Annotated retinal optical coherence tomography images (AROI) database for joint retinal layer and fluid segmentation

Martina Melinščak \(^a\,b\), Marin Radmilović \(^c\), Zoran Vatavuk \(^c\) and Sven Lončarić \(^b\)

\(^a\)Department of Mechanical Engineering, Karlovac University of Applied Sciences, Karlovac, Croatia; \(^b\)Department of Electronic Systems and Information Processing, Faculty of Electrical Engineering and Computing, Zagreb, Croatia; \(^c\)Department of Ophthalmology, Sestre milosrdnice University Hospital Center, Zagreb, Croatia

**ABSTRACT**

Optical coherence tomography (OCT) images of the retina provide a structural representation and give an insight into the pathological changes present in age-related macular degeneration (AMD). Due to the three-dimensionality and complexity of the images, manual analysis of pathological features is difficult, time-consuming, and prone to subjectivity. Computer analysis of 3D OCT images is necessary to enable automated quantitative measuring of the features, objectively and repeatedly. As supervised and semi-supervised learning-based automatic segmentation depends on the training data and quality of annotations, we have created a new database of annotated retinal OCT images – the AROI database. It consists of 1136 images with annotations for pathological changes (fluid accumulation and related findings) and basic structures (layers) in patients with AMD. Inter- and intra-observer errors have been calculated in order to enable the validation of developed algorithms in relation to human variability. Also, we have performed the automatic segmentation with standard U-net architecture and two state-of-the-art architectures for medical image segmentation to set a baseline for further algorithm development and to get insight into challenges for automatic segmentation. To facilitate and encourage further research in the field, we have made the AROI database openly available.

**INTRODUCTION**

Age-related macular degeneration (AMD) is an acquired degeneration of the retina that causes significant central visual impairment in its late-stage and is the leading cause of irreversible blindness in people 50 years of age or older in the developed world [1]. The estimated prevalence of any and late-stage AMD in people aged 45–85 years is 8.69% and 0.37%, respectively [2]. The prevalence of late-stage AMD sharply rises to 7.1% in those 75 years or older [3]. Due to the aging of the population, by 2040 an estimated 288 million people will be affected by AMD [2].

AMD is a progressive disease. The early and intermediate stages are usually asymptomatic, characterized by the accumulation of yellow granular deposits beneath the outermost layer of the retina, the retinal pigment epithelium (RPE) (Figure 1(a)). Advanced or late AMD is defined either by the development of atrophy of the RPE and the overlying photoreceptors or by the development of new blood vessels (neovascular membranes) beneath or above the RPE (Figure 1(b)). These new vessels tend to leak or rupture, with subsequent exudation or haemorrhage accumulating in different retinal layers [4]. These two forms of advanced AMD, the former usually referred to as geographic atrophy (GA), and the latter as exudative, wet, or neovascular AMD (nAMD), can occur alone or together, either simultaneously or sequentially, and both forms can lead to significant visual impairment [5]. There is currently no approved treatment available for GA [6]. Intravitreal anti-vascular endothelial growth factor (anti-VEGF) therapy is the mainstay of nAMD treatment [7].

Optical coherence tomography (OCT) is an imaging technique invaluable for diagnosing AMD and guiding AMD treatment because it provides high-resolution, pseudohistological cross-sectional images of the retina and choroid. In nAMD, OCT is used to detect and visualize specific lesions such as intraretinal fluid (IRF), subretinal fluid (SRF), subretinal hyperreflective material (SRHM), and retinal pigment epithelial detachment (PED). These changes represent exudates (IRF, SRF, some cases of SRHM, some forms of PED), haemorrhage (some cases of SRHM, some forms of PED), or neovascular membranes and fibrosis (some cases of SRHM and some forms of PED) [8]. Previous studies have shown these lesions can serve as OCT biomarkers for the visual function or therapy response [9–14]. In comparison to two-dimensional analyses of central B-scans, volumetric analyses of these pathologic lesions in the entire macular area might more precisely predict these outcomes [13,14].

**CONTACT**

Martina Melinščak martinamelinscak@gmail.com Karlovac University of Applied Sciences, Karlovac 47000, Croatia

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Automatic segmentation of retinal layers and of the IRF, SRF, SRHM, and PED (from this point on altogether referred to as fluids) is crucial for detecting and characterization of AMD objectively and in a reproducible way. A reliable automatic OCT system for segmentation is crucial for further development of diagnosing retinal disease. Due to the quantitative analysis of pathological changes, the therapy effectiveness could be predicted [15,16]. The occurrence of intra- and subretinal fluid is an important biomarker that plays a major role in (re-)treatment decisions and is prognostic of visual rehabilitation [17]. Quantitative analysis also allows the prediction of the transition from the middle to the late phase and the prediction of whether in the case of one diseased eye, the disease will affect the other eye [18,19]. Recent studies confirm the correlation between individual fluids and the success of anti-VEGF therapy, making computer segmentation and quantitative analysis even more important [20–22].

Currently, commercially available OCT software algorithms automatically perform retinal layer segmentation to a variable extent and derive basic parameters such as inner or outer retinal thickness, RPE elevation, or central subfield thickness. Although helpful and routinely used to inform clinical decisions, issues of susceptibility to segmentation errors and limited inter-device reproducibility have been raised [23,24].

However, the main issue of these algorithms is the lack of detailed information, as these basic thickness and elevation parameters do not distinguish specific underlying lesions (e.g. retinal fluids in different compartments).

Despite great progress in computer vision and medical image segmentation, a major shortcoming is the lack of publicly available databases of annotated images. In medical image segmentation supervised or semi-supervised methods are still predominant, and their accuracy depends on the quality and scale of annotated data. Most of the published methods are developed on the datasets of images in which no significant pathological changes are present, so it is questionable to what extent the methods are applicable in the case of severe disease. Also, it is difficult to evaluate and compare different methods as there is a lack of publicly available databases of manually labelled images, and results depend significantly on the number of processed images, quality of images (related to the type of OCT device), present disease, etc.

In 2017, MICCAI “Retinal OCT Fluid Challenge (RETOUCH)” was organized [25]. Considering the same dataset and the same metric for evaluation of segmentation and detection, this was a notable improvement in the evaluation of different methods. All applied methods (eight teams were participating) for automatic
segmentation and detection of fluids were based on deep learning methods (segmentation was usually performed with the popular U-net architecture [26]) in combination with other machine learning (ML) methods and image analysis methods. Details can be found on the challenge website [25] and in the accompanying paper [27]. Besides the RETOUCH challenge database, the only publicly available dataset, which some authors use for evaluation of their methods, is the DUKE dataset [28] containing 110 B-scans of 10 DME (diabetic macular oedema) patients acquired with the Spectralis OCT. In a recent review paper, Khan et al. [29] gave an extensive overview of publicly available databases in ophthalmology (up to May 2020) in which authors raise serious concerns about “data poverty” and argue about challenges in data collection.

In this paper, we describe the creation of the openly available Annotated Retinal OCT Images (AROI) database. We give an overview and analysis of the current state of development of the AROI database, with plans for further improvements and new features. It currently consists of 1136 annotated B-scans (from 24 patients suffering from nAMD) and associated raw high-resolution images. The existing research indicates that the approach in which layers and fluids are jointly segmented aims to take advantage of the interdependence of fluids and layers and thus achieves the best possible segmentation [30–35]. Therefore, we have provided annotations for pathological changes (fluid accumulation and related findings) and basic structures (layers). Also, we have presented results for intra- and inter-observer errors to enable the validation of developed algorithms in relation to human variability. To set a baseline for deep learning (DL) methods we have presented results for automatic segmentation with standard U-net architecture and two state-of-the-art architectures (U-net-like and U-net++) in medical image segmentation. The results indicate that there are major challenges for automatic segmentation in the case of severe pathologies. To the best of our knowledge, there are no such publicly available datasets in terms of scale, and with such exhaustive annotations in patients with severe pathology and still, they are crucial for the introduction of automatic segmentation into clinical practice.

Materials and methods

AROI database

Data collection

For the purpose of this study, we collected the database of manually Annotated Retinal OCT Images (AROI database). In collaboration with Sestre milosrdnice University Hospital Center (Zagreb, Croatia), images were collected and annotated by an ophthalmologist. Selection criteria included patients aged 60 years and older diagnosed with nAMD, irrespective of their anti-vascular endothelial growth factor (anti-VEGF) therapy status, with no significant media opacities precluding adequate retinal imaging, and with no other retinal disorders. The concurrent presence of geographic atrophy (GA) of any extent was not an exclusion criterion, since nAMD and GA occur simultaneously or sequentially in a significant number of patients with advanced AMD. From April 2018 to June 2018, 24 consecutive patients were included in the study. Macular SD-OCT volumes were recorded with the Zeiss Cirrus HD OCT 4000 device. Image quality was checked by an ophthalmologist. Overall signal strength of 6/10 or more and absence of any focal shadow artefacts or out of register artefacts was a prerequisite for further analysis. Each OCT volume consisted of 128 B-scans, spaced 47.24 µm apart, with a resolution of 1024 × 512 pixels (pixel size 1.96 µm × 11.74 µm). Retinal fluids and layers were annotated for 1136 B-scans out of a total of 3072 B-scans for 24 patients (37% of B-scans were annotated). Annotations were not provided for each B-scan. The central 10 B-scans around the foveal centre were annotated for each patient, as visual acuity mostly depends on the pathological changes in this area, while more eccentric B-scans were annotated at the ophthalmologist’s discretion: in case the adjacent B-scans were deemed similar, the annotations were skipped (performed for every 2nd to 10th scan, depending on the extent and complexity of pathological changes). The average number of annotated scans per patient was 47.3 ± 25.7.

Images within the database are available in PNG format and organized so that each filename is associated with a patient number (1–24) and B-scan ordinal number (0–127) (e.g. patient12_123.png). All raw images, and not just labelled ones, are available to enable 3D automatic segmentation. At this phase, images are not divided into training and test sets.

Data collection adhered to the tenets of the Declaration of Helsinki and the standards of Good Scientific Practice of Sestre milosrdnice University Hospital Center (Zagreb, Croatia). All patients signed informed consent, the images are anonymized and do not contain any additional information about patients. The presented study was approved by the Ethics Committee of the Sestre milosrdnice University Hospital Center (EP-3272/18-11) and the Faculty of Electrical Engineering and Computing (Zagreb, Croatia).

Fluids and layers annotations

All annotations were done by one expert ophthalmologist. The lesions of interest and annotated in this study were the intraretinal fluid (IRF), subretinal fluid (SRF), subretinal hyperreflective material (SRHM), and retinal pigment epithelial detachment (PED) (Figure 2). Although some of these lesions can reflect processes such as fibrotic scars or fibrovascular membranes, as
they most often reflect exudation or haemorrhage, the term “fluids” was liberally used here to refer to all these lesions. Additionally, as SRF and SRHM share the same location characteristics, and often reflect the same exudative process, the distinction between SRF and SRHM was not performed in this study, and they were annotated jointly as SRF. In the future, development of the enhanced segmentation method is planned in order to discriminate between SRF and SRHM, as well as between different types of SRHM. As the range of SRHM in AMD and other macular disorders includes a number of different lesions, such as neovascular membranes, fibrosis, exudate, haemorrhage, and lipofuscin like material, a new, more extensive dataset supported with multimodal imaging is in preparation.

To detect IRF, SRF, and PED, the knowledge of their location within or outside specific retinal layers can be used to facilitate their detection and differentiation. The retina is histologically divided into 10 layers: (1) internal limiting membrane (ILM), (2) retinal nerve fibre layer (RNFL), (3) ganglion cell layer (GCL), (4) inner plexiform layer (IPL), (5) inner nuclear layer (INL), (6) outer plexiform layer (OPL), (7) outer nuclear layer (ONL), (8) external or outer limiting membrane (ELM or OLM), (9) photoreceptor layer, and (10) retinal pigment epithelium/Bruch’s membrane complex (RPE/BM). The layers visible in the OCT scan have been correlated to these histological layers, with the exception of a few additional zones observed in the photoreceptor layer for which the exact histological counterpart is not yet defined [36].

The IRF is a localized extracellular intraretinal fluid accumulation seen as a hyporeflective area located anywhere between ILM and ELM, the SRF is a subretinal fluid accumulation seen as a hyporeflective area between photoreceptor layer and RPE (the SRHM is seen in the same location but as a homogenously or inhomogenously hyperreflective area), and the PED is either a hyporeflective accumulation of fluid or hyperreflective accumulation of other material between RPE and BM (the RPE/BM complex is separated in the case of PED) [8]. However, as pathological processes such as those in AMD can lead to extensive changes in retinal structure, it is often impossible for a reader to determine all the layers in the OCT scan reliably. Therefore, we traced ILM, the inner boundary of RPE, BM, and the boundary between IPL and INL (IPL/INL), as these layers/boundaries could be readily determined in virtually all images (Figure 2). While ILM, the inner boundary of RPE, and BM were chosen as pertinent for IRF, SRF, and PED localization, the IPL/INL boundary was chosen as it could be used to locate the foveal centre in case of an eccentric scan (in patients with poor fixation) or loss of normal foveal depression (in patients with extensive foveal oedema, elevation, traction, or parafoveal atrophy). As the foveal centre consists only of ILM, ONL, photoreceptor layer, and RPE/BM, while other layers taper towards the foveal centre, the centre can be defined as the point of least distance between the ILM and any of these other layers, including the IPL/INL boundary. In our dataset all scans were centred at the fovea and this step was not needed.

Figure 2. From left to right: example of the image with annotated boundaries, an image with annotated fluids, and an image prepared for semantic segmentation (with eight classes).
Images prepared for semantic segmentation
As simultaneous segmentation of layers and fluids should give better results than separate segmentation of fluids only or layers only, images were prepared as shown in Figure 2 thus reducing the problem of segmentation to semantic segmentation with eight classes: area above ILM (vitreous), area between ILM and INL/IPL, area between IPL/INL and RPE, area below the BM (choroid), and three retinal fluids (PED, SRF, IRF).

Models for automatic segmentation
Architectures
Since its introduction in 2015, the U-net architecture [26] and its various modifications have been the most used architectures for medical image segmentation. The U-net architecture consists of an encoder (contractive path), decoder (expanding path), and skip connections which enable simultaneous capturing of context and localization as it is shown in Figure 3. We will not explain it in detail as it is a well-known architecture for medical image segmentation. We used standard U-net architecture to set a baseline for automatic segmentation.

Although many modifications of U-net architecture were proposed [34,37–39] we chose two state-of-the-art architectures to get further insights into challenges for automatic segmentation. The groundbreaking improvement in computer vision was achieved with ResNet architecture [40] and DenseNet architecture [41], where both reached breakthrough results in classification on the ImageNet dataset [42]. The logical step was to improve the U-net architecture in the ResNet and DenseNet style fashion. We opted for U-net-like architecture [43] and U-net++ [44] as two state-of-the-art architectures. The former combines the good sides of U-net and ResNet architectures and the latter is inspired with DenseNet architecture. A recent paper by Isensee et al. [45], where they proposed a nnU-net (out-of-the-box tool) for biomedical image segmentation that uses U-net-like architecture was another argument for our choice of architecture.

The U-net-like architecture is shown in Figure 4. In the down-sampling path, residual blocks contain convolutional filters (3 × 3) followed by batch normalization (BN) [46] and a ReLU activation function. Down-sampling is accomplished by max-pooling (MP) which reduces image size by half. Skip connection in the residual block is not just an identity connection but it contains a convolutional filter (1 × 1) with strides equal to 2 and in that way is achieved down-sampling (image size is reduced by half, same as with MP in the main branch of the residual block). At the end of the residual block, the outputs from the main branch (also called a layer) and the skip connection are summarized. Residual blocks in the decoder are implemented in a similar manner: in a layer (the main branch) there are transpose convolutions (3 × 3), BN, activation function (ReLU), and up-sampling (which double the size of the image). In the skip connection, there is a convolution (1 × 1) and up-sampling.

The U-net++ architecture (a nested U-net architecture for medical image segmentation) is shown in Figure 5. It is seen how U-net architecture is enhanced with dense blocks and convolution layers between the encoder and decoder. The purpose of modified skip pathways is to reduce the semantic gap between the feature maps of the encoder and decoder before merging. Each circle represents a set of convolution operations. The shaded part shows the original U-net architecture, while the rest (middle part) shows the difference from the original architecture: dense convolution blocks on the skip pathways and deep supervision [44].

Training
Original images size 1024 × 512 pixels were resized to 512 × 256 pixels. Categorical cross-entropy loss was
used to train all models. The batch size was set to 4 (we obtained worse results with a larger batch size). The AdaBound optimizer [47] was used (as it combines advantages of Adaptive Moment Estimation (Adam) and Stochastic Gradient Descent (SGD)). Number of trainable parameters for standard U-net, U-net-like, and U-net++ architectures are 7,764,744, 8,230,536, and 9,041,832, respectively. Early stopping was used to prevent overfitting.

K-fold cross-validation was used where each fold contains images from four patients (1st fold contains images from patients 1 to 4, 2nd fold contains images from patients 5 to 8 and so on). K equals 6 in our procedure since, in that way, the test set share is approximately 15% as is a recommendation and common practice in a small data regime. We do not recommend splitting the sets of images from the same patient across training, validation, and test set as adjacent B-scans are similar and that would lead to overestimated validation of the method.

The models were trained on Google Colab [48] with a GPU. The models were implemented in Python, using the Keras library with the TensorFlow backend.

Results

Inter- and intra-observer error

For the purpose of calculating inter- and intra-observer error, annotations were additionally made for 75 B-scans (randomly chosen from the existing dataset). To calculate intra-observer error, the same expert who made all annotations (1st expert) made a re-annotation with no reference to previous annotations, and with a time delay (3 months after finishing the first annotations) for the 75 B-scans. To calculate inter-observer error, annotations were made by another expert (2nd expert) for the same 75 B-scans. Figure 6 shows two examples: an example of good and bad matching. Differences of opinion among experts can be observed. This can partly be explained by the poor quality of images (a large amount of speckle-noise), and by the fact that in the face of extensive disturbance of normal retinal structure, certain OCT findings cannot be reliably discerned or localized without normal anatomic landmarks. Also, with suboptimal image quality, the annotated changes could be confounded with other pathological phenomena (hyperreflective foci, pseudodrusen, outer retinal tubulations, etc.).

Model prediction errors

Some examples of the automatic segmentation results are shown in Figure 7. It is visible that results are good in case there are no significant pathological changes and deformation in retinal structure. In case there are pathological changes, segmentation predictions are deficient in preserving the topology. Also, fluids segmentation should be enhanced. The main cause is the low representation of pixels belonging to these classes in the total number of pixels, especially when it comes to IRF (not present in all patients nor in all B-scans; in addition, it is regularly smaller than SRF and PED).
**Figure 6.** From left to right: raw image, reference (annotations from 1st expert), re-annotations from 1st expert with time delay, annotations from 2nd expert. In the first row, there is an example with good matching. In the second row, there is an example with some differences even between annotations from the same expert (in annotations of PED and SRF) and between two experts (different annotations for fluids). Images are cropped and only the ROI is visible.

**Figure 7.** Three examples of the segmentation results. First row: a case with less pronounced pathological changes. Second row: a case with more pathological changes (PED and SRF are present). Third row: a case with extensive pathological changes (PED, SRF, and IRF are present and there is a large distortion of layers). From left to right: raw image, expert annotation (mask), the prediction from the standard U-net architecture, the prediction from the U-net-like architecture, and the prediction from the U-net++ architecture. Images are cropped and only the ROI is visible.
Comparison of inter-observer error and model prediction error

We use the Dice score to evaluate results as it is a similarity measure often used as a metric in the segmentation of medical images. It is calculated according to Equation (1) where TP is true positive, FP false positive, and FN false negative.

\[
DSC = \frac{2TP}{2TP + FP + FN} \tag{1}
\]

In Table 1 there are reported Dice scores (mean and standard deviation) for each class and for the inter- and intra-observer error, as well as the prediction errors from the U-net, U-net-like, and U-net++ models. The same results are shown in Figure 8 with bar graphs. We calculated the Dice score for each patient in test fold (that means four patients) but only for those B-scans with a reference segmentation. In case fluids (either SRF or IRF) were not present on a single B-scan of the patient, the Dice score is exempted when calculating the mean and standard deviation of Dice score for that class and that patient. We found it more appropriate than setting it to zero or one as it would lead to over- or underestimating the metric value. In the case of patients in whom fluids were present on some of the B-scans but not all, for B-scans where they were not present the value for Dice score was set to zero (the Dice score is not defined in the case of zero division which is a common situation in the absence of the class). It can be observed that for the inter-observer case and in all cases of the automatic segmentation the biggest errors occur in class 3 (surface between RPE and BM) and in classes that represent fluids (PED, SRF, and IRF).

One of the factors that contribute to the complexities of automatic segmentation is significant class imbalance: the background (area above ILM and under BM) occupies as much as 83.26% in the total number of pixels, while IRF occupies only 0.12%; surface between RPE and BM occupies 1.07%, SRF 1.05%, PED 1.5%. Further on, out of a total of 1136 scans, PED, SRF, and IRF are present in 1014 (89.26%), 648 (57.04%), and 229 (20.16%) B-scans, respectively.

As Dice score is not an appropriate metric in case of high class imbalance (the Dice score for regions above the ILM and below BM will always be close to one), we also provided results for evaluation in case of converting the layer-segmentation task into a boundary detection problem. In Table 2 there are reported mean square errors (MSE) with belonging standard deviations in the inter-observer case, the intra-observer

| Table 1. The Dice score (mean and standard deviation) in the inter-observer case, the intra-observer case, for standard U-net model, U-net-like model, and U-net++ model. |
|-----------------|--------|--------|--------|--------|--------|--------|--------|--------|
|                 | Above ILM | ILM-IPL/INL | IPL/INL-RPE | RPE-BM | Under BM | PED | SRF | IRF |
| Inter-observer  | 0.982 (0.072) | 0.950 (0.111) | 0.948 (0.112) | 0.699 (0.129) | 0.989 (0.114) | 0.860 (0.301) | 0.876 (0.366) | 0.735 (0.280) |
| Intra-observer  | 0.998 (0.003) | 0.973 (0.008) | 0.970 (0.117) | 0.778 (0.092) | 0.998 (0.001) | 0.912 (0.242) | 0.924 (0.331) | 0.844 (0.140) |
| Standard U-net  | 0.995 (0.011) | 0.950 (0.028) | 0.923 (0.083) | 0.669 (0.129) | 0.988 (0.016) | 0.638 (0.173) | 0.513 (0.287) | 0.480 (0.241) |
| U-net-like      | 0.995 (0.004) | 0.899 (0.040) | 0.890 (0.066) | 0.476 (0.132) | 0.988 (0.014) | 0.333 (0.139) | 0.372 (0.293) | 0.037 (0.061) |
| U-net++         | 0.992 (0.011) | 0.944 (0.032) | 0.924 (0.064) | 0.641 (0.133) | 0.986 (0.017) | 0.622 (0.159) | 0.487 (0.280) | 0.419 (0.274) |

Figure 8. The Dice scores (mean and standard error of the mean) for inter-observer variability, for standard U-net model, U-net-like model, and U-net++ model.
Table 2. The evaluation of the layer-segmentation task as a boundary detection problem: the mean square error (MSE) with belonging standard deviations in the inter-observer case, the intra-observer case, for standard U-net model, U-net-like model, and U-net++ model.

|                  | ILM | IPL/INL | RPE | BM   |
|------------------|-----|---------|-----|------|
| **Inter-observer** | 5.87| 20.46   | 51.74| 12.77|
|                  | (3.68) | (47.71) | (148.15) | (17.79) |
| **Intra-observer** | 2.10| 5.93    | 8.28 | 5.17 |
|                  | (1.43) | (4.80)  | (9.33) | (6.68) |
| **Standard U-net** | 6.23| 32.55   | 60.22| 15.88|
|                  | (3.88) | (50.22) | (173.23) | (19.87) |
| **U-net-like**   | 6.51| 40.34   | 65.47 | 24.56|
|                  | (4.22) | (56.67) | (177.23) | (31.22) |
| **U-net++**      | 6.02| 37.13   | 61.55 | 17.11|
|                  | (4.01) | (54.55) | (165.33) | (20.12) |

case, for standard U-net model, U-net-like model, and U-net++ model. Values are shown in pixels where each pixel corresponds to 1.96 μm along the axial (Z) axis.

Our research suggests that more complex architectures result in only slightly enhanced outcomes, no enhancement at all, or worse outcomes. For classes that represent fluids, the results are noticeably worse, compared to human error. Better results are obtained with U-net++ architecture than with U-net-like architecture, probably because U-net-like architecture lacks skip connections between encoder and decoder (skip connections only exist in each residual block within encoder/decoder) while there are dense blocks and convolutional layers between encoder and decoder in U-net++ architecture. Preliminary, we could conclude that better segmentation accuracy cannot be obtained only with more complex architectures, but rather using some of the efficient techniques in case of distinct class imbalance and the need for preserving the topology. Also, due to the suboptimal quality of images, preprocessing could help in achieving better accuracy.

**Discussion**

The AROI database contains a large sample of 1136 B-scans with exhaustive annotations (both fluids and retinal layers were annotated by an expert ophthalmologist). As images are collected from patients suffering from nAMD where pathologic biomarkers and large distortion of the retinal structure are present, automatic segmentation of such images presents a significant challenge. Figure 9 shows the confusion matrix for inter-observer error, and for model (U-net, U-net-like, and U-net++) prediction error. In the case of automatic segmentation, it is observed that IRF is often misclassified as the surface between IPL/INL and RPE (class 2) as it is in that area. In case the IRF is smaller or does not differ significantly in intensity and texture from the surface between IPL/INL and RPE, the model does not recognize it as a separate class. Also, it is observed that in a similar way SRF is often misclassified as PED or surface between IPL/INL and RPE. PED is mostly misclassified as an area under BM, and with further inception of individual predictions, it is apparent that it happens when Bruch’s membrane is not clearly visible (due to geographic atrophy or some other pathological changes). These results of automatic segmentation for three models (standard U-net architecture and two state-of-the-art architectures) can serve as a baseline for further development of deep learning models.

Also, intra- and inter-observer errors were calculated to enable the validation of algorithms for automatic segmentation. However, it is still not clear what level of segmentation accuracy we need in clinical practice as manual segmentation is rarely performed in clinical practice. There is probably no universal rule, and the required accuracy of segmentation will depend on the purpose — whether it is a diagnosis or prediction of the outcome of anti-VEGF therapy or prediction of another eye disease.

We have limited ourselves to collecting images from only one type of OCT device and from patients suffering from one type of disease. However, developing algorithms on as large and as diverse databases as possible would provide for more robust algorithms that could be implemented in commercial OCT device software. We hope that open access will become a common practice for the majority of research groups in the future and that online image collections and repositories will contribute to building a single database covering various diseases, various types of devices, and various retinal structures annotated from different experts.

![Figure 9](image-url)
As a lack of publicly available databases is one of the major obstacles to introducing AI and deep learning to ophthalmology, we consider the development of the AROI database as a step forward to introducing automatic segmentation in clinical practice and thus enabling quantitative analysis and more successful diagnosis and therapy.

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Disclosure statement
No potential conflict of interest was reported by the author(s).

Data availability statement
Code is available on GitHub: https://github.com/mmelinscak/OCT-images-segmentation. Dataset is available at: https://ipg.fer.hr/ipg/resources/oct_image_database [49] and https://doi.org/10.17605/OSF.IO/5WYR3 [50].

ORCID
Martina Meliňščak http://orcid.org/0000-0001-5128-3213

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