Semi-automated quantification of geographic atrophy with blue-light autofluorescence and spectral-domain optical coherence tomography: a comparison between the region finder and the advanced retinal pigment epithelium tool in the clinical setting

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ABSTRACT.

Purpose: To compare inter- and intraobserver reliability and intermodality agreement on quantification of geographic atrophy, using two routinely available quantification tools, based on blue-light fundus autofluorescence (BAF) and spectral-domain optical coherence tomography (SD-OCT).

Methods: Quantifications of atrophic lesions within the central 5 mm of 30 eyes from 30 patients (mean age: 76.1 years) were independently performed by two clinicians on BAF images using the REGION FINDER (RF; Heidelberg Engineering) and on SD-OCT using the advanced retinal pigment epithelium tool (ARPET; Carl Zeiss Meditec) at baseline and follow-up (mean interval: 336 days). Inter- and intraobserver reliability was determined by intraclass correlation coefficients (ICC) and Bland–Altman plots. Additionally, graders rated the experienced difficulty of each measurement.

Results: Intraclass correlation coefficients (ICC) showed excellent inter- and intraobserver reliability with values between 0.994 and 0.998 for RF and slightly higher values for ARPET of 0.997 and 0.999. Bland–Altman plots showed smaller variability for ARPET. Mean interobserver differences (95% CI) for size measurements were −0.11 (−0.27; 0.05) (baseline) and −0.05 mm² (−0.18; 0.08) (follow-up) for RF and −0.04 (−0.14; 0.06) and −0.06 mm² (−0.14; 0.02) for ARPET. Measurements of lesions were on average 0.57 mm² (0.35; 0.79) or 7.6% larger in ARPET. Lesion size between graders did not differ significantly. There was no statistically significant difference in relative enlargement rates between methods. There was poor to moderate agreement between graders when rating the experienced difficulty of each measurement.

Conclusion: Semi-automated analysis of geographic atrophy with RF and ARPET is equally reliable and reproducible in clinical settings, despite both algorithms require frequent adjustment by users. The ARPET restricts size measurements to the central 5 mm, which limits its ability to fully track GA progression. Results of both tools are not interchangeable as measurements with ARPET result in larger lesion sizes.

Key words: advanced RPE tool – blue-light autofluorescence – geographic atrophy – interobserver reliability – intraobserver reliability – optical coherence tomography – outer retinal atrophy – region finder – semi-automated quantification
Introduction

Age-related macular degeneration (AMD) is a major burden for health care systems, and demographic trends predict a significant rise in its global prevalence, from an estimated 196 to 288 million patients between 2020 and 2040 (Wong et al. 2014; Colijn et al. 2017). Geographic atrophy (GA) presents as localized atrophy of the retinal pigment epithelium (RPE), the outer retina and the choriocapillaris in advanced AMD, causing central scotomas and permanent loss of visual acuity (Sayegh et al. 2017; Fleckenstein et al. 2018). While no approved treatment for GA is available at present, several studies and trials focus on understanding the progression patterns or altering the enlargement speed of atrophic areas (Bandello et al. 2017; Evans & Lawson 2017; Sacconi et al. 2017). Monitoring GA development and progression over time is therefore invaluable in both research and clinical settings. Various imaging modalities are being used for this matter, ranging from conventional fundus colour photography and near-infrared reflectance imaging (IR) over blue-light fundus autofluorescence (BAF) or greenlight fundus autofluorescence (GAF) to spectral-domain (SD), swept-source or polarization-sensitive optical coherence tomography (Schmitz-Valckenberg et al. 2011; Yehoshua et al. 2011). Additionally, factors with a possible influence on agreement between graders such as the presence of multifocal lesions were analysed. We further assessed if there was a difference in measured lesions size or lesion progression between the two software tools. Finally, we evaluated if graders were agreeing on the subjective difficulty of every measurement and if the used quantification tool had an influence on the difficulty rating.

Material and Methods

This retrospective longitudinal study comparing two tools for semi-automated quantification of GA lesions was approved by the Medical University of Vienna institutional review board and conducted in compliance with the Declaration of Helsinki.

Study cohort

Subjects under regular clinical surveillance due to GA were selected from the database of the outpatient clinic for macular diseases. Selection procedure was performed as followed: In total, thirty patients with the diagnosis ‘advanced nonneovascular AMD’ were found who had received same day BAF on a Heidelberg Spectralis imaging platform and SD-OCT imaging on a Cirrus HD-OCT platform with a follow-up of at least 6 months between 01 January 2015 and 31 December 2017. Patients were then pseudonymized with a random number generator Research Randomizer–Version 4.0, Urbaniak, & Plous, (2013), and sequentially, both eyes of each patient (n = 60 eyes) were screened regarding the inclusion and exclusion criteria: Eligible eyes had to be free from a history of neovascularization or other non-AMD related retinal pathologies and refractive errors >6 dioptries. Blue-light fundus (BAF) images had to be acquired with a Spectralis HRA + OCT platform (Heidelberg Engineering), operating with a 488 nm solid state laser and a transverse resolution of 10 μm in high-speed mode. Spectral-domain (SD)-OCT images had to be acquired with a Cirrus HD-OCT 5000 platform (Carl Zeiss Meditec, Dublin, OH, USA), as these imaging modalities and software tools are widely available and easily accessible (Schmitz-Valckenberg et al. 2011; Yehoshua et al. 2011). Additionally, factors with a possible influence on agreement between graders such as the presence of multifocal lesions were analysed. We further assessed if there was a difference in measured lesions size or lesion progression between the two software tools. Finally, we evaluated if graders were agreeing on the subjective difficulty of every measurement and if the used quantification tool had an influence on the difficulty rating.

The present study aims to provide such information by comparing inter- and intraobserver reliability of two proprietary semi-automated quantification tools in a clinical setting, outside reading centre conditions. The REGION FINDER (RF) software in conjunction with BAF using the Heidelberg Spectralis imaging platform (Heidelberg Engineering, Heidelberg, Germany) was compared with the advanced RPE tool (ARPET) in conjunction with SD-optical coherence tomography (SD-OCT) on a Cirrus HD-OCT platform (Carl Zeiss Meditec, Dublin, OH, USA), as these imaging modalities and software tools are widely available and easily accessible (Schmitz-Valckenberg et al. 2011; Yehoshua et al. 2011). Additionally, factors with a possible influence on agreement between graders such as the presence of multifocal lesions were analysed. We further assessed if there was a difference in measured lesions size or lesion progression between the two software tools. Finally, we evaluated if graders were agreeing on the subjective difficulty of every measurement and if the used quantification tool had an influence on the difficulty rating.
All images had been taken in mydriasis, achieved routinely by instillation of 1% tropicamide (Mydriaticum Agepha, Vienna, Austria) and 2.5% phenylephrine eye drops. Regarding the characteristics of the atrophic lesions, no exclusion criteria were used and eyes with uni- and multifocal lesions as well as well-demarcated and less well-demarcated lesion borders were included.

**Procedures**

Included images were anonymized and put into random order by a co-author (GW) not involved in the assessment of GA size. Measurements with the RF would have been possible over the whole 30° × 30° BAF image, while the ARPET limited measurements to a 5 mm circle on the SD-OCT image. To make the working area comparable between the two programs, the circular constraint tool of the RF was used by the co-author (GW) to draw a 5 mm circle on the BAF image analogous to the 5 mm circle of the ARPET, using vessels and GA borders as landmarks for placement (Fig. 1A,B,H). The positions of the 5 mm circular constraints on RF images and the 5 mm circles on the ARPET images were not altered for the rest of the study, ensuring that both graders performed measurements in the same area.

Grader 1 (AR) and grader 2 (MGK) performed measurements independently, on separate days and with the same image order to prevent uneven learning curves. Measurements were performed 1 week apart, beginning with ARPET and RF baseline images and followed by ARPET and RF follow-up images. For intraobserver reliability, one grader (AR) did a second measurement cycle. All measurements were performed directly on the same Cirrus HD-OCT and Spectralis HRA + OCT platforms.

For the ARPET, graders were instructed to start with the suggested quantification of the algorithm and to use the integrated tools (brush, eraser, bucket), as well as the integrated OCT B-scans to adjust GA lesion size but could not use any other information or imaging modalities (Fig. 1E–G).

For the RF, graders were instructed to start with the suggested quantification of the algorithm and to use the integrated tools (brush, eraser, bucket), as well as the integrated OCT B-scans to adjust GA lesion size but could not use any other information or imaging modalities (Fig. 1E–G).

For the RF, graders were instructed to start with the suggested quantification of the algorithm and to use the integrated tools (brush, eraser, bucket), as well as the integrated OCT B-scans to adjust GA lesion size but could not use any other information or imaging modalities (Fig. 1E–G).

**Fig. 1.** Steps of measuring atrophic areas with REGION FINDER (first row) and advanced retinal pigment epithelium tool (ARPET) (second row). (A) Blue-light fundus (BAF) image shows non-autofluorescent atrophic areas in dark grey to black. Borders of the spared fovea are hardly visible. (B) A circular constraint (red circle) was manually placed to limit the workspace of graders analogous to the 5 mm border of the ARPET (compare to the white circle in image H). (C) After planting a seed, a grader ‘filled’ the region by adjusting the threshold for grey values (blue area). Without additional constraints, the spared fovea is falsely included in the atrophic lesion by the algorithm. (D) Final result. Manual constraints were drawn by grader 1 to demarcate foveal sparing. The measured area is ‘capped’ at the 5 mm circular constraint. (E) Optical coherence tomography (OCT) fundus image. In contrast to BAF, atrophic areas appear bright and foveal sparing is clearly visible. (F) Sub-retinal pigment epithelium (RPE) slab (top) and corresponding OCT image (bottom). B-Scan position is shown by the blue line. Areas with abnormal beam penetration are depicted in bright white. (G) Sub-RPE illumination map allows graders to manually correct the algorithm’s suggested area of atrophy. The yellow area is erased or enlarged using the information of the sub-RPE slab and B-Scan (image F). (H) Final Result. Atrophic areas are shown in bright white with black borders. Only areas within the 5 mm circle are measured. There are subtle differences regarding the presence of atrophy between BAF and spectral-domain-OCT (compare the areas in images D and G).
Statistical analysis

Analysis of the main question
The main goal of the study was to analyse the inter- and intraobserver reliability for measuring GA lesion size with RF and ARPET. Interobserver reliability was investigated using Bland–Altman plots. Estimated mean differences with 95% confidence intervals are reported for each method and each visit separately. Estimates and 95% confidence intervals for the interclass correlation coefficient (ICC) were computed with the ‘ irr ’ software package for R (cran.r-project.org, M. Gamer), based on a single-rating, absolute agreement, two-way random-effects model.

Intraobserver reliability was analysed in a similar manner, whereby ICCs were calculated based on a single-rating, absolute agreement, two-way mixed-effects model. Based on 95% confidence intervals, values between 0.5 and 0.75, between 0.75 and 0.9, and >0.90 were considered as moderate, good, and excellent reliability (Koo & Li 2016).

Analysis of secondary questions
To examine if the presence of multifocal lesions, retained RPE fragments within the atrophy or atrophy borders within the foveal region had an influence on the agreement between graders, different symbols in Bland–Altman plots were used and analysed descriptively.

To analyse if the grader or the used quantification tool had an influence on measured GA size, a mixed model (SAS quantification tool had an influence on were used and analysed descriptively. To analyse if the visit, method and the presence of atrophy borders within the foveal region had an influence on the difficulty rating, an ordinal logistic regression model with patient number as repeated factor (SAS Proc genmod) was calculated. p-values based on the score test as well as odds ratios with 95% Wald confidence intervals are reported.

Statistical analysis was conducted with r 3.3.2 (r-project.org, RCore Team), sas 9.4 (SAS Institute, Cary, NC, USA) and spss 24 (IBM Corp, Armonk, NY, USA). p-values of <0.05 were considered statistically significant.

Results

Patients, eyes and GA lesions
Characteristics of the included patients, eyes and GA lesions are shown in Table 1. Regarding demarcation of lesions, GA areas were generally well demarcated in ARPET as the sub-RPE slab allowed for a high contrast between atrophic and nonatrophic regions. In RF, region borders were either well demarcated, which was the case for nine eyes with multifocal and seven eyes with monofocal lesions at baseline (eight of each at follow-up), or less well demarcated, which was the case for seven multifocal and seven monofocal lesions at baseline and follow-up. Mean confocal scanning laser detector sensitivity for BAF images was 78.4 (SD: 7.7) at baseline and 86.4 (SD: 7.8) at the follow-up visit. Mean signal strength for OCT images was 7.9 (SD: 1.1) at baseline and 7.9 (SD: 1.0) at the follow-up visit.

Interobserver reliability and GA lesion size
Geographic atrophy (GA) lesion sizes at baseline and follow-up for both graders as well as mean differences in measurements between graders are shown in Table 2. Plotting the differences between measurements against their respective mean value (Bland–Altman plot) revealed higher variability in differences for RF in comparison with ARPET (Fig. 2). Lesion size did not correlate visibly or statistically with the measurement-difference between graders (Spearman’s \( \rho = 0.164 \) and \(-0.023\) for RF and 0.012 and 0.225 for ARPET). Intraclass correlation coefficients (ICCs; 95% CI) showed excellent interobserver reliability for RF with values of 0.994 (0.987; 0.997) at baseline and 0.996 (0.993; 0.998) at follow-up. Intraclass correlation coefficients (ICCs) for ARPET were slightly higher with 0.997 (0.995; 0.999) at baseline and 0.998 (0.996; 0.999) at follow-up.

Intraobserver reliability and GA lesion size for repeated measurements
Geographic atrophy (GA) lesion sizes and mean differences in repeated measurements of Grader 1 are shown in Table 2. Similar to interobserver agreement, variability in differences was slightly larger for RF than for ARPET as shown in the Bland–Altman Plots (Fig. 3).

Intraclass correlation coefficients (ICCs; 95% CI) showed excellent intraobserver reliability for RF with values of 0.997 (0.994; 0.999) at baseline and 0.998 (0.995; 0.999) at follow-up. Intraclass correlation coefficients
Table 2. Size of atrophic area (mm²) within the central 5 mm, assessed with the REGION FINDER and advanced retinal pigment epithelium (RPE) tool,  

|                        | Baseline visit |                              | Follow-Up visit |                              |
|------------------------|----------------|-------------------------------|-----------------|-------------------------------|
|                        | Mean | SD   | Median | Min/max | Mean | SD   | Median | Min/max |
| REGION FINDER          |       |      |        |         |       |      |        |         |
| Grader 1               | 8.62 | 3.96 | 9.67   | 2.95/15.78 | 10.06 | 4.08 | 11.40  | 3.40/17.44 |
| Grader 2               | 8.73 | 3.93 | 9.82   | 3.07/15.76 | 10.11 | 4.11 | 11.66  | 3.65/17.32 |
| Grader 1R*             | 8.52 | 3.85 | 9.83   | 2.90/15.54 | 10.08 | 4.10 | 11.67  | 3.43/17.69 |
| Advanced RPE Tool      |       |      |        |         |       |      |        |         |
| Grader 1               | 8.90 | 3.66 | 9.70   | 3.20/14.20 | 10.35 | 3.76 | 11.50  | 3.50/16.0 |
| Grader 2               | 8.93 | 3.67 | 9.90   | 3.00/14.10 | 10.41 | 3.71 | 11.60  | 3.80/15.50 |
| Grader 1R*             | 8.86 | 3.66 | 9.70   | 3.10/14.40 | 10.33 | 3.74 | 11.70  | 3.50/16.0 |

* Repeated measurement of grader 1 for intraobserver reliability.

Effects of method and grader on measured GA size

Mixed model analysis showed that GA lesion size was not affected by the grader [mean difference (95% CI): 0.063 (−0.11; 0.23), p = 0.470] but by the used quantification tool [ARPET versus REGION FINDER: mean difference (95% CI): 0.19 (0.017; 0.37), p = 0.032] and the interaction between the used tool and the factor ‘lesion extending beyond central 5 mm’ (p < 0.0001). For the latter, subset analyses were performed to investigate the effect of the used tool separately for patients with lesions extending beyond the automatically (ARPET) or manually (RF) placed 5 mm margin (n = 12) and for patients with lesions that did not expand beyond the central 5 mm circle (n = 18), respectively. For patients with lesions that did not expand over the central 5 mm circle, GA measurements were on average significantly larger when performed with ARPET (mean difference (95% CI): 0.57 (0.35; 0.79), p < 0.0001) compared with the REGION FINDER. Mean size for these lesions at baseline was 6.22 mm² (SD: 2.92) for RF and 6.72 mm² (SD: 2.91) for ARPET and 7.67 mm² (SD: 3.32) and 8.25 mm² (SD: 3.25) at follow-up.

For patients with lesions that were capped by the 5 mm margin (‘lesions extending beyond central 5 mm = yes’), no statistically significant difference between methods was observed [ARPET versus RF: mean difference (95% CI): −0.19 (−0.46; 0.084), p = 0.18]. Regarding the enlargement rate, there was no statistically significant difference between the two methods (mixed model, p = 0.746).

To prevent false-positive region growth in RF, 17 images at baseline and 16 at follow-up required placement of constraints in the foveal region. Overall, constraints had to be placed in 55 from 60 BAF images by graders, with an average of 2.67 constraints (SD: 1.73) at baseline and 3.28 (SD: 2.52) at follow-up. Analysis of the Bland–Altman plots showed that neither multifocal lesions (Fig. 4), nor retained RPE fragments (not shown) or GA involvement of the foveal region (Fig. 2) influenced the variability in differences between graders.

(ICC) for ARPET were slightly higher with 0.999 (0.998; 1) at baseline and 0.999 (0.999; 1) at follow-up.

Interobserver performance regarding number of lesions and foveal involvement

There was total agreement between graders and methods when assessing if a patient had mono- or multifocal GA. Lesion numbers per image ranged between 1 and 23 (median = 2) at baseline and between 1 and 17 (median = 1.5) at follow-up and were similar for graders and methods.
Subjective difficulty for graders

Ordinal difficulty ratings were similarly distributed between graders with RF at baseline and follow-up. Both graders rated ARPET measurements slightly easier at baseline and at follow-up, but no statistically significant effect of grader, visit, method or the interaction between foveal constrains and method on the difficulty rating could be observed (ordinal logistic regression with repeated measurements: method: \( p = 0.34 \), visit: \( p = 0.24 \), grader: \( p = 0.76 \), interaction foveal constraints*: method: \( p = 0.95 \)). Blue-light fundus (BAF) measurements were rated as more difficult, when the graders had to place constraints in the central foveal area [foveal constraints yes versus no: odds ratio (95% CI) = 2.93 (1.22; 7.01), \( p = 0.0249 \)]. For the individual measurements, weighted kappa coefficients (95% CI) showed only low to moderate interobserver agreement [at baseline: RF: 0.153 (−0.132; 0.437), ARPET: 0.463 (0.211; 0.715); at follow-up: RF: 0.250 (−0.028; 0.528), ARPET: 0.391 (0.160; 0.622)] and moderate to substantial intraobserver agreement [at baseline: RF: 0.344 (0.069; 0.620), ARPET: 0.756 (0.546; 0.966); at follow-up: RF: 0.407 (0.156; 0.658), ARPET: 0.554 (0.323; 0.785)].

Discussion

The purpose of this study was to compare two commercially available options for semi-automated quantification of GA in a clinical setting: the RF in conjunction with BAF and the ARPET with SD-OCT.

The two-dimensional BAF images used with the RF are not true anatomical images, but distribution maps of fluorescent compounds contained in lipofuscin, a not degradable, highly oxidized aggregation of cross-linked proteins that can be found in several human tissues as a result of cellular ageing and oxidative stress and other fluorophores (Delori et al. 1995; von Ruckmann et al. 1995; Jung et al. 2007). The contrast difference of autofluorescent nonatrophic areas and nonautofluorescent atrophic lesions improves analysis over fundus colour photos (von Ruckmann et al. 1997). The RF utilizes these contrast differences to form regions of similar BAF intensity, a principle known as binary enlarged objects ("BLOB") detection, and calculates their size with a scaling factor registered during image acquisition, which has been reported to be more accurate than manual outlining of GA lesions (Schmitz-Valckenberg et al. 2002, 2011). Drawbacks emerge from the technique of BAF and the RF algorithm: First, the distinction between healthy and atrophic regions around the central fovea is known to be challenging due to physiological autofluorescence signal reduction in this area (Delori et al. 1995; Sayegh et al. 2015; Domalpally et al. 2016). This signal reduction results in false-positive region growth of the RF’s algorithm when grading foveal atrophic lesions, a problem we encountered frequently as nearly half of all BAF images required placement of central constraints (Fig. 1C), which had a negative impact on the difficulty ratings of the graders but no significant impact on agreement between graders. Second, wrong region growth of the algorithm into vessels and beyond less well-demarcated lesions borders has been described (Schmitz-
Valckenberg et al. 2002, 2011), which required constraint placement in 92% of BAF images in the present study, underlining that BLOB-based region-detection is highly prone to false-positive region growth. While the combination of BAF analysis with imaging techniques that show less to none foveal signal reduction, such as GAF or IR, can help in distinguishing healthy from pathologic regions, it does not solve the false-positive region growth, which is inherent to BLOB-based lesion detection.

In contrast to two-dimensional imaging techniques, the ARPET uses cross-sectional information of a three-dimensional volume, acquired by a high-definition SD OCT (Cirrus HD-OCT). Summing and merging the signal of each A-scan allows for creation of OCT fundus images (see Fig. 1E) which can be used to outline atrophic areas (Barelle et al. 2009; Yehoshua et al. 2011; Pilotto et al. 2015). In contrast to a conventional OCT fundus image, the ARPET utilizes the different light scattering properties of retinal tissue such as RPE, choriocapillaris or choroid to create a sub-RPE en-face image based on beam penetration, known as the sub-RPE slab (Fig. 1H) (Yehoshua et al. 2011; Augustin 2012). Although the ARPET algorithm detects GA areas in the sub-RPE slab automatically as opposed to manual seed planting with the RF, manual correction was necessary in all images, showing that the software regularly demarcates lesion borders incorrectly. Additionally, while en-face OCT images have advantages over BAF images such as a constant signal around the fovea or availability of depth information (Fig. 1A,E,F), they are prone to motion artefacts due to longer scanning duration and interpolated, meaning that resolution depends on the number of performed scans (Sayegh et al. 2011; Yehoshua et al. 2011; Pilotto et al. 2015).

Although both methods rely on different technical backgrounds and software algorithms, they showed excellent inter- and intraobserver reliability and consistent measurements were performed without a measurable effect of graders. For RF, the mean interobserver differences between GA size measurements were 0.11 mm² (baseline) and 0.05 mm² (follow-up), which is very similar to the results of professional graders, who reported interobserver differences from 0.00 to 0.44 mm² in 30 eyes from 30 patients but were additionally allowed to use blue reflectance images to obtain additional information on the lesion borders and could copy constraints and regions from baseline to follow-up visits (Schmitz-Valckenberg et al. 2011). A second study on 29 eyes from 20 GA patients assessed intra- and interobserver agreements with RF between nonprofessional graders and found mean interobserver differences from 0.02 to 0.27 mm² (Panthier et al. 2014). Mean intraobserver differences were smaller than interobserver differences in the mentioned studies (−0.02 to 0.32 mm² for professional graders (Schmitz-Valckenberg et al. 2011), −0.01 to −0.17 mm² for nonprofessional graders (Panthier et al. 2014)) as well as in the presented one (−0.02 and 0.10 mm²).

For ARPET, mean inter- and intraobserver differences in GA size were at the level of RF, with values around 0.05 mm². While no detailed reports for ARPET were found, two studies by Yehoshua et al. (2011, 2015) stated excellent inter- and intraobserver reliability when using a 6 x 6 mm square SD-OCT fundus image (not to be confused with the sub-RPE slab of ARPET) and measuring GA size with external proprietary software (ADOBE PHOTOSHOP CS2, Adobe Systems, San José, California, USA) and a stylus-driven digitizing tablet. A big advantage of RF and ARPET over such approaches is, that both programs are ready to use on either the Heidelberg Spectralis imaging platform or the Cirrus SD-OCT and measurements can be done easily on the spot, which is more important for clinical use than for specialised reading centres.

Bland–Altman plots revealed that variability between graders’ measurements was even smaller with ARPET. This finding is unexpected, as ARPET relies more on direct mouse-driven region border correction and seems therefore to be more prone to human error than utilization of the region growth algorithm of RF. However, this finding was consistent for all measurements. Bland–Altman plots for RF in the presented study showed even slightly less variability in comparison with other reports, indicating that graders of this study achieved the expected performance (Schmitz-Valckenberg et al. 2011; Panthier et al. 2014). Region size outputs between the tools differ, as RF rounds region size to three decimal places whereas ARPET to just one decimal place, but the impact in decimal places is too subtle to explain for the difference in variabilities. From the current point of view, a possible explanation could be the threshold-driven approach of the region growing algorithm, as local disagreement between graders still affects the whole perimeter of the lesion. Panthier et al. (2014) reported a strong rise in observer disagreement for GA lesions of a size above 15.75 mm². They addressed this issue to the region growing algorithm, as size changes by adjusting the grey-scale threshold is ultimately proportional to the current lesion size, thus making larger regions more prone to substantial disagreement. A small area with a low contrast between atrophic and nonatrophic regions on a BAF image can thus affect the measured size of the whole lesion.

Intraclass correlation coefficients (ICC) were between 0.994 and 0.998 for RF and between 0.997 and 0.999 for ARPET, indicating excellent inter- and intraobserver agreement. A recent paper comparing BAF and GAF for measuring GA (both performed with RF) reported ICCs of 0.995 (GAF) and 0.991 (BAF), which were marginally smaller than the ones found in this study (Pfau et al. 2017).

Regarding the influence of methods on GA size, measurements were on average about 7.6% or 0.5 mm² larger when performed with ARPET. This noticeable difference is in agreement with an abstract on 15 eyes with GA, reporting mean lesion size of 4.8 mm² with ARPET and 4.1 mm² with manual outlining on colour fundus photos (Sharma et al. 2011). Post hoc comparison of the measurements in the current study showed that demarcations of GA regions on sub-RPE slabs were smoother and slightly larger than on BAF images (Fig. 5). Regarding lesion borders, BAF images appeared more detailed. Occasionally, RPE presence between GA lesions was indicated by autofluorescence in BAF, whereas the same region on SD-OCT indicated atrophy and absence of RPE (Fig. 1B, D,F,H). The less detailed images found with ARPET are most likely a result of the lower transversal resolution of the
Cirrus OCT compared with the Heidelberg Spectralis confocal scanning laser ophthalmoscope. Additionally, the OCT en-face and sub-RPE slab images are interpolated, which results in a loss of information. Besides differences in area demarcation, the scaling factors for both imaging platforms are not identical and a difference in region size can also be caused by the underlying technique of image scaling. We did not find a difference between the methods regarding GA size for lesions that did expand over the 5 mm border, which can be explained as followed: The 5 mm margin acts as a smooth artificial lesion border in RF as well as ARPET, thus masking the differences in depiction of the real lesion borders between the two methods and concealing possible differences in lesion size.

Ordinal difficulty ratings for each measurement were not useful, as agreement between graders was moderate at best. Overall difficulty was slightly lower with ARPET in comparison with RF, but no statistical significant difference was found. The weak agreement on difficulty ratings had no effect on the excellent inter- and intraobserver reliability of size measurements, indicating that graders agree with a measurement even if they disagree with the difficulty of performing it.

To the best of our knowledge, this is the first study comparing GA analysis with RF against ARPET. However, several limitations arise: Due to its retrospective nature, typical design problems such as selection bias can occur. Carefully defined inclusion and exclusion criteria were used to minimize this risk, and all included images were acquired by the same team of qualified orthoptists and medical photographers. Despite the excellent levels of agreement, the clinicians in this study were nonprofessional graders and results of this study have to be regarded as such. We did not assess resource factors such as grading time duration, as these attributes are of more relevance for professional reading centres and should therefore be performed in these settings. Regarding image resolution, choosing BAF images in high-resolution mode (1536 × 1536 pixels) could have provided even more details than in high-speed mode (768 × 768 pixels), which is beneficial when analysis of highly magnified images (15° × 15°) is performed. However, this was not the case in our study and high-speed mode has successfully been used when quantifying GA with RF (Schmitz-Valckenberg et al. 2011).

Finally, the workspace of RF had to be manually adjusted with the circular constraint to overlap with ARPET, as the latter limits the workspace automatically to a 5 mm circle. However, not doing so would have led to selection bias, as the RF algorithm causes errors in inter- and intraobserver agreement for large lesions as stated above (Panthier et al. 2014).

Concluding, RF and ARPET are equally suitable for measuring GA size in clinical settings as they both reach excellent inter- and intraobserver reliability but require significant adjustments by graders. REGION FINDER (RF) allows for analysis of larger areas, compared with ARPET, which restricts the user to a 5 mm circle. This limits the use of the ARPET to lesions that are not extensively progressed, ultimately limiting its potential to fully track GA enlargement in every patient in clinical practice, which was the case in 40% of our subjects. However, it also has to be noted that size measurements of large lesions with the RF are less reliable, as mentioned above. The characteristic hypoautofluorescent area

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**Fig. 5.** Comparison of geographic atrophy (GA) measurements between graders with REGION FINDER (RF) and advanced retinal pigment epithelium tool (ARPET). (A,E) Blue-light fundus (BAF) and sub-retinal pigment epithelium (RPE) slab of monofocal GA. (B,C,F,G) Final results of grader 1 (B and F) are very similar to the results of grader 2 (C and G). Lesion size was 9.360 (grader 1) and 9.367 (grader 2) with RF and 10.7 and 10.7 with ARPET. (D,H) Magnification of corresponding areas (white rectangles in C and G) shows that region borders are more detailed in BAF while the sub-RPE slab shows smoother but larger areas.
of the central fovea in BAF images had no significant impact on reliability, but measurements were more challenging when GA borders were present at the foveal region. Intraclass correlation coefficients (ICCs) for size measurements are excellent, even if graders agree generally poorly on the subjective difficulty of the performed measurements.

Contrary to what one would expect, inter- and intraobserver agreement is even better with ARPET, although the clinical relevance of the difference is debatable. The choice of method has an impact on the measured GA size, as lesions were on average 7% larger when using ARPET in comparison with RF. As a result, size measurements are not interchangeable between methods and clinicians outside of clinical trials should stick to a single imaging modality for longitudinal observations or aim for a multimodal imaging approach. Future studies will be necessary to find the best combination of imaging modalities as recently suggested by the Classification of Atrophy Meetings group (Holz et al. 2017).

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