

Article

Finding the Differences in Capillaries of Taste Buds between Smokers and Non-Smokers Using the Convolutional Neural Networks

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Featured Application: The aim of this work is to strengthen patient awareness and willingness to quit smoking by presenting them with the diagnostic results obtained using the capillaroscopy-based deep-learning artificial intelligence methods.

Abstract: Taste function and condition may be a tool that exhibits a rapid deficit to impress the subject with an objectively measured effect of smoking on his/her own body, because smokers exhibit significantly lower taste sensitivity than non-smokers. This study proposed a visual method to measure capillaries of taste buds with capillaroscopy and classified the difference between smokers and non-smokers through convolutional neural networks (CNNs). The dataset was collected from 26 human subjects through the capillaroscopy with the low and high magnification directly; of which 13 were smokers, and the other 13 were non-smokers. The acquired dataset consisted of 2600 images. The results of gradient-weighted class activation mapping (grad-cam) enabled us to understand the difference in capillaries of taste buds between smokers and non-smokers. Through the results, it was found that CNNs gave us a good performance with 79% accuracy. It was discussed that there was a shortage of extracted features when the conventional methods such as structural similarity index (SSIM) and scale-invariant feature transform (SIFT) were used to classify.

Keywords: capillaries of taste buds; convolutional neural network; deep learning; grad-cam; smokers; non-smokers



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1. Introduction

A lot of studies have confirmed that smokers exhibit significantly lower taste sensitivity than non-smokers [1]. The taste sensitivity level can be measured by electrogustometric (EGM) thresholds from various parts of the tongue (locus) [2]. After smoking cessation, thresholds of EGM decrease progressively and reach the taste sensitivity range of non-smokers depending on locus and time. It is known that the recovery in the posterior loci is complete after 9 weeks, and the recovery in the dorsal loci is observed only after 2 months or more. Smoking cessation results in a rapid recovery of taste sensitivity among smokers, with different recovery times [1,3]. Thus, it is considered that the use of taste sensitivity could be explored as a motivation tool for smoking cessation.

The function of capillaries of taste buds is the exchange of material between the blood and tissue cells for gustatory sensitivity [4]. Tobacco users are generally unaware of the effects of tobacco on general health, oral health, etc. [3]. The effect on sensory perception and the demonstration of its deficit to the subject might reveal an actual threat the smoker may wish to avoid. The taste function and the taste condition may be a tool that exhibits a rapid deficit to impress the subject with an objectively measured effect of smoking on

his/her own body because smokers exhibit significantly lower taste sensitivity than non-smokers [1,5,6]. From this point, the authors have an interest in whether there is a difference in capillaries of taste buds between smokers and non-smokers or not.

There have been two measuring methods of the taste sensitivity; for the whole mouth and for some mouth regions [2]. The evaluation for the whole mouth can be done with the use of colorless solutions of sweet, bitter, sour, and salt [7]. Then, the simplest regional test is EGM which was introduced in the clinical assessment of taste sensitivity during the 1950s [8]. In addition, contact endoscopy (CE) allows for both in vivo and in situ observations of pathology in the superficial layer of the tongue, nasal mucosa, vocal cords in the larynx-microsurgery and nasopharynx [9,10]. However, these methods are not easy for non-medical practitioners to use in daily life. Furthermore, images from CE are difficult for normal users to understand features related to diseases.

On the other hand, nailfold capillaroscopy is a non-invasive, inexpensive, and reproducible imaging technique to evaluate micro-circulations [11,12]. The capillaries are so abnormally altered that they can be seen with the naked eye, although magnification is usually required. And this method is used for the diagnosis of vascular dysfunction. That is the reason we used the nailfold capillaroscopy as the first testing. In nailfold videocapillaroscopy (NVC) qualitative assessment, scleroderma patterns can distinguish between primary and secondary Raynaud's phenomenon (RP) and represent an essential and reliable parameter for the early, as well as very early, diagnosis of systemic sclerosis (SSc) [13,14]. Thus, the recent introduction of capillaroscopic assessment into the new 2013 American College of Rheumatology (ACR)/European League Against Rheumatism (EULAR) classification criteria for SSc reflects its pivotal role in the diagnosis of the disease [15]. However, most studies using capillaroscopy focused on images of nailfold capillaries.

For the nailfold capillary analysis, there have been three approaches: manual, semi-automated, and automated segmentation [16]. The manual method depends on human-recognizable features and requires experts to perform certain tasks, rendering it impractical for mass and widespread use. [17]. The semi-automated method requires initial human intervention to mark the outer and inner parts of each capillary and requires data analysis, which may cause bias and mistakes [12]. The automated method combine a local threshold and the Simultaneous Truth and Performance Level Estimation (STAPLE) algorithm to distinguish nailfold capillaries [18]. However, these methods are contaminated by noise and require post-processing such as morphological operations. Recently, convolutional neural networks (CNNs) have been proposed for semantic segmentation [19–22]. Although these methods have been effectively applied on medical image segmentation tasks, such as liver, pancreas, MRI, and multiorgan, no CNN has been proposed for capillaries of taste buds.

Therefore, this study proposed a visual real-time method to measure capillaries of taste buds with capillaroscopy and classify the difference between smokers and non-smokers through CNNs. After that, this study confirmed which extracted featured points should be used to classify two classes.

This paper is organized as follows: In Section 2, related works about details of capillaries microscope and the data choosing for the testing; And we considered the algorithms with handcrafted feature extraction, such as SSIM, SIFT and algorithms without handcrafted feature extraction, such as convolutional neural networks (CNNs). After that, we chose the best method for classifying capillaries images which is our proposed method; Section 3 provides the results of the experiment settings and results; Section 4 provides the discussion; lastly, Section 5 gives the summary and conclusions of this work.

2. Methods

This section describes the capillaries microscope which is the small and simple device system to capture the capillaries images. Subsequently, we considered when choosing the nailfold capillaries or the blood vessels on tongue and give the reason of our choice. Our proposed method was implemented after testing two methods: the algorithms with handcrafted feature extraction, such as SSIM, SIFT and algorithms without handcrafted

feature extraction, such as convolutional neural networks (CNNs) on the applied data and chose the best.

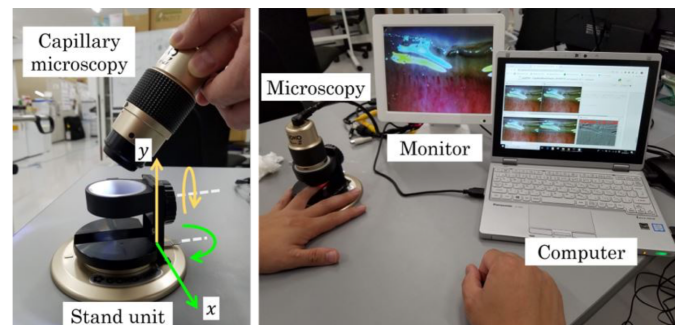
2.1. Experimental Environment and System

Table 1 shows pros and cons of different capillaroscopic devices [15]. Different devices can be used to perform capillaroscopy, as it is an *in vivo* imaging investigation that consists of a magnified view of the structural aspects of the microcirculation. The commercially available tools range from the wide-field microscope and videocapillaroscope to smartphone devices and are characterized by different portability, magnification, and costs. the USB-connected-typed microscopy is chosen because the necessary magnification for capillaries of taste buds in this study might be larger than $300\times$ and the device is simple enough for non-medical practitioners to use.

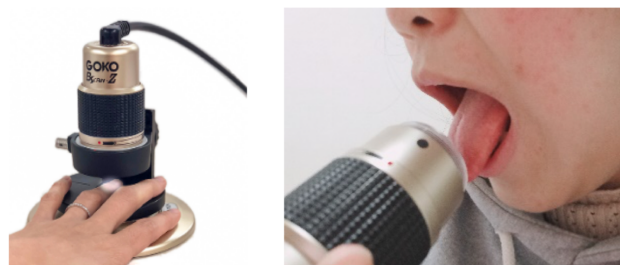
Table 1. Pros and cons of different capillaroscopic devices [15].

Device	Portability	Magnification	Costs
Stereomicroscopy	Not practical	Ranging between 10 and $200\times$	Medium to high
ophthalmoscopy	Very good	Up to $15\times$	Low
Dermatoscopy	Very good	Up to $10\times$	Low to medium
Smartphone devices	Very good	Up to $20\times$	Low
Digital USB microscopy	Good (laptop connection)	Up to $300\times$	Low to medium
Digital videocapillaroscopy	Good (laptop connection)	Raging between 50 and $500\times$	High

Figure 1a shows an overview of experimental environment and system consisted of the microscope with the low and high magnification. Table 2 [23] describes the specification of the experimental system of the microscopy with the low ($100\times$) and high ($410\times$) magnification.



(a) An overview of experimental environment and system consisted of the microscope with low and high magnification.



(b) Two measuring method according to two different measuring parts for nailfold capillaries of middle finger and taste bud capillaries of tongue.

Figure 1. (a) An overview of exp. environment consisting of the microscope with low and high magnification, (b) Measuring method for nailfold capillaries of finger on the left and measuring method taste bud capillaries of tongue on the right.

They are connected by cables and converter set to convert NTSC analogue images to uncompressed digital images for the real-time displaying on a monitor and saves them as video and still image. The microscope is a GOKO Bscan-Z (GOKO Imaging Devices, Kanagawa, Japan) with a vertical, cylindrical body and compact size. The body weight of the microscope is 150 g when the focus cap is attached; the size of the focus cap is diameter $\times$ length = $\phi 45 \times 10$ mm, and the body weight is 10 g. The diameter of the stand unit is $\phi = 120$, the camera holder is $\phi = 58$, and the height of the stand unit is 72.5 mm. The weight of the stand unit is 250 g. Because of the light body weight, it is easy to carry.

Table 2. Specification of microscope GOKO Bscan-Z.

Feature	Descriptions
2-ways-to-use	Desk and handheld.
Stand unit	Robust and is attached,
-	Reduces the tremors.
Zoom and focus	Powerful zoom,
-	Easy to focus
-	No need to change lenses.
Live video	Capture smooth without time lags.
Magnification	100 $\times$ to 410 $\times$ (on 14-inch monitor).
Dimensions	$\phi 45$ (external diameter) $\times$ 96 (length) mm
-	without attachment,
-	$\phi 45$ (external diameter) $\times$ 106 (length) mm
-	with attachment.

The range of the x-axis movement for the stand unit is 10 mm from the left to the right side, and that of the y-axis is 10 mm from the upper to the lower positions, as shown in Figure 1a. Although there is no range of the z-axis, a human subject can regulate his or her finger position. The real-time image of capillaries is projected onto a 14-inch monitor under a range of magnification from low to high magnification [11]. The user can spin the black middle part around to zoom out, zoom in and focus without any change of lens.

2.2. Data Acquisition

There is no opened data set to classify the difference of capillary distribution on nailfold and tongue between smokers and non-smokers in the world as we surveyed. Thus, it is necessary to measure the capillaries of the nailfold and tongue surface and make the data set directly. There are two measuring parts which had the contact between the tobacco and the human body while smoking. Figure 1b shows a description to measure capillaries of nailfold and taste bud by using the microscope. According to the measuring part, it is possible to attach or detach the microscope and stand unit.

Twenty-six human subjects (height: 172.2 ± 6.3 cm, weight: 68.3 ± 6.2 kg, Age: 24 ± 9 years old) were employed: 13 smokers in the university with the smoking careers of 5–10 years (class 1), and thirteen non-smokers who were university students without smoking careers (class 2). No subject from either class reported any health problem or a history of neurological disease, drug abuse, alcoholism, and medical constraints that might influence the experimental result. No major difference in body mass index (BMI) was observed between the two classes.

Figure 2 shows the location of the 9 recording loci on the surface of the tongue. The nine tongue loci were defined: Tip of the tongue middle (T), right (Tr) and left (Tl) where the density of fungiform papillae is highest, Dorsal right and left (Dr and Dl) where the density of fungiform papillae is lowest, Edge right and left (Er and El) on the foliate papillae, and Posterior right and left (Pr and Pl) just anterior to the circumvallate papillae.

After the authors explained the objectives and procedures of this study, the informed consent was obtained from all subjects. The experimental procedures were performed under the Declaration of Helsinki.

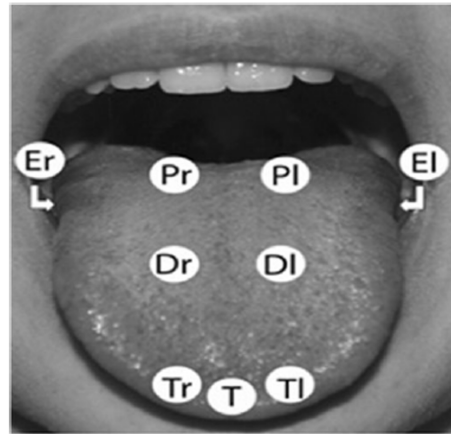


Figure 2. A location of the 9 recording loci on the surface of the tongue. T, tip; Tr and Tl, tip right and tip left; Er and El, edge right and edge left; Dr and Dl, dorsal right and dorsal left; Pr and Pl, fungiform papillae just anterior to the circumvallate papillae.

2.3. Data Training with Compound Model Scaling

Four different CNNs, with different different architectures (VGG [24], DenseNet [25], ResNet [26], and EfficientNet [27]) were trained on measured dataset. All training is performed using the Python programming language (version 3.8) on a workstation running on Jupyter Notebook with one Nvidia GeForce RTX 2080ti graphic cards (11 GB of RAM).

2.3.1. Model Scaling

All CNN [28] architectures follow the same general design principles of successively applying convolutional layers to the input, periodically downsampling the spatial dimensions while increasing the number of feature maps. While the classic network architectures (LeNet [29], AlexNet [30], and VGG) are comprised simply of stacked convolutional layers, modern architectures (Inception [31], ResNet, ResNeXt [32], DenseNet, and EfficientNet) explore new and innovative ways for constructing convolutional layers in a way which allows for more efficient learning. All these architectures are based on a repeatable unit which is used throughout the network.

For improving the performance, there are a lot of methods to scale up a CNN for different resource constraints. ResNet can be scaled up by regulating network depth which indicates layers, while WideResNet and MobileNet can be scaled by network width which indicates channels. It is also well-recognized that bigger input image size means bigger resolution that helps increasing accuracy with the overhead of more FLOPS (float point operations per second) that is a measure of computer performance. However, there is still the limitation to scale only one of the three dimensions (depth, width, and image resolution). That means that we should use only one scaling factor [27].

2.3.2. Compound Model Scaling

This study tries to use the CNN with compound model scaling, which means to use three scaling factors at the same time, and then evaluate the effect of compound model scaling on classification. The CNN layer i can be expressed as a function: $Y_i = F_i(X_i)$, where F_i is the operator, Y_i is output tensor, X_i is input tensor, with tensor shape $\langle H_i, W_i, C_i \rangle^1$, where H_i and W_i are spatial dimensions and C_i is the channel dimension [27,28]. A CNN N can be represented by a list of composed layers: $N = F_k \odot \dots \odot F_2 \odot F_1(X_1) = \odot_{j=1 \cdot \text{cot} \cdot k} F_j^{X_1}$, where $\odot$ means the connection between consecutive layers. CNN layers are often partitioned into multiple stages and all layers in each stage share the same architecture. Thus, the CNN can be defined as [27]:

$$N = \odot_{i=1 \dots s} F_i^{L_i}(X_{(H_i, W_i, C_i)}), \quad (1)$$

where $F_i^{L_i}$ denotes F_i of layer i is repeated L_i times in stage i , $\langle H_i, W_i, C_i \rangle$ denotes the shape of input tensor X of layer i .

When it is possible that all layers should be scaled with constant ratio, the problem for model scaling can be an optimization problem, which is to maximize the model accuracy for any given resource constraints. This is achieved by the following algorithm:

$$\begin{aligned} & \underset{d,w,r}{\text{maximize}} && \text{Accuracy}(N(d,w,r)) \\ & \text{subject to} && N(d,w,r) = \odot_{i=1 \dots s} \hat{F}_i^{d \cdot \hat{L}_i} (X_{\langle r \cdot \hat{H}_i, r \cdot \hat{W}_i, w \cdot \hat{C}_i \rangle}) \\ & && \text{Memory}(N) \leq \text{target} - \text{memory} \\ & && \text{FLOPS}(N) \leq \text{target} - \text{flops} \end{aligned} \quad (2)$$

where w, d, r are coefficients for scaling network width, depth, and resolution, and $\hat{F}_i, \hat{L}_i, \hat{H}_i, \hat{W}_i, \hat{C}_i$ are predefined parameters in baseline network.

The compound scaling method uses a compound coefficient ϕ to scale network width, depth, and resolution:

$$\begin{aligned} \text{depth:} & \quad d = \alpha^\phi \\ \text{width:} & \quad w = \beta^\phi \\ \text{resolution:} & \quad r = \gamma^\phi \\ \text{subject to} & \quad \alpha \cdot \beta^2 \cdot \gamma^2 \approx 2 \\ & \quad \alpha \geq 1, \beta \geq 1, \gamma \geq 1 \end{aligned} \quad (3)$$

where α, β, γ are constants that can be determined by a small grid search.

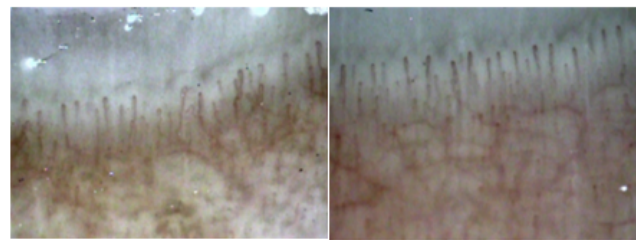
From the start with the baseline EfficientNet-B0, the compound scaling method is applied to scale it up with two steps. At first, it performs the search for α, β, γ through Equations (2) and (3) after fix $\phi = 1$. The values for EfficientNet-B0 are $\alpha = 1.2, \beta = 1.1$, and $\gamma = 1.15$ under the condition of $\alpha \cdot \beta^2 \cdot \gamma^2 \approx 2$. Then, scale up baseline network with different ϕ through Equation (3) after fix α, β, γ , to gain EfficientNet-B1 to B7.

3. Results

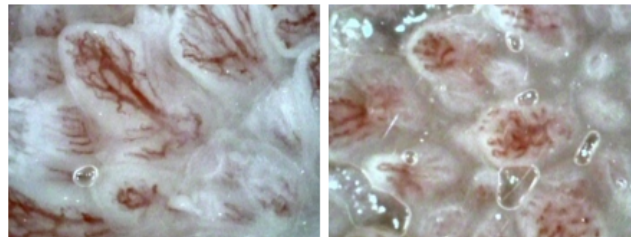
3.1. Example of Dataset

Figure 3 showed an example of the capillary image of nailfold (a) and taste bud (b) between smokers and non-smokers. The capillary image of the nailfold cannot give us the big difference between smokers and non-smokers. During daily activities such as washing, and doing something with hands, etc., the attached chemicals on the skin surface are eliminated.

On the other hand, the size of taste buds on the tongue for smokers seems to be smaller than that for non-smokers, thus, it looks like the condition of capillary distribution is different. The results of measurement as shown in Figure 3b was the dorsal for the middle of tongue (Dr and Dl) as shown in Figure 2.



(a) The nail fold image of no-smoker (left)/The nail fold image of smoker (right)

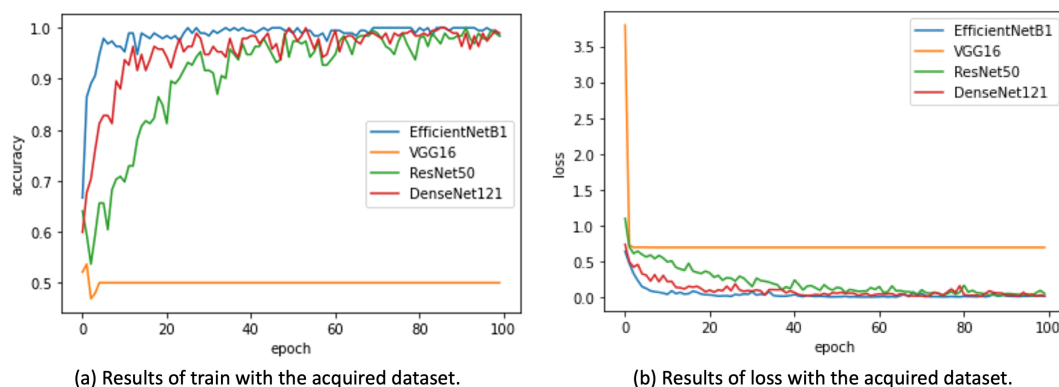


(b) The tongue image of non-smoker (left)/ The tongue image of smoker (right)

Figure 3. An example of the capillary image of nailfold (a) and tongue (b) between smokers and non-smokers.

3.2. Results of the Training and Prediction

Figure 4 shows the results of the training and loss for four applied models of CNNs: EfficientNet, VGG16, ResNet50, and DenseNet121. Although the results through VGG16 were not trained well, others showed good performance with nearly 80% accuracy. Our dataset includes 2 classes: smoker and non-smoker with 220 images per one class. We used rate (60/20/20) to split our dataset: 132 images in train set, 44 images in test set and validation set. Because the number of human subjects was small, the function of ImageDataGenerator Class in Keras was applied for increasing the number of images. It is well-known that CNN is only relevant when they are trained with a huge amount of data. In order to make the most of our few training examples, we can augment them via several random transformations. This helps to prevent overfitting and improves the generalization of the model. In Keras, this can be done by the class `keras.preprocessing.image.ImageDataGenerator`. The total number of images was 2600: 80% was used for the training and validation, and the left 20% was for the testing. We implemented experiment on four CNNs models: VGG, ResNet, DenseNet and EfficientNet with our dataset. And the result was showed on Tables 3 and 4.



(a) Results of train with the acquired dataset.

(b) Results of loss with the acquired dataset.

Figure 4. Results of training (a) and loss (b) for four models of CNNs: EfficientNet (blue color plot), VGG16 (orange color plot), ResNet50 (green color plot), and DenseNet121 (red color plot).

Table 3 represents the trainable and non-trainable parameters of four CNN models: EfficientNet-B1, VGG16, Resnet50 and DenseNet121. These results show that the total number of parameters of EfficientNet-B1 is the least of the four models, and is equal to the one fourth of the number of parameters of ResNet and half of VGG16 's. However, EfficientNet-B1 is the most effective, thus EfficientNet-B1 is considered to be the good choice in our data.

Table 3. The number of parameters for four models of convolutional neural networks (CNNs): EfficientNetB1, VGG16, ResNet50, and DenseNet121.

Model	Total Params	Trainable	Non-Trainable
EfficientNet-B1	6,576,513	6,514,465	62,048
VGG16	14,715,201	14,715,201	0
ResNet50	23,589,761	23,536,641	53,120
DenseNet121	7,038,529	6,954,881	83,648

Table 4 represents the results of prediction for four different algorithms of CNNs: EfficientNet, VGG16, ResNet50 and DenseNet121. It is shown that the results of prediction for EfficientNet indicates the best performance among four different CNNs.

Table 4. Results of prediction for four models of convolutional neural networks (CNNs): EfficientNet-B1, VGG16, ResNet50, and DenseNet121.

Model	Accuracy [%]	Loss
EfficientNet-B1	79.68	1.39
VGG16	50.00	0.69
ResNet50	60.94	0.89
DenseNet121	48.43	3.67

Compared to the result of algorithms with handcrafted feature extraction such as SSIM and SIFT, it was found that most of CNN algorithms without handcrafted feature extraction worked better for the image processing of capillaries, although VGG16 showed the failure.

3.3. Results of the Class Activation Map

The Class Activation Map (CAM) helps in the analysis of understanding as to what region of an input image influence the CNN's output prediction. The technique relies on the heat map representation which highlights pixels of the image that triggers the model to associate the image with a particular class.

Figure 5 compares the CAM images for three representative human subjects. Images are randomly picked from CNN validation set. The left image indicates the original image, the middle one indicates the results of CAM, and the right indicates the overlapped image with the original and CAM images. The results show that the CNNs tend to predict the non-smoker class by finding the large capillaries and white-colored taste buds. Additionally, it was also confirmed that the CNN model tended to predict the smoker class with the small capillaries and white-colored taste buds without capillaries [33].

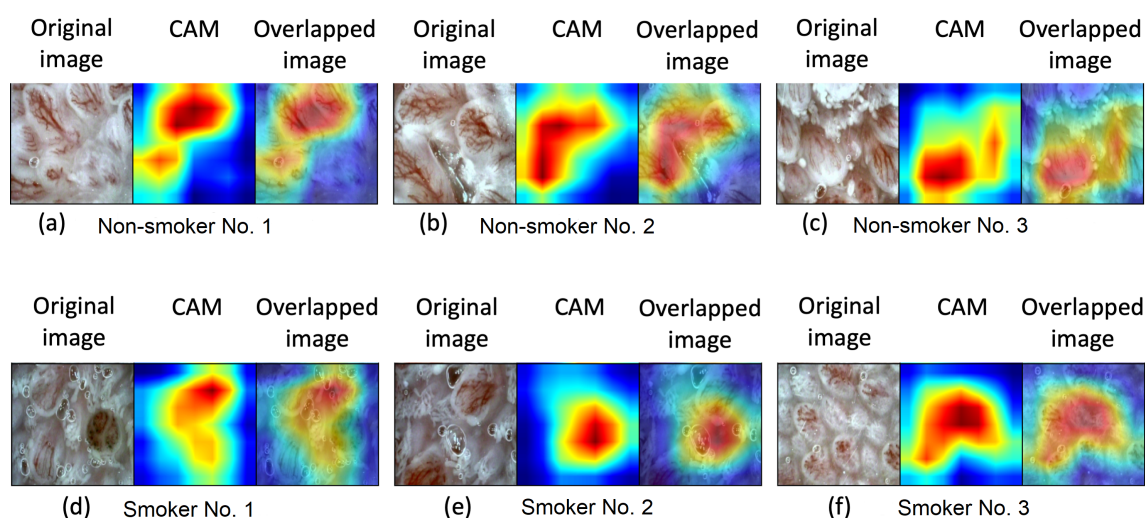


Figure 5. Results of Class Activation Map (CAM) visualization for Convolutional Neural Network (CNN).

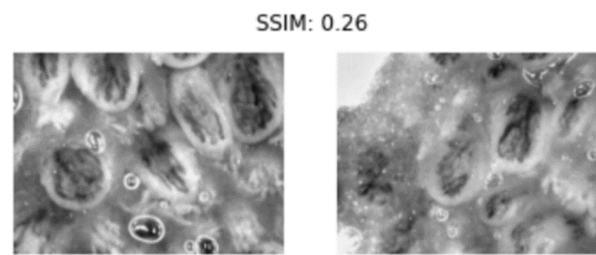
4. Discussion

We would like to analyze the CNNs model which is the best choice for classifying image data of capillaries. In the case, the algorithms with handcrafted feature extraction get the good result for capillaries images, there is no need to use the CNNs model without handcrafted feature extraction. However, the result of algorithms with handcrafted feature extraction gave us the bad score which showed in Figures 6 and 7.

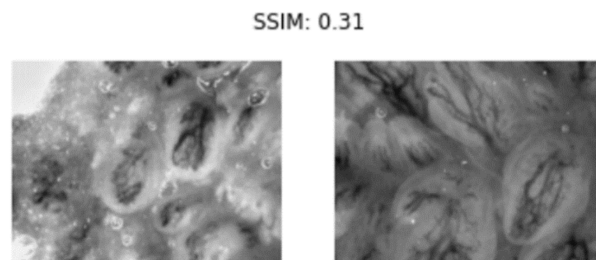
Figure 6 shows the results of the structural similarity index (SSIM) score [34] between two different tongue images for two smokers (a), and between two images for the smoker and non-smoker (b). SSIM score 1.00 represents that two images are same, and the 0.0 of that represents that those are fully different. The SSIM is a perceptual metric that quantifies image quality degradation based on the change in structural information, while also incorporating important perceptual phenomena, including both luminance masking and contrast masking terms. The difference to other methods, such as mean squared error (MSE) or Peak Signal-to-noise ratio (PSNR), is that these approaches estimate absolute errors. Structural information that pixels have strong inter-dependencies especially when they are spatially close. Through the results, it was confirmed that there were some difficulties in classifying the difference of tongue bud between two images of the smoker and the non-smoker through the SSIM score.

Figure 7 shows the results of the scale-invariant feature transform (SIFT) score [35] between two different tongue images for two smokers (a), and between two images for the smoker and non-smoker (b). The 100.0% of SIFT score represents that two images are same, and the 0.0% of that represents that those are fully different. SIFT keypoints of targets are first extracted from a set of reference images and stored in a database. A target is recognized in a new image by individually comparing each feature from the new image to this database and finding candidate matching features based on Euclidean distance of their feature vectors. Through the full set of matches, subsets of keypoints that agree on the target and its location, scale, and orientation in the new image are identified to filter out good matches. Through the results, it was confirmed that there were some difficulties in classifying the difference of tongue bud between two images of the smoker and the non-smoker through the SIFT score, because there was the shortage of good match points.

As a result, it was confirmed that there were some difficulties in classifying the difference of tongue bud between two images of the smoker and the non-smoker through the SSIM and SIFT score, because there was the shortage of good match points. It is necessary to have a lot of parameters when the feature from the image of capillaries is extracted. Thus, it is considered that algorithms with handcrafted feature extraction are not good for the image processing of capillaries.



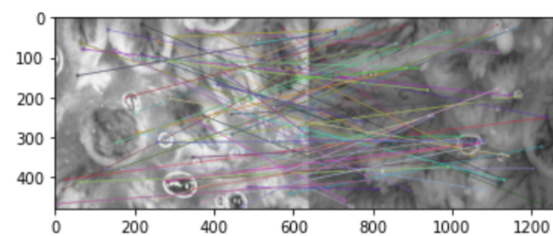
(a) Results of SSIM score between two tongue images for two smokers.



(b) Results of SSIM score between two tongue images for the smoker and the non-smoker.

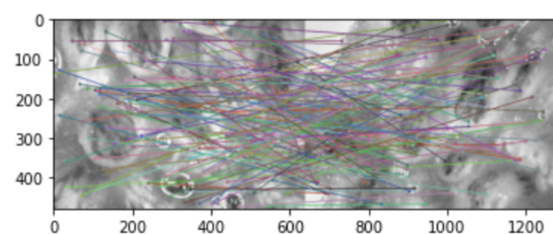
Figure 6. (a) The structural similarity index (SSIM) score between two tongue images for two different smokers, and (b) SSIM score between two images for the smoker and non-smoker.

The number of good match 1885
The score of SIFT is 8.06%



(a) Results of SIFT score between two tongue images for two smokers.

The number of good match 8267
The score of SIFT is 11.23%



(b) Results of SIFT score between two tongue images for the smoker and the non-smoker.

Figure 7. (a) The scale-invariant feature transform (SIFT) score between two tongue images for two different smokers, and (b) SIFT score between two images for the smoker and non-smoker.

5. Conclusions

In this study, some CNN models such as EfficientNet, ResNet, and DenseNet enabled us to train the data set of capillaries of taste buds, although two conventional methods such as structural similarity index (SSIM) and scale-invariant feature transform (SIFT) did not work well because of a shortage of extracted featured points. The results of class activation map (CAM) enabled us to understand a difference between smokers and non-smokers because CAM allowed us to know what were extracted featured points through CNNs.

The CNNs model without handcrafted feature extraction, especially EfficientNet with compound model scaling, proved the good performance to detect difference between smokers and non-smokers via tongue capillaries images which captured by the microscope, compared with conventional methods with handcrafted feature extraction. Then, the CAM enabled us to classify a difference in capillaries of taste buds between smokers and non-smokers. Our system with the bigger data, we can apply in hospital to support doctor in diagnosing disease, and it can be used daily for self-checking health and detecting the abnormal point of capillaries.

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Data Availability Statement: You can refer to the source code and the data set via this: <https://github.com/urgonguyen/efficientSmokingDetection>. The authors are waiting for any responses and comments.

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