Deep Learning Applied to Automated Segmentation of Geographic Atrophy in Fundus Autofluorescence Images

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Purpose: This study describes the development of a deep learning algorithm based on the U-Net architecture for automated segmentation of geographic atrophy (GA) lesions in fundus autofluorescence (FAF) images.

Methods: Image preprocessing and normalization by modified adaptive histogram equalization were used for image standardization to improve effectiveness of deep learning. A U-Net–based deep learning algorithm was developed and trained and tested by fivefold cross-validation using FAF images from clinical datasets. The following metrics were used for evaluating the performance for lesion segmentation in GA: dice similarity coefficient (DSC), DSC loss, sensitivity, specificity, mean absolute error (MAE), accuracy, recall, and precision.

Results: In total, 702 FAF images from 51 patients were analyzed. After fivefold cross-validation for lesion segmentation, the average training and validation scores were found for the most important metric, DSC (0.9874 and 0.9779), for accuracy (0.9912 and 0.9815), for sensitivity (0.9955 and 0.9928), and for specificity (0.8686 and 0.7261). Scores for testing were all similar to the validation scores. The algorithm segmented GA lesions six times more quickly than human performance.

Conclusions: The deep learning algorithm can be implemented using clinical data with a very high level of performance for lesion segmentation. Automation of diagnostics for GA assessment has the potential to provide savings with respect to patient visit duration, operational cost and measurement reliability in routine GA assessments.

Translational Relevance: A deep learning algorithm based on the U-Net architecture and image preprocessing appears to be suitable for automated segmentation of GA lesions on clinical data, producing fast and accurate results.

Introduction

Geographic atrophy (GA) is one of two end stages of age-related macular degeneration (AMD)—an age-associated disease of the macula that manifests in those aged 50 years and older. It is responsible for 8.7% of legal blindness globally, affecting approximately 5 million people worldwide, and 9 to 10 million cases are expected by the year 2040.¹,² The pathogenesis and etiology of AMD and its progression to GA is not completely understood, and drug therapies are currently not available.³–⁶ GA is characterized by death of the retinal pigment epithelium (RPE) and photoreceptor cells, as well as loss of the underlying choriocapillaris, which appear as sharply demarcated areas on retinal imaging.⁴ Vision loss occurs when atrophic lesions approach the central foveal area, and standard tasks such as reading and recognizing faces become increasingly difficult.⁷,⁸ In the absence of treatment, the condition continues to deteriorate over time, potentially leading to legal blindness (defined as 6/60 vision
in Australia). The rate of irreversible vision loss is also highly variable.8

Several imaging modalities are available for the assessment of GA, including fundus autofluorescence (FAF), color fundus photography (CFP), spectral domain optical coherence tomography (SD-OCT), and near-infrared FAF (IR). Currently, the growth of a GA lesion as seen on FAF is accepted as an outcome in clinical intervention trials by the United States Food and Drug Administration9; hence, there is much interest and need to accurately define borders and thus lesion size in a timely and efficient manner. Often, FAF is used with semiautomated software to help define the boundaries of a GA lesion.10 Such software relies heavily on a human user to correctly annotate and identify lesion boundaries within the image.

There are no clinically available complete automation software packages for the extraction of GA information from retinal images. Several groups have, however, developed artificial intelligence (AI)–based automation methods for isolation of GA lesions.11–30 These studies applied a process known as semantic segmentation—the labeling of each pixel within an image and the precise extraction of regions of interest. The range of algorithms tested included region-growing, interactive segmentation using watershed transform, level set approach, geometric active contour model, Fuzzy c-means, k-nearest neighbor (kNN), Chan-Vese model via local similarity factor, convolutional neural networks (CNN), sparse autoencoder deep networks, and an offline/self-learning model. The retinal images used in these GA-AI publications predominantly included FAF imaging. However, SD-OCT, combinations of SD-OCT/FAF, FAF/CFP, and FAF/IR were also used. Among these segmentation algorithms, sensitivity ranged from 0.47 to 0.983, specificity ranged from 0.93 to 0.99, accuracy ranged from 0.42 to 0.995, mean overlap ratio ranged from 0.659 to 0.899, correlation coefficient ranged from 0.82 to 0.998, and the Dice similarity coefficient (DSC) ranged from 0.66 to 0.89.

Although there are many metrics that can be used for the assessment of semantic segmentation algorithms, the DSC is the most suitable because it measures the overlap between machine-generated results and the ground truth (i.e., human annotated images).31 In the literature, four studies used this metric for evaluation of segmentation algorithms, with results ranging from DSC of 0.66, as reported by Liefers et al.,32 who described a U-Net–based encoder-decoder structure, to a study by Hu et al.,16 who reported a DSC of 0.89 for the application of the level set method on FAF images. In general, an overview of the literature suggests that results from algorithms applied to CFP images produce less promising results when compared with other, more grayscale-based FAF images. For example, sensitivities for CFP-based segmentation algorithms were in the range 0.47 to 0.65; however, for FAF-based algorithms, the range was 0.825 to 0.983. This is due to media opacities and low contrast between atrophic areas and the intact retina in CFPs, which make the detection of GA lesions and their boundaries difficult, even for highly qualified and experienced clinicians and graders.8,25,33 Furthermore, among the segmentation algorithms already applied in the GA-AI space, a majority of the publications reported relatively small image sample sizes. This was not unusual, given there are many constraints on accessing adequate medical data samples, including challenges on sharing data because of privacy or ethical concerns; the lack of equipment within healthcare systems that makes the sharing of available data challenging; and generally a lack of available cases which could be used to train an AI algorithm, especially in the case of deep learning.34–36

In this study, we applied the deep learning U-Net approach to FAF images obtained from GA-affected patients at various stages of progression. The U-Net is a modification of the CNN. It was designed to predict and classify each pixel within an image and thus create a more precise segmentation with fewer training images required.37 In the past, the U-Net architecture was used for GA segmentation by Wu et al.,28 who conducted segmentation on SD-OCT and synthesized FAF images, and by Schmidt-Erfurth et al.,30 who used a residual U-Net model to isolate hyperreflective foci voxels. Liefers et al.32 described segmentation on CFP using a deep learning model with an encoder-decoder structure with residual blocks, shortcut connections, and contracting-expanding pathways, citing the U-Net developers Ronneberger et al.37

Time-series segmentation captures a range of lesion sizes and shapes and provides added variability to reflect real-world clinical settings. For example, it is common for many patients to approach an optometrist or ophthalmologist several months after disease onset, when lesions may have already appeared in a variety of spatial patterns.38 Therefore an automated segmentation method with the capability of detecting GA lesions at all stages of the disease would be invaluable in a clinical setting.

In this article, the aim was to use the U-Net deep learning approach to isolate lesions in FAF images together with suitable image normalization and preprocessing to address image quality issues. Image segmentation is the first step in the automation of GA assessment because extraction of GA area is required.
Study Design and Participants

The study was approved by the Human Research Ethics Committee of the Royal Victorian Eye and Ear Hospital. The study was conducted in accordance with the International Conference on Harmonization Guidelines for Good Clinical Practice and tenets of the Declaration of Helsinki. Ethics approval was provided by the Human Research Ethics Committee (HREC: Project No. 95/283H/15) by the Royal Victorian Eye and Ear Hospital.

Subjects included in this retrospective analysis were AMD participants involved in macular natural history studies from the Centre for Eye Research Australia or from a private ophthalmology practice diagnosed with GA. Cases were referred from a senior medical retinal specialist (R.H.G.) and graded in the Macular Research Unit grading center. Inclusion criteria included being over the age of 50 years, having a diagnosis of AMD (on the basis of the presence of drusen greater than 125 μm) with progression to GA in either one of both eyes. An atrophic lesion was required to be in the macular and not extend beyond the limits of the FAF image at the first visit (i.e., baseline). Participants were required to have foveal centered FAF images and at least three visits recorded over a minimum of two years, with FAF imaging of sufficient quality. Good-quality images were classified as those with minimal or correctable artefacts (e.g., by correction of illumination with pre-processing techniques), and images should encompass the entire macular area and part (i.e., around half) of the optic disc. No minimum lesion sizes were set, because the objective of the study was to be able to automate all lesion sizes. Images contained both unifocal and multifocal lesions. Sampling in the training phase for the algorithm was augmented by time-series segmentation without limitation on lesion size.

Exclusion criteria included participants with neovascular AMD and macular atrophy from causes other than AMD, such as inherited retinal dystrophies, including Stargardt’s disease. These patients were excluded based on the determination of a retinal specialist (R.H.G.). Also excluded were patients who had undergone any prior treatment or participated in a treatment trial for AMD. Peripapillary atrophy was not included in the analysis and all participants required atrophy in the FAF image to be included.

Methods

Image Normalization and Preprocessing

Deep learning algorithms require very large datasets for training on raw images to cover for image acquisition problems that can affect performance. This potential challenge can be addressed by image preprocessing for image standardization, which can greatly reduce the sample size necessary. Problems during FAF image acquisition include illumination (poor uniformity of intensity in the image plane), blurred vision (from involuntary eye movements), physical discomfort (from viewing the blue-light beam), and dark contrasts or “shadowing” (because of vitreous opacities, incorrect adjustments of the camera, or the position of the patient relative to the camera). According to the Heidelberg Engineering’s HRA+OCT Spectralis Manual, the built-in real-time eye-tracking system was designed to minimize eye movement artefacts. The manual recommends obtaining and averaging between six to 24 scans to obtain a good-quality FAF image. Poor image quality may create a misinterpretation or might even render the FAF image uninterpretable. Although standardized protocols like the one stipulated by Heidelberg Engineering do exist, noises
and artefacts may still appear during image acquisition.\textsuperscript{40,41}

It is possible to correct for camera characteristics or other issues with image acquisition quality using preprocessing techniques.\textsuperscript{42} For this dataset, images with extreme artefacts that could not be salvaged with image preprocessing techniques were removed from the dataset. Extreme artefacts here were defined as the presence of overwhelming artefacts that skewed the extraction of information from the image, such as extreme darkness, blurriness or graininess. For the remaining set of images, residual artefacts were removed using the Contrast Limited Adaptive Histogram Equalization (CLAHE) technique.\textsuperscript{43} The CLAHE corrects for different illumination and contrast conditions, as well as improving the edges of objects within the image.\textsuperscript{43} Figure 1 illustrates the conversion of an image using CLAHE. Images in their original form (Fig. 1a) can, technically, be trained on a U-Net architecture. However, the resultant model would be flawed, producing contrast-related errors and outputs that do not accurately isolate lesions. By using CLAHE, it is easier to train the algorithm and for the algorithm to produce high quality outputs in its prediction of lesion areas.

Learning Algorithm

A deep learning model was developed using the U-Net architecture together with appropriate image contrast normalization, hyperparameters and training data. The basic U-Net architecture is illustrated in Figure 2 and consists of contracting (left side) and expansive (right side) pathways. The foundation of the architecture is the Fully Convolutional Network. For the contracting pathway, there is a repetitive pattern of two $3 \times 3$ convolutions, a rectified linear unit (ReLU) and a $2 \times 2$ max pooling operation. At every downsampling step, the number of convolution filters is doubled from the previous step (e.g., step 1 begins with 64 filters, which increases to 128 by step 2). For the expansive pathway, each repeated upsampling step consists of $2 \times 2$ convolutions that halve the number of filters, a concatenation with the correspondingly cropped feature map from the contracting path, and finally two $3 \times 3$ convolutions followed by a ReLU. A $1 \times 1$ convolution is used at the final layer to map each 64-component feature vector to the desired number of classes. The contracting and expansive pathways form a “U” shape, thus aptly giving this architecture its name.\textsuperscript{37} In our implementation, the ReLU-aware He Normal initialization was used. He Normal derives from the research of Glorot and Bengio,\textsuperscript{44} who used a scaled uniform distribution for initialization and assumed activations are linear. Proposed by He et al.,\textsuperscript{45} this initializer is considered to be more sound for ReLU activation, and involves each layer’s weight being initialized in accordance with the size of previous layers.

The adaptive learning rate optimization algorithm, ADAM, was used for stochastic gradient descent and was used with a learning rate of $LR = 3 \times 10^{-5}$. The ADAM optimizer includes bias corrections on both first- and second-order moments.\textsuperscript{46} The learning rate is an optimization hyperparameter that adjusts the weight of the algorithm during training. The learning rate was evaluated by assessment of the Dice loss and quality of segmentation outputs. For the dataset, we found larger learning rates, such as $LR = 10^{-4}$,
Figure 2. U-Net architecture. The U-Net architecture was created for biomedical image segmentation. The U-Net is aptly named because of the arrangements of the filters in a “U” shape. The contracting pathway (left) consists of two $3 \times 3$ convolutions, ReLU activation and $2 \times 2$ max pooling. The expansive pathway (right) consists of $2 \times 2$ convolution, a concatenation with the correspondingly cropped feature map from the contracting pathway, two $3 \times 3$ convolutions and ReLU activation (see Ronneberger et al.37).

produced suboptimal outputs that did not distinctly characterize GA lesion areas.

The batch size was set to 32, the steps per epoch was set to 175, and the number of epochs selected was 80. Batch sizes typically range from 32 to 512. Note that U-Net designers favor small batch sizes to minimize overhead and maximize graphics processing unit memory.37 The regularization effect of small batch sizes contributes to the ability to generalize.47,48 We found 80 epochs to be sufficient in reaching peak model performance and minimizing loss. Learning curves were created using Python’s ggplot (http://ggplot.yhathq.com/).

The hardware implementation of the U-Net was carried out on an operating system with an Intel Core i7-7820HQ CPU @ 2.90GHz. All training, testing, and statistics were performed using Keras (https://keras.io/) and Tensorflow (https://www.tensorflow.org/) using NVIDIA Quadro M1200 Graphics Processing Unit.

Training and Validation

The algorithm performance was evaluated by five-fold cross-validation,49 which is widely used in classification for model assessment and provides estimates of

Figure 3. Fivefold cross-validation. This cross-validation was chosen because it has been empirically shown to yield test error rates not excessively influenced by high bias and variances.

Performance Metrics

The following metrics were used for evaluating the performance of the U-Net algorithm: the DSC, DSC loss, sensitivity, specificity, mean absolute error (MAE), accuracy, recall, and precision.19,55,56
Bland-Altman plots and coefficient of repeatability (CR) were used to measure the difference between ground truth and segmentation results both visually and numerically, respectively. Bland-Altman plots were created using Python’s pyCompare (https://pypi.org/project/pyCompare/) and the unit of measure used was pixels. We further compared ground truth with automation segmentation using Spearman’s correlation coefficient ($\rho$) and plotted an appropriate regression line using Python’s ggplot.

Although DSC was our primary focus as the metric for segmentation performance, the additional evaluation metrics included were used so that results were comparable with other GA-AI findings. For example, in semantic segmentation, a lower sensitivity would suggest undersegmentation where lesion boundaries would not be captured in detail. Conversely, a lower specificity could indicate oversegmentation where lesions are resolved with too much detail possibly caused by noise or artefacts.

The metrics were computed from pixel-level values for true positives (TP), true negatives (TN), false positives (FP) and false negatives (FN). In the context of this study, TP was defined as correctly segmented GA lesion pixels, TN was the correctly identified background pixels, FP was the background pixels mistakenly segmented as GA lesion pixels, and FN was the GA lesion pixels mistakenly identified as background pixels.

The DSC is a spatial overlap index and a validation metric for reproducibility. It measures the agreement between results obtained using ground truth (such as human annotation) and machine-predicted results.

$$\text{DSC} = \frac{2TP}{2TP + FP + FN}$$

DSC loss is simply denoted as

$$\text{DSC}\_\text{loss} = 1 - \text{DSC}$$

Sensitivity (also known as recall) is defined as the proportion of TP pixels found within the lesions.

$$\text{Sensitivity} = \frac{TP}{TP + FN}$$

Specificity is the measure of diagnostic test accuracy and is defined as the proportion of TN pixels found within the background.

$$\text{Specificity} = \frac{TN}{TN + FP}$$

The MAE measures closeness of predictions (observed vs. predicted) and is expressed as

$$\text{MAE} = \frac{1}{n} \sum_{i=1}^{n} |y_i - x_i|$$

where $y_i$ is the prediction and $x_i$ is the true value.

The accuracy of the algorithm is its ability to distinguish different classes (i.e., GA lesion pixel or background pixel).

$$\text{Accuracy} = \frac{TP + TN}{TP + TN + FP + FN}$$

Precision represents the proportion of pixels correctly classified as GA lesions.

$$\text{Precision} = \frac{TP}{TP + FP}$$

Finally, the coefficient of repeatability (CR) measured the difference between ground truth and automated segmentation outcomes.

$$CR = 1.96 \times \sqrt{\frac{\sum (d_2 - d_1)^2}{n}}$$

These metrics were evaluated for every cross-validation fold and compared predicted outcomes to that of human-annotated ground truths. The metrics DSC, sensitivity, specificity, accuracy, and precision all have an outcome range of 0 to 1; the closer to 1 the result, the better the outcome. Conversely, the metrics DSC\_loss along with MAE should ideally be as close to 0 as possible to indicate that loss and error is minimized.

### Qualitative Assessment

While human subjectivity should be accounted for, combining a qualitative assessment along with a quantitative assessment strengthens the evaluation of model prediction. For the purposes of this study, qualitative assessment involves (a) the speed of the algorithm as compared to its human counterpart, and (b) the human visually evaluating machine-generated outcomes and determining whether outputs graphically appear to be accurate.

### Results

A total of 702 FAF images from 51 patients with GA secondary to AMD were included in the study, and whose images were manually annotated. The cohort of images and patients was quite large and diverse.
Figure 4. Learning curves with training/validation loss and accuracy across all fivefolds of cross-validation. (A) Cross-validation 1. (B) Cross-validation 2. (C) Cross-validation 3. (D) Cross-validation 4. (E) Cross-validation 5. The learning curve illustrates a consistent outcome of high accuracy and low loss throughout all fivefolds.
Figure 5. Learning curves for DSC and DSC_{loss} across all fivefolds of cross-validation. (A) Cross-validation 1. (B) Cross-validation 2. (C) Cross-validation 3. (D) Cross-validation 4. (E) Cross-validation 5. The learning curve illustrates a consistent outcome of high DSC and low loss throughout all 5-folds. DSC, Dice similarity coefficient.
Table 1. Geographic Atrophy U-Net Training Results

| Training Set | DSC   | DSC_loss | Sensitivity | Specificity | MAE    | Accuracy | Precision |
|--------------|-------|----------|-------------|-------------|--------|----------|-----------|
| Training Set 1 | 0.9752 | 0.0248   | 0.9949      | 0.8904      | 0.0443 | 0.9904   | 0.9943    |
| Training Set 2 | 0.9918 | 0.0082   | 0.9948      | 0.8925      | 0.0141 | 0.9903   | 0.9939    |
| Training Set 3 | 0.9916 | 0.0084   | 0.9951      | 0.884       | 0.0147 | 0.99     | 0.9934    |
| Training Set 4 | 0.9931 | 0.0069   | 0.9958      | 0.8913      | 0.0122 | 0.9917   | 0.9949    |
| Training Set 5 | 0.9855 | 0.0145   | 0.9967      | 0.785       | 0.0267 | 0.9935   | 0.9963    |

Mean ± SD  0.9874 ± 0.0067  0.0126 ± 0.0067  0.9955 ± 0.0007  0.8686 ± 0.0419  0.0224 ± 0.0121  0.9912 ± 0.0013  0.9946 ± 0.0010

SD, standard deviation.

Table 2. Geographic Atrophy U-Net Validation Results

| Validation Set | DSC   | DSC_loss | Sensitivity | Specificity | MAE    | Accuracy | Precision |
|----------------|-------|----------|-------------|-------------|--------|----------|-----------|
| Validation Set 1 | 0.9512 | 0.0488   | 0.9901      | 0.8529      | 0.0818 | 0.9702   | 0.974     |
| Validation Set 2 | 0.9887 | 0.0113   | 0.9887      | 0.871       | 0.0207 | 0.9838   | 0.9935    |
| Validation Set 3 | 0.9869 | 0.0131   | 0.9938      | 0.6055      | 0.0232 | 0.9824   | 0.9864    |
| Validation Set 4 | 0.9947 | 0.0053   | 0.998       | 0.5557      | 0.0103 | 0.9921   | 0.9938    |
| Validation Set 5 | 0.9678 | 0.0322   | 0.9932      | 0.7452      | 0.0569 | 0.979    | 0.9832    |

Mean ± SD  0.9779 ± 0.0161  0.0221 ± 0.0161  0.9928 ± 0.0032  0.7261 ± 0.1273  0.0386 ± 0.0267  0.9815 ± 0.0071  0.9862 ± 0.0073

SD, standard deviation.

Table 3. Geographic Atrophy U-Net Test Results

| Test Set | DSC   | DSC_loss | Sensitivity | Specificity | MAE    | Accuracy | Precision |
|----------|-------|----------|-------------|-------------|--------|----------|-----------|
| Test Set 1 | 0.9835 | 0.01645  | 0.9937      | 0.8504      | 0.0309 | 0.9883   | 0.9938    |
| Test Set 2 | 0.9916 | 0.0084   | 0.9904      | 0.8613      | 0.0160 | 0.9878   | 0.9970    |
| Test Set 3 | 0.9783 | 0.0217   | 0.9928      | 0.7207      | 0.0390 | 0.9666   | 0.9706    |
| Test Set 4 | 0.9820 | 0.0180   | 0.9824      | 0.7104      | 0.0309 | 0.9738   | 0.9880    |
| Test Set 5 | 0.9548 | 0.0452   | 0.9922      | 0.6060      | 0.0715 | 0.9703   | 0.9693    |

Average ± SD  0.9780 ± 0.0124  0.0220 ± 0.0124  0.9903 ± 0.0041  0.7498 ± 0.0955  0.0376 ± 0.0184  0.9774 ± 0.0090  0.9837 ± 0.0116

SD, standard deviation.

as compared to others in the GA-AI segmentation space.11 The cohort consisted of 99 eyes, 49 left eyes (49.5%) and 50 right eyes (50.5%). A total of 359 images were for the left eye and 343 images were for the right eye. The cohort consisted of 38 females (74.5%) and 13 males (25.5%) with an average age of 76.7 ± 8.9 years. Total follow-up time was 61.5 ± 25.3 months.

The intraclass correlation coefficient for consistency between the two graders was 0.9855 (95% confidence interval [CI]: 0.9298, 0.9971), showing close agreement between the graders. To suit the requirements of the cross-validation, the images were divided into four parts of 140 images and one part of 142 images by random allocation, with a mix of fast and slow progressors.

Learning curves across all five folds with training/validation loss and accuracy are presented in Figure 4 and for DSC and DSC_loss in Figure 5. For quantified training outcomes (Table 1), DSC ranged from 0.9752 to 0.9931, DSC_loss ranged from 0.0069 to 0.0248, sensitivity ranged from 0.9948 to 0.9967, specificity ranged from 0.785 to 0.8925, MAE ranged from 0.0122 to 0.0443, accuracy ranged from 0.99 to 0.9935, and precision ranged from 0.9934 to 0.9963.

For quantified validation outcomes (Table 2), DSC ranged from 0.9512 to 0.9947, DSC_loss ranged from 0.0053 to 0.0488, sensitivity ranged from 0.9887 to 0.998, specificity ranged from 0.5557 to 0.871, MAE ranged from 0.0103-0.0818, accuracy ranged from 0.9702 to 0.9921, and precision ranged from 0.974 to 0.9938.

For quantified test outcomes (Table 3), DSC ranged from 0.9548 to 0.9916, DSC_loss ranged from 0.0084 to 0.0452, sensitivity ranged from 0.9824 to 0.997, specificity ranged from 0.6060 to 0.8613, MAE ranged from 0.0160 to 0.0712, accuracy ranged from 0.9666 to 0.9883, and precision ranged from 0.9693 to 0.9970.

The Bland-Altman plots and CRs (Fig. 6) illustrate graphically that there are minimal differences between the ground truth and segmentation results. The Bland-Altman plots the difference between the ground truth and segmentation output measurements vs. the mean of the two measurements. The Bland-Altman showed a bias of (A) −1238.05 pixels (95% CI agreement: −5052.40, 2576.31), (B) −615.99 pixels (95% CI
Figure 6. Bland-Altman plots and coefficient of repeatability across all fivefolds of cross-validation (in units of pixels).

agreement: $-10788.48, 9556.50$, (C) 1451.87 pixels (95% CI agreement: $-10857.75, 13761.49$), (D) $-1876.28$ pixels (95% CI agreement: $-6607.24, 2854.68$), and (E) $-1115.70$ pixels (95% CI agreement: $-7306.82, 5075.42$). The coefficients of repeatability were (A) 4520.79 pixels (95% CI: 4030.03, 5148.73), (B) 10243.88 pixels (95% CI: 8941.18, 11994.43), (C) 12634.26 pixels (95% CI: 10961.46, 14914.32), (D)
5992.17 pixels (95% CI: 5334.85, 6835.69), (E) 6565.97 pixels (95% CI: 5818.85, 7534.94).

The Spearman’s correlation coefficient and regression line (Fig. 7) reveals that there is a strong positive correlation between ground truth and segmentation measurements. The Spearman’s correlation coefficients were (A) $\rho = 0.9936$ ($P < 0.001$), (B) $\rho = 0.9808$ ($P < 0.001$), (C) $\rho = 0.9249$ ($P < 0.001$), (D) $\rho = 0.9977$ ($P < 0.001$), (E) $\rho = 0.9984$ ($P < 0.001$).

Further to the quantifiable results of the algorithm, we evaluated visually the outputs generated by the U-Net to confirm that lesions were being extracted accurately. Figures 8 and 9 illustrate four sample cases of the U-Net GA lesion output of preprocessed FAF images. The presented cases showed extremely well-outlined lesions. The time in which GA lesions were extracted was compared between humans and the automation method. The average time it took a human, using RegionFinder, to annotate GA lesions was on average 1.04 minutes across all 702 images. The U-Net GA automation takes 6.06 seconds. The qualitative and quantitative assessment coupled illustrate a good performance of the algorithm.

In the current study, the 702 images from 51 persons are comparable with other recent studies using deep learning for GA image segmentation. See, for example, Liefers et al., who used 409 images for model development and evaluation from two cohorts: 87 images
Figure 8. Qualitative assessment of model prediction outcomes. Test cases A and B. In addition to assessing the performance of U-Net quantitatively, we evaluated the performance by visually assessing the degree of accuracy of U-Net-based lesion segmentation. The test cases presented demonstrate a good segmentation outcome.

Table 4. Geographic Atrophy U-Net Patient-Level Segmentation Results

|        | DSC    | DSC_loss | Sensitivity | Specificity | MAE    | Accuracy | Precision |
|--------|--------|----------|-------------|-------------|--------|----------|-----------|
| Training | 0.9931 | 0.0069   | 0.9953      | 0.9634      | 0.0116 | 0.9919   | 0.9951    |
| Validation | 0.9892 | 0.0108   | 0.9895      | 0.9318      | 0.0192 | 0.9831   | 0.9916    |
| Test   | 0.9793 | 0.0207   | 0.9792      | 0.8041      | 0.0382 | 0.9633   | 0.9818    |

from the BMES study (26 participants, 43 eyes) and 322 images from the RS study (149 participants, 195 eyes). The average time patients were observed in our cohort was quite long (61.5 months), which increased sample variability for model training.

The possibility of overfitting was addressed by (a) appropriate selection of hyperparameters, (b) early stopping in training, (c) batch size selection, and (d) its absence confirmed by results from the fivefold cross-validation—which would have revealed significant disparities between training and test results if the model was overfitted. For example, good results for the training phase, but poor results for the testing phase would be indicative of overfitting. Segmentation and cross-validation were carried out at the patient-level in addition to the image-level, using one image per eye for 99 eyes (Table 4). This is an additional check against overfitting, subject to certain statistical assumptions.61,62 We found similar outputs and results with patient-level and image-level segmentation. No evidence of overfitting was found.

The current study and experimental results included a number of features that, in combination, distinguish it from other studies: (a) it is a retrospective case study, under clinical conditions, (b) application of the U-Net deep learning architecture to GA segmentation with hyperparameters tuned to a new set of clinical data, (c) use of FAF imagery from the Heidelberg Spectralis instrumentation, and (d) data preprocessing using an optimal normalization process based on CLAHE. This combination of features does not appear to have been reported elsewhere in the research literature.
The automation of GA lesion segmentation is described using a deep learning algorithm based on the U-Net architecture. The algorithm was assessed both qualitatively and quantitatively. Quantitatively, performance was evaluated against the metrics DSC, DSC loss, sensitivity, specificity, MAE, accuracy, recall, and precision. The U-Net performance on FAF images coupled with preprocessing was successful in GA lesion segmentation. Training, validation, and testing scores were very high, particularly for the main metric of interest—the DSC—where the average DSC for training, validation and testing was 0.9874, 0.9779, and 0.9780, respectively. The highest DSC reported in the literature for GA-AI semantic segmentation was 0.89 for the study by Hu et al., using a level set method on FAF images. The test DSC score of 0.9780 ± 0.0124 produced here compares favorably with other U-Net-based algorithms. Wu et al. reported a DSC score of 0.872 ± 0.66 on SD-OCT and synthesized FAF images, whereas Liefers et al. reported an average DSC score of 0.72 ± 0.26 on CFP. Differences in performance can be ascribed to differences in data quality, different settings for hyperparameters, and the design of image normalization methods.

When comparison is made on the basis of accuracy (the more commonly used outcome evaluation), the algorithm compares favorably with scores reported by Hu et al. (i.e., 0.97 accuracy using kNN with FAF images) and Ji et al. (i.e., 0.986 and 0.995 accuracies using sparse autoencoders with SD-OCT scans). Our accuracies were 0.9912, 0.9815, and 0.9774 for training, validation, and testing, respectively. Similarly, the sensitivity of the algorithm was very good, with values of 0.9955, 0.9928, and 0.9903 for training, validation and testing, respectively. The highest sensitivity performance reported in the literature was 0.983, by Lee et al. (i.e., watershed transform algorithm with FAF images). In this study, qualitative evaluation by visual assessment of machine-generated outputs provided augmentation of scores based on objective metrics. Visually, the U-Net GA automation appears to capture all lesions visually accessible to the human grader. The speed of the automation was far greater

Figure 9. Qualitative assessment of model prediction outcomes. Test cases A and B. In addition to assessing the performance of U-Net quantitatively, we evaluated the performance by visually assessing the degree of accuracy of U-Net-based lesion segmentation. The test cases presented demonstrate a good segmentation outcome.
than the speed of human judgement. For the 702 images in the dataset, the speed of automation was \(~6.06\) seconds to complete each task, whereas the human grader averaged 1.04 minutes.

Metric values (whether DSC, sensitivity, specificity or accuracy) of 0.7–0.8 are typically considered acceptable, whereas 0.8 to 0.9 are considered excellent. Specificities for GA-AI segmentation algorithms ranged from 0.93 to 0.99 in the literature. For this algorithm, the specificities were 0.8686, 0.7261, and 0.7498 for training, validation and testing, respectively. With the results for specificity indicating good performance rather than excellent, there may be a possibility of oversegmentation occurring. Oversegmentation may be due to correctly detecting boundaries of interest within an image (lesion boundaries), but also insignificant boundaries as well. Visually, this may appear as the segmented areas being split up more than necessary. This is in contrast to undersegmentation, where individual segments are merged into singular segments. Future work to address this issue could involve combining the U-Net segmentation algorithm with other AI tools, such as texture discrimination, for improved resolution in spatial analysis.

Some limitations in the study included constraints on the use of the FAF imaging modality, such as image artefacts (e.g., blurriness, shadowing, and poor contrast), discomfort for the patient associated with the blue-light beam, low signal strength, and its potential for toxicity for the retina\(^{41}\). Preprocessing techniques were used to standardize FAF images to address some of these issues. Augmentation of machine learning performance in future may be possible by including other imaging modalities, such as gray-scale SD-OCT images during training, which have been used in the past to quantify atrophy of photoreceptors\(^{63–65}\). In the current study, the cross-validation approach was used to evaluate performance under controlled experimental conditions. In future, more extensive training and application to external datasets are possible using a model-to-data approach, also known as federated learning, as demonstrated by Mehta et al.\(^{36,66}\) This process involves exporting the partially trained deep learning model to different institutions for incremental training, while preserving local data privacy.

Conclusion

Estimation of GA area in FAF images is required to evaluate the severity and rate of progression of GA in clinical presentations. To automate this process, a deep learning approach was developed for semantic segmentation and applied to FAF images, with image preprocessing and normalization by CLAHE. The algorithm produced very high accuracy with very high DSC scores, matching or exceeding human performance in all metrics.

The automation results presented in this article are based on application to clinical data and therefore provide support that the algorithm is suitable for application in clinical settings. Automation of diagnostics for GA has the potential to provide savings with respect to patient visit duration, operational cost, and measurement reliability in routine GA assessments.

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