Genetic U-Net: Automatically Designed Deep Networks for Retinal Vessel Segmentation Using a Genetic Algorithm

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Abstract—Recently, many methods based on hand-designed convolutional neural networks (CNNs) have achieved promising results in automatic retinal vessel segmentation. However, these CNNs remain constrained in capturing retinal vessels in complex fundus images. To improve their segmentation performance, these CNNs tend to have many parameters, which may lead to overfitting and high computational complexity. Moreover, the manual design of competitive CNNs is time-consuming and requires extensive empirical knowledge. Herein, a novel automated design method, called Genetic U-Net, is proposed to generate a U-shaped CNN that can achieve better retinal vessel segmentation but with fewer architecture-based parameters, thereby addressing the above issues. First, we devised a condensed but flexible search space based on a U-shaped encoder-decoder. Then, we used an improved genetic algorithm to identify better-performing architectures in the search space and investigated the possibility of finding a superior network architecture with fewer parameters. The experimental results show that the architecture obtained using the proposed method offered a superior performance with less than 1% of the number of the original U-Net parameters in particular and with significantly fewer parameters than other state-of-the-art models. Furthermore, through in-depth investigation of the experimental results, several effective operations and patterns of networks to generate superior retinal vessel segmentations were identified.

Index Terms—Convolutional neural networks (CNNs), Genetic algorithms (GAs), Retinal vessel segmentation, Neural architecture search (NAS)

I. INTRODUCTION

The retinal vascular system can be observed non-invasively in vivo in humans [11]. In addition, changes in the vasculature are often associated with certain diseases, leading ophthalmologists and other physicians to consider fundus examination a routine clinical examination [2, 3]. Many diseases can be diagnosed and tracked [2] by observing the retinal vascular system. Pathological changes in retinal vessels can reflect either ophthalmology diseases or other systemic diseases, such as wet age-related macular degeneration and diabetes [4]. Diabetic retinopathy can lead to the growth of new blood vessels, and atherosclerosis [5] associated with wet age-related macular degeneration can cause the narrowing of blood vessels. Moreover, the retinal vascular system of each eye is unique. Without pathological changes, it does not alter throughout the lifetime. Hence, observation of the retinal vascular system can also be applied in biometrics [6, 7]. Through retinal vessel segmentation, relevant morphological information of retinal vascular trees (such as the width, length, and curvature of blood vessels) can be obtained [8]. Consequently, precise retinal vessel segmentation is significant. However, owing to the complexities of retinal vascular structures, manual inspection is subjective, time-consuming, and laborious [9, 10]. Therefore, developing an effective algorithm for the automated segmentation of retinal vessels to support ophthalmologists in clinical assessment has been of great interest.

Due to the complexities of fundus images, automated segmentation of retinal vessels is challenging. First, in fundus images, retinal vessels are difficult to distinguish from the background because of the subtle difference between the vascular region and background. Second, the structures of vascular trees are also complicated, with many cross-connected and minuscule vessels. Third, other factors such as pathological exudates and uneven illumination render segmentation difficult. Compared with the methods [11–15] based on traditional image processing or handcrafted features, the deep convolutional neural network (DNN) methods [16–20] offer advantages in dealing with these complications. In particular, U-Net [21] and its variants [8, 10, 22–26] have recently become the mainstream models used in retinal vessel segmentation.

However, these U-Net variants are designed manually and have two main shortcomings that constrain them in clinical applications. First, they still have deficiencies in extracting the features of vascular trees from complicated fundus images, which limits their capability of dealing with challenging cases (e.g., lesions areas, low contrast micro-vessels). Second, to improve segmentation performance, these manually designed U-Net variants tend to have many parameters. However, because constructing vessel segmentation datasets is laborious and highly dependent on expertise, only a small amount of annotated data is available. A network architecture with too many parameters is prone to overfitting when using insufficient training data and usually has high computational complexity. Moreover, the original U-Net and its variants utilize one or two types of identical blocks to form the architecture. However, different blocks in a U-shaped network are required to process different patterns of features in fundus images. Therefore, having the same structures for different blocks may not be optimal, as it limits the feature extraction ability and parameter...
Conversely, designing improved network architectures manually requires rich domain-specific knowledge and considerable time, especially when only a small number of parameters is expected. Therefore, this study developed a Neural Architecture Search (NAS)-based approach to automatically discover network architectures tailored for retinal vessel segmentation. Compared with natural images, fundus images have unique features \[26\]; therefore, the success of NAS in natural image segmentation \[28\] is not immediately transferable to retinal vessel segmentation. Several attempts have been made to deploy NAS in 2D medical image segmentation \[29, 30\], but the search spaces of these methods are highly constrained and usually include the repeated stacking of identical blocks with no topology optimization. The generated model retains many parameters, which renders it unsuitable for retinal vessel segmentation, which, with only a small amount of labeled data available is much more sensitive to overfitting. For NAS, the search space designed by human experts is crucial for the performance of the discovered architectures \[31, 32\].

In the proposed Genetic U-Net, we first define a specific search space and then realize the automated design of network architectures with an improved genetic algorithm (GA). When designing the search space, two key aspects are considered 1) defining a condensed but flexible search space, and 2) restricting the total number of architecture-based parameters. Since it is based on a U-Net backbone architecture, the search space is condensed to prevent the algorithm from having to search in an open-ended search space. It is flexible because different blocks of the U-Net architecture are diversified and optimized individually. As diverse optimized internal block topologies can be found in the overall architecture of U-Net, a model with superior performance yet far fewer parameters can be identified. Restricting the total number of architecture-based parameters of the possible architectures in the search space can reduce the computational requirements (e.g., GPU memory) and facilitate the acquisition of high-performance network architectures with even fewer parameters. Moreover, to improve the search efficiency of NAS, an improved GA is suggested with enhanced crossover and selection operations.

The importance of this research is described as follows: First, to our knowledge, this is the first instance of evolutionary NAS being applied to retinal vessel segmentation. Second, the proposed method exploits the potential of the U-shaped encoder-decoder structure and investigates whether an exceptional DNN can be designed with very few parameters and without using complex mechanisms (e.g., multiple encoders, multiple decoders, cascade structures, attention mechanisms). Finally, compared with previous CNNs that have been applied to retinal vessel segmentation, the proposed method can reduce the multiplicity of redundant parameters, which is important when training data are limited. Extensive experiments were performed to show that Genetic U-Net can design compact architectures that perform better than state-of-the-art models.

- We propose a novel automated design method for the U-shaped CNN architecture based on a specific search space and an improved GA, which enables us to acquire compact network architectures that outperform existing ones in retinal vessel segmentation.
- Through observation and analysis of the discovered architectures, we observed that some patterns and operations can significantly improve the performance of retinal vessel segmentation. The findings may lead to new insights into the design of network architecture.
- Compared with state-of-the-art models, the found models offered superior performance on several public datasets with the least parameters (less than 1% of the original U-Net).

II. RELATED WORK

A. Retinal Vessel Segmentation

Originally, researchers used traditional image processing techniques to segment retinal vessels, such as thresholding segmentation \[12\] or certain morphological operations \[11, 13\]. Under different circumstances, many of these methods’ hyperparameters had to be re-adjusted to achieve satisfactory segmentation results. Later, several learning-based methods \[14, 15\] incorporating handcrafted features were applied to this task. Since these handcrafted features do not have sufficient generalization ability to represent characteristics in a variety of complicated fundus images, these methods are misled by extreme cases (e.g., lesions areas and low contrast microvessels). Compared with the above methods, the DNNs-based methods \[16–20\] demonstrate particular advantages in dealing with complexity in fundus images, where the features are learned directly from the training data.

New state-of-the-art methods \[6, 10, 22, 26, 33\] for retinal vessel segmentation are dominated by deep learning models, especially U-Net variants. Yan et al. \[35\] adopted a joint loss to provide supervision information for U-Net, with two parts responsible for pixel-wise loss and segment-level loss. The joint loss can improve the capability of the model to balance the segmentation between thick and thin vessels. To better capture the diverse morphologies of vascular trees, Jin et al. \[3\] proposed DU-Net that replaces traditional convolution with deformable convolution. Wu et al. \[3\] designed a novel inception-residual block for a U-shape network and introduced four supervision paths with different convolution kernel sizes to utilize multi-scale features. Alom et al. \[24\] proposed R2U-Net with a recursive residual layer based on cyclical convolutions to capture features. Mou et al. \[9\] embedded dense dilated convolutional blocks between encoder and decoder cells at the same levels of a U-shape network and used a regularized walk algorithm to post-process model predictions. Dual U-Net proposed by Wang et al. \[10\] had two encoders. One encoder path was for extracting spatial information and the other was for extracting context information. A novel module was also proposed to combine the information of the two paths. Laibacher et al. \[23\] utilized pre-trained components of MobileNetV2 \[34\] on ImageNet \[35\] as the encoder of U-shape network and introduced novel contractive bottleneck blocks for the decoder, which achieved better performance, less computational cost, and faster inference speed. Gu et al. \[25\] proposed CE-Net by adopting multiple
convolution branches with different receptive fields to capture more high-level information and preserve spatial information for segmentation. Mou et al. [26] included a self-attention mechanism in the U-shape encoder-decoder to improve the hierarchical representation capture ability of the model. The abovementioned networks, which are unusually complex and have many parameters, were manually designed. To increase the feature extraction capability, some networks must be pre-trained with external datasets. In this study, we automatically design the network architectures that can perform well in extracting features from fundus images but require fewer parameters and no pre-training.

B. Neural Architecture Search

Neural architecture search (NAS) is an effective technique that assists end-users to design effective deep networks automatically. At present, depending on the optimization methods used, the three main categories of NAS are: (1) The methods based on reinforcement learning [36–38] that formulate NAS as a Markov decision process. A controller is used to sample the architecture and learn to generate improved architectures from a process of continuous trial and error; (2) The methods based on evolutionary algorithms [39, 40] that formulate NAS as an optimization problem and encode the architectures. Increasingly competitive architectures are generated by applying some genetic operations (e.g., crossover, and mutation) and will be retained as offspring in the next generation. The architectures are optimized from generation to generation until those with satisfactory performance are obtained; (3) For differentiable neural architecture search [41, 42], each operation option is assigned a weight coefficient. The parameter weights of the architecture and the weights of the operation options are optimized alternately by gradient descent. NAS has achieved great success in the natural image and medical image analysis [29, 30, 43–46], including some works that apply NAS to medical image segmentation. In the works of [30, 43–44], for medical image segmentation, the hyperparameters and operations of each layer of the building blocks were optimized, but the topology of the block was relatively fixed. Additionally, in the works [29, 46], the structure and operations of one or two types of building blocks were optimized, and then the architecture was constructed by repeatedly stacking them. However, in our work, the topology and operations of each block could be different yet simultaneously be optimized flexibly.

C. Genetic Algorithms

Genetic Algorithms (GAs) [47], one of the evolutionary algorithms, are metaheuristics inspired by evolution. They are widely used to find high-quality solutions for optimization problems (e.g., TSP [48], neuroevolution [49, 50], NAS [39]) by executing operators such as mutation, crossover, and selection. In addition, GAs have been applied to many medical image analysis tasks, such as segmentation [51], registration [52], disease detection [53], and image denoising [45]. Fan et al. [51] used GA to overcome numerical instability of the active model-based volumetric segmentation of brain images. Matsopoulos et al. [52] employed GA to optimize the parameters of affine transform to better register multimodal retinal images. Quellec et al. [53] adopted GA to optimize the parameters of wavelet transform for the detection of micro-aneurysms in retina photographs. Recently, Liu et al. [45] utilized GA to find optimal hyperparameters and CNN architectures for medical image denoising. However, to our knowledge, there are still no works on applying GAs to optimize CNN architectures for vessel segmentation in medical image analysis.

III. THE PROPOSED METHOD

In this section, we present the proposed method in detail. We first introduce the search space of the architectures, then explain the method of encoding an architecture into a binary string, and finally explain the genetic algorithm with evolutionary operations (e.g., crossover, mutation, and selection) searching for competitive architectures.

A. The search space and encoding

1) Backbone of the Search Space: As shown in Fig. 1(a), U-Net is composed of an encoder E and a decoder D. Both encoder E and decoder D contain several blocks, such as eᵢ (i = 0, 1, 2, 3) and dⱼ (j = 0, 1, 2). From top to bottom, U-Net is divided into different stages Sₖ (k = 0, 1, 2, 3), and the feature dimensions are constant at each stage. Skip connections are adopted in all, except for the last, stages to provide features with different semantic information extracted by the encoder to the decoder, which strengthens the connections between the encoder and the decoder and alleviates the vanishing gradient problem [54, 55] in model training. Decoder D must fuse features from the skip connections and up-samplings, and the two commonly used feature fusion operations are: concatenation or element-wise addition. Although the original U-Net employs concatenation for feature fusion, some U-Net variants [56] achieve good results using element-wise addition. Fig. 1(b) illustrates their main differences. Compared with element-wise addition, concatenation generates larger feature maps, which increase computational complexity. To mitigate such complexity in this study, we selected the addition operation for feature fusion.

The successful applications of U-Net and its variants reveal that the U-shaped encoder-decoder structure is highly applicable. Therefore, herein, we used the U-shaped encoder-decoder structure as the backbone. As shown in Fig. 1(a), it consists of seven blocks and four stages. Using this structure, satisfactory architectures can be found by adjusting the internal structures of these building blocks.

2) The Building Blocks and Their Encoding: In original U-Net, the internal structure of each block is composed of two basic layers (3 × 3 conv + ReLU). Usually, the U-Net variants [24, 25, 57, 58] improve their performance by adjusting the internal structures of blocks (e.g., ResNet block [59], DenseNet block [27], and InceptionNet block [60]), which illustrates the importance of the internal structures of blocks. The internal structures of the building blocks are represented similarly to Genetic CNN [39], which is sufficiently flexible to
The proposed approach has four fundamental differences from Genetic CNN [39]. First, the studied task is different. Genetic CNN is applied to natural image classification, while the proposed method is designed for retinal vessel segmentation. Second, the search space is different. The U-shaped encoder-decoder structure is employed as the backbone in our work but not utilized in Genetic CNN. In addition, the operations of the nodes can be optimized instead of a fixed operation in Genetic CNN. In Genetic CNN is applied to natural image classification, while the proposed method is designed for retinal vessel segmentation. The differences between these operational sequences are reflected by the convolutional kernel size, activation functions, activation types (pre-activation or post-activation), and normalization types (e.g., whether instance normalization is utilized). Therefore, four-bit binary encoding is utilized to represent the operational sequences shown in Table I. We assume that each node in one block has the same operation sequence; therefore, each block gene consists of an operation gene with four bits (to represent sixteen different operational sequences) and a connection gene (as shown in Fig. 3(b)). Overall, seven block genes together constitute the genotype of architecture (as shown in Fig. 3(b)), which indicates that the different blocks can have diverse structures and operations. Therefore, the designed search space is highly flexible.

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parameters. Last, an improved GA is proposed to promote search efficiency.

3) Number of Architecture-based Parameters: The number of the architecture parameters in the search space is closely related to the number of stages in the U-shape backbone, number of intermediate nodes in each block, and number of channels for convolution operations in the nodes. After selecting the backbone adopted for the search space to constrain the number of the architecture parameters and maintain architectural flexibility, the maximum allowed number of intermediate nodes in each block and the number of channels for convolution operations in the nodes are set to 5 and 20, respectively. Most network architectures in this search space have fewer than 0.4 Million parameters, while normal architectures used for retinal vessel segmentation have more than several Million or even dozens of Million parameters. By restricting the number of architecture-based parameters of the possible architectures within a range of smaller values allows better architectures with greater parameterization efficiency to be found.

B. Evolutionary Algorithm

Genetic U-Net follows an iterative evolutionary process to evolve a continuously improved population. In the population, each individual stands for architecture, and its fitness depends on the performance of the corresponding architecture in particular applications. The Genetic U-Net flowchart is summarized in Fig. 4 and the pseudocode of the proposed method is detailed in Algorithm 2.

Algorithm 1: Framework of the Proposed Method

Input: The population size $N$, the maximal generation number $T$, the crossover probability $p_c$, the mutation probability $p_m$, the mutation probability $p_b$, each bit.

Output: The discovered best architectures.

1) Crossover Operation: Crossover is intended to exchange information between individuals, and effective exchange can improve the performance of the algorithm. Many studies [66, 67] have shown the benefits of crossover operators involving a higher number of crossover points; consequently, a multi-point crossover is adopted herein. To improve search efficiency, we design a method named *difference-guided crossover* to choose two relatively different parent individuals for crossover. Algorithm 2 shows the details of the crossover operation. Initially, two individuals, $p_1$ and $p_2$, are chosen by binary tournament selection [68]. Next, the Hamming distance [69] between $p_1$ and $p_2$ is calculated and normalized to $[0,1]$ and denoted $diff$. If $diff$ is larger than the pre-defined threshold $\mu$, $p_1$ and $p_2$ are designated as parents. Otherwise, $p_1$ and $p_2$ will be re-selected by again using binary tournament selection. If ten re-selections of $p_1$ and $p_2$ do not meet the requirement, the last selection will be designated as parents. After that, the designated parents will mate with the probability $p_c$.

2) Mutation: Mutation can promote population diversity and prevent the algorithm from being trapped in a local optimum. In the proposed method, the offspring generated via crossover have the probability of $p_m$ for mutation, and each bit has the probability of $p_b$ for flipping independently. $p_b$ is a relatively small value (e.g., 0.05), which prevents an individual from being altered too much by mutation.

| ID | Operation sequence | ID | Operation sequence |
|----|-------------------|----|-------------------|
| 0  | $3 \times 3$ conv $\rightarrow$ ReLU | 8  | ReLU $\rightarrow$ $3 \times 3$ conv |
| 1  | $3 \times 3$ conv $\rightarrow$ Mish | 9  | Mish $\rightarrow$ $3 \times 3$ conv |
| 2  | $3 \times 3$ conv $\rightarrow$ IN $\rightarrow$ ReLU | 10 | IN $\rightarrow$ ReLU $\rightarrow$ $3 \times 3$ conv |
| 3  | $3 \times 3$ conv $\rightarrow$ IN $\rightarrow$ Mish | 11 | IN $\rightarrow$ Mish $\rightarrow$ $3 \times 3$ conv |
| 4  | $5 \times 5$ conv $\rightarrow$ ReLU | 12 | ReLU $\rightarrow$ $5 \times 5$ conv |
| 5  | $5 \times 5$ conv $\rightarrow$ Mish | 13 | Mish $\rightarrow$ $5 \times 5$ conv |
| 6  | $5 \times 5$ conv $\rightarrow$ IN $\rightarrow$ ReLU | 14 | IN $\rightarrow$ ReLU $\rightarrow$ $5 \times 5$ conv |
| 7  | $5 \times 5$ conv $\rightarrow$ IN $\rightarrow$ Mish | 15 | IN $\rightarrow$ Mish $\rightarrow$ $5 \times 5$ conv |

IN represents the instance normalization. ReLU indicates Rectified Linear Unit, and Mish is a self-regularized non-monotonic neural activation function.
Algorithm 2: Difference-guided Crossover Operation

Input: The population $P_t$, the probability for crossover operation $p_c$, the difference threshold of crossover operation $\mu$.

Output: Two offspring $o_1$, $o_2$.

1. $o_1$, $o_2 \leftarrow \emptyset$
2. for $j = 1$ to $10$ do
3.   $p_1, p_2 \leftarrow$ Select two individuals from $P_t$ by binary tournament selection;
4.   diff $\leftarrow$ Compute the difference between $p_1$ and $p_2$;
5.   if diff $> \mu$ then
6.     break;
7. end
8. $r \leftarrow$ Randomly generate a number from $(0, 1)$;
9. if $r < p_c$ then
10.   len $\leftarrow$ Compute the length of $p_1$ and $p_2$;
11.   ints $\leftarrow$ Randomly choose ten different integers from $[0, \text{len}]$ and sort them;
12.   $(i_0, i_1), (i_2, i_3), (i_4, i_5), (i_6, i_7), (i_8, i_9) \leftarrow$ Divide ints into five pairs in order;
13. for $k = 1$ to $5$ do
14.   $p_1, p_2 \leftarrow$ Exchange $p_1[i_{2k-2} : i_{2k-1}]$ and $p_2[i_{2k-2} : i_{2k-1}]$;
15. end
16. $o_1, o_2 \leftarrow p_1, p_2$;
17. else
18.   $o_1, o_2 \leftarrow p_1, p_2$;
19. end
20. return $o_1, o_2$.

3) Environmental Selection: Typically, GAs select the next population by tournament or roulette selection. Both selection methods may miss the best individual, resulting in performance degradation [72], which considerably impacts the final optimization results. Conversely, if we explicitly select the top $N$ individuals as the next generation, a premature phenomenon [70, 71] is more likely to trap the algorithm in a local optimum because of the heavy loss of population diversity. Hence, when choosing the next population, both elitist and non-elitist individuals are selected, thereby improving the convergence of the algorithm while maintaining the population diversity.

Algorithm 3 shows the process of environmental selection for the algorithm. First, given the current population $P_t$ and the generated offspring population $Q_t$, the top five best individuals are selected from $P_t \cup Q_t$. Second, $|P_t| - 5$ individuals are selected from $P_t \cup Q_t$ using the binary tournament selection, and then placed into $P_{t+1}$. The size of the next population $P_{t+1}$ is thus kept to that of the current population $P_t$.

Algorithm 3: Environmental Selection

Input: The parent population $P_t$, the offspring population $Q_t$.

Output: The parent population $P_{t+1}$.

1. $P_{t+1} \leftarrow \emptyset$;
2. $P_{\text{best}} \leftarrow$ Select the best five individuals from $P_t \cup Q_t$;
3. $P_t \cup Q_t \leftarrow$ Remove $P_{\text{best}}$ from $P_t \cup Q_t$;
4. $P_{t+1} \leftarrow P_{t+1} \cup P_{\text{best}}$;
5. while $|P_{t+1}| < |P_t|$ do
6.   $p \leftarrow$ Select an individual from $P_t$ by binary tournament selection;
7.   $P_{t+1} \leftarrow P_{t+1} \cup p$;
8. end
9. return $P_{t+1}$.
4) *Fitness Evaluation:* In Genetic U-Net, the fitness of an individual is the $F1$ (as explained in section IV-C) based on the architecture the individual represents and the data for validation because the $F1$ is a comprehensive metric for retinal vessel segmentation that helps resolve the problem of imbalanced samples. Algorithm 4 summarizes the procedure for evaluating individuals in the population. For evaluation, each individual transforms itself into an architecture, which is an inverse binary encoding process. Before training, He initialization [73] is used to initialize the architecture’s weights. Then, on the training data, the architecture is trained by Lookahead [74], which uses Adam [75] as the base optimizer. After 80 training epochs, the validation data are used to validate the trained architecture after each epoch until the $130^{\text{th}}$ epoch, and the best $F1$ during this process is set as the fitness of the corresponding individual.

**Algorithm 4: Evaluate Fitness**

**Input:** The population $P_i$ for fitness evaluation, training data $D_{train}$, validation data $D_{valid}$.

**Output:** The population $P_i$ with fitness.

```plaintext
foreach individual in $P_i$ do
    arch ← Transform the individual to its corresponding architecture;
    Apply weight initialization to arch;
    $F1_{best} ← 0$;
    for epoch = 1 to 130 do
        Train arch on $D_{train}$ by gradient descent for an epoch;
        if epoch > 80 then
            $F1 ←$ Evaluate the trained arch on $D_{valid}$;
            if $F1 > F1_{best}$ then
                $F1_{best} ← F1$;
            end
        end
    end
    Set $F1_{best}$ as the fitness of individual;
end
return $P_i$.
```

IV. MATERIALS FOR EXPERIMENTS

A. Loss Function

In fundus images, the ratio of vessel pixels is less than 0.1, and most pixels belong to the non-vessel class. Therefore, the problem of imbalanced samples must be resolved. For this purpose, focal loss [76] originally proposed to alleviate the sample imbalance problem in object detection is utilized as the loss function in this work, which is given as

$$
\text{Loss} = - \sum_{n=1}^{m} (\alpha y_n (1 - \hat{y}_n) \omega log \hat{y}_n + (1 - \alpha) (1 - y_n) \hat{y}_n^\alpha log (1 - \hat{y}_n))
$$

where $y$, $\hat{y}$, $n$, $m$, $\alpha$, and $\omega$ respectively indicate the ground truth, model prediction, $n^{th}$ sample, sample total, balance factor between positive and negative samples, and balance factor between simple and hard samples.

B. Datasets

Four public datasets DRIVE [11], CHASE_DB1 [77], STARE [12], and HRF [78] were used in our experiments. Descriptions of these datasets are given in Table II. The images of these datasets were captured by different devices and with different image sizes. DRIVE, STARE, and CHASE_DB1 have different annotations from two experts but only the annotations of the first expert were taken as the experimental ground truth. HRF has annotations from only one expert. The training and test datasets were split using the same method as in [18, 24, 33]. DRIVE has an officially determined training-testing split, 20 images for training and 20 for testing. For STARE, leave-one-out cross-validation was used to divide the dataset. For CHASE_DB1, the first 20 images were used for training, and the remaining 8 images were assigned to the test set. The 45 images in HRF are divided into three categories; healthy, diabetic retinopathy, and glaucomatous, each of which contains 15 images. The first five images of every category were used for training and the remainder were used for testing. Binary field of view (FOV) masks are offered for the DRIVE and HRF datasets. FOV masks for the STARE and CHASE_DB1 datasets were generated following [16] because they were not provided.

**TABLE II**

| Dataset     | Quantity | Resolution | training-testing split |
|-------------|----------|------------|------------------------|
| DRIVE       | 40       | 565×584    | 20/20                  |
| STARE       | 20       | 700×605    | leave one out          |
| CHASE_DB1   | 28       | 999×960    | 20/8                   |
| HRF         | 45       | 3504×2336  | 15/30                  |

C. Evaluation Metrics

Retinal vessel segmentation is a binary classification problem that classifies each pixel in the fundus image as either a vessel or a non-vessel. The output of the model is a probability map, which to each pixel assigns the probability of belonging to the class of vessels. In the experiments, the probability threshold was set to 0.5 to obtain the results. If a vessel pixel is correctly classified, it is a true-positive (TP); if not, it is a false-positive (FP). If a non-vessel pixel is precisely classified, it is a true-negative (TN); if not, it is a false-negative (FN). As shown in Table III, five metrics were selected for evaluation. Unless otherwise specified, we only used pixels inside FOVs to calculate the performance metrics.

**TABLE III**

| Metrics     | Description                                      |
|-------------|--------------------------------------------------|
| ACC (accuracy) | $AC = (TP + TN) / (TP + TN + FP + FN)$          |
| SE (sensitivity) | $SE = TP / (TP + FN)$                           |
| SP (specificity) | $SP = TN / (TN + FP)$                           |
| $F1$ (F1 score) | $F1 = (2 \times TP) / (2 \times TP + FP + FN)$ |
| AUROC       | Area Under the ROC Curve.                       |
V. Experiments

The experiments in the study consisted of two stages. In the first stage, the neural architectures were sought by GA. In the second stage, the found architectures were trained from scratch (validated) to establish their performances on retinal vessel segmentation. In this section, the stages of the experiments are explained and their results analyzed.

A. Experimental Setup

Dataset for searching: We sought the architectures using DRIVE. In the architecture search stage (first stage), the last five images of its training set were selected for validation, while the remaining fifteen images were for training. We also tested the architectures found using DRIVE on other datasets (STARE, CHASE_DB1, and HRF) in the second stage.

Genetic U-Net hyper-parameters: The generic upsampling and down-sampling operations (max pooling and transpose convolution) were adopted for the original U-Net [21]. The probability of crossover and mutation operations ($p_c$ and $p_m$) was set to 0.9 and 0.7 respectively, and the difference threshold $\mu$ was set to 0.2. The probability $p_h$ in the mutation process was 0.05. The population size $N$ was 20, and the number of the generations $T$ was 50.

Network training during search: For data argumentation, horizontal flipping, vertical flipping, and random rotation from [0°, 360°] were employed to increase the training data, which prevented the models from overfitting. The pixels of the images in HRF were randomly cropped out of a region 1000 × 1000 in size and used as the architecture input but the complete images were adopted as the input during testing. Additionally, due to the GPU memory limitation, during training, the images in HRF were randomly cropped out of a region 1000 × 1000 in size and used as the architecture input but the complete images were adopted as the input during testing.

B. Experimental Results of the Discovered Model

1) Comparison with Existing Methods: We report the test results of the architectures searched for on DRIVE on all four public datasets (DRIVE, STARE, HRF, and CHASE_DB1) and compare them with other existing methods (mostly are CNN-based methods).

The experimental results are summarized in Tables IV and VII. In these tables, the results of existing methods except CE-Net [25] and CS²-Net [26] are obtained from the original papers. The results of CE-Net and CS²-Net are obtained using the same training settings as the discovered architecture. The existing methods marked with stars calculate the performance metrics with FOVs, while those without stars were not explicitly claimed in the papers. Here, we report the performance of the discovered architecture both with FOVs and without FOVs. For fairness, our method used the same data split as the other methods. For all the datasets, the proposed method dominates all the methods of comparison on two comprehensive metrics (F1 and AUROC), which means that the discovered architecture achieves the best overall performance without post-processing or special pre-processing. Additionally, the proposed method outperformed the other methods in SE (sensitivity) by some margin, which indicates that the discovered architecture is more capable of detecting vessels. ACC is a trade-off between SE and SP. Since the non-vessel region is much larger than the vessel region in fundus images, ACC is more susceptible to SP. The ACC of the proposed method also compares well against the other methods. Moreover, the parameters of the discovered architecture are the least among these methods (0.27 M). We utilized only a few data augmentations such as flipping and rotation that are simpler and less diverse than those used by most other methods. The architecture searched for on DRIVE also successfully transferred to the other three datasets, STARE, HRF, and CHASE_DB1.

| Methods                         | Year  | ACC   | SE    | SP    | Fl   | AUROC | Params(M) |
|---------------------------------|-------|-------|-------|-------|------|--------|-----------|
| Vega et al. [23]                | 2015  | 0.9412| 0.7444| 0.9612| 0.6884| N/A    | N/A       |
| Li et al. [24]                  | 2015  | 0.9527| 0.7569| 0.9816| N/A  | 0.9738 | N/A       |
| Orlando et al. [25]             | 2016  | N/A   | 0.7897| 0.9684| 0.7857| N/A    | N/A       |
| Fan and Mo [26]                 | 2016  | 0.9612| 0.7814| 0.9784| N/A  | 0.9785 | N/A       |
| Lukowski et al. [27]            | 2016  | 0.9555| 0.7811| 0.9807| N/A  | 0.9790 | 48.00     |
| Mo and Zhang [28]               | 2017  | 0.9521| 0.7779| 0.9798| N/A  | 0.9782 | 7.63      |
| Yan et al. [29]                 | 2018  | 0.9542| 0.7653| 0.9818| N/A  | 0.9792 | 31.35     |
| Alam et al. [30]                | 2019  | 0.9556| 0.7792| 0.9815| N/A  | 0.9794 | 1.07      |
| Jin et al. [31]                 | 2019  | 0.9566| 0.7963| 0.9800| 0.8227| 0.8902 | 0.88      |
| Bo Wang et al. [32]             | 2019  | 0.9567| 0.7940| 0.9816| 0.8270| 0.9772 | N/A       |
| YiCheng Wu et al. [33]          | 2019  | 0.9578| 0.8038| 0.9802| N/A  | 0.9821 | 1.70      |
| Mou Lei et al. [34]             | 2019  | 0.9594| 0.8126| 0.9798| N/A  | 0.9796 | 56.03     |
| CE-Net [35]                     | 2019  | 0.9545| 0.8276| 0.9735| 0.8243| 0.9794 | 15.28     |
| CS²-Net [36]                    | 2021  | 0.9553| 0.8154| 0.9757| 0.8228| 0.9784 | 8.91      |

Genetic U-Net w/o FOVs - 0.9777 0.8300 0.9756 0.8314 0.9823 0.27
Genetic U-Net w/o FOVs 0.9707 0.8300 0.9843 0.8314 0.9885 0.27

*W/’ and ‘W/o’ mean ‘with’ and ‘without’, respectively.

| Methods                         | Year  | ACC   | SE    | Fl   | AUROC | Params(M) |
|---------------------------------|-------|-------|-------|------|--------|-----------|
| Vega et al. [23]                | 2015  | 0.9483| 0.7019| 0.9671| 0.9614| N/A       |
| Li et al. [24]                  | 2015  | 0.9628| 0.7726| 0.9444| N/A   | 0.9879   |
| Orlando et al. [25]             | 2017  | N/A   | 0.7480| 0.9738| 0.7644| N/A       |
| Fan and Mo [26]                 | 2016  | 0.9654| 0.7834| 0.9799| N/A   | 0.9785   |
| Lukowski et al. [27]            | 2016  | 0.9566| 0.7867| 0.9754| N/A   | 0.9785   |
| Mo and Zhang [28]               | 2018  | 0.9674| 0.8147| 0.9844| N/A   | 0.9885   |
| Yan et al. [29]                 | 2018  | 0.9612| 0.7581| 0.9946| N/A   | 0.9801   |
| Alam et al. [30]                | 2019  | 0.9712| 0.8292| 0.9862| 0.8475| 0.9914   |
| Jin et al. [31]                 | 2019  | 0.9641| 0.7959| 0.9878| 0.8143| 0.9832   |
| CE-Net [35]                     | 2019  | 0.9565| 0.8406| 0.9813| 0.8363| 0.9871   |
| CS²-Net [36]                    | 2021  | 0.9670| 0.8396| 0.9813| 0.8420| 0.9875   |

Genetic U-Net w/ FOVs 0.9719 0.8658 0.9846 0.8630 0.9921
Genetic U-Net w/o FOVs 0.9792 0.8658 0.9886 0.8630 0.9942

2) Comparison with the Baseline Models: As this work uses the U-shaped encoder-decoder structure as the backbone,
TABLE VI
Comparison with existing methods on the CHASE_DB1 dataset.

| Methods                  | Year | ACC  | SE   | SP   | FI   | AUROC |
|--------------------------|------|------|------|------|------|-------|
| Li et al. [13]           | 2015 | 0.9581 | 0.7507 | 0.9793 | N/A | 0.9716 |
| Fan and Mo [27]          | 2016 | 0.9573 | 0.7656 | 0.9704 | N/A | 0.9781 |
| Yan et al. [47]          | 2018 | 0.9610 | 0.7633 | 0.9809 | N/A | 0.9781 |
| Akom et al. [14]         | 2019 | 0.9634 | 0.7756 | 0.9820 | 0.7928 | 0.9815 |
| Bo Wang et al. [10]      | 2019 | 0.9661 | 0.8074 | 0.9821 | 0.8037 | 0.9812 |
| Yicheng Wu et al. [22]   | 2019 | 0.9661 | 0.8132 | 0.9814 | N/A | 0.9860 |
| CE-Net [25]              | 2019 | 0.9641 | 0.8093 | 0.9797 | 0.8054 | 0.9834 |
| CS²-Net [26]             | 2021 | 0.9651 | 0.8329 | 0.9784 | 0.8141 | 0.9851 |
| Genetic U-Net w/o FOVs   | -    | 0.9667 | 0.8463 | 0.9645 | 0.8223 | 0.9880 |
| Genetic U-Net w/o FOVs   | -    | 0.9769 | 0.8463 | 0.9857 | 0.8223 | 0.9914 |

TABLE VII
Comparison with existing methods on the HRF dataset.

| Methods                  | Year | ACC  | SE   | SP   | FI   | AUROC |
|--------------------------|------|------|------|------|------|-------|
| Orlando et al. [14]      | 2016 | N/A  | 0.7794 | 0.9650 | 0.7341 | N/A   |
| Yan et al. [33]          | 2018 | 0.9437 | 0.7881 | 0.9592 | N/A | 0.9831 |
| Jin et al. [37]          | 2019 | 0.9651 | 0.7464 | 0.9874 | N/A | 0.9831 |
| CE-Net [25]              | 2019 | 0.9613 | 0.7805 | 0.9798 | 0.7895 | 0.9766 |
| CS²-Net [26]             | 2021 | 0.9618 | 0.7890 | 0.9795 | 0.7935 | 0.9758 |
| Genetic U-Net w/o FOVs   | -    | 0.9667 | 0.8220 | 0.9818 | 0.8179 | 0.9872 |
| Genetic U-Net w/o FOVs   | -    | 0.9715 | 0.8220 | 0.9839 | 0.8179 | 0.9891 |

TABLE VIII
Comparison with U-Net and FC-Densenet on four datasets.

| Datasets      | Models | ACC  | SE   | SP   | FI   | AUROC |
|---------------|--------|------|------|------|------|-------|
| DRIVE         | U-Net  | 0.9550 | 0.8089 | 0.9720 | 0.8191 | 0.9795 |
|               | FC-Densenet | 0.9555 | 0.8231 | 0.9743 | 0.8325 | 0.9799 |
|               | Genetic U-Net | 0.9577 | 0.8300 | 0.9758 | 0.8314 | 0.9823 |
| STARE         | U-Net  | 0.9654 | 0.8187 | 0.9798 | 0.8273 | 0.9862 |
|               | FC-Densenet | 0.9673 | 0.8276 | 0.9805 | 0.8369 | 0.9872 |
|               | Genetic U-Net | 0.9719 | 0.8658 | 0.9846 | 0.8630 | 0.9921 |
| CHASE_DB1     | U-Net  | 0.9650 | 0.8298 | 0.9829 | 0.8092 | 0.9859 |
|               | FC-Densenet | 0.9654 | 0.8307 | 0.9789 | 0.8144 | 0.9862 |
|               | Genetic U-Net | 0.9667 | 0.8463 | 0.9818 | 0.8223 | 0.9880 |
| HRF           | U-Net  | 0.9647 | 0.8212 | 0.9794 | 0.8101 | 0.9849 |
|               | FC-Densenet | 0.9652 | 0.8098 | 0.9811 | 0.8125 | 0.9851 |
|               | Genetic U-Net | 0.9667 | 0.8220 | 0.9818 | 0.8179 | 0.9872 |

C. Experimental Analysis of Architecture Search

The evolutionary trajectory of Genetic U-Net is the blue line shown in Fig. 8. The fitness of the best individual gradually increases from the first generation and converges at approximately the 50th generation. Thus, we terminate the architecture search after the 50th generation and select the best individual from the last population as the result. Fig. 12 displays the discovered architecture decoded from the selected individual.

1) Observations and Findings: The results of the evolutionary algorithm usually contain some useful information for us to further improve our work. To find some patterns for more efficient architecture design, we observe and analyze the top five architectures of the last generation.

Topology structure: We observe the topology within each block of these architectures displayed in the Appendix. Almost all these blocks have the allowed maximum number of nodes, and their internal structures are relatively complex. Several skip connections can be found between the nodes. In addition, all blocks have two or three parallel branches within that resemble InceptionNet [60] blocks.

Operations and operational sequences: We obtain the statistics of the frequency of occurrence of all operational sequences listed in Table 1. Fig. 7 shows that the operation sequence with ID 11 has the highest frequency. To further verify the effect of this operation sequence, we use it to replace the basic convolutional layer (3 × 3 conv + ReLU) of the U-Net [21] blocks, which almost do not change parameters and MACs of U-Net, and conduct experiments on DRIVE. The result is reported in the eighth row of Table X. As expected, we comprehensively compared the discovered architecture with the baseline models, the original U-Net [21], and FC-Densenet [58], which is also a U-shaped network. For a fair comparison, we trained the original U-Net and FC-Densenet with the same parameter settings as the discovered architecture. Table VIII reveals that the discovered architecture outperformed the original U-Net and FC-Densenet on all four datasets.

Some examples of the results are also presented in Fig. 5, Fig. 6, and Fig. 9. The blue pixels in the images indicate false negatives, which are from the vessel regions not detected. It shows that there are more blue pixels in the results of the original U-Net and FC-Densenet, in both the overall view and the locally magnified view. It further shows that the original U-Net and FC-Densenet have limitations in extracting complicated structural features, while the discovered architecture extracts them more effectively.

The computational efficiency of the discovered architecture is also analyzed, and the results are provided in Table IX. Based on Table IX the parameters in the discovered architecture total approximately 0.27 Million, which is a 115x/34x reduction compared to the 31.03 Million/9.32 Million parameters for U-Net/FC-Densenet. The model size of the discovered architecture is 1.2 MB, which is approximately a 100x/32x reduction compared to the 120MB/38MB of U-Net/FC-Densenet. The execution time and Multiplication Accumulation operations (MACs) results on DRIVE were obtained with an input dimension 3×565×584. The total number of MACs in the discovered model in one forward propagation is 41 Billion, which is a 3.2x/2.0x reduction compared to the 142 Billion/83 Billion MACs in U-Net/FC-Densenet. For the execution time, the discovered architecture also achieved an approximately 1.3x/2.9x reduction compared to U-Net/FC-Densenet.
after U-Net used this operation sequence, its performance of retinal vessel segmentation on DRIVE significantly improved. Surprisingly though, its performance surpassed existing methods listed in Table IV except the discovered architecture using the proposed approach in this study. It is important to explore the reasons why only changing some operations of U-Net blocks can lead to such a significant improvement of its performance. After replacing the original basic convolution layer in U-Net with the operation sequence with ID 11, U-Net has a different activation function (Mish [64]) and activation type (pre-activation) and instance normalization, which further boosts its performance. To discover which factor or factors will have a greater impact on U-Net, we conducted six additional sets of experiments with the results listed in the second to the seventh row in Table X. From the data in the second to the fourth rows, adding instance normalization or using pre-activation is shown to improve the performance somewhat. In particular, the effect of adding instance normalization is apparent. U-Net is not improved by only changing the activation function to Mish. Furthermore, the data of the fifth to the eighth rows in Table X show that adding instance normalization or using pre-activation is useful for promoting better performance. When instance normalization and pre-activation are employed together, the effect is more notable. In addition, pre-activation with Mish is slightly more effective than pre-activation with ReLU.

Fig. 5. Overall view visualization of the segmentation results. The green pixels indicate true positive, red pixels indicate false positive, and blue pixels indicate false negative.
Genetic U-Net can obtain multiple competitive network architectures from these evolutionary results. We can extract knowledge and several principles of network architecture design for retinal vessel segmentation through the observations and the statistical analyses of these network architectures, which may potentially be important knowledge bases to further improve our future work.

**D. Ablation Study**

**Difference-guided crossover:** To verify the effect of utilizing the difference-guided crossover, we compare the performances of the algorithms searching with and without the difference-guided crossover. We conduct this experiment with the same settings for both algorithms. We obtain the fitness of the best individual in the population of each generation. As shown in Fig. 8, the difference-guided crossover improves the performance.

**Environmental selection:** To improve search efficiency and simultaneously prevent premature convergence, we adopted a selection scheme that considers both elitism (size=5) and diversity. Here, we demonstrate the benefits of this scheme by comparing it with the top $N$ selection [79] and elitist selection (size=1) [80]. The experiments were conducted with the same initial population and other settings (e.g., the same crossover operation.). Fig. 8 also clearly reveals the advantage of the proposed selection scheme. The top $N$ selection results in premature convergence of the algorithm.

**VI. DISCUSSION**

**A. Performance on Challenging Cases**

The images provided in the third and fourth columns of Fig. 5 and the second column of Fig. 9 all show lesions, and the image in the third column of Fig. 5 is disrupted by severely uneven illumination. In these cases, fewer red pixels (false positive or misidentified vessel regions) of lesion areas are seen in the resulting output images of the discovered architecture than those of the other models. This shows that the discovered architecture is less sensitive to lesions and uneven illumination. Similarly, as shown in the magnified view of Fig. 6, there are fewer blue pixels (false-negative or undetected vessel regions) of the cross-connected and tiny vessel areas in
the resulting output images of the discovered architecture than those of the other models. This demonstrates that the discovered architecture can capture more cross-connected vessels and low-contrast tiny vessels than the experimental competitors. Overall, among the methods compared in Fig. 5, Fig. 9 and Fig. 6 the vascular trees segmented by the discovered architecture best match the ground truth. This indicates that the discovered architectures are beneficial to the segmentation of these challenging regions.

B. Performance on High-Resolution Fundus Images

Compared with STARE, DRIVE, and CHASE_DB1, the HRF dataset includes fundus images with much higher resolutions. Several experimental results are displayed in Fig. 9. Both qualitative and quantitative results on HRF demonstrate that the discovered architecture achieves better segmentation results than the state-of-the-art methods and the baseline models (the original U-Net and FC-DenseNet). This indicates that the discovered architecture can also achieve promising performance on high-resolution fundus images.

C. Adjusting the Number of Architecture-based Parameters

To explore the effect of model capacity further, we adjusted the channels of convolutional operations $c$ within the nodes of the discovered architecture from 20 to 2, 3, 4, 5, 10, and 30 (thus changed the number of architecture-based parameters) and trained the adjusted models from scratch. We evaluated their performance on DRIVE and CHASE_DB1 and analyzed their computational complexity. The performance of the adjusted models is reported in Table XI. When $c$ is increased from 20 to 30, compared with the original model, the adjusted model has more parameters and higher computational complexity (approximately a 125% increase on MACs and Params). However, its performance was almost not improved (On DRIVE, there was an approximately 0.04% increase in F1 and almost no increase in AUROC). When $c$ is decreased, the adjusted models have fewer parameters and lower computational complexity, and their performance does not degrade distinctly. For instance, when $c = 5$ or 10, there is approximately a 94% or 75% reduction in MACs and Params, respectively. On DRIVE, however, F1 only degenerates by approximately 0.20% or 0.91%, and AUROC by approximately 0.13% or 0.15%. When $c = 5$, the adjusted model with fewer architecture-based parameters (about 0.06%/0.19% of the original U-Net/FC-DenseNet) can still achieve competitive performance with the original U-Net, FC-DenseNet, and the other methods listed in Table IV and Table VI. Even if $c = 2$ and the model only contains 0.0028 Million (2.8 K) parameters, the performance is still comparable with some existing methods. These results confirm that decreasing the capacity of the discovered architecture does not significantly influence its performance, which is beneficial for practical clinical applications. This reveals that the discovered architecture has strong parameter efficiency and is not sensitive to reductions in the number of architecture-based parameters.

D. Performance using Reduced Training Data

To verify the performance of the discovered architecture after reducing the training data, we evaluated it on the reduced training data of DRIVE and CHASE_DB1. In the previous data split, both DRIVE and CHASE_DB1 had 20 images for model training. Here, we reduced the number of images used...
for model training from 2 to 2, 5, and 10 in both datasets and conducted multiple experiments on the discovered architecture and the baseline models (the original U-Net and FC-Densenet). The F1-score showed a degradation as the number of training images decreased. On DRIVE and CHASE_DB1, both the discovered architecture and the baseline models suffered performance degradation; however, for the baseline models, the degradation is more apparent. For example, when the training data on DRIVE/CHASE_DB1 consisted of only 5 images, the F1 of the discovered architecture decreased from 0.8314/0.8223 to 0.8270/0.8063 (0.52%/1.94% degradation), the F1 of the original U-Net decreased from 0.8190/0.8092 to 0.8043/0.7881 (1.80%/2.61% degradation), and the F1 of FC-Densenet decreased from 0.8235/0.8144 to 0.8073/0.7827 (1.97%/3.89% degradation). Even with only 5 training images, the performance of the discovered architecture is still comparable with the existing methods listed in Table [V] and Table [VI]. This confirms that the discovered architecture has more stable performance under reduced training data, which is crucial when using the limited annotated data available for retinal vessel segmentation.

### Table XI

| Model configs | Params | MACs | F1 | AUROC | F1 | AUROC |
|---------------|--------|------|----|-------|----|-------|
| c = 20 | 0.2724 M 41.35 B | | 0.8314 | 0.9823 | 0.8223 | 0.9880 |
| c = 30 | 0.6117 M 92.77 B | | 0.8317 | 0.9824 | 0.8228 | 0.9880 |
| c = 10 | 0.0685 M 10.42 B | | 0.8306 | 0.9815 | 0.8214 | 0.9871 |
| c = 5 | 0.0173 M 2.65 B | | 0.8297 | 0.9810 | 0.8179 | 0.9863 |
| c = 4 | 0.0111 M 1.71 B | | 0.8275 | 0.9806 | 0.8134 | 0.9856 |
| c = 3 | 0.0063 M 0.97 B | | 0.8229 | 0.9791 | 0.8103 | 0.9833 |
| c = 2 | 0.0028 M 0.44 B | | 0.8138 | 0.9746 | 0.7914 | 0.9804 |

The MACs are calculated based on an input size of 512 × 512.

### Table XII

| Models | Training samples | F1 | AUROC | F1 | AUROC |
|--------|------------------|----|-------|----|-------|
| DRIVE  | CHASE_DB1        |    |       |    |       |
| Genetic U-Net | 20 | 0.8314 | 0.9823 | 0.8223 | 0.9880 |
|           | 10 | 0.8303 | 0.9813 | 0.8154 | 0.9861 |
|           | 5  | 0.8270 | 0.9786 | 0.8063 | 0.9831 |
|           | 2  | 0.8122 | 0.9730 | 0.7866 | 0.9792 |
| FC-Densenet | 20 | 0.8235 | 0.9799 | 0.8144 | 0.9862 |
|           | 10 | 0.8213 | 0.9799 | 0.8057 | 0.9828 |
|           | 5  | 0.8073 | 0.9717 | 0.7827 | 0.9764 |
|           | 2  | 0.7789 | 0.9603 | 0.7601 | 0.9604 |
| U-Net    | 20 | 0.8191 | 0.9795 | 0.8092 | 0.9859 |
|           | 10 | 0.8146 | 0.9794 | 0.7996 | 0.9824 |
|           | 5  | 0.8043 | 0.9700 | 0.7881 | 0.9806 |
|           | 2  | 0.7597 | 0.9514 | 0.7687 | 0.9707 |

### Table XIII

| Models | ACC | SE | SP | F1 | AUROC |
|--------|-----|----|----|----|-------|
| DRIVE  |     |    |    |    |       |
| FC-Densenet | 0.9112 | 0.8179 | 0.9401 | 0.8097 | 0.9587 |
| CE-Net | 0.9099 | 0.8253 | 0.9233 | 0.8161 | 0.9645 |
| CS²-Net| 0.9109 | 0.8263 | 0.9272 | 0.8163 | 0.9634 |
| Genetic U-Net | 0.9171 | 0.8376 | 0.9446 | 0.8212 | 0.9666 |

### E. Generalization Ability

To further evaluate the generalization ability of the discovered architecture, we tested it (searched on DRIVE) on the challenging task of cell boundary segmentation, which is an application similar to vessel segmentation. In this experiment, the regions of the cell membranes (boundaries) are the segmented areas (positive). The adopted EM dataset [SI] contains 30 training images and 30 test images 512 × 512 in size, but the ground truth of the 30 test images is not public.

### Table XIV

| Models | Model performance on DRIVE/CHASE_DB1 |    |       |    |       |
|--------|-------------------------------------|----|-------|----|-------|
| DRIVE  | F1-AUROC                            |    |       |    |       |
| FC-Densenet | 0.8122 | 0.8270 | 0.8213 | 0.8154 | 0.8154 |
| CE-Net | 0.8013 | 0.8273 | 0.8213 | 0.8154 | 0.8154 |
| CS²-Net| 0.8109 | 0.8263 | 0.8272 | 0.8163 | 0.9634 |

### F. Future work

Even though the network architecture obtained by the proposed method is capable of segmenting vessels closer to the ground truth than other models, it may fail in some very challenging cases. To improve the performance of the...
proposed approach further, a better design of the search space will be investigated. We must also extend the application of the proposed method to other scenarios to further verify its efficacy and generality.

VII. CONCLUSION

In this study, a novel neural architecture search (NAS) for retinal vessel segmentation, named Genetic U-Net, is proposed based on the U-shaped encoder-decoder structure. The existing CNN methods for retinal vessel segmentation can make hardly any further improvements and usually have profuse architecture-based parameters. Genetic U-Net employs an improved GA to evolve a model that outperforms existing mainstream methods in retinal vessel segmentation using a condensed but flexible search space that limits the number of architecture-based parameters within a range of small values. Therefore, the resulting architecture achieves a significant reduction in the number of parameters and the computational cost. This indicates that it is feasible to design an exceptional network architecture with fewer parameters for vessel segmentation that would facilitate its deployment for clinical applications and overcome the insufficiency of training data. Furthermore, by analyzing the evolved results, we found that utilizing several effective operations and patterns in the model’s building blocks greatly enhanced the performance in vessel segmentation, which provides instrumental domain knowledge for use in future studies.

ACKNOWLEDGMENT

This work was supported by the Key Lab of Digital Signal and Image Processing of Guangdong Province, by the Key Laboratory of Intelligent Manufacturing Technology (Shantou University), Ministry of Education, by the Science and Technology Planning Project of Guangdong Province of China under grants (180917144960530, 2019A050519008, 2019A050520001), by the State Key Lab of Digital Manufacturing Equipment & Technology under grant DMETKF2019020.

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VIII. APPENDIX

A. Visualization of Discovered Architectures

![Diagram of the first best architecture](image1)

Fig. 12. The first best architecture.

![Diagram of the second best architecture](image2)

Fig. 13. The second best architecture.
Fig. 14. The third best architecture.

Fig. 15. The fourth best architecture.

Fig. 16. The fifth best architecture.