## AI BASED RECTAL CANCER AND STAGE PREDICTION OVER WEB IN REAL - TIME

**A PROJECT REPORT**

***Submitted by***

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***in partial fulfillment for the award of the degree***

*of*

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IN

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**(An Autonomous Institution, Affiliated to Anna University, Chennai)**

**MAY 2022**

# PANIMALAR ENGINEERING COLLEGE

###### (An Autonomous Institution, Affiliated to Anna University, Chennai)

**BONAFIDE CERTIFICATE**

Certified that this project report **“AI BASED RECTAL CANCER AND STAGE PREDICTION OVER WEB IN REAL-TIME”** is the bonafide work of **“BOGGADA MOUNIKA (211418104041), Y.EESHA SAI SRI (211418104057) and PANCHETI DIVIJA REDDY (211418104184)”** who carried out the project work under my supervision.

|  |  |
| --- | --- |
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Certified that the above mentioned students were examined in End Semester project viva-voice held on \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_.

### INTERNAL EXAMINER EXTERNAL EXAMINER

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**DECLARATION BY THE STUDENT**

### We BOGGADA MOUNIKA(211418104041), Y.EESHA SAI SRI(211418104057) and PANCHETI DIVIJA REDDY (211418104184) hereby declare that this project report titled“ AI BASED RECTAL CANCER AND STAGE PREDICTION OVER WEB IN REAL-TIME”, under the guidance of DR. SANGEETHA , M.E.,Ph.D is the original work done by us and we have not plagiarized or submitted to any other degree in any university by us.

**BOGGADA MOUNIKA**

**Y. EESHA SAI SRI**

**PANCHETI DIVIJA REDDY**

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### BOGGADA MOUNIKA

**Y. EESHA SAI SRI** **PANCHETI DIVIJA REDDY**

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**ABSTRACT**

Now-a-days, with the development of targeted therapies, many treatments are based on molecular studies, which require sampling tumor tissue from paraffin blocks for sequencing. An automated solution could potentially reduce the workload of the pathologists by acting as a screening device and may reduce the subjectivity in diagnosis. In tissue-based diagnostics, most of the work still needs to be done manually by a pathologist using a microscope to examine stained slides. The foundation of such tasks is to accurately distinguish cancer/malignant cells from normal/benign cells. However, the determination of tumor content is poorly reproducible with significant variation. As the size of tumor regions can be very small, pathologists are often required to use high magnification for detecting tumor cells. This requirement significantly increases the workload for pathologists. As digital pathology datasets have become publicly available and have opened up the possibility of evaluating the feasibility of applying deep learning techniques to improving the efficiency and quality of histologic diagnosis. In this project we introduce an application to detect Colorectal cancer based on the Convolutional Neural Network and Ranking algorithm. Here we will collect the tissue from lab or hospital and we will train the image and do data processing with segmentation and morphological filtering. Now we will store that in Azure ML server. In prediction website we will select the image and we will predict that one. The result will be displayed with ranking.

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### LIST OF ABBREVATIONS

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **S. NO.** | | | **ABBREVATION** | **EXPANSION** |
| 1 | | | CNN | Convolutional Neural Network |
| 2 | | | LSTM | Long Short-Term Memory |
| 3 | | | MRI | Magnetic Resonance Imaging |
| 4 | | | MATLAB | Matrix Laboratory |
| 5 | | | ResNet | Residual Network |
| 6 | | | CRC | Colorectal Cancer |
| 7 | | | viii  LGN | Lateral Geniculate Nucleus |
| 8 | | | SVF | Stromal Vascular Fraction |
| 9 | | | WAT | White Adipose Tissue |
| 10 | | | BAT | Brown Adipose Tissue |
| 11 | | | API | Application Program Interface |
| 12 | | | PIM | Protocol Independent Multicast |
| 13  14 | | | ROLM  SIANN | Randomized On-Line Matching  Space Invariant Artificial Neural |
| 15 | RNN Recurrent Neural Network | | | |
| 16 | EC2 Elastic Compute Cloud | | | |
| 17 | AZ Azure Web Services | | | |
| 18 | AMI Azure Machine Image | | | |
| 19 | EBS Elastic Block Store | | | |
| 20 | IP Internet Protocol | | | |
| 21 | IPv4 Internet Protocol Version 4 | | | |
| 22 | VPC Virtual Private Cloud | | | |

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## CHAPTER 1

## INTRODUCTION

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### CHAPTER 1 INTRODUCTION

In the modern era, cancer is the most spreading complex disease. Identifying cancer without biopsy at an early stage is further imperative. Also, taking a biopsy is not good for health also. In general, cancer has been caused by hereditary instability and accumulation of multiple molecular alterations. It is also caused by cellular genes abnormal activation that controls cell growth or cell mitosis. Colorectal cancer is cancer from uncontrolled cell growth in the colon or rectum. This was the third most commonly diagnosed cancer in the world. Colorectal cancer is also known as colon cancer, bowel cancer or colorectal adenocarcinoma. The main negative aspect of cancer is its diagnosis and treatment too late. Due to this problem, cancer has overtaken heart disease as the leading cause of death for any age on. Therefore, early detection of cancer is important. The images are collected and manually annotated for image processing. These images represent controlled imaging conditions and a wide variety in patient demographics. Each image has a dimension ranging from [155x240] to [960x1280] pixels with storage size of 10kB to 252kB per image.

**1.1 OVERVIEW**

With the development of targeted therapies, many treatments are based on molecular studies, which require sampling tumor tissue from paraffin blocks for sequencing. An automated solution could potentially reduce the workload of pathologists by acting as a screening device and may reduce the subjectivity in diagnosis. In tissue-based diagnostics, most of the work still needs to be done manually by a pathologist using a microscope to examine stained slides. The foundation of such tasks is to accurately distinguish cancer/malignant cells from normal/benign cells. However, the determination of tumor content is poorly reproducible with significant variation.

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As the size of tumor regions can be very small, pathologists are often required to use high magnification for detecting tumor cells. This requirement significantly increases the workload for pathologists. As digital pathology datasets have become publicly available and have opened up the possibility of evaluating the feasibility of applying deep learning techniques to improving the efficiency and quality of histologic diagnosis. In this project we introduce an user facing AI based application to detect and predict the stage of Rectal cancer based on CNN with Attention mechanism and Ranking algorithm.

**1.2 PROBLEM DEFINITION**

Colorectal cancer is also known as colon cancer, bowel cancer or colorectal adenocarcinoma. The main negative aspect of cancer is its diagnosis and treatment too late. Due to this problem, cancer has overtaken heart disease as the leading cause of death for any age on. Therefore, early detection of cancer is important. With the development of targeted therapies, many treatments are based on molecular studies, which require sampling tumor tissue from paraffin blocks for sequencing. An automated solution could potentially reduce the workload of pathologists by acting as a screening device and may reduce the subjectivity in diagnosis. As datasets have become publicly available and have opened up the possibility of evaluating the feasibility of applying deep learning techniques to improving the efficiency and quality of histologic diagnosis. In this project we introduce an application to detect Rectal cancer based on Convolutional Neural Network and Ranking algorithm.

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### CHAPTER 2

### LITERATURE SURVEY

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### CHAPTER 2 LITERATURE SURVEY

**2.1- TITLE : MACHINE LEARNING FOR COLORECTAL CANCER**

**RISK PREDICTION**

**AUTHOR :** Ling Zheng, Elijah Eniola, Jiacun Wang

**YEAR :**2021

**DESCRIPTION :**

Colorectal cancer is the third most prevalent cancer and the second most common cause of cancer deaths in the United States. Screening is one of the most powerful based on history of colorectal cancer and age. To facilitate a more effective screening of colorectal cancer, this paper explores the feasibilityof machine learning algorithms for the colorectal cancer risk colorectal cancer risk prediction. The longitudinal Pancreatic, Lung, Colorectal, Ovarian Cancer dataset from the National Cancer Institute was utilized for the training and testing of eight machine learning algorithms. The experiment results show that the gradient boosting model has the largest area under the Receiver Operating Characteristics curve 0.82, and the random forest model has the highest accuracy 0.75, highest recall 0.76 and highest F1score 0.75. The two optimal models were also used to evaluate the importance of top risk factors, which are helpful for a more effective screening recommendation.

**METHODOLOGY USED :**

This paper explores the feasibility of machine learning algorithm. The machine learning algorithms are used for training and testing.

**MERITS :**

The two optimal methods used are top risk factors and helpful for a more effective

screening recommendation.

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**DEMERITS:**

Dataset for the training and testing requires of eight machine learning algorithms.

**2.2 – TITLE : TWO STAGE CLASSIFICATION WITH CNN FOR**

**COLORECTAL CANCER DETECTION**

**AUTHOR :** Pallabi Sharma, Kangkana Bora, Kunio Kasugaiand Bunil

Kumar

**YEAR :**2020

**DESCRIPTION :**

In this paper, it addresses the current problem in medical image processing, the detection of colorectal cancer from colonoscopy videos. According to worldwide cancer statistics, colorectal cancer is one of the most common cancers. The process of screening and the removal of precancerous cells from the large intestine is a crucial task to date. The traditional manual process is dependent on the expertise of the medical practitioner. In this paper, a two-stage classification is proposed to detect colorectal cancer. In the first

stage, frames of colonoscopy video are extracted and are rated as significant if it contains a polyp, and these results are then aggregated in a second stage to come to an overall decision concerning the final classification of that frame to be neoplastic and non-Neoplastic. In doing so, a comparative study is being made by considering the applicability of deep learning to perform this two-stage classification. The CNN models namely VGG16, VGG19, Inception V3, Xception, GoogLeNet, ResNet50, ResNet100, DenseNet, NASNetMobile, MobilenetV2, InceptionResNetV2 and fine-tuned version each model. It is observed that the VGG19 model is the best deep learning method for colonoscopy image diagnosis.

**METHODOLOGY USED :**

The CNN models namely VGG16, VGG19, inception V3,Xception, GoogleNet, resnet50, Resnet 100, densenet, NASnet mobile, mobilenetV2, inception Resnet V2 and

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fine-tuned version of each model is evaluated.

**MERITS :**

As the process of screening and the removal of pre-cancerous cells from the large intestine is a crucial task by using these, we can reduce the task.

**DEMERITS:**

It fine-tuned version of each model is evaluated.

**2.3** – **TITLE**  **:** **GRADING OF COLORECTAL CANCER USING**

**HISTOLOGY IMAGES.**

**AUTHOR :** Namita Sengar; Neeraj Mishra, Malay Kishore Dutta, Jiri

Prinosil, Radim Burget

**YEAR :** 2020

**DESCRIPTION :**

This paper proposed an automated system for grading of colorectal cancer using image processing methods. Almost half a million people die every year due to colon cancer. Histopathological tissue analysis is a common method for its detection, which needs an expert pathologist. Screening for this cancer is effective for prevention as well as early detection. The method proposed segment the glands automatically by using intensity-based thresholding and organizational properties for classification. In existing literature, the majority of studies are based on gland segmentation in healthy or benign samples, but rarely on intermediate or high grade cancer. Unlike most of the existing methods this system is fully automated and grades the images as benign healthy, benign adenomatous, moderately differentiated malignant and poorly differentiated malignant. The proposed method achieves overall accuracy of 81% when tested on 165 histology images.

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**METHODOLOGY USED :**

Image processing methods.

**MERITS:**

We can predict at its earlier stag**e.**

**DEMERITS:**

It gives less accuracy result and its long process.

**2.4 – TITLE : AUTOMATIC CLASSIFICATION OF NON-INFORMATIVE**

**FRAME IN COLONOSCOPY VIDEOS**

**AUTHOR :**  Ballesteros, Trujillo and C. Mazo

**YEAR :** 2020

**DESCRIPTION:**

Colonoscopy is the most recommended test for prevention of colorectal cancer. Nowadays, digital videos are recorded during colonoscopy procedures and used for training machine learning algorithms. Machine learning algorithms are used for automatically recognizing lesions based on supervised learning. Moreover, annotation of lesions is a difficult and time consuming process that is manually made by gastroenterologists. Those annotations may contain frames that have not useful information, called non-Informative frames. The presence of non-Informative frames in a group of frames labelled as lesion affects the accuracy of machine learning algorithms. In this paper, a method based on edge detection is proposed to automatically classify a frame -from a colonoscopy video - into either Informative and Non-Informative. Non- Information Frames usually do not contain many edges. However, brightness regions produce false edges. Therefore, the proposed method includes a technique for brightness segmentation to identify false edges. The proposed method is evaluated using videos

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annotated by gastroenterologists. Elimination of No - Informative frames may reduce

significantly the number of frames to be annotated by gastroenterologists and may improve the accuracy of machine learning algorithms. Experimental evaluation showed that the accuracy and the precision of the proposed method is over 95%.

**METHODOLOGY USED:**

A random forest classifier was used for classification. An enhanced edge detection-based method was proposed.

**MERITS:**

It includesa technique for brightness segmentation to have accurate false edges.

**DEMERITS:**

1. Presence of Non-Information frames in a group of frames labelled as lesion affects the accuracy of machine learning algorithms.
2. Time consuming process.

**2.5 – TITLE : NON-INFORMATIVE FRAME CLASSIFICATION IN**

**COLONOSCOPY VIDEOS USING CNN**

**AUTHOR :** A. B. M. R. Isla, A. Alammari, W. Tavanapong, J. wong and P.

C.de groen.

**YEAR :** 2019

**DESCRITION :**

In the US, colorectal cancer is the second leading cause of cancer-related deaths behind lung cancer, causing about 49,000 annual deaths. Colonoscopy is currently the gold standard procedure for colorectal cancer screening. However, recent data suggest that there is a significant (4-12%) miss-Rate for the detection of even large polyps and

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cancers. To address this, we have been investigating an 'automated feedback system' which measures quality of colonoscopy automatically by analyzing colonoscopy video

frames in order to assist the endoscopist to improve the quality of the actual procedure being performed. One of the fundamental steps analyzing colonoscopy video frames for the automated quality feedback system is to distinguish non-informative frames from informative ones. Most methods to detect and classify these non-informative frames are based on the hand-engineered features. However, it is very tedious to design optimal hand-engineered features. In this paper, we explore the effectiveness of Convolutional Neural Network (CNN) to detect and classify these non-informative frames. The experimental results show that the proposed approaches are promising.

**METHODOLOGY USED:**

A CNN model was used with random trained dataset.

**MERITS:**

Easy for implementation and reduces the hand engineered work.

**DEMERITS:**

It is very tedious to design optimal hand-engineered features.

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**CHAPTER 3**

**SYSTEM ANALYSIS**

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**CHAPTER-3**

**SYSTEM ANALYSIS**

**3.1 EXISTING SYSTEM**

In the existing system, the concept of data mining systems supported on cancer prophecy system merging the prediction scheme with mining tools. The categorization algorithms used in the existing system is called decision tree. The user enters into the cancer prophecy scheme, and then required to retort the queries, connected to genetic and non-genetic skin textures. In that case the prediction structure allots the hazard rate to both query bases on the client retorts. One time the exposure significance is estimated, the series of the coercion preserve is resolute by the forecast structure. Research data shows that the accuracy of cancer prediction system is about 73%.

* 1. **PROPOSED SYSTEM**

With the development of targeted therapies, many treatments are based on molecular studies, which require sampling tumor tissue from paraffin blocks for sequencing. An automated solution could potentially reduce the workload of pathologists by acting as a screening device and may reduce the subjectivity in diagnosis. In tissue-based diagnostics, most of the work still needs to be done manually by a pathologist using a microscope to examine stained slides. The foundation of such tasks is to accurately distinguish cancer/malignant cells from normal/benign cells. However, the determination of tumor content is poorly reproducible with significant variation. As the size of tumor regions can be very small, pathologists are often required to use high magnification for detecting tumor cells. This requirement significantly increases the workload for pathologists. As digital pathology datasets have become publicly available and have opened up the possibility of evaluating the feasibility of applying deep learning techniques to improving the efficiency and quality of histologic diagnosis.

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As digital pathology datasets have become publicly available and have opened up the possibility of evaluating the feasibility of applying deep learning techniques to improving the efficiency and quality of histologic diagnosis. In this project we introduce an user facing AI based application to detect and predict the stage of Rectal cancer based on CNN with Attention mechanism and Ranking algorithm.

* 1. **FEASIBILITY STUDY**

Reduce the workload of pathologists by acting as a screening device and also

reduce the subjectivity in diagnosis.

Inclusion of feature in Scanning Devices for Quick analysis.

Possibility of evaluating the feasibility of applying deep learning techniques to

improving the efficiency and quality of histologic diagnosis.

**3.3.1 AREAS OF FEASIBILITY**

**Economic Feasibility:**

The financial cost related to this project it feasible as it only requires trained Model and system with good processing power.

* Total number of lines of code(LOC) = 2000K
* KLOC = 2000/1000 = 2
* Effort = 2.4\*(2)^1.05 = 4.969 person-month
* Development time = 2.5(4.969)^0.38 = 4.597 months
* Average staff size = 4.969/4.597 = 1.0809 person
* Productivity = 2/4.969 = 0.402 KLOC/person-month
* P = 402 LOC/person-month

Hence, it’s clear that this project is economically feasible

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**Technical Feasibility:**

* It is related to the feasibility of training the model and implementing it in an web application.
* Since the system implementation relies on processing power a decent machine with good processing capability is required.
* This project is based on machine learning algorithms and the technologies are :

1. Machine learning algorithm – CNN

2. Artificial Intelligence

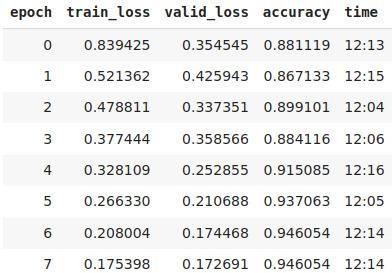
3. Azure ML

4. IDE : visual studio

**Schedule Feasibility:**

Based on the designed timeline chart the proposed system only requires 2-3 months

for developing it without any delay.



Using 7 epochs we are getting is 94% accuracy.

Fig. No. 3.1 Training with Epoch

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Fig. No. 3.2 Accuracy with Epoch

**3.4 HARDWARE ENVIRONMENT**

Processor - Pentium –IV

Speed - 1.1 Ghz

RAM - 4GB RAM

Hard Disk - 20 GB

Key Board - Standard Windows

Mouse - Two or Three Button Mouse

Monitor - ANY

**3.5 SOFTWARE ENVIRONMENT**

Operating System - Unix/Linux/XP/7/8/8.1/10

Coding Language - Python >= 3.8.0

Flask

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**CHAPTER 4**

**SYSTEM DESIGN**

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**CHAPTER 4**

**SYSTEM DESIGN**

**4.1 ER Diagram**

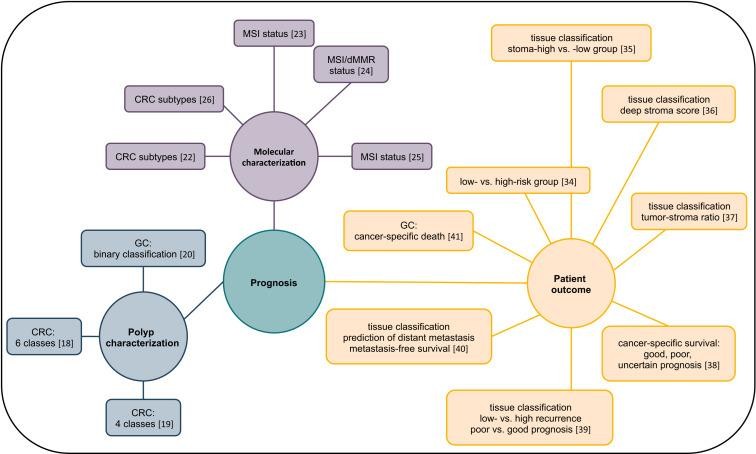


Fig. No. 4.1 ER Diagram

In the above diagram, the Prognosis connects with patient outcome, Molecular

characterization and Polyp characterization.

An Entity Relationship (ER) Diagram is a type of flowchart that illustrates how “entities”

such as people, objects or concepts relate to each other within a system.

**4.2** **DATA DICTIONARY**

The diagram depicts the data dictionary of the project. The entity ‘pixel’ has the attribute image, numpyis the data type. The entity ‘dataset’ has the attribute class with string as data type.

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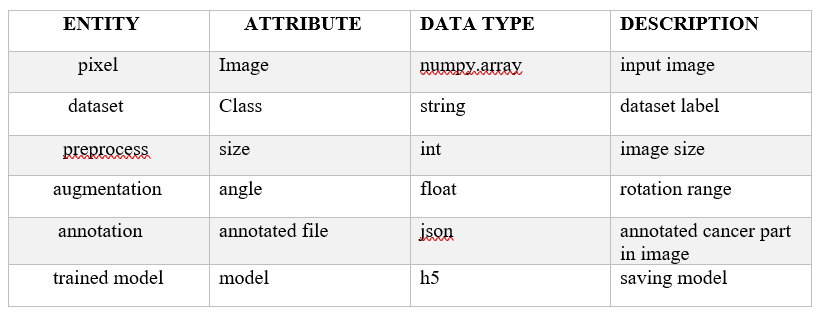


Table No.4.1 Data Dictionary

The entity ‘pre-processor’ has the attribute size and integer as data type. The entity ‘augmentation’ has angle as attribute and float as data type. The entity ‘annotation’ has annotated file as attribute and json as data type. The final entity ‘trained model’ has attribute model and h5 as data type.

* 1. **DATABASE DIAGRAM**

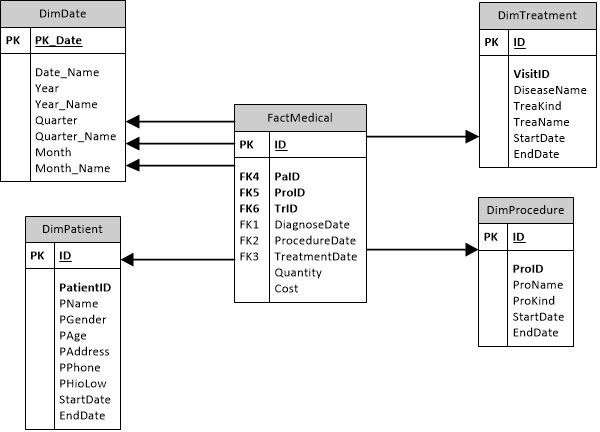


Fig. No. 4.2 Database Diagram

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In the above diagram, the class is static diagram and it is used to model the static view of the system. The static view describes the vocabulary of the system.

**4.4 DATA FLOW DIAGRAM**

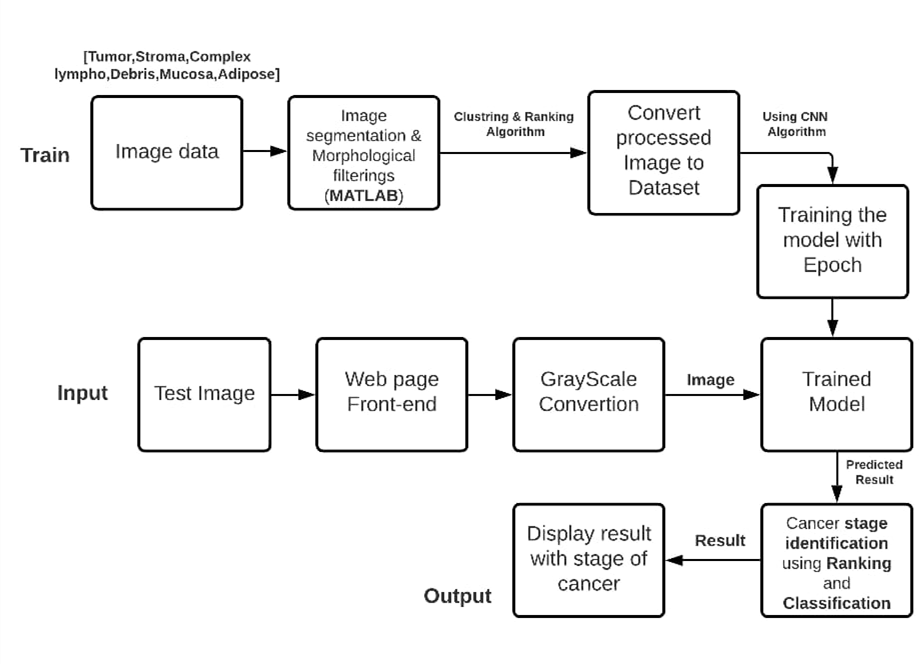


Fig. No.4.3 Data flow Diagram

* In training phase, we collect tissue datasets and apply morphological filtering’s.
* Second stage is clustering and ranking algorithm
* Clustering is used to group the images that comes under tumor, stroma etc., separately
* Ranking is used to rank the image
* Then we train the dataset using CNN algorithm, we train using epoch

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**4.5 UML DIAGRAM**

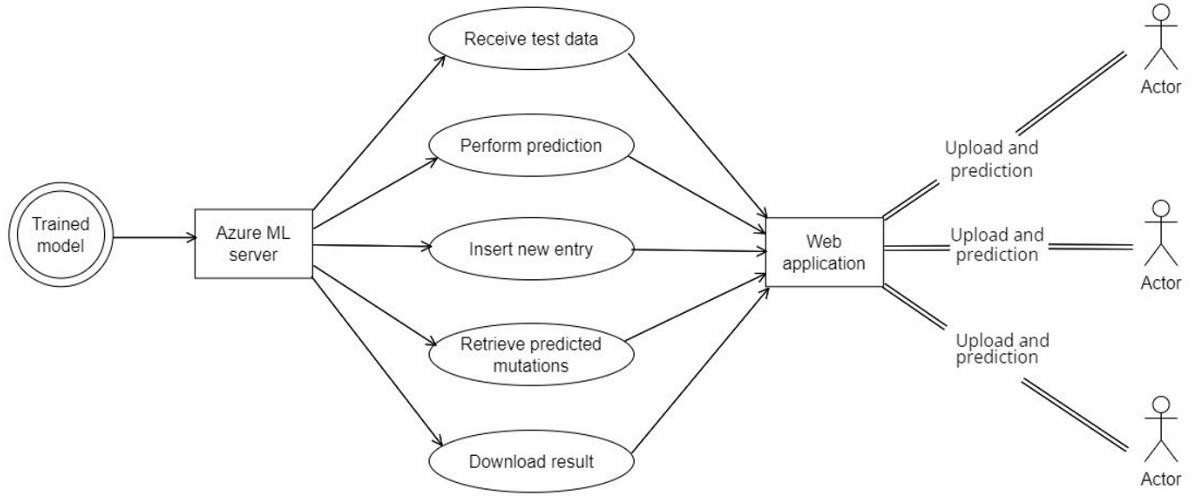
****

Fig. No. 4.4 UML Diagram

* In the above diagram, the images are trained and we get the trained model.
* The trained model is stored in Azure ML server.
* Though that we will perform prediction, insert new entry, retrieve predicted mutations and download result in web application.
* Through the web application, we will receive test data, then perform prediction, insert new entry, retrieve predicted mutual and download result in web application. In web application we will upload and predict then we will send to actor .

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**4.6 UI DIAGRAM**

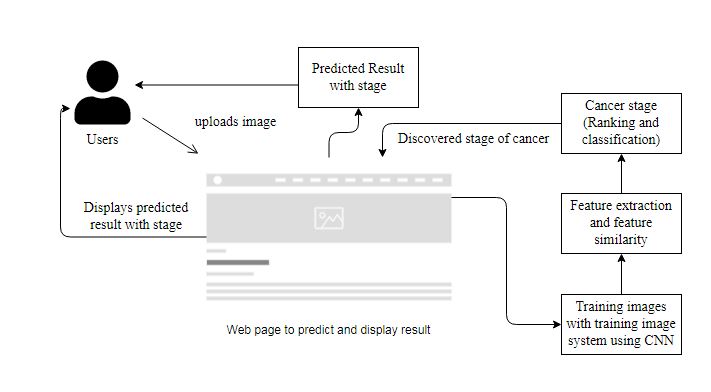


Fig. No.4.5 UI Diagram

* Here, the user uploads image in the webpage for prediction.
* The uploaded images are trained using CNN algorithm and the cancer stage is predicted.
* The predicted result with stage is displayed to the user in the webpage.

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**CHAPTER 5**

**SYSTEM ARCHITECTURE**

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**CHAPTER 5**

**SYSTEM ARCHITECTURE**

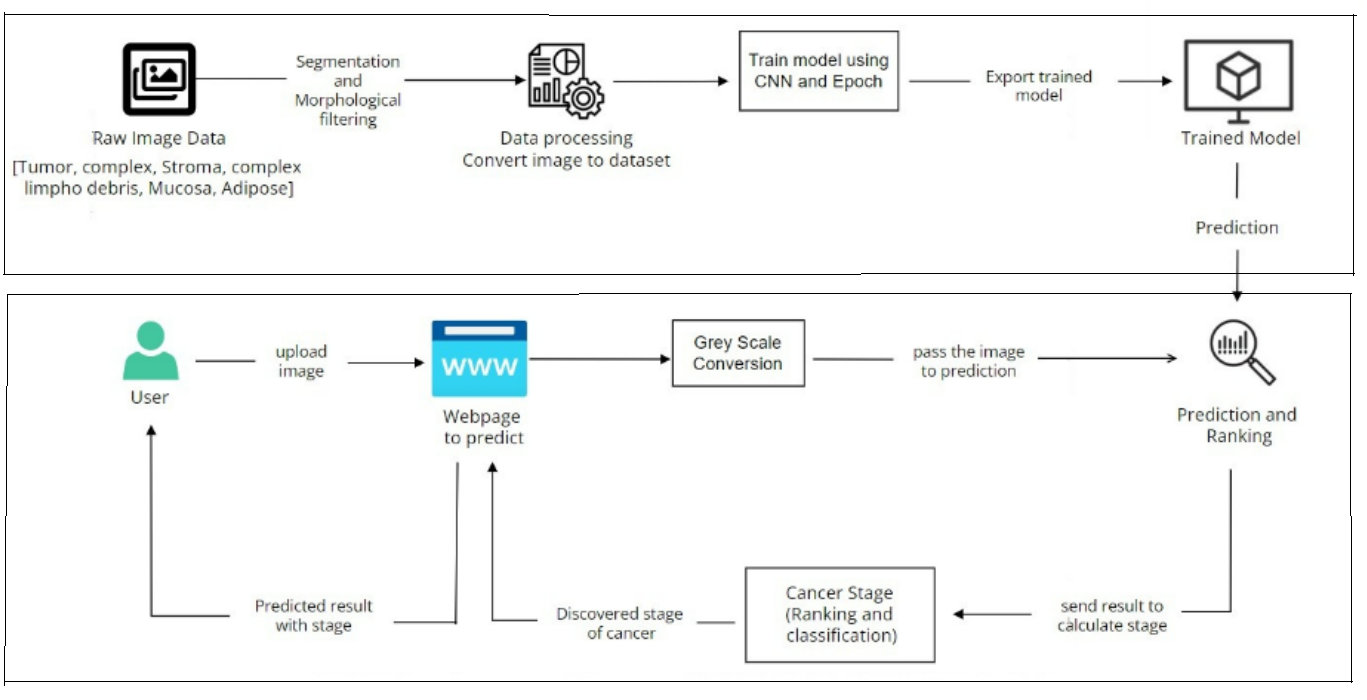


Fig. No. 5.1 Architecture Diagram

**5.1 MODULE DESIGN SPECIFICATION**

There are 4 modules:

1.Dataset preparation and pre-processing.

2.Dataset splitting.

3.Modelling.

4.Model deployment over web.

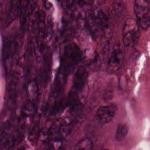
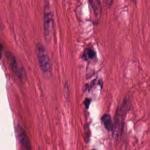
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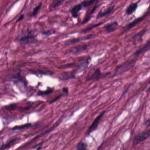
1. **Dataset preparation and pre-processing:**

Data is the foundation for any machine learning project. The second stage of project implementation is complex and involves data collection, selection, preprocessing, and transformation which include:

1. **Data collection:**

The image is collected from the external source via endoscopic ultrasound which uses ultrasound imaging and endoscopy to determine abnormalities in the colon. A special endoscope uses high-frequency sound waves to produce detailed images of the lining and walls of your digestive tract and chest. We have collected around 5000 images i.e., under each tissue there are 645 images. Parameters Of the image such as brightness, contrast and exposure are maintained such that the minor variability in them does not cause any deviation in the final output. The images are examined individually as it may contain unwanted noise which may be removed using image segmentation and morphological filtering.





Tumor Stroma Complex

1. **Data preprocessing:**

Here the collected tissues are given as raw data set to train the model for the prediction. After image conversion the images undergo image segmentation.

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Image segmentation involves converting an image into a collection of regions of pixels that are represented by a mask or a labeled image. By dividing an image into segments, you can process only the important segments of the image instead of processing the entire image.

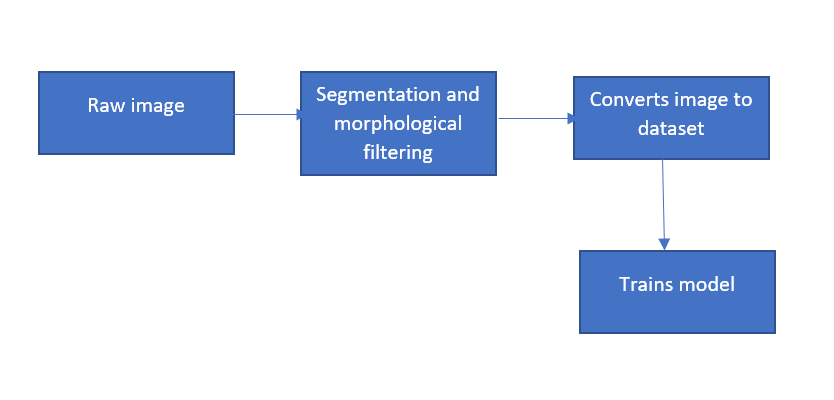


Fig. No. 5.2 Data Processing

1. **Data transformation:**

After training the data user has to select the image for the prediction using CNN algorithm.

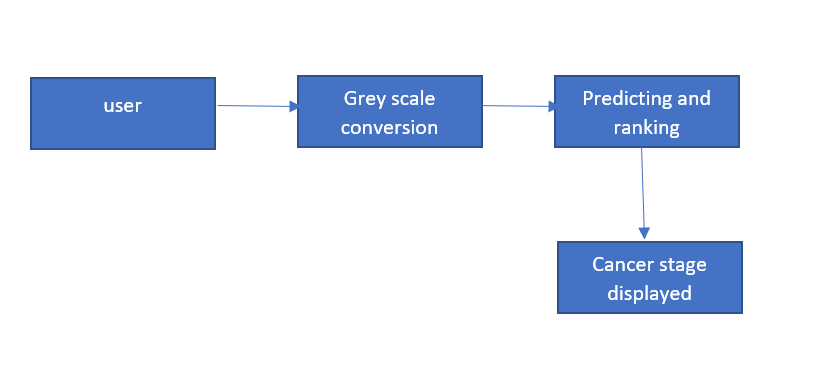


Fig. No. 5.3 Data Transformation

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1. **Data visualization:**

Here the data is represented the chart as we split the types of cancer in the form of distributed data.

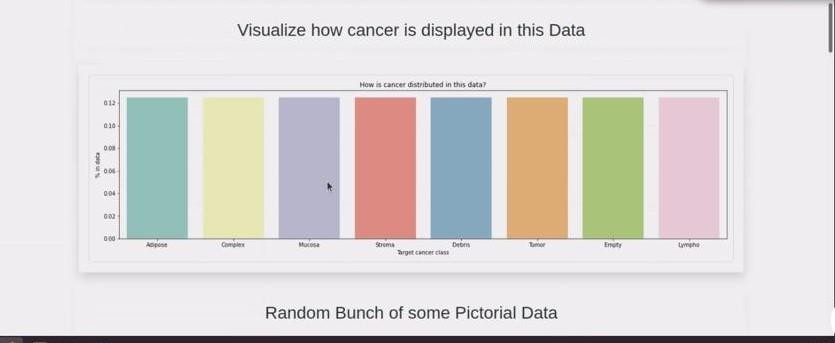


Fig. No. 5.4 Data Visualization

1. **Data splitting:**

**i) Training set:**

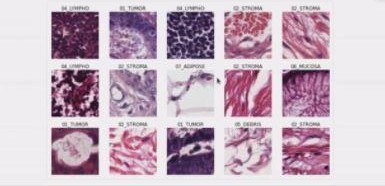
A data scientist uses a training set to train a model and define it optimal parameters it has to learn from data.

Fig. No. 5.5 Training set

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**ii) Test set:**

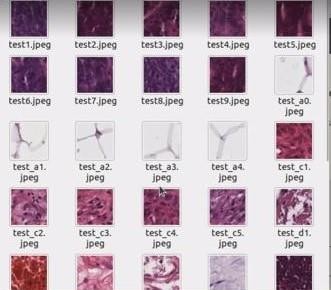
A test set is needed for an evaluation of the trained model and its capability for generalization.

Fig. No. 5.6 Test set

1. **Modelling:**

During this stage we train numerous models to define which one of them provides the most accurate prediction.

1. **Model training:**

Here the human tissues collected from the lab/hospitals are to be drained and converted to datasets.

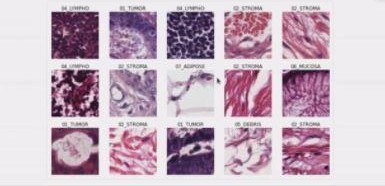


Fig. No. 5.7 Model Training

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1. **Model evaluation and testing:**

This process is done by the user as the trained images are available and need to be selected and the stage of the cancer is predicted. This is done in the web page.

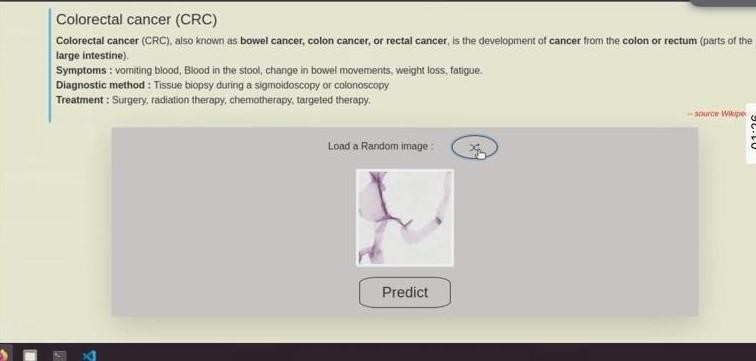


Fig. No. 5.8 Model evaluation and testing

1. **Model deployment over web:**

After selecting the image by the result as the cancer prediction and ranking is done over the web.

**5.2. ALGORITHMS**

**5.2.1 CLUSTERING**

Cluster analysis or clustering is the task of grouping a set of objects in such a way that objects in the same group (called a cluster) are more similar (in some sense) to each other than to those in other groups (clusters). Cluster analysis itself is not one specific algorithm, but the general task to be solved. It can be achieved by various algorithms that differ significantly in their understanding of what constitutes a cluster and how to efficiently find them.

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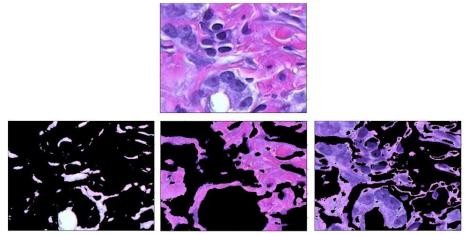


Fig. No. 5.9 Image Segmentation using Clustering

* It is a main task of exploratory data analysis, and a common technique image
* analysis and machine learning.
* The dataset contains a number of images. Clustering is used to group the images
* in the dataset under tumor, stroma etc.
* It is necessary to modify data preprocessing and model parameters until the

result achieves the desired properties.

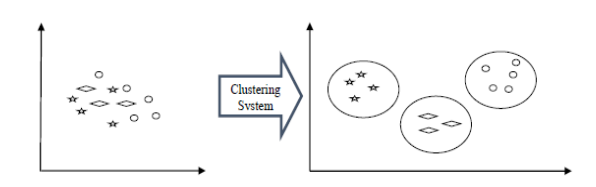


Fig. No. 5.10 Clustering of objects

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**5.2.2 RANKING ALGORITHM**

Randomized On-Line Matching, a representative of a class of algorithms, is a sequential algorithm that exploits a randomized efficient on-line matching algorithm that calculates maximal matchings in bipartite graphs, named the Ranking algorithm, as its basis. The Ranking algorithm makes a matching decision considering one output after the other. Specifically, during every switch cycle, the Ranking algorithm calculates the (maximal) matching, incrementally with the following steps:

S1: Calculate a random permutation *π*(*In*) (ordering) of inputs, which is the same for all outputs

S2: Consider outputOut[0] and identify the requests to it (i.e., the first input in *π*(*In*) that has a request for Out[0]; the requests of the selected input are deleted from the graph)

S3: MatchOut[0] to the eligible input (if any) of highest rank S4:

Repeat Steps S2 and S3 for all remaining outputs.

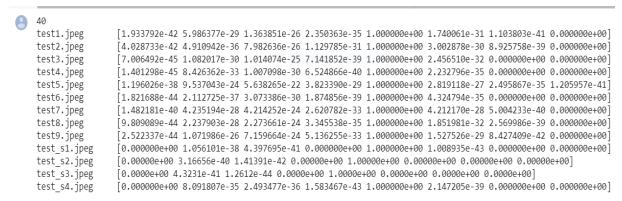


Fig. No. 5.11 Output for Ranking Algorithm

There are seven different types of cancer cells are as follows:

∙ Tumor

∙ Stroma

∙ Complex

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∙ Lymph

∙ Debris

∙ Mucosa

∙ Adipose

The ranking algorithm is used to rank the images.The above-mentioned tissues are allotted a rank number each and the images in the dataset is ranked based on their group.

For example, tumor – rank no 1, stroma – rank no 2, and so on.

* + 1. **CNN ALGORITHM**

In deep learning, a convolutional neural network (CNN, or ConvNet) is a class of deep neural networks, most commonly applied to analyzing visual imagery. They are also known as shift invariant or space invariant artificial neural networks (SIANN), based on the shared-weight architecture of the convolution kernels that shift over input features and provide translation equivariant responses. Counter-intuitively, most convolutional neural networks are only equivariant, as opposed to invariant, to translation. They have applications in image and video recognition, recommender systems, image classification, Image segmentation, medical image analysis, natural language processing, brain-computer interfaces, and financial time series are regularized versions of multilayer perceptron.

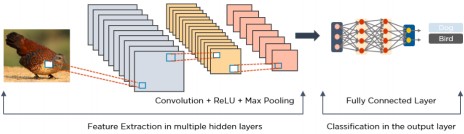


Fig. No. 5.12 Image processed via CNN

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Multilayer perceptron usually means fully connected networks, that is, each neuron in one layer is connected to all neurons in the next layer. The "full connectivity" of these networks makes them prone to overfitting data. Typical ways of regularization, or preventing overfitting, include: penalizing parameters during training (such as weight decay) or trimming connectivity. CNNs take a different approach towards regularization: they take advantage of the hierarchical pattern in data and assemble patterns of increasing complexity using smaller and simpler patterns embossed in their filters. Therefore, on a scale of connectivity and complexity, CNNs are on the lower extreme.

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**CHAPTER 6**

**SYSTEM INPLEMENTATION**

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**CHAPTER 6**

**SYSTEM IMPLEMENTATION**

**6.1. TRAINING CNN AND LSTM**

from fastai import \*

from fastai.vision import \*

import numpy as np

import pandas as pd

import seaborn as sns

import matplotlib.pyplot as plt

from sklearn.metrics import auc,roc\_curve

import os

print (os.listdir("/content/drive/My Drive/Colab Notebooks"))

% Matplotlib inline

class\_names = {1: "Tumor", 2: "Stroma", 3: "Complex", 4: "Lympho",

5: "Debris", 6: "Mucosa", 7: "Adipose", 8: "Empty"}

class\_numbers = {"Tumor": 1, "Stroma": 2, "Complex": 3, "Lympho": 4,

"Debris": 5, "Mucosa": 6, "Adipose": 7, "Empty": 8}

class\_colors = {1: "Red", 2: "Orange", 3: "Gold", 4: "Limegreen",

5: "Mediumseagreen", 6: "Darkturquoise", 7: "Steelblue", 8: "Purple"}

label\_percentage = df.label.value\_counts() / df.shape[0]

class\_index = [class\_names[idx] for idx in label\_percentage.index.values]

plt.figure(figsize=(20,5))

sns.barplot(x=class\_index, y=label\_percentage.values, palette="Set3");

plt.ylabel("% in data");

plt.xlabel("Target cancer class");

plt.title("How is cancer distributed in this data?");

tfms=get\_transforms(flip\_vert=True, max\_warp=0.)

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tfms=get\_transforms(flip\_vert=True, max\_warp=0.)

data = (ImageList.from\_folder(path)

.split\_by\_rand\_pct()

.label\_from\_folder()

.transform(tfms, size=150)

.databunch(num\_workers=2, bs=32))

learner= cnn\_learner(data, models.resnet34, metrics=[accuracy], model\_dir='/content/drive/My Drive/Colab Notebooks')

# Train the model on 4 epochs of data at the default learning rate

#learner.fit\_one\_cycle(4)

## Fit the model over 8 epochs

lr=5e-3 ## uncomment this

learner.fit\_one\_cycle(8, lr) ## uncomment this

#save the model

learner.save('/content/drive/My Drive/Colab Notebooks/level-1')

#print(os.listdir("./drive/My Drive/Colab Notebooks"))

#load the model

#learner.load('level-1')

#save the model

learner.save('level-2') ## uncomment this

#load the model

#learner.load('level-1')

# intrepting most confused

interp.most\_confused()

# ROC curve

fpr, tpr, thresholds = roc\_curve(lb.numpy(), preds.numpy()[:,1], pos\_label=1)

# ROC area

pred\_score = auc(fpr, tpr)

print(f'ROC area is {pred\_score}')

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plt.figure()

plt.plot(fpr, tpr, color='green', label='ROC curve (area = %0.2f)' % pred\_score)

plt.plot([0, 1], [0, 1], color='red', linestyle='--')

plt.xlim([-0.01, 1.0])

plt.ylim([0.0, 1.01])

plt.xlabel('False\_Positive\_Rate')

plt.ylabel('True\_Positive\_Rate')

plt.title('Receiver\_Operating\_Characteristic')

plt.legend(loc="lower right")

####/\*\*\*\*\*\*\*\*\*\*\*\*\*\*Testing and prediction (load level-2)\*\*\*\*\*\*\*\*\*\*\*\*\*\*/###

#learner.load("level-2")

learner.load("/content/drive/My Drive/Colab Notebooks/Kather\_texture\_2016\_image\_tiles\_5000/old\_level-1")

####/\*\*\*\*\*\*\*\*\*\*\*\*\*\*Testing and prediction \*\*\*\*\*\*\*\*\*\*\*\*\*\*/###

#learner.load("level-2")

# lets save our model with two formats: pkl and pth

#learner.export('pkl\_colorectal\_CNN\_model.pkl')

#learner.save('pth\_colorectal\_CNN\_model')

imageC1=random.choice(os.listdir("/content/drive/My Drive/Colab Notebooks/Kather\_texture\_2016\_image\_tiles\_5000/04\_LYMPHO/"))

#read = cv2.imread("/content/drive/My Drive/Colab Notebooks/test.jpeg")

#test\_image=cv2.imwrite("/content/drive/My Drive/Colab Notebooks/test\_tif.tif",read)

**6.2. EXPORT MODEL AND PERFORM UNIT TESTING**

print(imageC1)

# test case 1:

#159A9\_CRC-Prim-HE-07\_022.tif\_Row\_901\_Col\_151.tif ; 1EAE\_CRC-Prim-HE-10\_029.tif\_Row\_1\_Col\_451.tif [tumor or debris]

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#4B46\_CRC-Prim-HE-07.tif\_Row\_301\_Col\_601.tif [tumor debris adipose]

#test\_image=plt.imread("/content/drive/My Drive/Colab Notebooks/Kather\_texture\_2016\_image\_tiles\_5000/01\_TUMOR/"+imageC1)

# test case 2:

test\_image=plt.imread("/content/drive/My Drive/Colab Notebooks/Kather\_texture\_2016\_image\_tiles\_5000/04\_LYMPHO/"+imageC1)

#test case 3:

#test\_image=plt.imread("/content/drive/My Drive/Colab Notebooks/Kather\_texture\_2016\_image\_tiles\_5000/03\_COMPLEX/17D73\_CRC-Prim-HE-01\_034.tif\_Row\_451\_Col\_301.tif")

#print(os.listdir("/content/drive/My Drive/Colab Notebooks/Kather\_texture\_2016\_image\_tiles\_5000/02\_STROMA"))

#test\_image=plt.imread("/content/drive/My Drive/Colab Notebooks/Kather\_texture\_2016\_image\_tiles\_5000/02\_STROMA/11385\_CRC-Prim-HE-06\_003.tif\_Row\_601\_Col\_151.tif")

plt.imshow(test\_image)

file\_name=[]

predictions=[]

from PIL import Image as PImage

import cv2

#from fastai.vision import \*

#-----check for lympho---

lympholist=os.listdir("/content/drive/My Drive/Colab Notebooks/test\_images/")

print(len(lympholist))

for i in range(0,len(lympholist)):

if(lympholist[i].endswith(".jpeg")):

test\_image=plt.imread("/content/drive/My Drive/Colab Notebooks/test\_images/"+lympholist[i])

#------check end for lympho-----

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frame = cv2.cvtColor(test\_image,cv2.COLOR\_BGR2RGB)

pil\_im = PImage.fromarray(frame)

x = pil2tensor(pil\_im ,np.float32)

preds\_num = learner.predict(Image(x))[2].numpy()

#print(preds\_num)

if True:# preds\_num[4]!=0 and preds\_num[4]==max([preds\_num[0],preds\_num[1],preds\_num[2],preds\_num[3],preds\_n um[4],preds\_num[5],preds\_num[6],preds\_num[7]]) :

#print(l,"\n",class\_names)

file\_name.append(lympholist[i])

predictions.append(preds\_num)

#print(lympholist[i])

#print(preds\_num)

#break

#from sklearn.externals import joblib

#print(os.listdir("./drive/My Drive/Colab Notebooks/Kather\_texture\_2016\_image\_tiles\_5000"))

#classifer = joblib.load("./drive/My Drive/Colab Notebooks/Kather\_texture\_2016\_image\_tiles\_5000//drive/My Drive/Colab Notebooks/pkl\_colorectal\_CNN\_model.pkl")

for fn,pre in zip(file\_name,predictions):

print(fn,"\t",pre)

count=0

pridict=[]

for i in range(len(preds\_num)):

if(preds\_num[i]!=0):

count+=1

#print(class\_names[i+1]," ---> ",preds\_num[i])

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pridict.append(class\_names[i+1])

#print(class\_names[i+1],"\t")

print("According to our dataset the scan matches with:\n"," and ".join(pridict),"type of colorectal cancer")

**6.3. IMPLEMENT MODEL TO PREDICT OVER WEB**

from flask import Flask,render\_template,request,flash,url\_for,redirect

from werkzeug.utils import secure\_filename

import json

import random

import tablib

l=learner.load\_learner("./models/level1.pth")

app=Flask(\_\_name\_\_)

app.secret\_key = 'h432hi5ohi3h5i5hi3o2hi'

#create a route

@app.route('/')

def home():

return render\_template('index.html')

@app.route('/prediction',methods=['GET','POST'])

def result():

if request.method == 'POST':

#flash(" ".join(request.form.keys()))

f=request.form['img\_file'].split("/")

#-----------------------------------#

#result=jsonify(l.predict(f)) #

#json.dump(result,"testfile.json") #

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#-----------------------------------#

with open("testfile.json") as jfile:

dicl=json.load(jfile)

ifile=f[len(f)-1]

if ifile in dicl.keys():

result=dicl[ifile]

furl="/test\_images/"+f[len(f)-1]

stage="Can't identify"

if 'Mucosa' in result:

stage="S0"

if 'Lympho' in result:

stage="S1"

elif 'Debris' in result:

stage="S2"

if 'Stroma' in result or 'Complex' in result:

stage="S3I"

elif 'Lympho' in result:

stage="S3"

if 'Stroma' in result or 'Complex' in result:

stage="S3I"

#print(cell)

#print(result1)

return render\_template('index.html',isindex=True,imagef=str(url\_for("static",filename=furl)),result=result,stage=stage)

else:

return redirect(url\_for('home'))

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@app.route('/model')

def model():

return render\_template('model.html')

**PIP lock file:**

{

"\_meta": {

"hash": {

"sha256": "d5e270fef618e43f481bda408d839a4499afea70db57b11d01d3c414e4b94b4f"

},

"pipfile-spec": 6,

"requires": {

"python\_version": "3.7"

},

"sources": [

{

"name": "pypi",

"url": "https://pypi.org/simple",

"verify\_ssl": true

}

]

},

"default": {

"beautifulsoup4": {

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41 "sha256:84729e322ad1d5b4d25f805bfa05b902dd96450f43842c4e99067d5e1369eb25",

"sha256:fff47e031e34ec82bf17e00da8f592fe7de69aeea38be00523c04623c04fb666"

],

"index": "pypi",

"version": "==4.9.3"

},

"click": {

"hashes": [

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"sha256:dacca89f4bfadd5de3d7489b7c8a566eee0d3676333fbb50030263894c38c0dc"

],

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42

"sha256:8a4fdd8936eba2512e9c85df320a37e694c93945b33ef33c89946a340a238557"

],

"index": "pypi",

"version": "==1.1.2"

},

"itsdangerous": {

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"sha256:b12271b2047cb23eeb98c8b5622e2e5c5e9abd9784a153e9d8ef9cb4dd09d749"

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"version": "==1.1.0"

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"sha256:a6d58433de0ae800347cab1fa3043cebbabe8baa9d29e668f1c768cb87a333c6"

],

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"markers": "python\_version >= '2.7' and python\_version not in '3.0, 3.1, 3.2, 3.3, 3.4'",

"version": "==2.11.3"

},

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"hashes": [

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"sha256:9add70b36c5666a2ed02b43b335fe19002ee5235efd4b8a89bfcf9005bebac0d",

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"sha256:b1dba4527182c95a0db8b6060cc98ac49b9e2f5e64320e2b56e47cb283197 8c7",

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"sha256:e8313f01ba26fbbe36c7be1966a7b7424942f670f38e666995b88d012765b9be",

"sha256:ba59edeaa2fc6114428f1637ffff42da1e311e29382d81b339c1817d37ec93c6",

"sha256:84dee80c15f1b560d55bcfe6d47b27d070b4681c699c572af2e3c7cc90a3b8e0",

"sha256:d53bc011414228441014aa71dbec320c66468c1030aae3a6e29778a3382d9 6e5",

"sha256:acf08ac40292838b3cbbb06cfe9b2cb9ec78fce8baca31ddb87aaac2e2dc3bc2",

"sha256:195d7d2c4fbb0ee8139a6cf67194f3973a6b3042d742ebe0a9ed36d8b6f0c07f",

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45 "sha256:13d3144e1e340870b25e7b10b98d779608c02016d5184cfb9927a9f10c689f42",

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"sha256:cdb132fc825c38e1aeec2c8aa9338310d29d337bebbd7baa06889d09a60a1 fa2",

"sha256:6788b695d50a51edb699cb55e35487e430fa21f1ed838122d722e0ff0ac5ba15",

"sha256:1027c282dad077d0bae18be6794e6b6b8c91d58ed8a8d89a89d59693b9131db5",

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**CHAPTER 7**

**PERFORMANCE ANALYSIS**

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**CHAPTER 7**

**PERFORMANCE ANALYSIS**

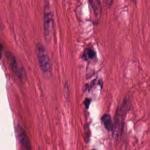
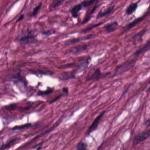
**7.1 RESULTS & DISCUSSIONS**

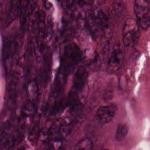
The training starts with dataset preparation and pre-processing and the data is separated using data splitting followed by modelling and then finally model deployment over web.

**Dataset preparation and pre-processing:**

Data is the foundation for any machine learning project. The second stage of project implementation is complex and involves data collection, selection, preprocessing, and transformation which include:

**Data collection:**

The image is collected from the external source via endoscopic ultrasound which uses ultrasound imaging and endoscopy to determine abnormalities in the colon. We have collected around 5000 images i.e., under each tissue there are 645 images.



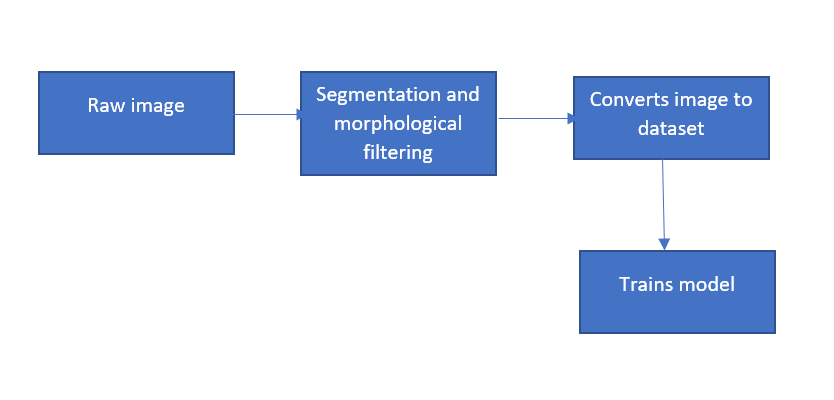


Fig. No. 7.1 Data Processing

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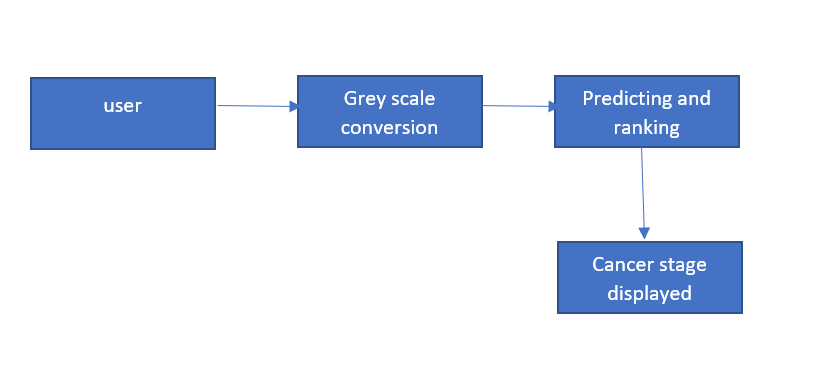


Fig. No. 7.2 Data Transformation

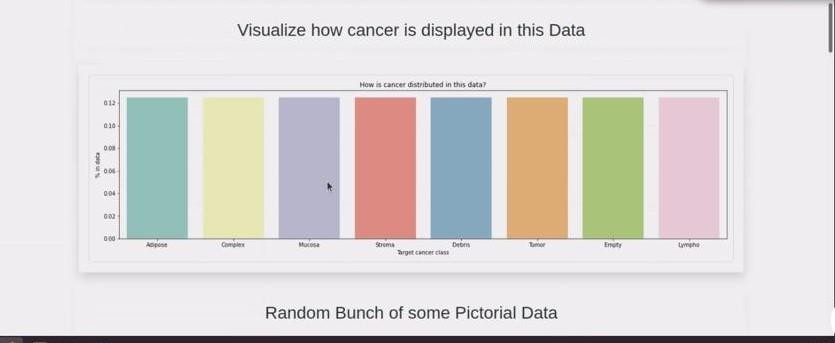


Fig. No. 7.3 Data Visualization

**Data splitting:**

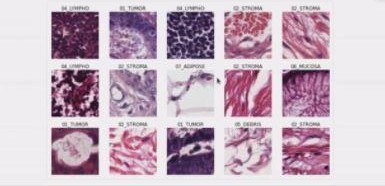
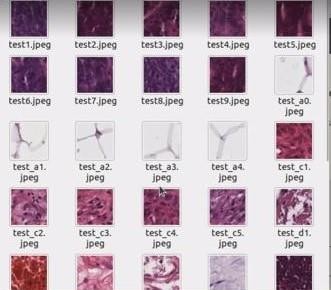


Fig. No. 7.4 Training set Fig. No. 7.5 Test set

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**Modelling:**

During this stage we train numerous models to define which one of them provides the most accurate prediction.

**Model training:**

Here the human tissues collected from the lab/hospitals are to be drained and converted to datasets.

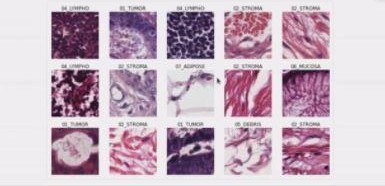


Fig. No. 7.6 Model Training

**Model evaluation and testing:**

This process is done by the user as the trained images are available and need to be selected and the stage of the cancer is predicted. This is done in the web page.

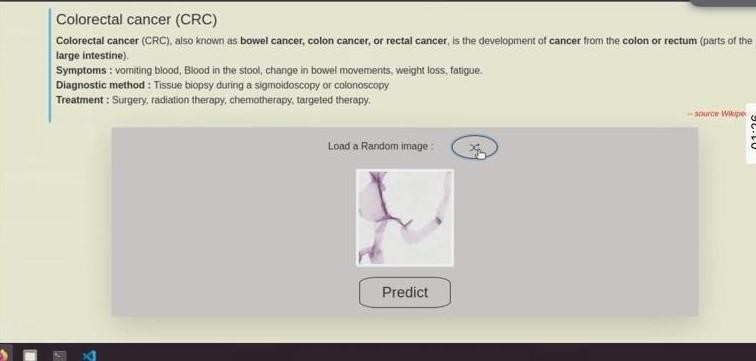


Fig. No. 7.7 Model evaluation and testing

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**Model deployment over web:**

After selecting the image by the result as the cancer prediction and ranking is done over the web.

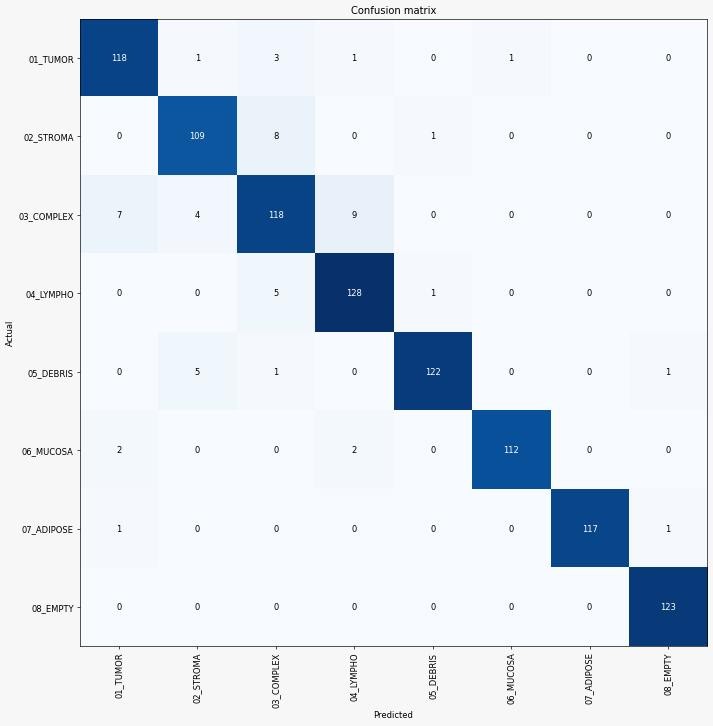


Fig. No. 7.8 Confusion Matrix

* 1. **ACCURACY**

**Accuracy :** It gives you the overall accuracy of the model, meaning the fraction of the total samples that were correctly classified by the classifier. To calculate accuracy, use the following formula: ***(TP+TN)/(TP+TN+FP+FN)***.

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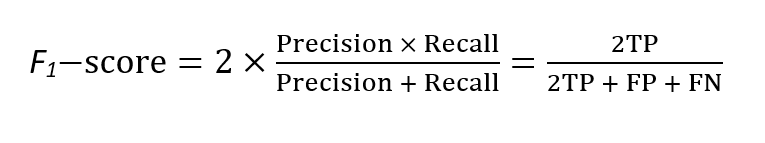
**Misclassification Rate :** It tells you what fraction of predictions were incorrect. It is also known as Classification Error. You can calculate it using ***(FP+FN)/(TP+TN+FP+FN)*** or ***(1-Accuracy)***.

**Precision:** It tells you what fraction of predictions as a positive class were actually positive. To calculate precision, use the following formula: ***TP/(TP+FP)***.

**Recall:** It tells you what fraction of all positive samples were correctly predicted as positive by the classifier. It is also known as True Positive Rate (TPR), Sensitivity, Probability of Detection. To calculate Recall, use the following formula: ***TP/(TP+FN)***.

**Specificity:**It tells you what fraction of all negative samples are correctly predicted as negative by the classifier. It is also known as True Negative Rate (TNR). To calculate specificity, use the following formula: ***TN/(TN+FP)***.

**F1-score:** It combines precision and recall into a single measure. Mathematically it’s the harmonic mean of precision and recall. It can be calculated as follows:



Now, in a perfect world, we would want a model that has a precision of 1 and a recall of 1. That means a F1-score of 1, i.e. a 100% accuracy which is often not the case for a machine learning model. So what we should try, is to get a higher precision with a higher recall value.

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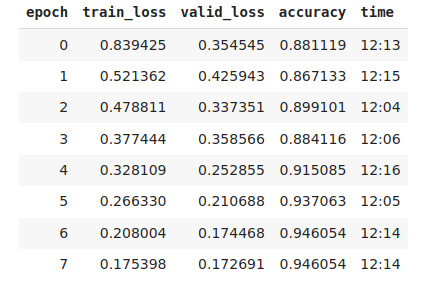


Table 7.1 Accuracy with Epoch

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**CHAPTER 8**

**CONCLUSION AND FUTURE WORK**

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**CHAPTER 8**

**CONCLUSION AND FUTURE WORK**

Early detection of cancer is very important in the medical field. In this work, we present an image-based feature extraction, segmentation and training approaches for classification and screening of cancer tissues. The previous work focused on k-mean clustering which is less efficient and accuracy was 73%. In our work we have increased the accuracy to 94% by training the model with EPOCH. LSTM is used for fast processing and stores the best result for comparison in the future. Analysis is done by training a Neural network using the processed set of images in predicting the future output from input given to model. Future works Will Be Directed Towards Analysis of additional data sets acquired under controlled imaging conditions. Since the datasets under analysis in this work represent a huge variety of imaging condition variabilities, the observations from the experimental analysis are more generalizable yet limited in classification capabilities.

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**APPENDICES**

**A.1 SAMPLE SCREENS**

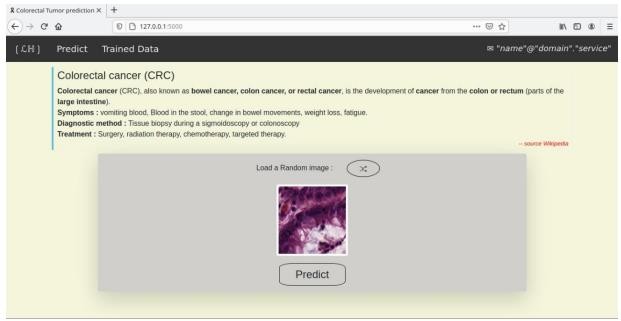


Fig 8.1 Uploading the image

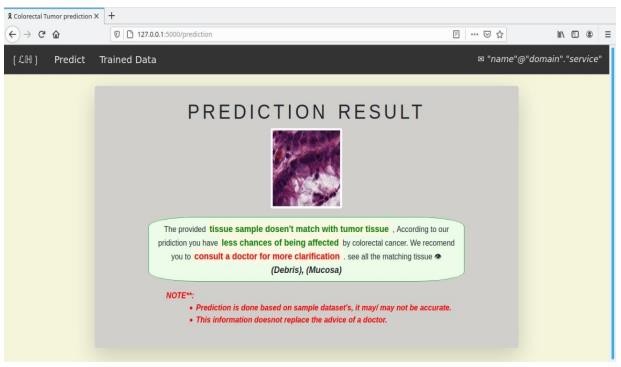


Fig 8.2 Result with less chance of colorectal cancer

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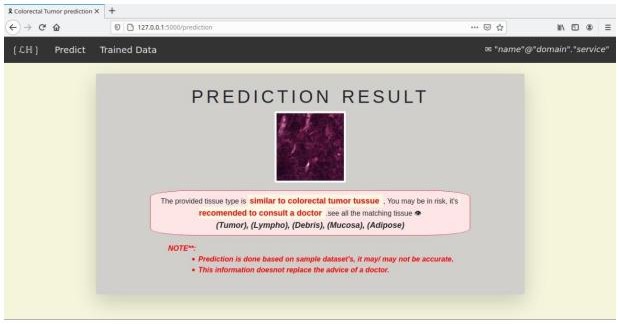


Fig 8.3 Result more chance of colorectal cancer

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