

# Tissue Artifact Segmentation and Severity Analysis for Automated Diagnosis Using Whole Slide Images

by

# Galib Muhammad Shahriar Himel (21-92094-2)

Master of Science in Computer Science (Intelligent Systems)

Department Computer Science
Faculty of Science and Technology
American International University-Bangladesh

September 2022

# **Declaration**

I declare that this thesis is my original work and has not been submitted in any form for another degree or diploma at any university or other institute of tertiary education. Information derived from the published and unpublished work of others has been acknowledged in the text and a list of references is given.

\_\_\_\_\_

**Galib Muhammad Shahriar Himel** 

21-92094-2

# **Approval**

The thesis titled "Tissue Artifact Segmentation and Severity Analysis for Automated Diagnosis Using Whole Slide Images" has been submitted to the following respected members of the board of examiners of the department of computer science in partial fulfilment of the requirements for the degree of Master of Science in Computer Science on 15.09.2022 and has been accepted as satisfactory.

Dr. Hossain Md. Shakhawat Dr. Md. Sakir Hossain Assistant Professor & Supervisor Associate Professor & External Department of Computer Science Department of Computer Science American International University-Bangladesh American International University-Bangladesh \_\_\_\_\_ Dr. Dip Nandi Dr. Tabin Hasan Professor & Director (Computer Science) Professor & Head (Graduate Program) Department of Computer Science Department of Computer Science American International University-Bangladesh American International University-Bangladesh Dr. Tafazzal Hossain Dr. Carmen Z. Lamagna

Professor & Dean (Computer Science)

Department of Computer Science

American International University-Bangladesh

Vice Chancellor

American International University-Bangladesh

## Acknowledgement

First of all, I would like to express my gratitude to my thesis supervisor Dr. Hossain Md. Shakhawat, Assistant Professor, Department of Computer Science, American International University-Bangladesh (AIUB) for his adequate guidance and motivation in all aspects of my thesis work.

I would like to express my gratitude to my external supervisor, Dr. Md. Sakir Hossain, Associate Professor, Faculty of Science and Technology, American International University Bangladesh (AIUB) for his comments which were instrumental in the implementation phase of the thesis.

Finally, I would also like to express my gratitude towards 'SHINSHU MEDICAL UNIVERSITY', as the dataset needed for my research was collected from that institute.

#### **Abstract**

Pathological specimens are scanned using a whole slide imaging (WSI) scanner to create digital images for monitor-based diagnosis and analysis. The slide needs to be rescanned if the image quality is occasionally unsatisfactory because of focus-error or noise. An evaluation method for reference less quality was previously proposed, although some artifacts (such as tissue folds and air bubbles) were identified as false positives. Those artifacts must be disregarded when deciding whether or not rescanning is required because slide preparation, not scanning, is what causes them. By differentiating between the sources of quality degradation—the focus-error or noise created by the scanner and the artifact occurred during the slide preparation—this research suggests a strategy for a more practical approach to assess WSI quality. In the method, two UNet based architecture: DoubleUNet and ResUNet++ is used to segment the area in which the artifact is located. Besides the segmentation, the artifact severity is also analyzed using ensemble learning. To do the ensemble learning, at first transfer learning was implemented using various pre-trained model and then, from the generated model best models are selected for ensemble learning. In case of artifact segmentation above 97% accuracy is achieved and in the case of severity analysis 99.99% accuracy is gained which is almost perfect.

# Table of Contents

1.Introduction	9
2.Literature Review	13
3.Methodology	16
3.1. Overview of The Proposed Method	17
3.2. Artifact Detection Method	
3.3. Artifact Severity Classification	19
4.Experimment	21
4.1. Dataset	21
4.2. Experimental Setup	23
4.2.1. Experimental setup for artifact segmentation	23
4.2.2. Experimental setup for severity classification	24
5.Results and Discussion	26
5.1. Artifact Segmentation Result	26
5.2. Severity Analysis/ Classification Result	45
5.3. Critical Analysis	55
6.Conclusion	56
Bibliography	57
Author Information	60

# **List of Figures**

Figure 1: (a) focus blur affected (b) noisy (c, d) under graded region of interest [7]	10
Figure 2: Tissue artifacts (Tissue-fold and Air-bubble)	11
Figure 3: Proposed method	12
Figure 4: WSI system [42]	16
Figure 5: Proposed method for artifact segmentation and severity analysis	17
Figure 6: DoubleUNet and ResUNet++ architecture	18
Figure 7: Ensemble learning	19
Figure 8: Dataset for segmentation	21
Figure 9: Dataset for severity classification	22
Figure 10: Best results of ResUNet++ on Tissue-fold dataset	27
Figure 11: Testing comparison (on Tissue-fold using ResUNet++)	28
Figure 12: Best results of DoubleUNet on Tissue-fold dataset	30
Figure 13: Testing comparison (on Tissue-fold using DoubleUNet)	31
Figure 14: Comparison graph for ResUnet++ and DoubleUNet for Tissue-fold dataset	32
Figure 15: Best results of ResUNet++ on Air-bubble dataset	34
Figure 16: Testing comparison (on Air-bubble using ResUNet++)	35
Figure 17: Best results of DoubleUNet on Air-bubble dataset	37
Figure 18: Testing comparison (on Air-bubble using DoubleUNet)	38
Figure 19: Comparison graph of ResUNet++ and DoubleUNet for Air-bubble dataset	39
Figure 20: ROC comparison	42
Figure 21: Comparison graph of ResUNet++ and DoubleUNet	44

# **List of Tables**

Table 1: Hyper parameter tuning for segmentation experiment	23
Table 2: Hyper parameter optimization	24
Table 3: Pre-trained models; their image dimensions	24
Table 4: Best segmentation result for Tissue-fold using ResUNet++	26
Table 5: Best segmentation result for Tissue-fold using DoubleUNet	29
Table 6: Best segmentation result for Air-bubble using ResUNet++	33
Table 7: Best segmentation result for Air-bubble using DoubleUNet	36
Table 8: Accuracy and IOU for Tissue Fold dataset.	40
Table 9: Accuracy and IOU for Air Bubble dataset	40
Table 10: 6-Fold Cross validation for Tissue fold dataset	41
Table 11: 6-Fold Cross validation for Air bubble dataset	41
Table 12: Over all comparison of segmentation models	43
Table 13: Final ROC based comparison of segmentation models	43
Table 14: Transfer Learning for base model selection	45
Table 15: Top 6 models description	53
Table 16: Ensemble Learning Combinations using 10 meta learners	54

#### 1.Introduction

The development of a whole slide imaging (WSI) scanner made it possible to digitize pathological works [1][2][3]. Glass slide specimens are converted into high-resolution digital images by the scanner for use in computational pathology and pathology practice. In hospitals and pathology labs, it enables us to undertake picture analysis and diagnosis using automated algorithms or human viewing on a computer screen. It pushes the boundaries of light microscopy and has great promise not just for research and education but also for primary and secondary diagnoses [4]. However, it is important to take into account a number of factors before using a WSI scanner as a standard medical device, including image quality, color variability, inadequate consistency of slide preparation, and image format [3][5]. In this research, we set our focus on mainly tissue artifact area segmentation and artifact severity analysis which are primarily base on image quality.

In research labs, hospitals, medical colleges and pathology departments a whole Slide Imaging System would be used to scan biological specimens on a regular basis [6]. WSI scanners may produce images of insufficient quality. Images of poor quality can seriously hinder diagnosis and render analysis ineffective. The operator is currently required by the WSI system to guarantee the accuracy of the scanned image. However, such manual evaluation is expensive and unreliable since it is susceptible to fatigue, inconsistent results, and prejudice. In order to provide access to high-quality photographs, it is important to analyze the quality of scanned images automatically [7].

Failures during scanning might result in focus blur or noise being introduced into the WSI scanner, which is one of the main reasons of image quality loss. For scanning the glass slides, focus points can be manually or automatically chosen. When focus points are chosen from an area with a different focus depth than the normal tissue area, the surrounding area is rendered out of focus. The independent random value that differs from adjacent image pixels is known as noise. Figure 1 demonstrates the noise, focus blur and under graded tissue area.

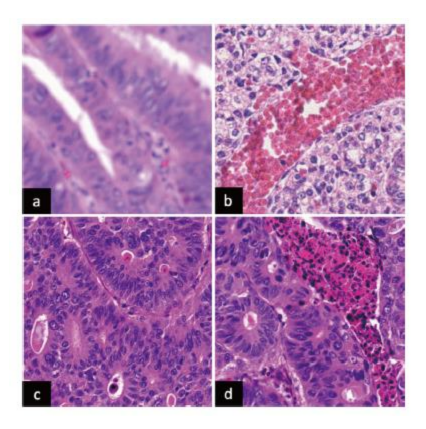


Figure 1: (a) focus blur affected (b) noisy (c, d) under graded region of interest [7].

It is important to rescan the slide if the focus error or noise significantly affects the scanned image. Hashimoto et al. previously suggested a reference less quality evaluation approach (RQM) to assess the scanned image's quality in order to identify slides that needed to be rescanned. The major causes of quality failure in the WSI scanner are sharpness and noise measurement, which is how this approach assesses the quality of images.

Tissue artifacts are produced during the glass slide preparation stage, which is another reason why image quality fails [9]. The two main artifacts discovered in WSI are a tissue fold and an air bubble [10], as illustrated in fig. 2. Tissue artifacts deceive analysis and diagnosis by concealing or changing important information. Furthermore, focusing errors are brought on by such objects [11]. They present various focus depths, and if focus locations are chosen from these artifacts, the surrounding area is rendered out of focus. Tissue artifacts like air bubbles and tissue fold would also be found using the prior RQM approach [8]. If a slide has a scanning problem, rescanning it can restore the image's quality but not remove tissue artifacts. It is useless to rescan the slide if the WSI quality is low due simply to tissue artifacts. For both analysis and quality assessment, artifacts

should be found and disregarded. Therefore, to apply the judgment's outcome in the workflow and address the quality failure, an efficient system must identify the reason of quality failure.

# **Tissue Fold** Air Bubble

Figure 2: Tissue artifacts (Tissue-fold and Air-bubble).

But, simply removing all the artifacts is not a feasible solution. There can be such slides those contain minor artifacts while contain major tissue specimen which can be proven important to identify disease. So, those kind slides can generate important features for machine learning. But, due to presence of minor artifacts if those images are disregarded then again the same problem which is wrong diagnosis of disease can occur. That is why the severity of the artifact is to be determined before removing from the diagnosis process.

In this research, we suggest adding the tissue artifact detection stage and the severity analysis of the artifacts, as shown in fig. 3. The suggested method uses a UNet-based image segmentation methodology to identify artifacts, then classifies the images based on their severity before deciding whether or not to exclude them from the diagnosis process.

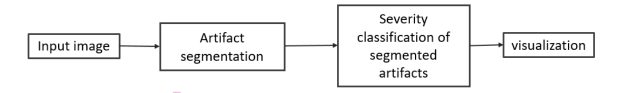


Figure 3: Proposed method.

#### 2.Literature Review

Although reduced-reference methods are available in some situations, methods for evaluating image quality can be roughly categorized into two groups: those that use reference images and those that do not. However, because a perfect reference picture is not available, reference-based evaluation cannot be used to identify a quality problem in the WSI scanner system. For the purpose of evaluating the quality of applications involving cameras, displays, video transmission, and codecs, references-less quality evaluation methods (RQMs) have been created [12][13][14][15][16][17][18][19][20].

Focus inaccuracy and noise are the key problems in WSI applications for managing the quality of digital pathology [8][21][22][23]. The RQMs have been used in this way, taking specific distortion into account. The method put out by Hashimoto et al. based on sharpness and noise assessment predicts both a subjective quality score as well as an objective score, such as a root mean square error, because computational image analysis is a crucial issue in both visual diagnostics and digital pathology. In order to gauge the reproducibility of the scanner, Shrestha et al. provided a method to assess the WSI's quality [21]. This approach, however, is unsuitable for practical applications because it depends on a reference image for evaluation. Another technique used deep learning to find out-of-focus areas, but it ignored noise-induced distortion [22]. Because it is effective and simple, the system suggested in this work is based on the method developed by Hashimoto et al. To distinguish between the tissue artifacts and scanning errors, it is vital to take into account the source of degradation while using the WSI scanning procedure in digital pathology.

In several areas of computer vision, object detection or categorization in an image is carried out. Both supervised and unsupervised methods exist. The supervised method works well for the proposed system's air-bubble and tissue-fold detection because the unsupervised method could mix up artifacts with other anomalies that could have clinical significance. Template matching is a well-known and established supervised method for object detection. Statistical techniques like SVM and neural network classifiers have also been frequently used to identify objects in an image. Handcrafted features have been employed in this type of supervised classification, including local binary patterns, the gray level co-occurrence matrix (GLCM), and scale-invariant feature

transform. It is well known that employing the sequential feature selection strategy when choosing the features for a classifier can increase performance [24][25].

More recently, non-handcrafted features that CNN gathered from the data were made possible by the development of deep learning technology. CNN can also be applied directly to an image. It has been demonstrated that a deep learning strategy, like CNN, performs better than a handmade feature based approach, while requiring a larger training dataset. When applied to a small dataset, it frequently becomes over fitted [26]. Transfer learning is typically employed in such instances to alleviate the problem of data limitation [27][28]. A network can avoid overfitting by using dropout and data augmentation [29][30]. In this work, we compared the segmentation method based on DoubleUNet and ResUNet++ with a VGG19 model as decoder. As for image classification process for severity analysis, we used various transfer learning methods for base model selection and lastly used several meta learners for stacking based ensemble learning.

In histopathology, object detection using segmentation and image classification methods were used in previous researches. A review paper on importance of artifact detection and possibility of wrong disease diagnosis due to presence of artifact is published by Syed Ahmed Taqi at el.in the year 2018 [31]. In the year 2019, Kay R.J. Oskal at el. Published a paper regarding a UNet based tissue segmentation method in WSI system [32]. An automated blur detection method is proposed by Xavier et al. in the year 2013 [33]. Samar khan published a paper regarding the probability of wrong diagnosis due to artifact in the year 2014 [34]. Ozan Oktay et al. introduced a unique attention gate (AG) model for medical imaging in 2018 that automatically learns to focus on target structures of various sizes and forms. Models trained with AGs intuitively learn to emphasize prominent features that are helpful for a particular task while suppressing irrelevant regions in an input image. Because of this, they can do away with the need for explicit external tissue/organ identification modules in cascaded convolutional neural networks (CNNs) [35]. A SVM based approach was taken by Hossain Md. Shakhawat in the year 2020 to evaluate image quality in WSI systems [7]. In the year 2019, Morteza Babaie et al. proposed a unique approach that mimerges the sym with dense layer 201 model to enhance the feature extraction process for medical images [36]. In the year, 2009, Pinky A. Bautista had done a research regarding artifact detection in WSI systems using basic object detection model [37]. In the year 2015, Hang Wu proposed a method regarding detecting blur artifacts in WSI [38]. Sonal Kothari published a research article on

eliminating tissue fold artifact in the year 2013 [39]. Wei Lu et al. in the year 2013 researched on the presence of air bubble artifact in slides [40]. Pinky A. Bautista again in the year 2010 published another paper regarding enhancing the visualization and detection of tissue fold artifacts in whole slide imaging system [41].

So, from the literature, it is apparent that, continuous researches are being done by the researchers to further improve the accuracy of the detection of artifacts and also how to eliminate the artifacts automatically using different approaches.

#### **Problem Statement**

Existing methods only detects the tissue artifacts but does not analyze the severity of a particular artifact.

But, severity analysis is important otherwise, significant amount of tissue can be eliminated from the specimen.

Till now no work has been done to analyze the severity of the artifact. That's why we focused our research on mainly two things:

- Improve the accuracy of artifact segmentation.
- Analyze the artifact severity.

#### 3. Methodology

The WSI scanner is used for scanning whole slide images. It is hard to determine the region of interest using human eye. So various approaches are being taken to enhance the automatic artifact detection. Among those approaches there are several machine learning approaches applied for advanced image processing.



Figure 4: WSI system [42].

Pathological specimens are a bit complex to extract meaningful image data from. Because from a lot of images which are apparently similar, important distinguishable features have to be extracted. For this purpose, we have proposed a method which will extract necessary information which will be used for successful feature extraction; ultimately leading to efficient pathological analysis and disease diagnosis. In our proposed method, we have applied image annotation, data pre-processing, segmentation, classification and visualization respectively in an organized manner.

#### 3.1. Overview of The Proposed Method

In the preliminary stage, the images are captured using WSI system. The whole image will be divided into many small parts and segmentation is done for each part.

The proposed method takes input images then detect the artifact using UNet based image segmentation approach. This segmentation process will identify the tissue artifact and will highlight that part while separating it from the background. After the segmentation process, the artifacts are being classified in terms of severity. For this classification task, Ensemble learning is applied. Ensemble learning requires several methods. For selecting those methods transfer learning is applied using different combinations of parameters. After the severity classification being done, the final output is generated and sent for visualization.

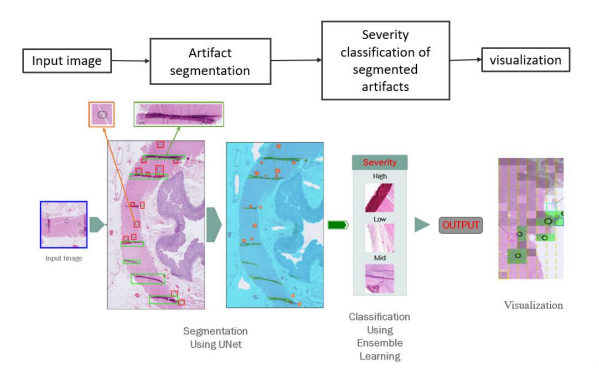


Figure 5: Proposed method for artifact segmentation and severity analysis.

In the visualization screen, those identified artifacts along with their severity level will be shown and pathologists will be able to successfully pinpoint the affected region.

#### 3.2. Artifact Detection Method

For, artifact detection we have used UNet based architecture for image segmentation. At first we used basic UNet. But the results were not good. Then we used 'DoubleUNet' and 'ResUNet++' as they have become quite popular in recent times for medical image segmentation tasks.

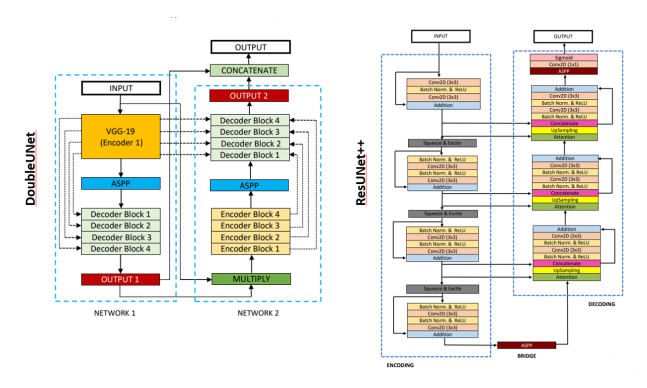


Figure 6: Double UNet and Res UNet++ architecture.

DoubleUnet architecture uses VGG-19 model as encoder. The first layer of UNet generates an output which is multiplied with the primary input and that multiplied output is given as input to the second layer of UNet. The primary output and the secondary output concatenate and generates the final output.

The ResUNet++ architecture follows a basic ResNet architecture and builds a bridge between the ResNet and UNet architecture. In simple words, it is a UNet architecture which follows ResNet architecture for segmentation.

#### 3.3. Artifact Severity Classification

After the artifacts are being identified using segmentation techniques, the images need to be classified. For classification, we primarily used transfer learning using various pre-trained model. After getting the results we have compared the results according to the accuracy and loss values. Then we selected the best models to ensemble the results using stacking method.

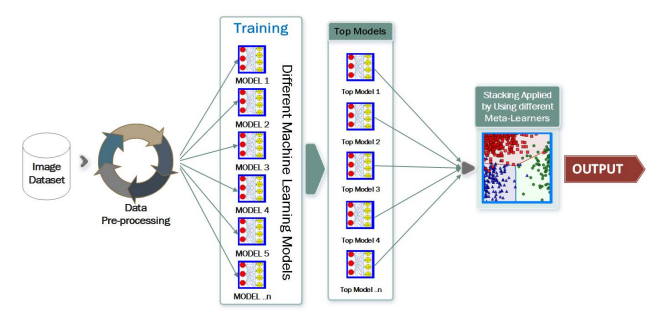


Figure 7: Ensemble learning.

For implementing ensemble learning, we tested 10 different meta learners:

- Logistic Regression
- K Nearest Neighbor
- Support Vector Machine
- Decision Tree
- Random Forest
- Ada Boost Classifier
- XGB Classifier
- Gradient Boost Regressor
- Gradient Boost Classifier
- Gaussian Naïve Bias

Severity analysis requires extensively complex calculation process for selecting differentiable features for image classification. The transfer learning models we used are not able to give perfect results from every perspective. Each model focuses on particular type of feature extraction policy. That's why, to combine those strategies, ensemble learning is used.

Within the ensemble learning itself there are approaches those are specific to different areas. Meta learners used in stacking policy acts in different ways. That's why several meta learners assessed to find out which meta learner generates the best results.

#### 4.Experimment

#### 4.1. Dataset

Several slides are used in this experiment, which included the tissues of different organs including kidney, liver, heart, lunge, brain etc. So the dataset is organ independent.

The slides were collected from various biomedical laboratories of the world.

#### **Segmentation task**

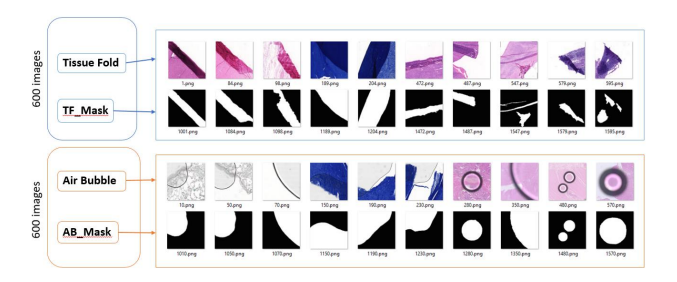


Figure 8: Dataset for segmentation.

Each category consists of 600 images. Data augmentation was applied for data balancing. Every images were segmented manually in adobe Photoshop by using binary masking function.

#### **Classification task**

Taking images from the same source the dataset was categorized in three categories:

- High severity
- Low severity
- Mid severity

The dataset was again augmented for balancing purpose. This time the dataset contains 1380 images. This data annotation was done by certified expert.

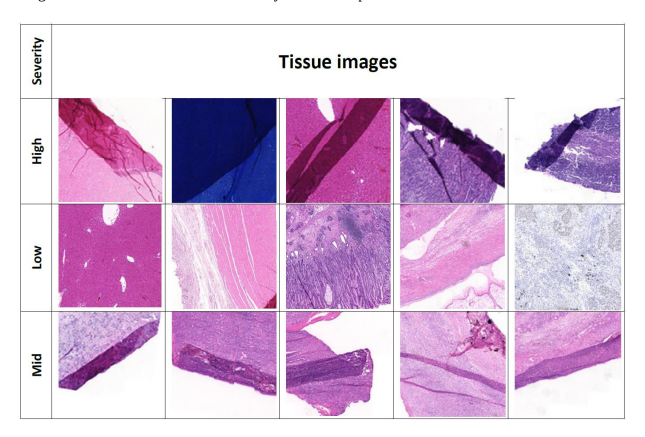


Figure 9: Dataset for severity classification.

#### 4.2. Experimental Setup

Our experiment is divided in two parts: a) artifact segmentation; b) severity analysis/ classification.

#### 4.2.1. Experimental setup for artifact segmentation

For the segmentation experiment the hyper parameters were tuned in different combinations.

Table 1: Hyper parameter tuning for segmentation experiment.

Train images	480
Valid images	60
Test images	60
Batch size	8
UNet architecture	DoubleUNet, ResUNet++
Optimizer	Adam, Adamax, <b>RMSprop</b> , SGD
Loss function	Dice Coef Loss, binary cross entropy
Learning Rate	Starting from 1e <sup>-4</sup>
ReduceLROnPlateau	monitor='val_loss', factor=0.1, patience=4
EarlyStopping	monitor='val_loss', patience=10

Different epochs were tried but the result was not always uniformly tuned and that's why we used early stopping function to prevent overfitting and gain a reliable value for segmentation.

#### 4.2.2. Experimental setup for severity classification

For classification task we used the following combinations described in table 2 and table 3.

Table 2: Hyper parameter optimization.

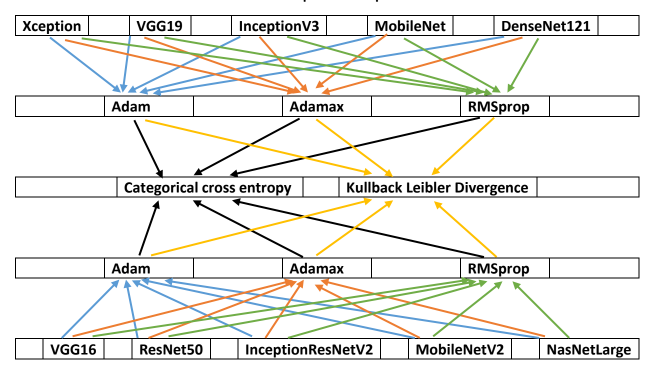
Hyper parameter	Optimization Space
Epoch	25
Batch size	32
Learning rate	0.0001
Optimizer	'Adam', 'Adamax', 'RMSprop'
Loss function	[Categorical cross entropy], [Kullback Leibler Divergence]
Class mode	categorical

For implementing ensemble learning, we had to run transfer learning several times using different pre-trained model in various combinations. We used the most famous 10 pre-trained model from the official website of 'keras'.

Table 3: Pre-trained models; their image dimensions.

Da	ta Augr	menta	tion	Image inpu	ıt dimension
Parameter		Value		Xception	224×224
Rescale		1./255		VGG16	224×224
Zoom range		0.3		VGG19	224×224
Rotation range	•	15.		ResNet50	224×224
Horizontal flip		True		InceptionV3	224×224
Data set		InceptionResNetV2	224×224		
Γotal images	Train in	nages	Test images	MobileNet	224×224
High	32	:0	140	MobileNetV2	224×224
_OW	32	.0	140	DenseNet121	224×224
Mid	32	.0	140	NasNetLarge	331×331
Grand Total	96	0	420		331331

Total 60 combinations of code were run to complete the experiment.



<b>Pretrained Model</b>	Optimizer	Loss function	calculation	Total combination
10	3	2	10 × 3 × 2	60

\*\*\*Note: "Accuracy Graph, ROC Curve and Confusion Matrices are generated for every combination but, for the paper writing convenience only the Accuracy graph, ROC curve and Confusion matrices of the Top 6 Models are mentioned in the Result section".

#### 5. Results and Discussion

#### 5.1. Artifact Segmentation Result

#### Tissue Fold: ResUNet++

Table 4 represents the hyper-parameter combination for the best result for tissue fold artifact using ResUNet++ architecture. RMSprop optimizer performs better than any other optimizers. The learning rate was set to 0.0001. But, the 'ReduceLROnPlateau' function reduces the learning rate at every step where validation loss increases. The training was stopped when Ir was  $1e^{-9}$ .

Table 4: Best segmentation result for Tissue-fold using ResUNet++.

Train images	480
Valid images	60
Test images	60
Batch size	8
Epoch	38 Epoch
Optimizer	RMSprop
Loss function	Dice Coef Loss
Learning Rate	$1e^{-4} > 1e^{-5} > 1e^{-6} > 1e^{-7} > 1e^{-8} > 1e^{-9}$
ReduceLROnPlateau	monitor='val_loss', factor=0.1, patience=4
EarlyStopping	monitor='val_loss', patience=10

loss	-0.9736	Val_loss	-0.9791
recall	29.65%	Val_recall	34.97%
precision	99.99%	Val_ precision	99.97%
Mean-IOU	85.81%	Val_mean-IOU	88.88%
IOU	94.87%	Val_IOU	96.05%
Dice Coef	97.36%	Val_Dice Coef	97.98%

Dice coef loss is defined as negative value that's why the loss values are negative.

#### Graph

Figure 10 represents the graph of mean IOU, Loss curve, precision curve, recall curve, dice-coefficient and the IOU. The last two graphs which are the dice-coefficient and IOU curve defines the actual performance of the model.

In the next page, figure 11 represents some of the example comparison of the above mentioned model.

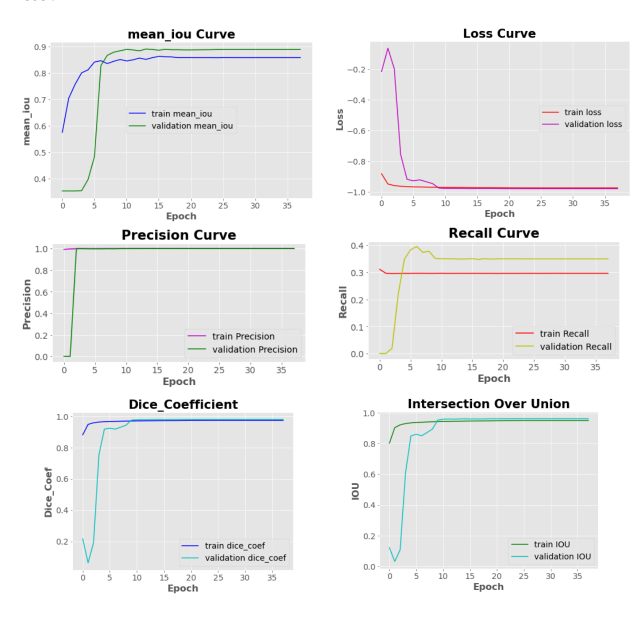


Figure 10: Best results of ResUNet++ on Tissue-fold dataset.

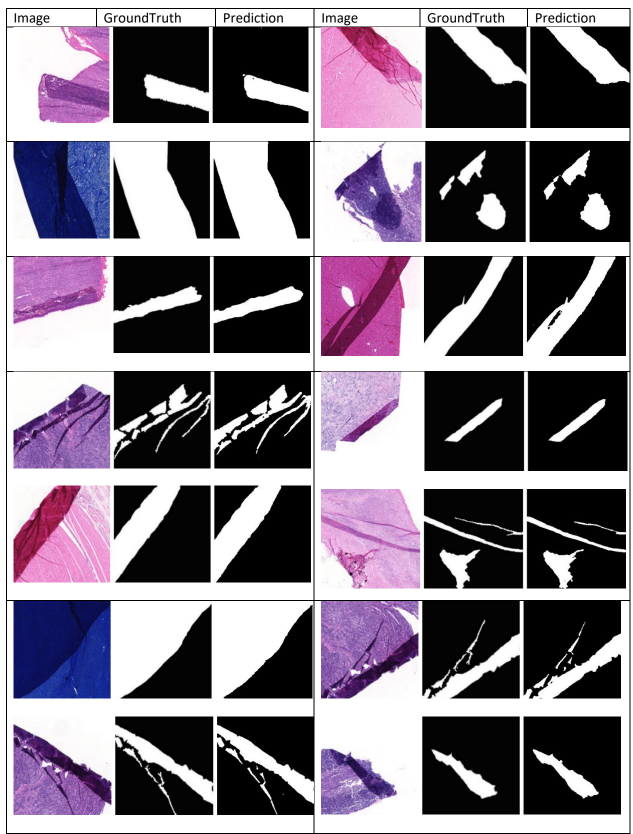


Figure 11: Testing comparison (on Tissue-fold using ResUNet++).

#### **Tissue Fold: DoubleUNet**

Table 5 represents the hyper-parameter combination for the best result for tissue fold artifact using DoubleUNet++ architecture. RMSprop optimizer performs better than any other optimizers. The learning rate was set to 0.0001 and was consistent throughout the whole training process. The result was achieved at epoch 50.

Table 5: Best segmentation result for Tissue-fold using DoubleUNet.

Train images	480
Valid images	60
Test images	60
Batch size	8
Epoch	50 EPOCH
Optimizer	RMSprop
Loss function	Dice Coef Loss
Learning Rate	1e <sup>-4</sup>
ReduceLROnPlateau	monitor='val_loss', factor=0.1, patience=10
EarlyStopping	monitor='val_loss', patience=10

loss	-0.9692	Val_loss	-0.9762
recall	29.64%	Val_recall	35.26%
precision	99.98%	Val_ precision	99.97%
mean-IOU	41.03%	Val_mean-IOU	35.12%
IOU	94. 03%	Val_ IOU	95.50%
Dice Coef	96.92%	Val_Dice Coef	97.69%

Dice coef loss is defined as negative value that's why the loss values are negative.

#### Graph

Figure 12 represents the graph of mean IOU, Loss curve, precision curve, recall curve, dice-coefficient and the IOU. The last two graphs which are the dice-coefficient and IOU curve defines the actual performance of the model.

In the next page, figure 13 represents some of the example comparison of the above mentioned model.

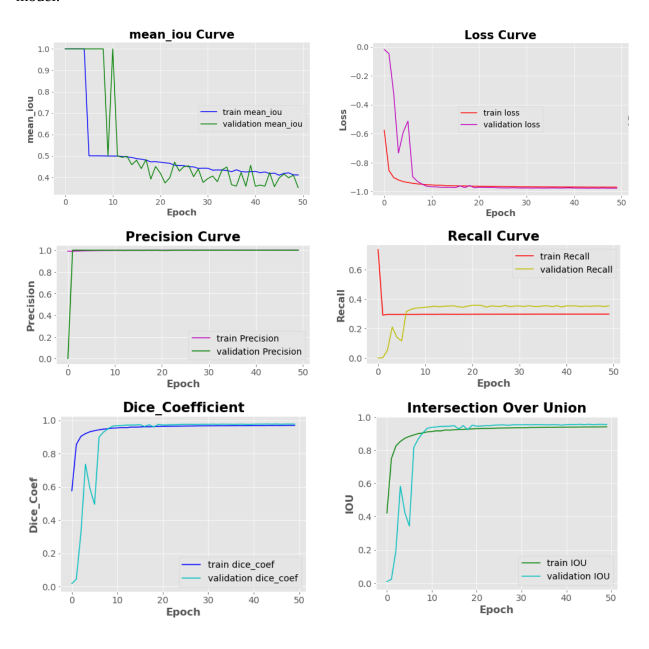


Figure 12: Best results of DoubleUNet on Tissue-fold dataset.

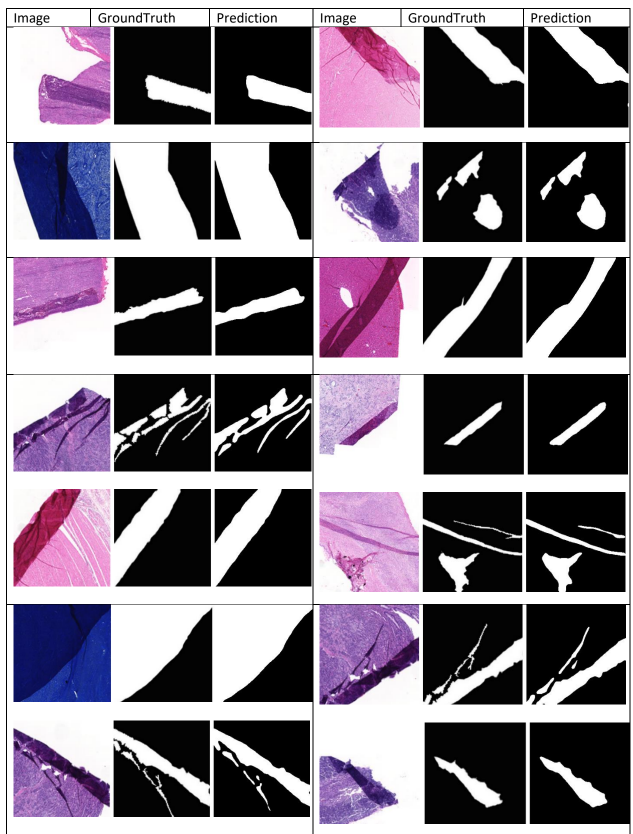
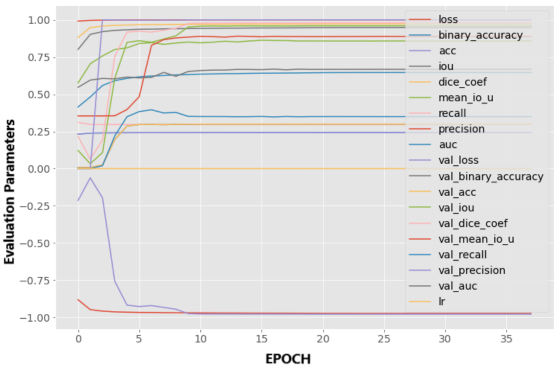


Figure 13: Testing comparison (on Tissue-fold using DoubleUNet).

#### Overall comparison of Tissue-Fold detection between ResUNet++ and DoubleUNet

#### Comparison of ResUNet++ on Tissue Fold Data set



### **Comparison of DoubleUNet on Tissue Fold Data set**

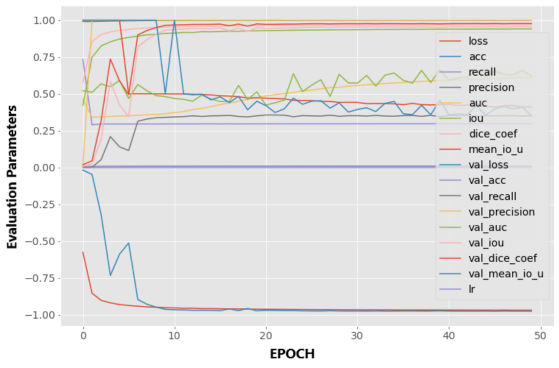


Figure 14: Comparison graph for ResUnet++ and DoubleUNet for Tissue-fold dataset.

#### **Air Bubble: ResUNet++**

Table 6 represents the hyper-parameter combination for the best result for air bubble artifact using ResUNet++ architecture. RMSprop optimizer performs better than any other optimizers. The learning rate was set to 0.0001. But, the 'ReduceLROnPlateau' function reduces the learning rate at every step where validation loss increases. The training was stopped when lr was  $1e^{-13}$ .

Table 6: Best segmentation result for Air-bubble using ResUNet++.

Train images	480	
Valid images	60	
Test images	60	
Batch size	8	
Epoch	56 EPOCH	
Optimizer	RMSprop	
Loss function	Dice Coef Loss, binary cross entropy	
Learning Rate	$1e^{-4} > 1e^{-5} > 1e^{-6} > 1e^{-7} > 1e^{-8} > 1e^{-9} > 1e^{-10} > 1e^{-11} > 1e^{-12}$	
ReduceLROnPlateau	monitor='val_loss', factor=0.1, patience=4	
EarlyStopping	monitor='val_loss', patience=10	

loss	-0.9858	Val_loss	-0.9873
recall	40.62%	Val_recall	45.50%
precision	99.99%	Val_ precision	99.99%
mean-IOU	89.96%	Val_mean-IOU	91.32%
IOU	97.21%	Val_ IOU	97.45%
Dice Coef	98.58%	Val_Dice Coef	98.71%

Dice coef loss is defined as negative value that's why the loss values are negative.

#### Graph

Figure 15 represents the graph of mean IOU, Loss curve, precision curve, recall curve, dice-coefficient and the IOU. The last two graphs which are the dice-coefficient and IOU curve defines the actual performance of the model.

In the next page, figure 16 represents some of the example comparison of the above mentioned model.

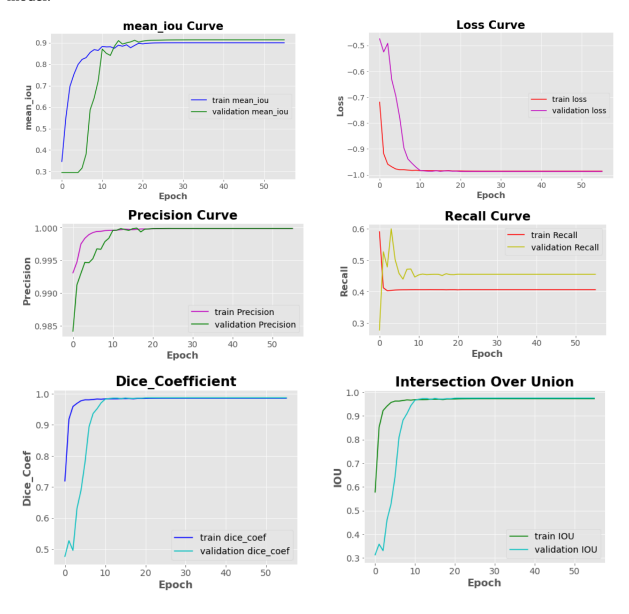


Figure 15: Best results of ResUNet++ on Air-bubble dataset.

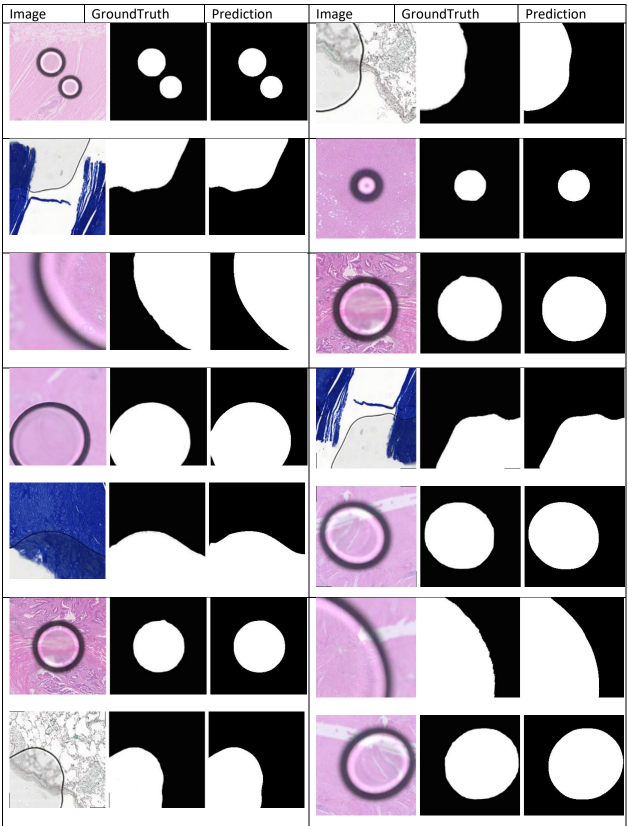


Figure 16: Testing comparison (on Air-bubble using ResUNet++).

#### Air Bubble: DoubleUNet

Table 7 represents the hyper-parameter combination for the best result for tissue fold artifact using DoubleUNet++ architecture. RMSprop optimizer performs better than any other optimizers. The learning rate was set to 0.0001 and was consistent throughout the whole training process. The result was achieved at epoch 50.

Table 7: Best segmentation result for Air-bubble using DoubleUNet.

Train images	480	480				
Valid images	60	60				
Test images	60	60				
Batch size	8	8				
Epoch	50 EPOCH	50 EPOCH				
Optimizer	RMSprop	RMSprop				
Loss function	Dice Coef Loss	Dice Coef Loss				
Learning Rate	1e <sup>-4</sup>	1e <sup>-4</sup>				
ReduceLROnPlateau	monitor='va	monitor='val_loss', factor=0.1, patience=10				
EarlyStopping	monitor='va	monitor='val_loss', patience=10				
loss	-0.9817	Val_loss	-0.9841			
recall	40.60%	Val_recall	45.45%			
precision	99.97%	Val_ precision	99.96%			
mean-IOU	32.97%	Val_mean-IOU	30.82%			
IOU	96.41%	Val_ IOU	96.83%			
Dice Coef	98.17%	Val_Dice Coef	98.39%			

Dice coef loss is defined as negative value that's why the loss values are negative.

#### Graph

Figure 17 represents the graph of mean IOU, Loss curve, precision curve, recall curve, dice-coefficient and the IOU. The last two graphs which are the dice-coefficient and IOU curve defines the actual performance of the model.

In the next page, figure 18 represents some of the example comparison of the above mentioned model.

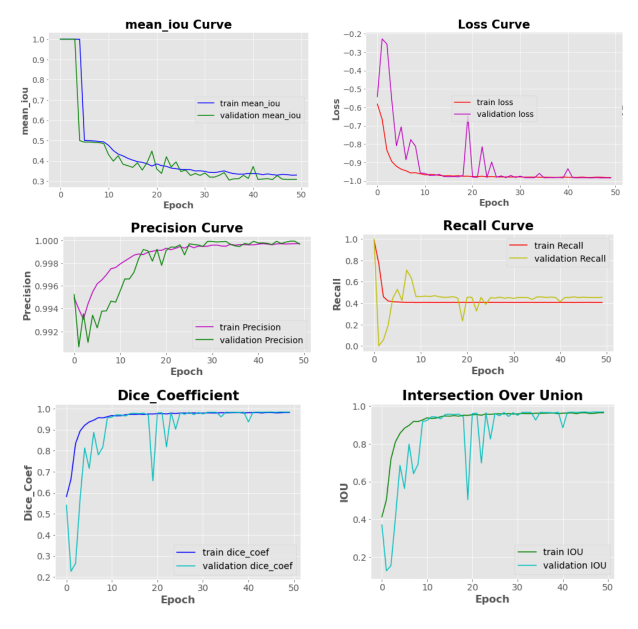


Figure 17: Best results of DoubleUNet on Air-bubble dataset.

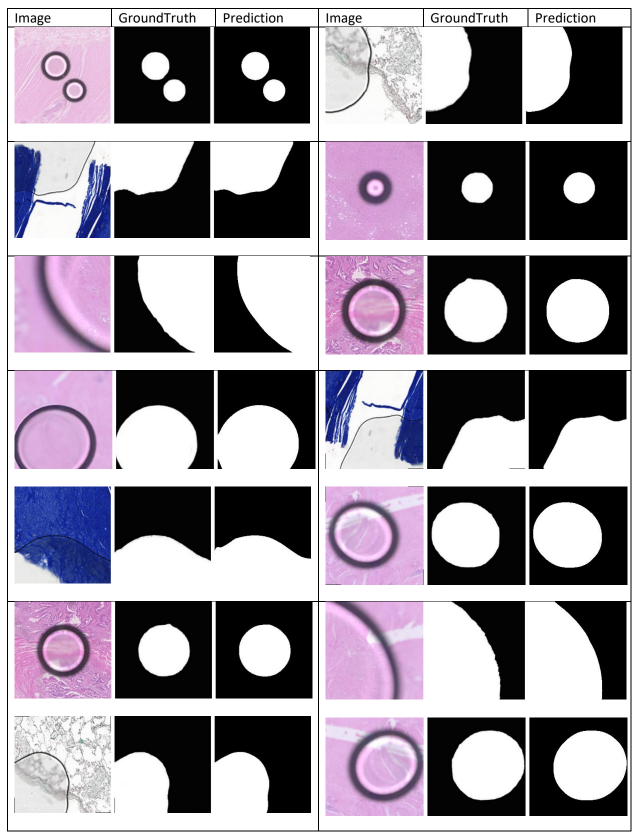
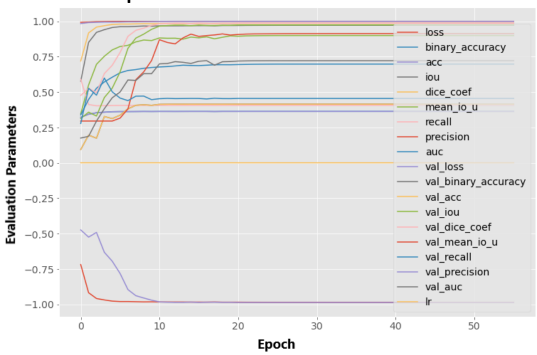


Figure 18: Testing comparison (on Air-bubble using DoubleUNet).

## Overall comparison of Air-bubble detection between ResUNet++ and DoubleUNet

# Comparison of ResUNet++ on Air Bubble Data set



# **Comparison of DoubleUNet on Air Bubble Data set**

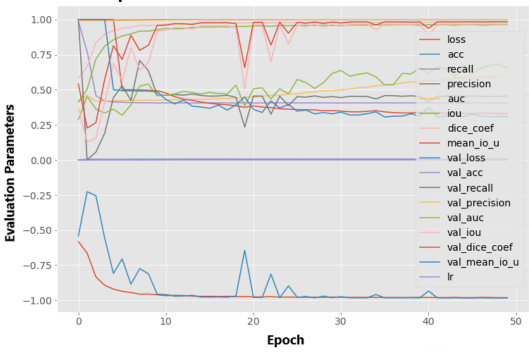


Figure 19: Comparison graph of ResUNet++ and DoubleUNet for Air-bubble dataset.

### **Different Optimizers Tested for comparison**

"AVERAGE TEST IOU" is calculated using the IOU value of the test set.

Here, Test data set contains 60 images.

\*AVG TEST IOU = 
$$\frac{\text{IOU of 1st image + IOU of 2nd image + IOU of 3rd image + . . . . . + IOU of 60th image}}{60}$$

Test Accuracy is determined by using different IOU threshold values.

If Value > threshold, then append (1: True) If Value < threshold then append (0: False)

[ \*For example, If, 58 out of 60 are true then test accuracy is (58/60) \*100 %= 96.66 %]

Table 8: Accuracy and IOU for Tissue Fold dataset.

UNet	Optimizer	AVERAGE Test IOU	Test Accuracy (IOU Threshold = 90%)	Test Accuracy (IOU Threshold = 85%)
41	Adamax	96.54 %	90.00 %	95.00 %
ble let	Adam	97.18 %	95.00 %	98.33 %
Double	RMSprop	98.02 %	98.33 %	100 %
	SGD	28.24 %	0 .00%	0.00 %
;	Adamax	95.74 %	88.33 %	96.66 %
ığ,	Adam	96.95 %	96.66 %	100 %
ResUNet ++	RMSprop	97.72 %	96.66 %	100 %
~	SGD	79.62 %	36.66 %	50.00 %

Table 9: Accuracy and IOU for Air Bubble dataset.

UNet	Optimizer	AVERAGE Test IOU	Test Accuracy (IOU Threshold = 90%)	Test Accuracy (IOU Threshold = 85%)
<b>a</b> )	Adamax	99.08 %	100 %	100 %
Souble	Adam	91.35 %	68.33 %	81.66 %
Double	RMSprop	99.11 %	100 %	100 %
	SGD	45. 96 %	0.00 %	0.00 %
<del>,</del>	Adamax	97.43 %	96.66 %	98.33 %
ž +	Adam	98.80 %	100 %	100 %
ResUNet ++	RMSprop	98.15 %	96.66 %	96.66 %
~	SGD	28.29 %	0.00 %	0.00 %

After doing all the combinations, it was found that "<u>RMSprop</u>" performs better than the other three optimizers. And SGD generates the poorest result.

So, in the next steps, for K-fold cross validation, "<u>RMSprop</u>" is used for both 'DoubleUNet' & 'ResUNet++'.

# **6-Fold Cross validation**

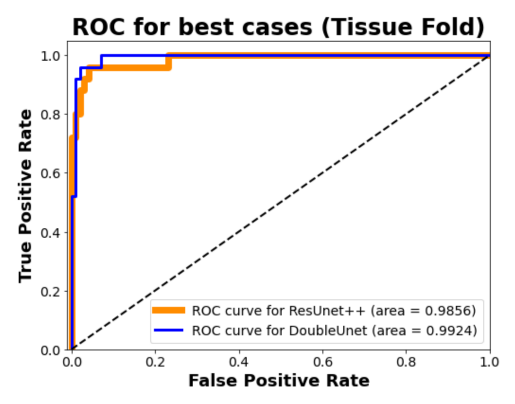
Table 10: 6-Fold Cross validation for Tissue fold dataset.

	Fold no.	Training Set	Testing Set	Test Acc (IOU Threshold = 90%)
	Fold 1	A, B, C, D, E	F	98.33 %
	Fold 2	B, C, D, E, F	Α	96.66 %
DoubleUNet	Fold 3	C, D, E, F, A	В	95.00 %
Doubleonet	Fold 4	D, E, F, A, B	С	98.33 %
	Fold 5	E, F, A, B, C	D	100 %
	Fold 6	F, A, B, C, D	E	100 %
	Average			97.96%
	Fold 1	A, B, C, D, E	F	95.00 %
	Fold 2	B, C, D, E, F	Α	96.66 %
DocUMetri	Fold 3	C, D, E, F, A	В	96.66 %
ResUNet++	Fold 4	D, E, F, A, B	С	95.00 %
	Fold 5	E, F, A, B, C	D	98.33 %
	Fold 6	F, A, B, C, D	E	96.66 %
	Average			96.38 %

Table 11: 6-Fold Cross validation for Air bubble dataset.

	Fold no.	Training Set	Testing Set	Test Acc (IOU Threshold = 90%)
	Fold 1	A, B, C, D, E	F	100 %
	Fold 2	B, C, D, E, F	Α	100 %
DoubleUNet	Fold 3	C, D, E, F, A	В	98.33 %
Doubleonet	Fold 4	D, E, F, A, B	С	100 %
	Fold 5	E, F, A, B, C	D	100 %
	Fold 6	F, A, B, C, D	E	100 %
	Average			99.72 %
	Fold 1	A, B, C, D, E	F	96.66 %
	Fold 2	B, C, D, E, F	Α	100 %
ResUNet++	Fold 3	C, D, E, F, A	В	100 %
Resolvet++	Fold 4	D, E, F, A, B	С	98.33 %
	Fold 5	E, F, A, B, C	D	100 %
	Fold 6	F, A, B, C, D	E	100 %
	Average			99.16 %

# **ROC Curve**



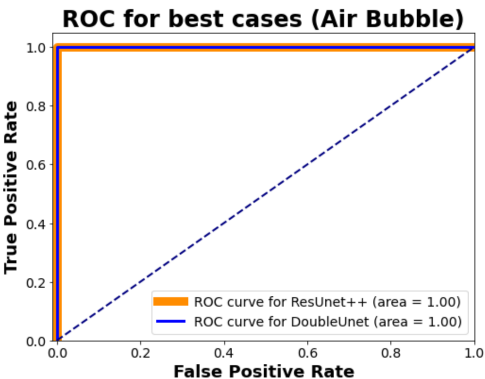


Figure 20: ROC comparison.

Table 12: Over all comparison of segmentation models.

Artifact	UNet	Train Loss	Val Loss	Train Recall	Val Recall	Train Precision	Val Precision	Train IOU	Val IOU	Train Dice- Coef	Val Dice- Coef
iue Id	Double UNet	-0.9692	-0.9762	0.2964	0.3526	0.9998	0.9997	0.9403	0.9550	0.9692	0.9769
Tissue Fold	ResUNet ++	-0.9736	-0.9791	0.2965	0.3497	0.9999	0.9997	0.9487	0.9605	0.9736	0.9798
r ble	Double UNet	-0.9817	-0.9841	0.4060	0.4545	0.9997	9666:0	0.9641	0.9683	0.9817	0.9839
Air Bubble	ResUNet ++	-0.9858	-0.9873	0.4062	0.4550	0.9999	0.9999	0.9721	0.9745	0.9858	0.9871

Table 13: Final ROC based comparison of segmentation models.

Tissue Artifact Type	UNet Architecture	ROC Score
Tissue Fold	DoubleUNet	99.24%
rissue roid	ResUNet++	98.56%
Air Bubble	DoubleUNet	1.00
Air Buddle	ResUNet++	1.00

Table 12 demonstrates the accuracy (IOU and dice-coefficient), loss, precision and recall comparison for both train and validation dataset.

From table 13 we can see that, for air-bubble dataset, both model gains 100% accuracy in terms of area under curve or ROC score while in the case of tissue fold dataset, DoubleUNet outperforms ResUNet++.

#### **Comparison Graph**

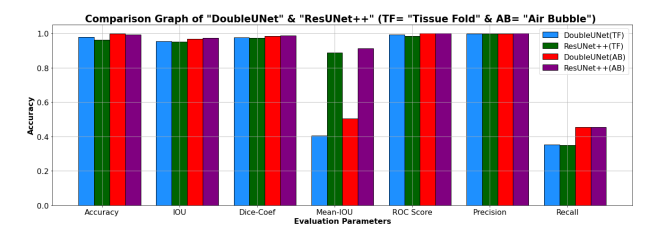


Figure 21: Comparison graph of ResUNet++ and DoubleUNet.

Over all The DoubleUNet out performs the ResUNet++. But the results are almost similar. Only the mean IOU value is greater in case of ResUNet++. But, Overall accuracy of DoubleUnet is better.

For the segmentation experiment, we can conclude that, our model works better for air-bubble dataset. Both models generate better result for air- bubble. While running 6-fold cross validation, in one or two cases, the result of ResUNet++ was better but overall the DoubleUNet works better which was explicitly clear throughout the experiment.

But one interesting findings was for both dataset resUNet++ generates better Mean-IOU than the DoubleUNet when the actual assessment criteria for our result was IOU and dice-coefficient.

Another finding from our segmentation experiment is that, the precision is over 99.9% while the recall is between 33%-45%. It means that, false positive rate is greater than false negative.

# 5.2. Severity Analysis/ Classification Result

Table 14 represents the accuracy and loss results of transfer learning based on 10 models.

Table 14: Transfer Learning for base model selection.

Pretraind Model	Opimizer	Loss Function	Learning Rate	Validation Loss	Test Accuracy	ROC Score
	Adam	CCE	0.0001	0.0170	99.52%	99.99%
Xception	Auaiii	KLD	0.0001	0.0222	99.05%	99.98%
	Adamay	CCE	0.0001	0.0304	99.29%	99.99%
cep	Adamax	KLD	0.0001	0.0287	99.52%	99.99%
×	RMSprop	CCE	0.0001	0.0103	99.76%	99.99%
	KiviSprop	KLD	0.0001	0.0093	99.52%	100%
	Adam	CCE	0.0001	0.0444	98.81%	99.99%
10	Adam	KLD	0.0001	0.0480	98.57%	99.98%
VGG16	A da	CCE	0.0001	0.1173	97.86%	99.93%
99/	Adamax	KLD	0.0001	0.1192	97.86%	99.92%
_	DMCman	CCE	0.0001	0.0298	99.05%	99.99%
RM	RMSprop	KLD	0.0001	0.0224	99.52%	100%
	A al a	CCE	0.0001	0.0441	99.05%	100%
	Adam	KLD	0.0001	0.0435	99.05%	99.99%
319		CCE	0.0001	0.1259	97.38%	99.95%
VGG19	Adamax	KLD	0.0001	0.1145	98.33%	99.93%
	RMSprop	CCE	0.0001	0.0217	99.76%	100%
		KLD	0.0001	0.0252	99.05%	99.99%
	Adam	CCE	0.0001	0.4082	86.19%	96.57%
00	Adam	KLD	0.0001	0.4418	78.81%	96.53%
ResNet50	Adamax	CCE	0.0001	0.5723	80.71%	94.05%
Nsa		KLD	0.0001	0.5385	78.10%	94.07%
Re	RMSprop	CCE	0.0001	0.4903	78.57%	95.77%
RMSprop	KiviSprop	KLD	0.0001	0.5587	80.71%	95.12%
	Adam	CCE	0.0001	0.0422	99.05%	99.96%
<b>N</b> 3	Audili	KLD	0.0001	0.0457	99.05%	99.96%
ior	Adamax	CCE	0.0001	0.0492	99.05%	99.89%
ept	Auailiax	KLD	0.0001	0.0523	99.05%	99.84%
InceptionV3	DMCnron	CCE	0.0001	0.0315	99.29%	99.98%
	RMSprop	KLD	0.0001	0.0317	99.29%	99.98%
	Adam	CCE	0.0001	0.0299	98.81%	99.99%
on 72	Auam	KLD	0.0001	0.0269	99.05%	99.98%
Inception ResNetV2	A domes:	CCE	0.0001	0.0631	98.81%	99.92%
cek	Adamax	KLD	0.0001	0.0441	98.81%	99.95%
In Re	DMC	CCE	0.0001	0.0368	98.57%	99.99%
	RMSprop	KLD	0.0001	0.0152	99.05%	100%

Adam							
RMSprop   Adamax   CCE   0.0001   0.0296   99.29%   99.99%   99.99%	Adam	Adam	CCE	0.0001	0.0631	98.81%	99.92%
RMSprop   KLD   0.0001   0.0093   99.76%   100%	let	Name Adam	KLD	0.0001	0.0229	99.29%	99.99%
RMSprop   KLD   0.0001   0.0093   99.76%   100%	i e N	A domest	CCE	0.0001	0.0296	99.29%	99.98%
RMSprop   KLD   0.0001   0.0093   99.76%   100%	idc	Adamax	KLD	0.0001	0.0263	99.52%	99.97%
Adamax   CCE   0.0001   0.0093   99.76%   100%	Š	Σ nus	CCE	0.0001	0.0174	99.76%	99.99%
Adam   KLD   0.0001   0.0119   99.76%   99.99%		KIVISProp	KLD	0.0001	0.0093	99.76%	100%
Adam   CCE   0.0001   0.0207   99.76%   100%	7	A da	CCE	0.0001	0.0148	99.52%	99.99%
Adam   CCE   0.0001   0.0207   99.76%   100%	, Ç	Adam	KLD	0.0001	0.0119	99.76%	99.99%
Adam   CCE   0.0001   0.0207   99.76%   100%	Ž	A domest	CCE	0.0001	0.0169	99.52%	100%
Adam   CCE   0.0001   0.0207   99.76%   100%	e	Adamax	KLD	0.0001	0.0245	99.29%	99.99%
Adam   CCE   0.0001   0.0207   99.76%   100%	10	10k	CCE	0.0001	0.0024	99.76%	100%
Adamax   KLD   0.0001   0.0212   99.29%   99.99%		KiviSprop	KLD	0.0001	0.0207	99.76%	100%
Adamax   CCE   0.0001   0.0171   99.52%   99.99%	1	مر مام ۸	CCE	0.0001	0.0296	98.81%	99.99%
Adamax   CCE   0.0001   0.0171   99.52%   99.99%	112	Audili	KLD	0.0001	0.0212	99.29%	99.99%
Adamax   CCE   0.0001   0.0171   99.52%   99.99%	let let	Adamay	CCE	0.0001	0.0280	99.52%	99.99%
Adamax   CCE   0.0001   0.0171   99.52%   99.99%	seľ	Adamax	KLD	0.0001	0.0406	98.81%	99.96%
Adamax   CCE   0.0001   0.0171   99.52%   99.99%	en	DMCnron	CCE	0.0001	0.0035	99.76%	100%
Adamax KLD 0.0001 0.0971 98.81% 99.97%  CCE 0.0001 0.0442 98.81% 99.93%  KLD 0.0001 0.0328 99.05% 99.99%  CCE 0.0001 0.0799 99.05% 99.93%		△ KIMSprop	KLD	0.0001	0.0171	99.52%	99.99%
RMSprop   RMSprop   RLD   0.0001   0.0971   98.81%   99.97%	a)	Adam	CCE	0.0001	0.1059	98.81%	99.92%
Adamax         CCE         0.0001         0.0442         98.81%         99.93%           KLD         0.0001         0.0328         99.05%         99.99%           RMSprop         CCE         0.0001         0.0799         99.05%         99.93%           KLD         0.0001         0.0387         99.29%         99.99%	rge	80 Adam	KLD	0.0001	0.0971	98.81%	99.97%
KLD   0.0001   0.0328   99.05%   99.99%	tLa	Adamay	CCE	0.0001	0.0442	98.81%	99.93%
RMSprop CCE 0.0001 0.0799 99.05% 99.93% KLD 0.0001 0.0387 99.29% 99.99%	Ne Se	Adamax	KLD	0.0001	0.0328	99.05%	99.99%
KLD 0.0001 0.0387 99.29% 99.99%	las	DMSpren	CCE	0.0001	0.0799	99.05%	99.93%
		KIVISprop	KLD	0.0001	0.0387	99.29%	99.99%

This table represents the results of different combinations of transfer learning. This table is used for base model selection for ensemble learning. The base models are selected comparing the test accuracy. There are 7 models which have same test accuracy. Among those 7 models, 1 model has relatively higher validation loss. So, we excluded that model from our base model list. So, we finally selected the top 6 models as our base model for applying ensemble learning. The top models are described in details in the next page.

Every base model selected for ensemble learning is described in details in terms of their loss, accuracy, miss-classification rate and ROC score in their respective pages along with the confusion matrix.

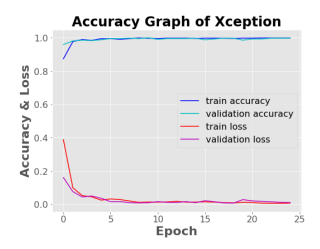
#### **Best Models**

#### Model 1

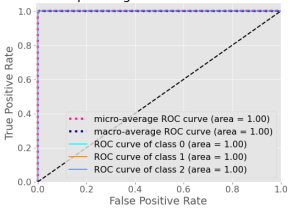
Model	XCEPTION
Epoch	25
Batch size	32
Learning rate	0.0001
Optimizer	RMSprop
Loss function	Categorical cross entropy (CCE)

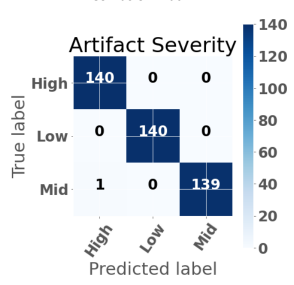
Loss	Test accuracy	ROC Score	Mismatch
0.0103149469	0.9976190328	0.9999914965	1 of 420 (1 mid predicted as high)

## Graph



## Receiver operating characteristic to multiclass

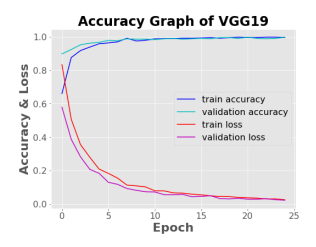




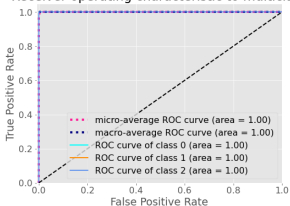
Model	VGG19
Epoch	25
Batch size	32
Learning rate	0.0001
Optimizer	RMSprop
Loss function	Categorical cross entropy (CCE)

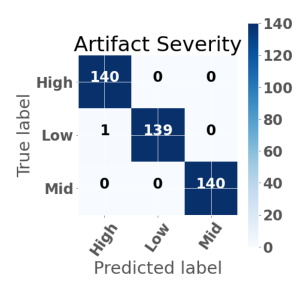
Loss	Test accuracy	ROC Score	Mismatch
0.0216745790	0.9976190328	1.0000000000	1 of 420 (1 low predicted as high)

#### Graph



### Receiver operating characteristic to multiclass

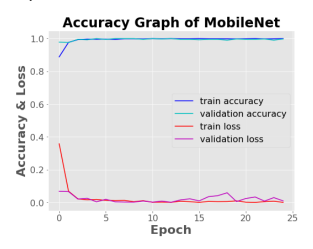




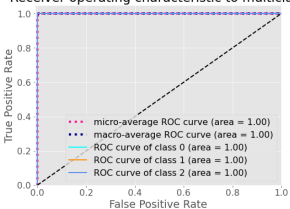
Model	MobileNet
Epoch	25
Batch size	32
Learning rate	0.0001
Optimizer	RMSprop
Loss function	Kullback Leibler Divergence (KLD)

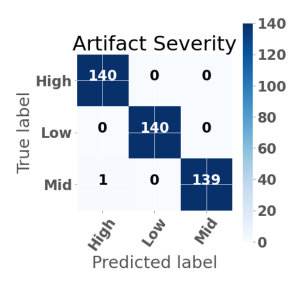
Loss	Test accuracy	ROC Score	Mismatch
0.0092747136	0.9976190328	1.0000000000	1 of 420 (1 mid predicted as high)

#### Graph



### Receiver operating characteristic to multiclass

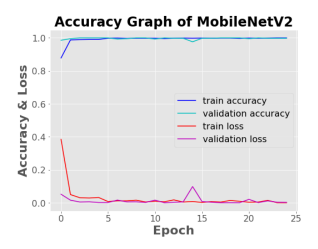




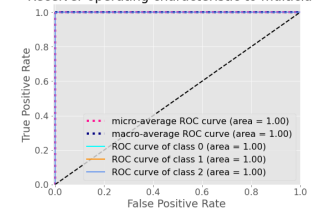
Model	MobileNetV2
Epoch	25
Batch size	32
Learning rate	0.0001
Optimizer	RMSprop
Loss function	Categorical cross entropy (CCE)

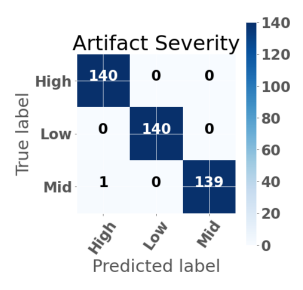
Loss	Test accuracy	ROC Score	Mismatch
0.0024069594	0.9976190328	1.0000000000	1 of 420 (1 mid predicted as high)

#### Graph



## Receiver operating characteristic to multiclass

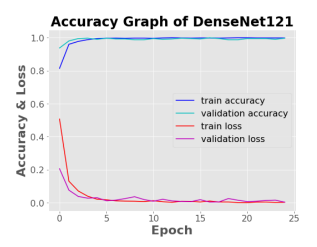




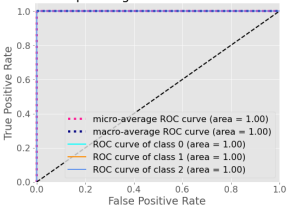
Model	DenseNet121
Epoch	25
Batch size	32
Learning rate	0.0001
Optimizer	RMSprop
Loss function	Categorical cross entropy (CCE)

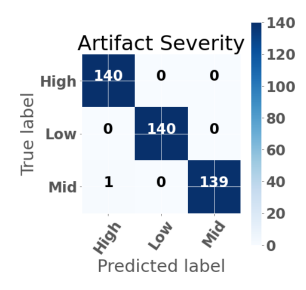
Loss	Test accuracy	ROC Score	Mismatch
0.0034660110	0.9976190328	1.0000000000	1 of 420 (1 mid predicted as high)

#### Graph



## Receiver operating characteristic to multiclass

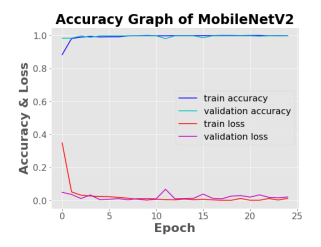




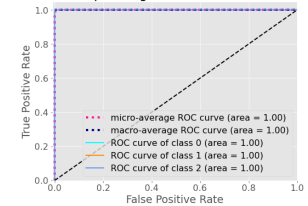
Model	MobileNetV2
Epoch	25
Batch size	32
Learning rate	0.0001
Optimizer	RMSprop
Loss function	Kullback Leibler Divergence (KLD)

Loss	Test accuracy	ROC Score	Mismatch
0.0206717420	0.9976190328	1.0000000000	1 of 420 (1 mid predicted as high)

#### Graph



#### Receiver operating characteristic to multiclass



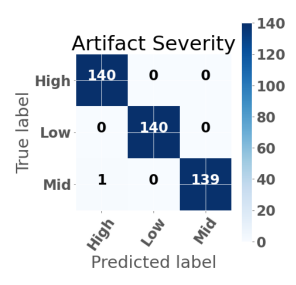


Table 15: Top 6 models description

		Combination		Accuracy	
Model name	Pretrained Model	Optimizer	Optimizer Loss function		
Model 1	XCEPTION	RMSprop	CCE	0.0103149469	0.9976190328
Model 2	VGG19	RMSprop	CCE	0.0216745790	0.9976190328
Model 3	MobileNet	RMSprop	KLD	0.0092747136	0.9976190328
Model 4	MobileNetV2	RMSprop	CCE	0.0024069594	0.9976190328
Model 5	DenseNet121	RMSprop	CCE	0.0034660110	0.9976190328
Model 6	MobileNetV2	RMSprop	KLD	0.0206717420	0.9976190328

For Ensemble Learning various methods can be applied such as stacking, voting etc. For our experiment, we have used stacking.

We used various combinations of models as follows:

- ➤ Using top 6 models
- Using top 5 models
- ➤ Using top 4 models
- ➤ Using top 3 models
- Using top 2 models

And for every combination, we used Logistic Regression, KNN, SVM, Decision Tree, Random Forest, Ada Boost Classifier, XGB Classifier, GB Classifier, GB Regressor and Gaussian Naïve bias as meta learners and compared the results. The comparison table is mention in the next page.

Table 16: Ensemble Learning Combinations using 10 meta learners.

						Accura	icy (%)				
Models	Combination	Logistic Regression	KNN	MVS	Decision Tree	Random Forest	AdaBoost Classifier	XGB Classifier	GB Regressor	GB Classifier	Gaussian NB
Top 2	Model (4, 5)	6666'66	6666'66	6666'66	99.8211	6666'66	6666'66	6666'66	6666'66	6666'66	6666'66
Top 3	Model (3, 4, 5)	6666666	99.9999	99.9999	99.9999	99.9999	99.9999	99.9999	99.9999	6666.66	99.9999
Top 4	Model (1, 3, 4, 5)	6666666	6666.66	99.9999	99.6441	99.8211	6666.66	99.8217	99.8217	6666.66	99.9999
Top 5	Model (1, 2, 3, 4, 5)	6666666	6666.66	99.9999	99.6415	6666.66	6666.66	99.8211	6666.66	99.8211	99.9999
Тор б	Model (1, 2, 3, 4, 5, 6)	6666666	6666.66	6666.66	99.8211	6666.66	6666.66	6666.66	99.8211	99.8217	99.8217

From the table it is clear that,

Logistic Regression, K Nearest Neighbor and Ada Boost Classifier performs better than any other models for our experiment. And from the model perspective, using the top 3 models generates the best result.

## 5.3. Critical Analysis

We have so far covered the results and related discussion on both the segmentation and classification tasks. In terms of the segmentation part, the segmentation accuracy is assessed based on some particular IOU threshold value instead of pixel wise accuracy. If the value is above 90% we considered it to be as a successful identification case. Based on this assessment criterion the segmentation result we got is quite good which is over 97% (while using DoubleUNet) for air bubble and over 96% (while using DoubleUNet) for tissue fold. Due to manual binary masking, some masking imperfection occurred as it couldn't have been avoided altogether. But, if perfect binary masking was possible, then we could get even better results in terms of segmentation accuracy.

As for classification task, the levels of severity were determined by human eye. For this, the result can very if the annotation is done by another human. But, looking at the result, which is 99.99%, we can easily say that, our model will definitely achieve 99% accuracy in terms of severity classification even if done by different experts providing that the binary masking should be close to perfect and the data annotation process should follow standard annotation techniques.

Another thing is that, for severity analysis, regression is better than classification as no hard and fast rules applies to the severity labels, but in that case, the manual observation will be required even after visualization of the artifacts and human decision will be necessary. In our case, to avoid that scenario we implemented classification instead of regression.

# 6.Conclusion

In this research, we worked on improving the tissue artifact segmentation accuracy to further enhance the detection mechanism of the whole slide imaging scanner. We have managed to achieve our goal using DoubleUNet and ResUNEt++ architecture; the DoubleUnet architecture performs better in most of the cases. We have also proposed an ensemble learning based classification model which analyze the severity of the detected artifact. Our model was tested on a dataset annotated by experts in this field. This will help the WSI to take decision on removing artifacts from diagnosis process according to the severity detected by our model. This proposed model will enhance the WSI experience and lead the digital pathology to a new level, making the diagnosis process easier than ever before and also more accurate.

Our future plan is to apply regression for severity analysis instead of classification and also to reconstruct the artifact containing region and to check whether the reconstructed area can be used in meaningful diagnosis or not. If these can be done successfully, the digital pathology will be elevated to a new step and the future of pathological image analysis will be brighter than ever before.

# **Bibliography**

- 1. Gilbertson, John R., et al. "Primary histologic diagnosis using automated whole slide imaging: a validation study." *BMC clinical pathology* 6.1 (2006): 1-19.
- 2. Ho, Jonhan, et al. "Use of whole slide imaging in surgical pathology quality assurance: design and pilot validation studies." *Human pathology* 37.3 (2006): 322-331.
- 3. Ghaznavi, Farzad, et al. "Digital imaging in pathology: whole-slide imaging and beyond." *Annu Rev Pathol* 8.1 (2013): 331-359.
- 4. Pantanowitz, Liron. "Digital images and the future of digital pathology." *Journal of pathology informatics* 1 (2010).
- 5. Farahani, Navid, Anil V. Parwani, and Liron Pantanowitz. "Whole slide imaging in pathology: advantages, limitations, and emerging perspectives." *Pathol Lab Med Int* 7.23-33 (2015): 4321.
- 6. Thorstenson, Sten, Jesper Molin, and Claes Lundström. "Implementation of large-scale routine diagnostics using whole slide imaging in Sweden: Digital pathology experiences 2006-2013." *Journal of pathology informatics* 5.1 (2014): 14.
- 7. Shakhawat, Hossain Md, et al. "Automatic Quality Evaluation of Whole Slide Images for the Practical Use of Whole Slide Imaging Scanner." *ITE Transactions on Media Technology and Applications* 8.4 (2020): 252-268.
- 8. Hashimoto, Noriaki, et al. "Referenceless image quality evaluation for whole slide imaging." *Journal of pathology informatics* 3.1 (2012): 9.
- 9. Jimson, S. U. D. H. A., et al. "Artifact in histological section." *Biomedical and Pharmacology Journal* 9.2 (2016): 843-845.
- 10. Ekundina, V. O., and G. Eze. "Common artifacts and remedies in histopathology (a review)." *African Journal of Cellular Pathology* (2015): 1-7.
- 11. Yagi, Yukako, and John R. Gilbertson. "A relationship between slide quality and image quality in whole slide imaging (WSI)." *Diagnostic pathology*. Vol. 3. No. 1. BioMed Central, 2008.
- 12. Choi, Min Goo, Jung Hoon Jung, and Jae Wook Jeon. "No-reference image quality assessment using blur and noise." *International Journal of Computer Science and Engineering* 3.2 (2009): 76-80.
- 13. Konstantinides, Konstantinos, Balas Natarajan, and Gregory S. Yovanof. "Noise estimation and filtering using block-based singular value decomposition." *IEEE transactions on image processing* 6.3 (1997): 479-483.
- 14. Suthaharan, Shan. "No-reference visually significant blocking artifact metric for natural scene images." *Signal Processing* 89.8 (2009): 1647-1652.

- 15. Sazzad, ZM Parvez, Yoshikazu Kawayoke, and Yuukou Horita. "No reference image quality assessment for JPEG2000 based on spatial features." *Signal Processing: Image Communication* 23.4 (2008): 257-268.
- 16. Babu, R. Venkatesh, Sundaram Suresh, and Andrew Perkis. "No-reference JPEG-image quality assessment using GAP-RBF." *Signal Processing* 87.6 (2007): 1493-1503.
- 17. Ferzli, Rony, and Lina J. Karam. "A no-reference objective image sharpness metric based on the notion of just noticeable blur (JNB)." *IEEE transactions on image processing* 18.4 (2009): 717-728.
- 18. Narvekar, Niranjan D., and Lina J. Karam. "A no-reference perceptual image sharpness metric based on a cumulative probability of blur detection." 2009 International Workshop on Quality of Multimedia Experience. IEEE, 2009.
- 19. Varadarajan, Srenivas, and Lina J. Karam. "An improved perception-based no-reference objective image sharpness metric using iterative edge refinement." 2008 15th IEEE international conference on image processing. IEEE, 2008.
- 20. Liu, Lixiong, et al. "No-reference image quality assessment based on spatial and spectral entropies." *Signal processing: Image communication* 29.8 (2014): 856-863.
- 21. Shrestha, Prarthana, et al. "A quantitative approach to evaluate image quality of whole slide imaging scanners." *Journal of pathology informatics* 7.1 (2016): 56.
- 22. Senaras, Caglar, et al. "DeepFocus: detection of out-of-focus regions in whole slide digital images using deep learning." *PloS one* 13.10 (2018): e0205387.
- 23. Hosseini, Mahdi S., et al. "Focus quality assessment of high-throughput whole slide imaging in digital pathology." *IEEE transactions on medical imaging* 39.1 (2019): 62-74.
- 24. Marcano-Cedeño, Alexis, et al. "Feature selection using sequential forward selection and classification applying artificial metaplasticity neural network." *IECON 2010-36th annual conference on IEEE industrial electronics society*. IEEE, 2010.
- 25. Khaire, Utkarsh Mahadeo, and R. Dhanalakshmi. "Stability of feature selection algorithm: A review." *Journal of King Saud University-Computer and Information Sciences* (2019).
- 26. Yamashita, R., et al. "Convolutional neural networks: An overview and application in radiology. Insights into Imaging." (2018): 611-629.
- 27. Pan, Sinno Jialin, and Qiang Yang. "A survey on transfer learning." *IEEE Transactions on knowledge and data engineering* 22.10 (2009): 1345-1359.
- 28. Zhao, Wei. "Research on the deep learning of the small sample data based on transfer learning." *AIP Conference Proceedings*. Vol. 1864. No. 1. AIP Publishing LLC, 2017.

- 29. Srivastava, Nitish, et al. "Dropout: a simple way to prevent neural networks from overfitting." *The journal of machine learning research* 15.1 (2014): 1929-1958.
- 30. Perez, Luis, and Jason Wang. "The effectiveness of data augmentation in image classification using deep learning." *arXiv preprint arXiv:1712.04621* (2017).
- 31. Taqi, Syed Ahmed, et al. "A review of artifacts in histopathology." *Journal of oral and maxillofacial pathology: JOMFP* 22.2 (2018): 279.
- 32. Oskal, Kay RJ, et al. "A U-net based approach to epidermal tissue segmentation in whole slide histopathological images." *SN Applied Sciences* 1.7 (2019): 1-12.
- 33. Moles Lopez, Xavier, et al. "An automated blur detection method for histological whole slide imaging." *PloS one* 8.12 (2013): e82710.
- 34. Khan, S., et al. "Artifacts in histopathology: A potential cause of misinterpretation." *Res Rev J Dent Sci* 2 (2014): 23-31.
- 35. Oktay, Ozan, et al. "Attention u-net: Learning where to look for the pancreas." *arXiv preprint arXiv:1804.03999* (2018).
- 36. Babaie, Morteza, and Hamid R. Tizhoosh. "Deep features for tissue-fold detection in histopathology images." *European Congress on Digital Pathology*. Springer, Cham, 2019.
- 37. Bautista, Pinky A., and Yukako Yagi. "Detection of tissue folds in whole slide images." 2009 Annual International Conference of the IEEE Engineering in Medicine and Biology Society. IEEE, 2009.
- 38. Wu, Hang, et al. "Detection of blur artifacts in histopathological whole-slide images of endomyocardial biopsies." 2015 37th annual international Conference of the IEEE Engineering in Medicine and biology society (EMBC). IEEE, 2015.
- 39. Kothari, Sonal, John H. Phan, and May D. Wang. "Eliminating tissue-fold artifacts in histopathological whole-slide images for improved image-based prediction of cancer grade." *Journal of pathology informatics* 4.1 (2013): 22.
- 40. Lu, Wei, et al. "Evidence for trapped surface bubbles as the cause for the twinkling artifact in ultrasound imaging." *Ultrasound in medicine & biology* 39.6 (2013): 1026-1038.
- 41. Bautista, Pinky A., and Yukako Yagi. "Improving the visualization and detection of tissue folds in whole slide images through color enhancement." *Journal of pathology informatics* 1.1 (2010): 25.
- 42. <a href="http://www.globalengage.co.uk/pathology/docs/PosterShakhawatetal.pdf">http://www.globalengage.co.uk/pathology/docs/PosterShakhawatetal.pdf</a>

# **Author Information**



**Galib Muhammad Shahriar Himel** 

ID: 21-92094-2

MSCS, Dept. of Computer Science, AIUB

ORCID: https://orcid.org/0000-0002-2257-6751

LinkedIn: https://www.linkedin.com/in/galib-muhammad-shahriar-himel-3763b0138/

Email: galib.muhammad.shahriar@gmail.com