Abstract: Objective and quantitative assessment of skin conditions is essential for cosmeceutical studies and research on skin aging and skin regeneration. Various handcraft-based image processing methods have been proposed to evaluate skin conditions objectively, but they have unavoidable disadvantages when used to analyze skin features accurately. This study proposes a hybrid segmentation scheme consisting of Deeplab v3+ with an Inception-ResNet-v2 backbone, LightGBM, and morphological processing (MP) to overcome the shortcomings of handcraft-based approaches. First, we apply Deeplab v3+ with an Inception-ResNet-v2 backbone for pixel segmentation of skin wrinkles and cells. Then, LightGBM and MP are used to enhance the pixel segmentation quality. Finally, we determine several skin features based on the results of wrinkle and cell segmentation. Our proposed segmentation scheme achieved a mean accuracy of 0.854, mean of intersection over union of 0.749, and mean boundary F1 score of 0.852, which achieved 1.1%, 6.7%, and 14.8% improvement over the panoptic-based semantic segmentation method, respectively.

Keywords: Image segmentation; skin texture; feature extraction; dermoscopy image

1 Introduction

Skin texture is the outermost indicator of the skin condition or the progression level of skin aging. Skin texture depends on various intrinsic and extrinsic factors that affect the health of the skin layers [1–5]. For instance, prolonged exposure to sunlight and excessive smoking can adversely affect skin layers, as the skin loses moisture and becomes dry, which is then reflected in the skin texture. Therefore, tracking and observing changes in skin texture can help intuitively understand changes in skin health.

However, objective assessment of skin condition has been regarded as challenging because dermatologists have traditionally diagnosed the skin condition by examining the skin texture with the naked eye, making the diagnostic results subjective. To diagnose the skin condition objectively, two general approaches have been proposed: three-dimensional topography analysis (3DTA) and two-dimensional image analysis (2DIA).
3DTA mainly analyzes depth-related skin features such as maximal, mean, minimum depth of roughness, and smoothness [6–9]. This usually requires silicon replicas of the skin surface and optical equipment. Therefore, 3DTA suffers from significant problems such as long processing time, complexity of analysis, and high cost. In contrast, most 2DIA methods focus on analyzing the structural shapes of skin texture using dermoscopy devices [10–17]. The 2DIA methods aim to extract diverse representative visual features from dermoscopy images, such as irregular polygons and borderlines that are regarded as cells and wrinkles, respectively. Skin features include wrinkle width, wrinkle length, and the density of cells that can be used as essential criteria to evaluate skin conditions or estimate the degree of skin aging [12–14]. For decades, 2DIA methods have generally been used for skin condition diagnosis in terms of time and cost efficiency, but 2DIA methods, too, suffer from various limitations:

- The image preprocessing steps of 2DIA methods, such as contrast enhancement, noise removal, and color histogram equalization, have been designed differently for different datasets.
- Most 2DIA methods have opted for a handcrafted feature-based method, but handcrafted features are generally not robust and are computationally intensive due to their high dimensionality.
- Traditional 2DIA methods must choose which features are important in a given dataset. Therefore, substantial engineering experience and dermatology knowledge are required to obtain successful results.

Deep learning is one alternative technique to handcrafted feature-based approaches because it can learn visual features automatically to solve a specific task, such as object classification and target segmentation. In many research fields, a convolutional neural network (CNN)-based model, which is a deep learning technique, has been reported to be extremely effective compared with handcraft-based approaches. We believe that applying a deep learning model overcomes the disadvantage of the 2DIA methods in accurate skin feature extraction. In this study, we address three major issues, with the following main contributions:

- We performed wrinkle and cell segmentation using Deeplab v3+ [18] with an Inception-ResNet-v2 backbone [19], which is a popular CNN model. By using this approach, we overcome the serious limitations of traditional handcrafted feature-based methods.
- We improved the quality of wrinkle and cell segmentation by using LightGBM [20] and morphological processing (MP), which can accurately extract skin features.
- We demonstrated the effectiveness of our proposed scheme by extracting various skin features from the segmentation results and comparing them with other existing methods in terms of accuracy.

The remainder of this paper is organized as follows. Related works of skin image analysis are introduced in Section 2. The major differences between previous works and our proposed scheme are described in Section 3. Finally, the experimental results and conclusions are presented in Sections 4 and 5, respectively.

2 Related Works

This section presents several studies that have been conducted to assess skin conditions. Since the 2000s, most skin image analyses have been performed through molding replica analysis based on the 3DTA method [21–23]. Masuda et al. [21] proposed a 3D morphological characterization method for molding replica analysis. Using 3D measurement equipment and a surface analysis
method, they extracted and analyzed several skin texture features such as depth, width, and length of skin furrows. Friedman et al. [22] conducted a 3D skin topology analysis, a non-invasive approach for identifying facial rhytides and acne scars. They utilized a micro-topology imaging system and a charge-coupled device (CCD) camera to record the skin surface topology. They concluded that 3D skin topology analysis could allow fast and quantitative assessment of skin conditions. Molding replica and 3D topology analysis methods have been regarded as useful approaches, but their requirements for expensive equipment and substantial analytic experience make their widespread application difficult.

Recently, 2DIA-based approaches have attracted much attention [15–17,24–28]. Zou et al. [17] suggested an objective modeling approach for skin surface characterization. They developed new measurement parameters, including polygons on the skin texture and the average area of the polygons detected. They argued that skin surface analysis using image processing techniques is useful for quantitatively expressing the skin condition. Cula et al. [24] proposed an automatic wrinkle detection algorithm to extract the orientation of polygon borders and bidirectional histogram features of illumination. Facial wrinkle features were extracted using contrast stretching and Gabor filtering, and the wrinkle types were classified to identify the subjects. Tanaka et al. [25] applied the cross-binarization method to detect skin wrinkles. Image binarization was first conducted for accurate wrinkle detection, and subsequently, the straight-line matching algorithm was applied to measure the length of skin wrinkles. Razalli et al. [26] proposed a wrinkle detection method to determine the relationship between aging and changes in wrinkle shape. The Hessian-based filter (HBF) was applied to extract the facial features, and then, the method was evaluated using the FG-NET database. Choi et al. [27] proposed a framework for skin texture extraction and skin aging trend estimation. The depth of wrinkles, wrinkle length and width, and the number of closed polygons were calculated as a feature set. Then, a support vector machine model was utilized to estimate the trend of skin aging. A skin aging estimation accuracy of more than 90% was achieved in a dermatologist's blind test.

More recently, using the versatility of machine learning techniques, breakthrough attempts have been made to detect or diagnose critical diseases such as melanoma, lesions, and COVID-19 [29–33]. Afza et al. [29] proposed an optimal color feature selection scheme for skin lesion classification. They used contrast stretching, color feature extraction, and entropy-controlled feature optimization. To improve the performance of skin lesion classification, they proposed a hierarchical 3-step super-pixels with deep learning–based framework that includes a pre-trained deep learning model (ResNet-50) and a new optimization technique called Grasshopper [30]. Khan et al. [31] proposed an intelligent framework that includes Mask R-CNN, ResNet with feature pyramid network (FPN), and 24-layered CNN for localizing and classifying multi-class skin lesions. They used pre-trained DenseNet, entropy-controlled least square support vector machine and extreme learning machine (ELM) techniques to improve the accuracy of skin lesion classification [32].

To summarize, 3DTA, 2DIA, and machine learning methods have been used to objectively assess skin conditions and diagnose critical diseases such as melanoma and skin lesions. 3DTA is an excellent approach to accurately analyze skin conditions, but it requires expensive equipment and dermatological knowledge and experience. Although 2DIA is a cost-effective alternative to 3DTA, it is difficult to guarantee its applicability as most 2DIA methods follow a handcrafted feature-based method that makes it difficult to obtain robust features and rely heavily on an engineer’s experience and knowledge [34]. To overcome the shortcomings of the 3DTA and 2DIA
methods, we propose a scheme that performs image segmentation based on a deep learning model and extracts accurate skin features from the segmentation result.

3 Methodology

3.1 Comparison with Previous 2DIA Methods

In this section, we show the difference between the traditional 2DIA approaches and the proposed scheme. Traditional 2DIA methods for skin feature extraction first perform some preprocessing steps to eliminate light interference, noise, and distortion to obtain skin features of reasonable quality [10–17].

The skin image is then converted into a binary image, which is regarded as the groundwork for identifying wrinkle and cell areas. Fig. 1 presents the typical structure of traditional 2DIA methods with the preprocessing steps in the yellow box. Image cropping eliminates the harmful effects of vignetting because a luminous source of the skin image from a dermoscopy device is typically concentrated around its center [10–13]. Contrast stretching and histogram equalization can minimize light interference. Generally, noise filtering and morphological transformation improve the effectiveness of preprocessing. The watershed transform method [35] performs wrinkle and cell segmentation by calculating the intensity difference in a grayscale image and draws borderlines based on the intensity difference. Finally, wrinkles and cells are obtained using the watershed transform.

![Figure 1: Typical structure of traditional 2DIA methods for skin feature extraction](image)

Fig. 2 presents an illustration of our scheme. Unlike traditional approaches, our scheme adopts a CNN, LightGBM, and MP as preprocessing units. The CNN is organized by multiple convolutional layers that perform image classification and segmentation tasks. Convolutional layers reduce the images into a form that is easier to process without losing features. This is critical for achieving good predictions. Each convolutional layer extracts valuable visual features, considering image distortion, noise, and brightness. Therefore, well-designed convolutional layers can extract reasonable visual features that represent the semantic information of the image. For this, the
Deeplab v3+ with Inception-ResNet-V2 backbone model, a popular CNN model, was used to replace several preprocessing steps in the traditional approach. In addition, to enhance the quality of binarization, LightGBM and MP were used together. Manually, the ground truth was made based on the annotated wrinkle and cell class for every pixel in the dermoscopy image, and these features were utilized to train the Inception-ResNet-V2 model [19]. Subsequently, we applied watershed transformation and performed wrinkle and cell feature extraction.

![Figure 2: Overall structure of the proposed scheme for skin feature extraction](image)

![Figure 3: Segmentation examples (a) Input images (b) Ground truth](image)
3.2 Wrinkle and Cell Segmentation

In this section, we present details of the wrinkle and cell segmentation process with the organized CNN architecture and LightGBM. Figs. 3a and 3b present the input skin images captured by a commercial dermoscopy device and their ground truth, respectively. We defined that the black pixels represent the wrinkle class and the white pixels indicate the cell class.

Fig. 4 presents the overall flow of our proposed scheme. In this study, a pre-trained Inception-ResNet-v2 model obtained from the MathWorks repository was used. Inception-ResNet-v2 showed the best performance in the ImageNet Large Scale Visual Recognition Challenge (ILSVRC) 2015, and its performance in object recognition and segmentation is better than other inception models, such as Inception-ResNet-v1, Inception-v3, and Inception-v4.

![Figure 4: Overall flow of our proposed scheme](image)

Inception-ResNet-v2 is a combination of residual connections and an inception module. The front module of Inception-ResNet-v2 is a stem layer, which generates a 35 × 35 × 256 feature map using five convolutions of 3 × 3, single max-pooling of 3 × 3, and one convolution of 1 × 1 from the 299 × 299 × 3 input image. In general, the stem layer is used to maximize the performance of inception neural networks. In the figure, the pink boxes indicate the various types of inception modules. Each inception module acts as multiple convolution filters, such as 1 × 1 and 3 × 3, which are designed for efficient visual feature extraction. Fig. 5 compares the various schemas of the Inception-ResNet-A, Inception-ResNet-B, and Inception-ResNet-C modules. Each module extracts general and local visual features simultaneously and reduces the computation resources. This is because the inception modules replace any n × n convolution by a 1 × n convolution followed by an n × 1 convolution. This form can significantly reduce computational cost as n increases. The green box represents the reduction modules of Inception-ResNet-v2. Fig. 6 shows detailed schemas of the reduction-A and reduction-B modules. The reduction-A module reduces the 35 × 35 × 256 feature map to a 17 × 17 × 896 feature map, and reduction-B reduces the 17 × 17 × 896 feature map to an 8 × 8 × 1792 feature map. These two modules reduce the size of the features from the
previous layer and pass them to the next layer. Atrous spatial pyramid pooling (ASPP) extracts the high-level features from the results of the Inception-ResNet-C module and fed them to the decoder module. The decoder infers the semantic information of each pixel by using the extracted representative features. The DeepLab v3+ decoder module was used to infer the wrinkle and cell area of the skin texture image. The DeepLab v3+ decoder performs up-sampling twice, which restores the dimensions to the input image size. During up-sampling, the encoder module first concatenates the corresponding low-level features from the stem layer. Then, the encoder module up-samples by a factor of four to infer the pixel segmentation. The last layer is the soft-max layer, which outputs a vector that represents the probability distribution of the potential outcomes. This layer takes the output of multiple dimensional vectors and rescales them in the range from 0 to 1, with a sum of one. Through the arguments of the maxima (argmax) function, the segmentation results for all pixels can be obtained. If the argmax function calculates the probability as zero, it is predicted as a cell class. In contrast, if the argmax function calculates the probability as one, it is predicted as a wrinkle class.

Figure 5: Detail schema of Inception-ResNet-v2 network (a) Inception-RestNet-A module (b) Inception-ResNet-B module (c) Inception-ResNet-C module

Figure 6: Detailed schema of Inception-ResNet-v2 network (a) Reduction-A module (b) Reduction-B module
To verify the segmentation performance of Inception-ResNet-v2, the soft-max activation layer was visualized, as presented in Fig. 7c. The white pixels belong to the cell class and black pixels belong to the wrinkle class. In the figure, the red boxes indicate false segmentation candidates.

![Figure 7](image_url)

**Figure 7:** Example of soft-max activation and prediction result (a) Input image (b) Ground truth (c) soft-max activation (d) Prediction result from the Deeplab v3+ with Inception-ResNet-v2 backbone

To alleviate the false segmentation problem, a tree-based ensemble framework, namely, LightGBM, was used. LightGBM has obvious advantages, such as high training speed, low memory usage, and high prediction accuracy, when processing large-scale data. In this case, LightGBM was used to improve the segmentation quality by considering the output probabilities of the soft-max layer as input. For this, we sequentially calculated the probabilities of the soft-max layer extracted from the training images. The overall probability values were then accumulated and reshaped to a flattened form. If one image has $299 \times 299 \times 2$ soft-max probabilities, the probabilities were reshaped to $89,401 \times 2$ (class probabilities). Using class labels of the ground truth, we trained all reshaped-probabilities as the input of LightGBM. The segmentation improvement is shown in the experimental section.

### 3.3 Morphological Processing

Although CNN and LightGBM show good performance in wrinkle and cell segmentation, their results are not always satisfactory. For instance, the segmentation includes areas wherein it is unclear whether they contain wrinkles or cells, which degrades the reliability of the segmentation results [36–39]. Another problem is false segmentation pixels, which appear as noise. Most skin-related 2DIA studies attempted to eliminate noise using median and linear filters, such as Gaussian and Wiener [40]. These filtering methods often lead to structural changes. To avoid this, an MP method was used. Morphological processing has the advantage of removing a single-pixel line, dot, or tiny pixel structures while maintaining the structural shape. MP performs shrinking and growing processes. The shrinking process rounds large structures and the growing process removes small structures. To achieve this, the dilation and erosion processes were used together. For instance, for the segmentation result illustrated in Fig. 8a obtained by means of CNN and LightGBM methods, Fig. 8b shows the result of MP, in which false segmented areas were removed or merged.
3.4 Skin Texture Feature Extraction

After segmentation, various skin features are extracted from the segmentation results. In this study, four skin features that are essential for evaluating skin condition and aging were considered. They are the length of wrinkle lines, wrinkle width, number of cells, and area of detected cells. The length of wrinkle lines is easily identified using the skeleton pixels classified as wrinkle class. To calculate the wrinkle width, we used Algorithm 1. The number of cells and their area were calculated using the polygon mesh detection algorithm (PMDA), which is presented in Algorithm 2 [12]. Fig. 9 depicts the steps for skin texture feature extraction.

Algorithm 1: Wrinkle feature extraction

Input: Segmented wrinkle $SW$

Output: Wrinkle width $W_w$, Wrinkle length $W_l$

Binary wrinkle skeleton $S \leftarrow$ watershed_transform($SW$);
Wrinkle contour $WC \leftarrow$ canny_edge_detection($SW$);
Angle $A \leftarrow$ PCA_angle($S$);

for each point $p$ in $S$ and each angle $a$ in $A$
do
  Measurement line $ML \leftarrow$ find_orthogonal($a, p$)
  [Start point $s$, end point $e$] $\leftarrow$ find_start_end_points($ML, WC$);
  localWrinkleWidth $+= euclidean_distance(s, e)$;
endo

$W_l = cnt(S)$;
$W_w = localWrinkleWidth/cnt(ML)$;

return $W_w, W_l$

Algorithm 2: Polygon mesh detection

Input: Segmented wrinkle $SW$

Output: Number of cell $C_n$, Average area of detected cell $C_{avg_area}$

Binary wrinkle skeleton $S \leftarrow$ watershed_transform($SW$);

(Continued)
for each point $p$ in $S$
    do
        recursively_seek_4_neighbors();
        localCellArea++;
        if $p$ belongs to image border then
            localCellArea = 0;
        end if
        $C_n$++;
        totalCellArea += localCellArea;
    while not reaching the borderlines
end for
$C_{\text{avg\_area}} = \text{totalCellArea}/C_n$;
return $C_n$, $C_{\text{avg\_area}}$

Figure 9: Steps for skin texture feature extraction (a) Input image (b) Segmentation result (c) Wrinkle-related feature extraction (d) Cell-related feature extraction using PMDA

4 Results and Discussion

4.1 Dataset and Model Configuration

To construct a skin image dataset, $50 \times$ magnified facial images from 365 healthy subjects were collected. All measurements were conducted in a room at a temperature of $23 \pm 3\,^{\circ}\text{C}$ and humidity of $50 \pm 10\%$. Fig. 10 shows the sample images of our dataset. In the experiment, none of the subjects applied any makeup on the face.

To train the Inception-ResNet-v2 model, the resolution of the input images was first set to $299 \times 299$. Then, data augmentation was conducted using random vertical flip, random horizon flip, 10\% zooms, and 10\% shear changes. The stochastic gradient descent with a momentum (SGDM), learning rate of $1 \times 10^{-3}$, and L2 regularization were then used to configure the training strategy. For verification, 80\% of augmented facial images were used as training and validation sets, and the remaining 20\% as the test set. To train LightGBM, the following conditions were set: $1 \times 10^{-3}$ learning rate, 5,000 iterations with early stopping, 10 max depth branching, 1,024 max leaves, and the gradient-boosted decision trees (GBDT) method. 10-fold cross-validation was conducted to assess the LightGBM performance. The ratio of the training set, validation set, and test set is equal to that of the Inception-ResNet-v2.
4.2 Performance Evaluation of Segmentation

In the evaluation of segmentation performance, three metrics were used: mean accuracy (MA), mean of intersection over union (MIOU), and mean of boundary F1 score (MBF) [41]. MA, which can be defined by Eq. (1), indicates the percentage of correctly identified pixels as wrinkle class or cell class. For each class, accuracy is the ratio of precisely matched pixels to the total number of pixels in that class, according to the ground truth. In Eq. (2), $TP$, $FN$, and $FP$ represent the number of true positives, the number of false negatives, and the number of false positives, respectively. In addition, $i$ is the total number of images.

\[
MA = \frac{1}{i} \times \sum_{n=1}^{i} \frac{TP_n + TN_n}{TP_n + TN_n + FP_n + FN_n},
\]

(1)

MIOU is another popular metric for measuring the performance of image segmentation and can be expressed as Eq. (2). IOU indicates the overlap percentage between the prediction pixels and the ground truth pixels.

\[
MIOU = \frac{1}{i} \times \sum_{n=1}^{i} \frac{TP_n}{TP_n + FN_n + FP_n},
\]

(2)
The last metric is the MBF score between the segmentation result and the ground truth. MBF, which is usually used to evaluate the contour matching, indicates how close the boundary of the segmentation matches the boundary of the ground truth. Eq. (3) shows how to calculate the MBF. In Eq. (3), $P_c$ and $R_c$ indicate the precision and recall of class $c$, respectively, and $B_{cps}$ and $B_{cgt}$ indicate the boundary binary map of the predicted segmentation and the ground truth in class $c$, respectively. In Eq. (3), “[[]]” is the Iverson bracket notation, where $[[z]] = 1$ if $z = \text{true}$ and 0 otherwise. In addition, $d()$ represents the Euclidean distance measured in pixels. In Eq. (4), $BF$ is defined as the harmonic mean of the precision and recall values with a distance error tolerance. Finally, we obtained the MBF by averaging the per-image BF scores.

$$P_c = \frac{1}{|B_{cps}|} \sum_{z \in B_{cps}} \left[ \left[ d(z, B_{cgt}) < \theta \right] \right] \quad \text{and} \quad R_c = \frac{1}{|B_{cgt}|} \sum_{z \in B_{cgt}} \left[ \left[ d(z, B_{cps}) < \theta \right] \right]$$

$$BF_c = \frac{2 \times P_c \times R_c}{R_c + P_c},$$

In the experiment, we considered diverse segmentation methods including Choi’s method, U-Net, SegNet, Deeplab v3+ with ResNet backbone families, and Panoptic-Deeplab with ResNet-101 backbone for comparison. Tab. 1 compares their wrinkle and cell class segmentation performances. As is clear from the table, the higher the score, the better the segmentation result. Our proposed scheme showed the best segmentation performance compared with other methods. It means that our proposed scheme significantly improved the segmentation quality in terms of finding the borderlines and regions of wrinkle and cell. Choi’s method presented the worst performance because it is basically a handcraft-based method. In contrast, the Deeplab v3+ models with ResNet backbone families showed better performance than U-Net and SegNet. Panoptic-Deeplab scored higher than Deeplab v3+ models in MA evaluation, but it showed relatively lower accuracy in wrinkle and cell boundary matching than did Deeplab v3+ models. Fig. 11 presents the example of segmentation results. The first and second columns show the input skin texture image and the ground truth of the segmentation task, respectively. From the third to the last column, the segmentation results of different CNN models are presented. When the boundaries of the wrinkles and cells are difficult to distinguish in the image, SegNet and two DeepLab v3+ models showed relatively low segmentation performance compared with the proposed model.

| Model                                      | MA   | MIOU  | MBF   |
|--------------------------------------------|------|-------|-------|
| Choi et al. [8]                            | 0.664| 0.612 | 0.606 |
| U-Net [42]                                 | 0.672| 0.653 | 0.642 |
| SegNet [43]                                | 0.790| 0.688 | 0.726 |
| Deeplab v3+ (ResNet-18 backbone) [18]      | 0.798| 0.694 | 0.819 |
| Deeplab v3+ (ResNet-50 backbone) [18]      | 0.809| 0.698 | 0.823 |
| Deeplab v3+ (Inception-ResNet-v2 backbone) [18] | 0.842| 0.739 | 0.843 |
| Panoptic-Deeplab (ResNet-101 backbone) [44] | 0.847| 0.702 | 0.742 |
| Ours (w/o) MP                              | 0.851| 0.743 | 0.849 |
| Ours                                       | **0.854**| **0.749**| **0.852** |
Figure 11: Segmentation results of SegNet [43], Deeplab v3+ (Resnet-18 backbone), Deeplab v3+ (Resnet-50 backbone) [18], and the proposed scheme

4.3 Performance Evaluation for Skin Texture Feature Extraction

In this experiment, we evaluated the accuracy of skin feature extraction using four popular metrics. First, the accuracy of wrinkle line extraction is defined using Eq. (5). In this equation, $DWP$ and $WSEG$ represent the detected wrinkle pixels and ground truth pixels of the wrinkle, respectively. To calculate the accuracy, the equation first counts the correctly overlapped pixels between the $DWP$ and $WSEG$, and then, it divides them by the total number of detected wrinkle pixels.

$$\text{Accuracy}_{WL} = \frac{\#(DWP \cap WSEG)}{\#(DWP)} \times 100$$  (5)
To evaluate the accuracy of wrinkle width extraction, the mean absolute percentage error (MAPE) was used, which can be calculated using Eq. (6). In this equation, \( ML \) indicates the number of measurement lines, and \( AWW_i \) and \( EWW_i \) indicate the actual length and estimated length of the \( i \)th measurement line on the wrinkle width, respectively.

\[
MAPE_{WW} = \frac{100}{ML} \times \sum_{i=1}^{ML} \frac{|AWW_i - EWW_i|}{|AWW_i|}
\]  

(Eq. 6)

Eqs. (7) and (8) show how to count valid cells and calculate their accuracy. To count valid cells, the weighted distance was calculated between the detected cells \( DC \) and ground truth cells \( GC \). Then, the IOU was calculated between the \( DC \) and \( GC \). If the IOU between the \( i \)th matched \( DC \) and \( GC \) is greater than 0.6, it is counted as a valid cell.

\[
VC_i = \begin{cases} 
1, & \text{if } IOU(\text{DC}_i, \text{GC}_i) \geq 0.6 \\
0, & \text{otherwise}
\end{cases}
\]  

(Eq. 7)

\[
\text{Accuracy}_{VC} = \frac{\#\{VC\}}{\#\{GC\}} \times 100
\]  

(Eq. 8)

Finally, the MAPE of the valid cell area was measured using Eq. (9). Here, \( VC \) is the number of valid cells, and \( AGC_i \) and \( AVC_i \) are the areas of the \( i \)th ground truth cell and valid cell, respectively.

\[
MAPE_{AC} = \frac{100}{VC} \times \sum_{i=1}^{VC} \frac{|AGC_i - AVC_i|}{|AGC_i|}
\]  

(Eq. 9)

Tab. 2 shows the performance comparison of skin feature extraction. The same skin feature extraction methods were used for each segmentation model. CNN-based models such as SegNet, U-Net, Deeplab v3+, and Panoptic-Deeplab showed better performance than did Choi’s model, which is based on the handcrafted features. The proposed scheme showed the best performance in extracting skin features, and U-Net presented the worst performance. Despite the fact that Choi’s model used a handcrafted method, it performed better than the U-Net model because it was mainly designed to extract wrinkles and cells from dermoscopy images.

**Table 2: Performance comparisons of skin texture feature extraction**

| Model                           | Accuracy\(_{WL}\) | MAPE\(_{WW}\) | Accuracy\(_{VC}\) | MAPE\(_{AC}\) |
|--------------------------------|-------------------|----------------|-------------------|----------------|
| Choi et al. [8]                 | 63.242            | 7.160          | 61.517            | 6.184          |
| U-Net [42]                      | 62.118            | 7.319          | 61.154            | 6.519          |
| SegNet [43]                     | 68.176            | 7.461          | 63.517            | 5.342          |
| Deeplab v3+ (ResNet-18 backbone) [18] | 70.283           | 6.251          | 68.804            | 5.008          |
| Deeplab v3+ (ResNet-50 backbone) [18] | 71.721           | 6.130          | 70.760            | 4.859          |
| Deeplab v3+ (Inception-ResNet-v2 backbone) [18] | 72.498           | 5.821          | 73.499            | 4.802          |
| Panoptic-Deeplab (ResNet-101 backbone) [44] | 72.528           | 5.740          | 72.824            | 4.908          |
| Ours (w/o) MP                   | 73.312            | 5.625          | 74.118            | 4.716          |
| Ours                            | **74.204**        | **5.517**      | **74.833**        | **4.659**      |
5 Conclusions

In this study, a hybrid segmentation scheme for skin feature extraction using Deeplab v3+ with an Inception-ResNet-v2 backbone, LightGBM, and MP was proposed. As traditional 2DIA approaches for skin analysis use handcraft-based methods, it is difficult to obtain satisfactory results using them. To alleviate this problem, based on deep neural networks, which can perform efficient feature extraction from deep and complex networks, the hybrid approach was used to improve segmentation quality and skin feature extraction accuracy. To validate the effectiveness of the proposed scheme, extensive comparisons with other popular models were performed using diverse evaluation metrics. From the experimental results, the proposed scheme was confirmed to outperform the handcraft-based method and other popular CNN models under every evaluation metric. It is our belief that the proposed scheme can be used in diverse skin-related applications, such as skin damage estimation, skin condition assessment, and skin aging estimation.

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