°–1° and 3° degrees was significantly associated with drusen volume, whereas mean sensitivity at 5° and 7° was not associated with drusen volume after adjusting for AMD stage, age, and the presence...
### TABLE 2. Descriptive Analysis and Group Comparisons

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Early AMD</th>
<th>iAMD</th>
<th>Early AMD vs. iAMD</th>
<th>Early AMD vs. Control</th>
<th>iAMD vs. Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drusen load, mm³</td>
<td>0.000236</td>
<td>0.002718</td>
<td>0.155821</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>(0.0003033)</td>
<td>(0.0015339)</td>
<td>(0.1945323)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BCVA</td>
<td>87.6 (4.1)</td>
<td>83.5 (4.1)</td>
<td>81.6 (6.8)</td>
<td>0.306</td>
<td>0.016</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>LIVA</td>
<td>74.0 (4.6)</td>
<td>70.0 (6.4)</td>
<td>64.0 (9.3)</td>
<td>0.008</td>
<td>0.06</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>MAC-VA</td>
<td>68.4 (3.9)</td>
<td>64.1 (4.2)</td>
<td>60.1 (6.7)</td>
<td>0.026</td>
<td>0.006</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Contrast sensitivity</td>
<td>38.2 (2.9)</td>
<td>37.0 (3.6)</td>
<td>33.6 (3.1)</td>
<td>&lt;0.001</td>
<td>0.137</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>IReST</td>
<td>166.8 (21.9)</td>
<td>162.7 (23.9)</td>
<td>149.6 (28.1)</td>
<td>0.077</td>
<td>0.693</td>
<td>0.012</td>
</tr>
<tr>
<td>Mesopic microperimetry</td>
<td>26.2 (1.7)</td>
<td>24.9 (3.2)</td>
<td>23.0 (2.8)</td>
<td>0.016</td>
<td>0.056</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Dark-adapted microperimetry</td>
<td>22.7 (1.4)</td>
<td>23.1 (5.7)</td>
<td>20.5 (2.4)</td>
<td>0.004</td>
<td>0.715</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

*P values are based on the Wilcoxon rank-sum test. Bold numbers indicate statistical significance.

### TABLE 3. Linear Regression of Visual Function Measures Against Drusen Volume (in mm³)

<table>
<thead>
<tr>
<th></th>
<th>β Coefficient</th>
<th>P</th>
<th>Adjusted for AMD Stage and Age</th>
<th>β Coefficient</th>
<th>P</th>
<th>Adjusted for AMD Stage, Age, and Presence of SDD</th>
<th>β Coefficient</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>BCVA</td>
<td>-0.001</td>
<td>0.571</td>
<td>-0.001</td>
<td>0.581</td>
<td>0.001</td>
<td>-0.001</td>
<td>0.588</td>
<td>0.001</td>
</tr>
<tr>
<td>LIVA</td>
<td>-0.005</td>
<td>0.007</td>
<td>-0.003</td>
<td>0.091</td>
<td>0.005</td>
<td>-0.003</td>
<td>0.093</td>
<td>0.005</td>
</tr>
<tr>
<td>MAC-VA</td>
<td>-0.009</td>
<td>&lt;0.001</td>
<td>-0.007</td>
<td>0.008</td>
<td>&lt;0.001</td>
<td>-0.007</td>
<td>0.008</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Contrast sensitivity</td>
<td>-0.014</td>
<td>0.001</td>
<td>-0.007</td>
<td>0.148</td>
<td>0.008</td>
<td>-0.008</td>
<td>0.130</td>
<td>0.008</td>
</tr>
<tr>
<td>IReST</td>
<td>-0.001</td>
<td>0.125</td>
<td>0.000</td>
<td>0.368</td>
<td>0.000</td>
<td>0.368</td>
<td>0.349</td>
<td>0.000</td>
</tr>
<tr>
<td>Mesopic microperimetry</td>
<td>Global</td>
<td>-0.019</td>
<td>0.001</td>
<td>0.000</td>
<td>0.784</td>
<td>0.000</td>
<td>0.784</td>
<td>0.000</td>
</tr>
<tr>
<td></td>
<td>0°–1°</td>
<td>-0.022</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>3°</td>
<td>-0.018</td>
<td>&lt;0.001</td>
<td>0.015</td>
<td>0.015</td>
<td>0.006</td>
<td>0.015</td>
<td>0.015</td>
</tr>
<tr>
<td></td>
<td>5°</td>
<td>-0.012</td>
<td>0.033</td>
<td>-0.005</td>
<td>0.306</td>
<td>-0.007</td>
<td>0.195</td>
<td>0.012</td>
</tr>
<tr>
<td></td>
<td>7°</td>
<td>-0.007</td>
<td>0.130</td>
<td>-0.001</td>
<td>0.784</td>
<td>-0.001</td>
<td>0.609</td>
<td>0.000</td>
</tr>
<tr>
<td>Dark-adapted microperimetry</td>
<td>Global</td>
<td>-0.017</td>
<td>0.001</td>
<td>-0.011</td>
<td>0.022</td>
<td>-0.012</td>
<td>0.022</td>
<td>0.022</td>
</tr>
<tr>
<td></td>
<td>0°–1°</td>
<td>-0.016</td>
<td>&lt;0.001</td>
<td>-0.010</td>
<td>0.051</td>
<td>-0.010</td>
<td>0.037</td>
<td>0.037</td>
</tr>
<tr>
<td></td>
<td>3°</td>
<td>-0.021</td>
<td>&lt;0.001</td>
<td>-0.016</td>
<td>0.001</td>
<td>-0.017</td>
<td>0.001</td>
<td>0.001</td>
</tr>
<tr>
<td></td>
<td>5°</td>
<td>-0.010</td>
<td>0.029</td>
<td>-0.006</td>
<td>0.185</td>
<td>-0.006</td>
<td>0.166</td>
<td>0.166</td>
</tr>
<tr>
<td></td>
<td>7°</td>
<td>-0.005</td>
<td>0.270</td>
<td>-0.001</td>
<td>0.983</td>
<td>-0.000</td>
<td>0.951</td>
<td>0.951</td>
</tr>
<tr>
<td>Pigment changes</td>
<td>0.089</td>
<td>0.013</td>
<td>0.038</td>
<td>0.319</td>
<td>0.034</td>
<td>0.372</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Bold numbers indicate statistical significance.*

of SDD (Table 3). In univariate regression, the presence of pigment changes was significantly associated with drusen volume (P = 0.013), but adjusting for pigment changes in multivariate regression showed no effect on overall results.

### DISCUSSION

In our study, drusen volume was associated with visual impairment detected in functional tests under low luminance and challenging contrast conditions in early stages of AMD. Specifically, MAC-VA and both mesopic and dark-adapted microperimetry were significantly associated with drusen volume. These results indicate a structure–function relationship in early stages of AMD that may not be detectable using conventional high-luminance, high-contrast functional tests.

The association between drusen volume and microperimetric sensitivity found in this study is consistent with previous studies in early stages of AMD; however, in those studies only mesopic microperimetry was assessed. We also found global dark-adapted microperimetric sensitivity, as well as sensitivities at 0°–1° and 3° eccentricities but not at 5° and 7° eccentricities, to be associated with drusen volume. Our results revealed a significant association between MAC-VA and drusen volume. We also found that MAC-VA was the only functional test that differed significantly among all three groups. When comparing the iAMD group to controls, we found that the performance of all visual function tests decreased significantly in the iAMD group. These findings are comparable with previous studies that have also reported a reduced visual function in these tests. Chandramohan and colleagues also found BCVA, LIVA, and mesopic microperimetry significantly decreased in patients with iAMD compared to healthy controls. Similar results were found by Wu et al., who found that results of these tests were significantly reduced for all AMD groups except early AMD compared to controls, which is in line with our results. BCVA was on average four letters worse in eyes with early AMD, and this difference was statistically significant. This is comparable to the findings from Owseley et al. and Klein et al., who reported a significant difference of two letters between these two groups. We did not find a significant relationship between the presence...
of SDDs and drusen volume, and adjusting for SDDs in multivariate regression analysis demonstrated no changes in the results. Interestingly, we found a higher prevalence of SDDs in eyes with early AMD (31.3%) than in eyes with iAMD (19.4%), which is in contrast to many other studies and likely a spurious finding due to our small sample size.

We found that LLVA and contrast sensitivity were decreased in the iAMD group compared to controls and early AMD; however, we did not find a significant difference in these functional tests between early AMD and controls. Puell et al. showed that LLVA was impaired in early stages of AMD before changes in BCVA were observed. Decreased contrast sensitivity in early AMD compared to healthy controls has been reported by Feigl and coworkers.

In our study, drusen volume was found to be largest in the iAMD group and significantly lower in the early AMD group and controls. This is in accordance with several other studies demonstrating that drusen volume increases with increasing AMD stage and is predictive of progression to late AMD.

For early AMD, we calculated a mean drusen volume of 0.0027 mm³, which is lower than the values reported by Li and coworkers, who found a mean drusen volume of 0.03 (range, 0.00–0.28) mm³ in eyes with early AMD. This could also be explained by the small sample size in our early AMD group. The mean drusen volume measure we obtained for the iAMD group of 0.138 mm³ is comparable to the results of a study by Yehoshua et al., who reported drusen volume measures of 0.095 to 0.375 mm³ in the highest quintile for eyes with nonexudative AMD.

We showed that drusen volume is associated with visual impairment detected by functional tests. This is in line with previous studies that have found that functional tests under low lighting are correlated with retinal morphology in AMD.

Strengths of our study include the comprehensive panel of functional tests, including the relatively new MAC charts, for which few data are available, as well as the use of reading performance and dark-adapted microperimetry. All participants were phenotyped according to the current reference-standard retinal imaging in combination with a clinical examination. For drusen volume calculation, we used a new CNN-based approach that allows for the fully automated segmentation of drusen in OCT images. Gorgi Zadeh et al. demonstrated that the CNN-based approach yields much better results than a previous state-of-the-art method reported by Chen et al. and therefore allows for accurate automated assessment of drusen load in AMD. A limitation of our study is the relatively small sample size of the early AMD group and the controls, leading to less statistical power. As common with exploratory studies, no adjustment for multiple testing was done which might lead to an overestimation of statistical power. Another limitation of our study is the fact that the control group was younger than both AMD groups, although univariate linear regression revealed no significant association between age and drusen volume ($P = 0.642$), and we adjusted for age in the multivariate regression analyses.

In conclusion, our study showed that MAC-VA and mesopic and dark-adapted microperimetry are associated with drusen volume in early stages of AMD and thus might provide an indication of structural changes. Our findings suggest that these visual function tests might be useful measurements in monitoring and diagnosing early AMD and iAMD and could be used as functional endpoints in clinical studies. However, more research is warranted, and a longitudinal follow-up will be needed to evaluate the performance of these functional tests as intended, for example, by the MACUSTAR consortium.

Acknowledgments

We thank Jeany Q. Li, Matthias M. Mautsitz, Maximilian Pfau, Christopher A. Turau, and Gabriele N. Turski for their support in conducting the ophthalmic examinations.

This research was supported by the German Scholars Organization/Else Kröhner Fresenius Stiftung (GSO/EKFS 16) and BONFOR GEROK Program, Faculty of Medicine, University of Bonn (grant no. O-137.0028, MW). We are grateful for the technical support of Carlo Pellizzari from CenterVue SpA (Padua, Italy), which provided research material (S-MAIA) necessary to conduct this study. CenterVue had no role in the design or conduct of the experiments.

Disclosure: S.G. Ponderfer, Heidelberg Engineering (F), Optos (F), Carl Zeiss Meditec (F), CenterVue (F); M.W.M. Wintergerst, Heidelberg Engineering (F), Optos (F), Carl Zeiss Meditec (F), CenterVue (F), Heine Optotechnik (C, F), DigiSight Technologies (F), D-Eye (F); S. Gorgi Zadeh, None; T. Schultz, None; M. Heinemann, Heidelberg Engineering (F), Optos (F), Carl Zeiss Meditec (F), CenterVue (F); E.G. Holz, Heidelberg Engineering (F, C, R), Optos (F), Carl Zeiss Meditec (F, C), CenterVue (F), Allergan (F, R), Alcon/Novartis (F, R), Genentech/Roche (F, R), Bayer (F, R), Acucela (F, R), Boehringer Ingelheim (F, R); R.P. Finger, Heidelberg Engineering (F), Optos (F), Carl Zeiss Meditec (F), CenterVue (F), Bayer (C), Novartis (F, C), Santen (C), Opthea (C), Novelion (C), Retina Implant (C), Oxford Innovation (C)

References

10. Hageman GS, Luthert PJ, Victor Chong NH, Johnson LV, Anderson DH, Mullins RF. An integrated hypothesis...


