SPECIAL ISSUE ARTICLE





Deep learning in medical imaging: A brief review

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Abstract

Researchers have used deep learning methods for a human level or better disease identification and detection. This paper reports, in brief, the recent work in deep learning identification of diseases occurring at three unique parts of the human body: the skin, the thorax, and the eye. While earlier reviews reported on the theory, applications, and challenges of such research, what distinguishes this work from the others is the reporting and comprehensive analysis of the key results. In doing so, the paper not only summarizes the major conclusions drawn from them but also emphasizes their weaknesses. The hope is to help the researchers see the big picture in deep learning classification of the diseases of the skin, the thorax and the eye, and guide them to find the right future research direction.

1 | INTRODUCTION

Physicians use different imaging modality techniques, such as MRI, CT, ultrasound, and X-ray to help them detect, diagnose, and cure serious diseases. With help from radiologists, they can interpret a vast number of diseases occurring at distinct parts of a human body, ranging from brain to lungs to skin. Still, the tremendous diversity in disease pathology can make human errors possible.

In search of computer methods to aid in disease detection and recognition, researchers have experimented on convolutional neural networks (CNN) since the mid-90s.¹ Used for hand-written digit recognition at first,² CNN's popularity increased with the discovery of deep CNNs in 2012.³⁻⁸ Since then, a growing number of researchers are using CNNs for medical image analysis (such as References 9-17). Some even show that CNNs can give comparable performance to medical doctors.¹⁸

We may use the Internet of Things (IoT) to connect medical devices in hospitals for the collection of big data. Each hospital can join a central unit this way for the automated detection of diseases on images using CNN. IoT is a crucial tool for reaching data in an instant, and artificial intelligence is essential for modeling for disease detection on images. Together they may form what we call The Internet of Medical Things (IoMT).¹⁹

CNN's distinct architectures and vast application areas (such as this and References 20-22) have likewise led the research groups to publish a few review papers on using deep CNNs in medical imaging (eg, References 23-27). While these papers focus on the theory, applications, and challenges of deep CNNs, they do not detail the results discovered by the researchers.

This survey paper, yet, aims to fill that gap by reporting the recent advances of deep learning use in medical imaging and analyzing the key results. In doing that, it hopes to offer the reader an opportunity to see the big picture, including any strengths and weaknesses that such research might have. The intention is not to cover the entire application areas in medical imaging but focus on only a few parts of the human body and be thorough. Hence, this survey paper includes

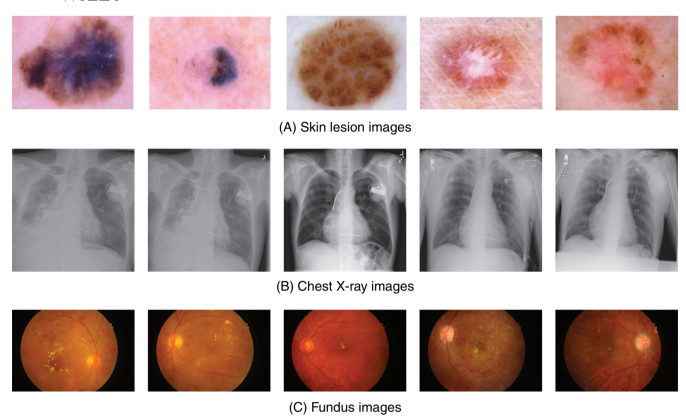


FIGURE 1 Sample images from each medical image category

the performance review of the deep learning methods for skin lesion classification, listing the earlier approaches, and comparing their accuracies. It reports the performances of the deep learning methods for the detection of throat diseases, again, listing the earlier deep learning approaches and comparing their performances. At last, it details the glaucoma recognition using deep learning methods and compares their performances.

Figure 1 shows the medical images of the three unique parts of the human body this paper is focusing on. The first is the human skin lesions. Figure 1A displays a variety of malignant and benign skin lesions. The second focus is on the human chest. Figure 1B displays a variety of chest X-ray images. The third and last focus is the human eye. Figure 1C shows distinct fundus images of the eye, in particular those that have glaucoma.

The organization of this review is in Section 2, we explain in brief a few CNN methods. In Section 3, we review the deep learning approaches for skin lesion classification and single, ensemble, and feature-based deep CNN models for skin lesion detection. Sections 4 and 5 summarize the deep learning methods for throat disease classification and report on the approaches for glaucoma classification. Section 6 gives the details of popular datasets used during the deep learning classification of skin, throat, and glaucoma diseases. Sections 7 and 8 report, analyze, and discuss the key results of researchers when they classified the three aforementioned diseases with deep learning. Section 9 concludes the review.

2 | DEEP LEARNING METHODS

We describe a few deep learning methods in this section.

2.1 | AlexNet deep network

AlexNet³ consists of eight layers. Input images go through five convolutional layers and then three fully connected layers. Images are convolved with 96 filters of size 11×11 in the first convolution layer. After, the output of the first convolutional

layer is used as input to the second convolutional layer. Then the pooling is performed on the output of the second convolutional layer. This input is filtered with 256 kernels of size 5×5 in the second convolutional layer. This convolution process is done for third and fourth and fifth convolutional layers. After, fully connected layers are constructed.

2.2 | VGG-16 deep network

VGG Net⁴ consists of 16 and 19 layers. The input images pass through two 64 filters with 3×3 sizes. Then, another two 128 sets of filters of 3×3 sizes are utilized to process the output of the first filtering. This convolution process is utilized with 256 and 512 sets of filters. Finally, fully connected layers are created.

2.3 | GoogleNet deep network

GoogleNet^{5,6} consists of 20 layers. This deep learning model is based on inception modules. Inception modules allow parallel filtering of the layers. This is network is much deeper than VGG and AlexNet deep learning models.

2.4 | ResNet deep network

ResNet-18, ResNet-32, ResNet-50, and ResNet-152⁷ are the mainly used models for medical image classification. These models are based on the residual learning method. Residual learning does not allow error accumulation on the convolution layers but enables a better representation of the content in the convolution layers.

3 | DEEP LEARNING IN SKIN LESION RECOGNITION

There are three different classification approaches for skin lesion classification. They are single deep learning methods, ensemble deep learning methods, and feature-based deep learning methods. The following sections review some of the recent research covering these methods.

3.1 | Single deep learning methods

Esteva et al²⁸ used GoogleNet Inception v3 deep learning model for skin lesion classification. They trained the deep learning method using 129 450 clinical images to recognize keratinocyte carcinomas and seborrheic keratoses, malignant melanomas, and benign nevi. The authors did two classification tasks. First, they performed keratinocyte carcinoma and seborrheic keratoses skin lesions classification. Second, they classified malignant melanomas and benign nevi detection lesions.

Han et al²⁹ deploy another model for twelve skin lesion classification. They based their proposed model on a fine-tuned ResNet-152 deep learning model. The created model allows basal cell carcinoma, squamous cell carcinoma, intraepithelial carcinoma, actinic keratosis, seborrheic keratosis, malignant melanoma, melanocytic nevus, lentigo, pyogenic granuloma, hemangioma, dermatofibroma, and warts classification.

Menegola et al³⁰ made use of ResNet-101 and Inception-v4 deep learning models for melanoma detection. They recognized melanoma among seborrheic keratoses and benign nevi skin lesions. The authors used several external datasets for creating deep learning models.

Zhang et al³¹ proposed attention-based residual convolution neural networks for melanoma recognition. This new deep network is ResNet-50 deep network modification. They enhanced residual learning by attention modeling. The performance evaluation of this model outperforms the classic ResNet-50 residual-based model.

Fujisawa et al³² used GoogLeNet CNN model for skin lesion classification. First, this method grouped malignant and benign types of skin lesions. Second, the method further divided malignant and benign types into epithelial and melanocytic groups. Likewise, they further divided each group into more subcategories. As a result, they could recognize twenty-one classes of skin lesions.

Rezvantalab et al³³ used DenseNet-210, ResNet-152, Inception v3, Inception ResNet v2 deep CNNs for melanoma classification. These models detect melanoma, basal cell carcinoma, actinic keratosis, benign keratosis, melanocytic nevi, dermatofibroma, vascular lesions, and atypical nevi. The ResNet-152 models outperform other models for melanoma and melanocytic nevi detection. The Inception v3 model outperforms the other models for actinic keratosis and benign keratosis classification. DenseNet-210 provided the best results for basal cell carcinoma and dermatofibroma classification.

Khamparia et al¹⁹ researched a framework driven by deep learning Internet of Health Things (IoHT) for skin lesion classification. As part of it, they used transfer learning and pretrained VGG19, Inception V3, ResNet50, and SqueezeNet models for benign and malignant skin lesion classification.

Sujitha et al²² presented a new image compression method for remote sensing using CNN. To balance image quality and the compression efficiency, they used two CNN, one at the encoding and the other at the decoding side.

3.2 | Ensemble deep learning methods

Matsunaga et al³⁴ used an ensemble of ResNet-50 CNNs for recognizing melanoma and seborrheic keratosis skin lesions. They based this approach on generating several ResNet-50 deep models. Each ResNet-50 model builds on different augmentation data. They fused each of the model probabilities for melanoma and seborrheic keratosis classes for skin class identification.

Harangi³⁵ used an ensemble of AlexNet, VGG, GoogleNet, and ResNet CNNs and joint model output probabilities for melanoma classification. Four different techniques lead the final probabilistic skin class prediction. The fusion strategies are the sum, product, maximal probabilities, and majority voting. This ensemble approach outperforms single AlexNet, VGGNet, GoogLeNet, and ResNet deep learning methods.

Nyiri and Kiss³⁶ used DenseNet, ResNet, VGG16, VGG19, Inception V3, and Xception CNNs for skin lesion classification. They predicted skin lesions using each of these CNN and again estimated lesion types by fusing output estimates of each of the models. The study presents two training strategies. One of them is to train the models using original images. Another is to train the models using cropped skin images around the region of interest. Joint CNNs outperform the single CNN for skin lesion recognition.

Nozdryn-Plotnicki et al³⁷ generated DenseNet, ResNet, Inception V3, and SEResNet deep learning models and fused these models for skin lesion prediction. The author's fusion approach led to higher accuracy of prediction than single CNN models.

Li and Shen³⁸ joint two ResNet models for melanoma detection. The authors based their approach on convolving images using two models and then combining saliency maps of the two models. At last, they estimated the skin lesions using joint output saliency maps.

Bi et al³⁹ used two ResNet models for recognizing melanoma among seborrheic keratosis and nevus. First, the authors predict melanoma using each of the ResNet models. Then they combine two ResNet architectures for the melanoma prediction. The performance results have shown that an ensemble of two ResNet models outperforms each of the single ResNet models.

3.3 | Feature-based deep learning methods

Mahbod et al⁴⁰ used AlexNet, VGGNet, and ResNet deep networks for feature extraction and then classified these futures using support vector machines. The authors likewise fused extracted features for skin lesion recognition. This study shows that fused features for lesion prediction outperform the characteristics of each of the deep learning models.

Hagerty et al⁴¹ used ResNet-50 network together with several feature images for melanoma recognition. Handcraft features and deep features of ResNet are joint for melanoma identification. The results have shown that fusing handcraft and deep features provided higher melanoma accuracy.

Serte and Demirel¹⁰ proposed Gabor wavelet-based deep learning to recognize malignant melanoma and seborrheic keratosis skin lesions. This method decomposed the skin image into seven directional Gabor band and then modeled each of the representations using AlexNet and ResNet-18 deep learning models.

Serte and Demirel¹¹ used a wavelet-based technique for melanoma classification. They based their approach on getting skin image into wavelet band and then modeling each image band using deep networks.

4 | DEEP LEARNING IN CHEST X-RAY RECOGNITION

Wang et al⁴² used AlexNet, GoogleNet, VGG-16, and ResNet-50 deep CNNs. Authors generated these models by ImageNet models kept on chest X-ray models. This work provided eight thorax disease classification and showed that the ResNet-50 deep learning model outperforms other deep learning models.

Yao et al⁴³ proposed multi-resolution-based multi-deep learning models. Authors use the ResNet model to reduce chest X-ray images and then use the DenseNet model to classify chest X-ray images in different resolutions.

Wang and Xia⁴⁴ used classification and attention-based deep learning models. The authors named this model a Chest-Net model. The classification part of the ChestNet model contained the ResNet-152 CNN model. The output of the classification map of this model is further modeled using convolutions for more exact classification. This work compared the proposed model with the classic ResNet model on the Chest X-ray 14 dataset. The performance evaluation of the ChestNet model showed that ChestNet outperforms the ResNet model.

Gundel et al⁴⁵ used the DenseNet-121 CNN for twelve chest x-ray image classification. The authors created this network by adapting the ImageNet pretrained DenseNet-121 model to chest images. The proposed method proved higher overall accuracy than AlexNet, GoogleNet, VGG-16, and ResNet-50 deep networks.

Rajpurkar et al⁴⁶ applied the DenseNet-121 CNN for the detection of pneumonia. The authors likewise used this model for the detection of 12 thorax diseases. Authors trained the DenseNet-121 model using Chest X-ray 14 dataset.⁴²

Li et al⁴⁷ used an attention-based ResNet CNN. Their approach applies to U-Net deep learning segmentation model and they classify the local regions of the X-ray images using the ResNet model.

Kermany et al⁴⁸ proposed transfer learning-based CNNs for pneumonia detection.

Varshni et al⁴⁹ joint DenseNet-169 model and support vector machines SVM. They represented chest X-ray appearances and related extracted feature vectors using DenseNet-169 deep networks. They got the image representations bypassing the architecture comprising 169 layers. Then they classified image appearances using SVM for health and pneumonia images.

Ayan and Unver⁵⁰ used Xception and VGG-16 deep networks pneumonia classification and healthy X-ray images. They base their approach on retraining existing models for X-ray images and use transfer learning-based models for detection pneumonia among pneumonia and healthy X-ray images. The VGG-16 model outperforms the Xception model for the classification task.

5 | DEEP LEARNING IN GLAUCOMA RECOGNITION

The following sections review in brief the earlier glaucoma research that used deep learning and other classification methods.

5.1 | Glaucoma classification using deep learning methods

Various experts have studied deep learning to detect glaucoma. Among them, Chen et al⁵¹ proposed a six-layer deep learning architecture for better characterization of glaucoma hidden patterns. In a separate work, Chen et al⁵² showed an automated deep learning feature learning method to detect glaucoma. For automated glaucoma detection, Orlando et al⁵³ used preprocessing and pretrained CNNs. Chai⁵⁴ joint domain knowledge with fundus images and deep learning for automatic glaucoma detection. Zilly⁵⁵ segmented optic cup and optic disc to detect glaucoma by using a convolutional learning architecture with ensemble learning and entropy sampling. Raghavendra et al⁵⁶ classified fundus images into normal and glaucoma classes using an 18-layer CNN.

5.2 | Glaucoma classification using other methods

Likewise, researchers have used a few other classifiers to detect glaucoma: sequential minimal optimization,⁵⁷ support vector machine (SVM),⁵⁸⁻⁶⁰ k-nearest neighbor,⁶¹ naïve Bayes (NB),⁶² least square-support vector machine (LS-SVM)^{63,64} and artificial neural network (ANN).^{65,66}

6 | DATASETS

This section describes the databases related to skin lesions, chest X-rays, and glaucoma imaging. Table 1 shows the number of images of skin databases. Table 2 shows the number of images of the different thorax diseases included in the databases, and Table 3 lists the number of fundus images in the databases related to types of glaucoma. Table 4 lists the abbreviations used in the datasets for each disease.

6.1 | Skin lesion datasets

6.1.1 | PH2

PH2⁶⁷ dataset contains images of two types of skin lesions: nevi and melanoma. Common nevus is observed on 80 of nevi skin lesion images and atypical nevus is observed on another 80. Malignant melanoma is observed on 40 skin lesion images.

6.1.2 | 2016 ISIC challenge

2016 International Skin Imaging Collaboration (ISIC) Skin Lesion Analysis Towards Melanoma challenge dataset⁶⁸ contains images of two skin lesion types: benign and malignant melanoma. Benign melanoma is observed on 388 skin lesion images and malignant melanoma is observed on 891.

6.1.3 | 2017 ISIC challenge

2017 ISIC challenge dataset⁶⁹ has images of three types of skin lesions: melanoma, seborrheic keratosis, and benign nevi. Of these images, melanoma is observed on 521 images, seborrheic keratosis on 386, and benign nevi on 1843 images.

TABLE 1 Skin lesion datasets and the number of images in each dataset category

Dataset	MEL	NV	ВСС	AK	SCC	SK	BK	DF	VASC
PH2 ⁶⁷	40	160	_	_	_	_	_	_	_
2016 ISIC ⁶⁸	1279	_	_	_	_	_	_	_	_
2017 ISIC ⁶⁹	521	1843	_	_	_	386	_	_	_
2018 ISIC ⁷⁰	1113	6705	514	327	_	_	1099	115	142
2019 ISIC ⁶⁹⁻⁷¹	4522	12875	3323	867	628	_	2624	239	253

TABLE 2 Chest X-ray datasets and the number of images in each dataset category

Dataset	NR	ATEL	CARD	EFF	INF	MS	NOD	PNA	PNX	CON	ED	EM	FB
Chest X-ray 8 ⁴²	84 312	5789	1010	6331	10 317	6046	1971	1062	2793	_	_	_	_
Chest X-ray 14 ⁴²	_	11 167	12 071	7646	16 316	142	6480	9836	4693	4544	8875	170	10 380

TABLE 3 Glaucoma datasets and the number of images in each dataset category

Dataset	Normal (Healthy)	Early Glaucoma	Glaucoma	Advanced Glaucoma	Total
HRF ⁷²	18	_	27	_	45
Drishti-GS1 ^{73,74}	31	_	70	_	101
RIM-ONE ⁷⁵	261	_	194	_	455
sjchoi86-HRF ⁷⁶	300	_	101	_	401
ACRIMA ⁷⁷	309	_	396	_	705
Dataset of ⁷⁸	786	289	_	467	1542

TABLE 4 Abbreviations for diseases.

Imaging	Abbreviation	Disease
Skin	MEL	Melanoma
Skin	NV	Melanocytic Nevi
Skin	BCC	Basal Cell Carcinoma
Skin	SCC	Squamous Cell Carcinoma
Skin	BK	Benign Keratosis
Skin	DF	Dermatofibroma
Skin	VASC	Vascular Skin Lesions
Chest	NR	Normal
Chest	ATEL	Atelectasis
Chest	CARD	Cardiomegaly
Chest	EFF	Effusion
Chest	INF	Infiltration
Chest	MS	Mass
Chest	NOD	Nodule
Chest	PPNA	Pneumonia
Chest	PNX	Pneumothorax
Chest	CON	Consolidation
Chest	ED	Edema
Chest	EM	Emphysema
Chest	FB	Fibrosis
Chest	PL	Pleural Thickening
Chest	HE	Hernia

6.1.4 | 2018 ISIC challenge

2018 ISIC challenge dataset⁷⁰ contains images of seven skin lesions. Melanoma is observed on 1113 images, melanocytic nevus on 6705, basal cell carcinoma on 514, actinic keratosis on 327, benign keratosis on 1099, dermatofibroma on 115 and vascular lesion on 42 images.

6.1.5 | 2019 ISIC challenge

This dataset⁶⁹⁻⁷¹ has images of eight different skin lesions. Melanoma is observed on 4522 images, melanocytic nevus on 12 875, basal cell carcinoma on 3323, actinic keratosis on 867, benign keratosis on 2624, dermatofibroma on 239, vascular lesion on 253 and squamous cell carcinoma images on 628 images.

6.2 | Chest X-ray datasets

6.2.1 | Chest X-ray 8 and chest X-ray 14

Chest X-ray 14 dataset⁴² is one of the largest datasets related to chest X-ray images. This dataset has images of 14 thorax diseases, including atelectasis, cardiomegaly, effusion, infiltration, mass, nodule, pneumonia, pneumothorax,

consolidation, edema, emphysema, fibrosis, pleural thickening, hernia. The dataset also contains images that are observed to have no thorax diseases.

Chest X-ray 8 dataset, on the other hand, contains images of eight different thorax diseases.

6.3 | Glaucoma datasets

6.3.1 | HRF

In this dataset,⁷² there are 45 fundus images. Diabetic retinopathy is seen on 15 images and glaucoma is seen on another 15. The rest of the images contain no eye disease.

6.3.2 | Drishti-GS1

This dataset^{73,74} contains 101 fundus images. Seventy-one of these images are glaucomatous and 30 images contain no eye disease.

6.3.3 | RIM-ONE

RIM-ONE r2 dataset⁷⁵ has 455 fundus images. The sign or suspicion of glaucoma is observed on 200 these images. The rest of the images contain no eye disease.

6.3.4 | sjchoi86-HRF

sjchoi86-HRF dataset⁷⁶ contains 401 fundus images. Of these, 101 are glaucomatous. Three hundred images contain no sign of eye disease.

6.3.5 | ACRIMA

This dataset⁷⁷ has 705 fundus images. Glaucoma is observed on 396 of them. There is no eye disease on 309 images.

7 | PERFORMANCE EVALUATION

The area under the receiver operating characteristic curve (AUC), accuracy, sensitivity, and specificity (SP) performance merits are used to test the accuracy of the methods. We can describe accuracy, sensitivity, and specificity as:

$$Accuracy = \frac{TP + TN}{TP + TN + FP + FN},$$
 (1)

Sensitivity =
$$\frac{TP}{TP+FN}$$
, (2)

Specificity =
$$\frac{TN}{TN+FP}$$
, (3)

where true positive, positive, true negative, false positive, and false negative are denoted as TP, TN, FP, and FN, respectively.

TABLE 5 AUC values of recently studied deep learning models for skin lesion detection

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Authors	Method	Network	Dataset	MEL	SK	NV	BCC	DF	AK	VASC
Serte and Demirel ¹¹	Wavelet trans.	ResNet-18	2017 ISIC	0.96	0.81	_	_	_	_	_
Serte and Demirel ¹¹	Wavelet trans.	ResNet-50	2017 ISIC	0.74	0.80	_	_	_	_	_
Serte and Demirel ¹¹	Wavelet coeff.	ResNet-18	2017 ISIC	0.82	0.82	_	_	_	_	_
Serte and Demirel ¹¹	Wavelet coeff.	ResNet-50	2017 ISIC	0.96	0.87	_	_	_	_	_
Serte and Demirel 11	Wavelet coeff.	ResNet-50-18	2017 ISIC	0.91	0.87	_	_	_	_	_
Serte and Demirel ¹⁰	Gabor	AlexNet	2017 ISIC	0.85	0.70	_	_	_	_	_
Serte and Demirel 10	Gabor	AlexNet	2017 ISIC	0.95	0.80	_	_	_	_	_
Serte and Demirel ¹⁰	Gabor	ResNet-18	2017 ISIC	0.84	0.73	_	_	_	_	_
Serte and Demirel 10	Gabor	ResNet-18	2017 ISIC	0.96	0.86	_	_	_	_	_
Harangi ³⁵	Images	Ensemble	2017 ISIC	0.85	0.93	_	_	_	_	_
Zhang et al ³¹	Images	Attention	2017 ISIC	0.86	0.95	_	_	_	_	_
Li and Shen ⁴⁷	Lesion Indexing	Ensemble	2017 ISIC	0.91	_	_	_	_	_	_
Matsunaga et al ³⁴	Images	Ensemble	2017 ISIC	0.87	0.95	_	_	_	_	_
Diaz ⁷⁹	Images	Ensemble	2017 ISIC	0.85	0.96	_	_	_	_	_
Menegola et al ³⁰	Images	Ensemble	2017 ISIC	0.87	0.94	_	_	_	_	_
Bi et al ³⁹	Images	Ensemble	2017 ISIC	0.87	0.92	_	_	_	_	_
Mahbod et al ⁴⁰	Deep features	Ensemble	2017 ISIC	0.87	0.95	_	_	_	_	_
Mahbod et al. ⁴⁰	Images and features	Ensemble	2017 ISIC	0.84	0.97	_	_	_	_	_
Hagerty et al ⁴¹	Features	Ensemble	2017 ISIC	0.90	_	_	_	_	_	_
Hagerty et al ⁴¹	Images and features	Ensemble	2017 ISIC	0.94	_	_	_	_	_	_
Rezvantalab et al ³³	Images	DenseNet-210	2018 ISIC	0.938	_	0.97	0.993	0.999	0.981	1
Rezvantalab et al ³³	Images	ResNet-152	2018 ISIC	0.944	_	0.973	0.991	0.998	0.976	1
Rezvantalab et al ³³	Images	Inception-v3	2018 ISIC	0.934	_	0.97	0.986	0.99	0.984	1
Rezvantalab et al ³³	Images	Inception-ResNet-v2	2018 ISIC	0.932	_	0.957	0.986	0.993	0.968	0.963

7.1 | Skin lesion classification

Table 5 shows the performance comparison of some of the newest research on skin lesion detection. The highest AUC values in each category are indicated in bold.

These recent studies have focused on the 2017 and 2018 ISIC challenge databases. The results in bold of studies that used the 2017 ISIC challenge database show that frequency-domain use of the images has improved the AUC performance for melanoma skin lesion classification. Using an ensemble of CNN has given a better AUC performance than image frequency domain use for seborrheic keratosis classification.

Rezvantalab et al³³ have used the 2018 ISIC challenge dataset for skin lesion classification. The results in bold of Table 5 show that all four CNN models used have almost equally shown high skin lesion detection results, with ResNet-152 architecture being the best of them.

7.2 | Chest X-ray classification

Table 6 reports the AUC performance comparison of unique CNN architectures on Chest X-ray 8 and Chest X-ray 14 datasets. Again, the highest AUC values are given in bold. We see from them that for both datasets the higher-layered CNN architectures have performed the best in chest X-ray classifications.

TABLE 6 Area under the receiver operating characteristic curve values of recently studied deep learning models for thorax disease detection

Authors	Network	Dataset	ATEL	CARD	EFF	INF	MS	NOD	PNA	PNX	CON	ED	EM	FB	PL	HE
Wang et al ⁴²	AlexNet	Chest X-ray 8	0.645	0.692	0.664	0.604	0.560	0.648	0.549	0.742	_	_	_	_	-	-
Wang et al ⁴²	GoogleNet	Chest X-ray 8	0.630	0.705	0.688	0.608	0.536	0.557	0.599	0.782	_	_	_	_	-	-
Wang et al ⁴²	VGG-16	Chest X-ray 8	0.628	0.708	0.650	0.589	0.510	0.655	0.510	0.751	_	_	_	_	-	-
Wang et al ⁴²	ResNet-50	Chest X-ray 8	0.706	0.814	0.736	0.612	0.560	0.716	0.633	0.789	_	_	_	_	-	-
Wang et al ⁴⁴	ResNet-152	Chest X-ray 14	0.743	0.874	0.811	0.677	0.783	0.697	0.695	0.809	0.725	0.832	0.822	0.804	0.751	0.899
Yao et al ⁸⁰	DenseNet	Chest X-ray 14	0.772	0.904	0.859	0.695	0.792	0.717	0.713	0.841	0.788	0.882	0.829	0.767	0.765	0.914
Yao et al ⁴³	ResNet-64	Chest X-ray 14	0.733	0.858	0.806	0.675	0.778	0.727	0.690	0.805	0.717	0.806	0.842	0.757	0.724	0.871
Gundel et al ⁴⁵	DenseNet-121	Chest X-ray 14	0.767	0.883	0.828	0.709	0.821	0.758	0.731	0.846	0.745	0.835	0.895	0.818	0.761	0.896

Method	Network	Dataset	AUC
Diaz-Pinto et al77, 77	Xception	HRF	0.83
Diaz-Pinto et al ⁷⁷	Xception	Drishti-GS1	0.80
Diaz-Pinto et al ⁷⁷	Xception	RIM-ONE	0.85
Diaz-Pinto et al ⁷⁷	Xception	sjchoi86-HRF	0.77
Diaz-Pinto et al ⁷⁷	Xception	ACRIMA	0.76
Ahn et al ⁷⁸	Average CNN	78	0.94
Ahn et al ⁷⁸	Inception v3	78	0.93
Chen et al ⁵¹	6 layer CNN	ORIGA	0.831
Shibata et al ⁸¹	ResNet	Private	0.965
Christopher et al ⁸²	ResNet-50	Private	0.91
Li et al ⁸³	AlexNet	ORIGA	0.8384
Li et al ⁸⁴	Inception v3	Private	0.986

TABLE 7 Area under the receiver operating characteristic curve (AUC) values of recently studied deep learning models for glaucoma classification

7.3 | Glaucoma classification

Table 7 shows the AUC comparison of different recent studies on glaucoma classification. The best performing single CNN architectures for this classification are the results in bold. We see from them that ResNet and Inception v3 are among the best performing architectures for glaucoma classifications.

8 | DISCUSSIONS

The results have shown us that for improved performance the studies have preferred to use an ensemble of CNNs or frequency domain representations on skin lesion images rather than single CNN architectures. However, the observation is that no unique way of improving the performance exists, that is, a method that works for the detection of one skin lesion might not work for the detection of the others. What we have further observed is that the researchers have not used ensemble or frequency domain methods so far for the detection of skin lesions included in the 2018 ISIC challenge dataset.

The results of thorax disease detection on two different datasets have indicated that high layered CNN architecture use has resulted in the best performance. To see how generalizable this observation is, we need further research that applies the thorax disease detection to other different datasets.

One observation of glaucoma classification is that if the training and testing datasets are different, it affects the performance (see Diaz Diaz-Pinto et al⁷⁷ vs the others in Table 6). Moreover, the researchers need to check if their results are generalizable by applying the detection architectures to people of different ethnic backgrounds.

9 | CONCLUSION

In this paper, our goal has been to review the newest research on deep learning methods for skin, chest, and eye disease identification. Unlike previous research, we have aimed to report, analyze, and discuss the results provided on these topics. To accomplish this, we have compared the performances of different deep learning methods used for the detection of skin lesions and chest and eye diseases. First, we described the disease datasets, and then we listed the number of images included in these datasets. After, we provided the performance evaluation of proposed deep learning methods that used these datasets.

Our analysis has shown that the performances of deep learning methods depend on their architectures and the number of layers. We likewise observed that for skin classification, the researchers need to apply ensemble or frequency domain methods to newer datasets. For thorax disease detection, they also need to apply higher layered CNN models to other datasets and for glaucoma classification, they should consider a more diverse population of patients.

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How to cite this article: Serte S, Serener A, Al-Turjman F. Deep learning in medical imaging: A brief review. *Trans Emerging Tel Tech.* 2020;e4080. https://doi.org/10.1002/ett.4080