PROJECT REPORT

TEAM ID:PNT2022TMID03542

1. Introduction

1.1 Project Overview

Now a day's people are suffering from skin diseases, more than 125 million people suffering from Psoriasis also skin cancer rate is rapidly increasing over the last few decades especially Melanoma is most diversifying skin cancer. If skin diseases are not treated at an earlier stage, then it may lead to complications in the body including spreading of the infection from one individual to the other.

To overcome the above problem, we are building a model which is used for the prevention and early detection of skin cancer, psoriasis. Basically, skin disease diagnosis depends on the different characteristics like colour, shape, texture etc. Here the person can capture the images of skin and then the image will be sent the trained model. The model analyses the image and detect whether the person is having skin disease or not.

1.2 Purpose

The diseases are not considered skin diseases, and skin tone is majorly suffered from the ultraviolet rays from the sun. However, dermatologists perform the majority of non-invasive screening tests simply with the naked eye, even though skin illness is a frequent disease for which early detection and classification are essential for patient success and recovery. The characteristic of the skin images is diversified so that it is a challenging job to devise an efficient and robust algorithm for automatic detection of skin disease and its severity. Automatic processing of such images for skin analysis requires quantitative discriminator to differentiate the diseases.

2. Literature Survey

2.1 Existing problem

A neglected public health problem Skin diseases are among the most common health problems in humans. Considering their significant impact on the individual, the family, the social life of patients, and their heavy economic burden, the public health importance of these diseases is underappreciated.

2.2 References

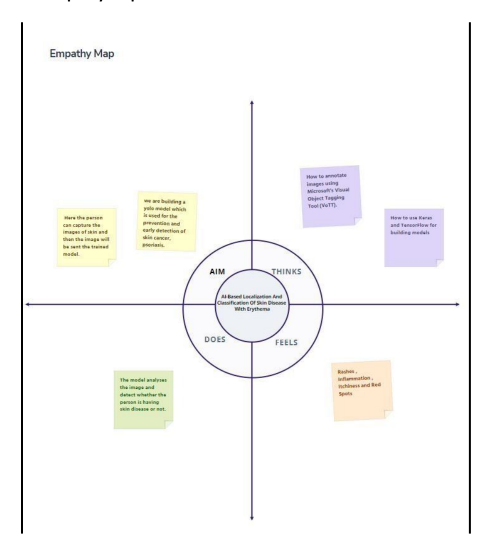
- [1] J. Kawahara and G. Hamarneh, "Multi-resolution-tract CNN with hybrid pretrained and skin-lesion trained layers," in International Workshop on Machine Learning in Medical Imaging, pp. 164–171, Springer, New York, NY, USA, 2016.
- [2] S. Verma, M. A. Razzaque, U. Sangtongdee, C. Arpnikanondt, B. Tassaneetrithep, and A. Hossain, "Digital diagnosis of Hand, Foot, and mouth disease using hybrid deep neural networks," IEEE Access, vol. 9, pp. 143481–143494, 2021.
- [3] P. P. Rebouças Filho, S. A. Peixoto, R. V. Medeiros da Nobrega´ et al., "Automatic histologically-closer classification of skin lesions," Computerized Medical Imaging and Graphics, vol. 68, pp. 40–54, 2018.

2.3 Problem Statement Definition

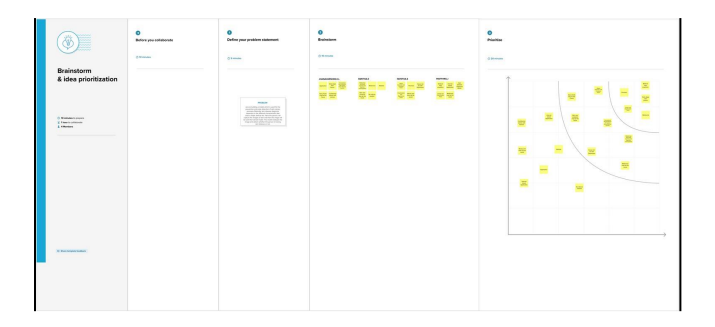
We're trying to find a solution to identify Skin Disease but Developed model is under training because given an image of skin, we can decompose, segment, and classify in a sequential manner which takes to Early detection of skin cancer, psoriasis.

3. Ideation and Proposed Solution

3.1 Empathy Map Canvas



3.2 Ideation and Brainstorming



3.3 Proposed Solution

Two-phase analysis model. The original image primarily enters a pre-processing stage, where normalization and decomposition occur. Afterwards, the first step is segmentation, where cluster of abnormal skin are segmented and cropped. The second step is classification, where each cluster is classified into its corresponding class. Developed Model is Still under training.

3.4 Problem Solution fit

Skin disease can appear in virtually any part of body and there is a lack of data required to form an association between the probability of a skin disease based on the body part. A Solution model used for the prevention and early detection of skin cancer and psoriasis by image analyses to detect whether the person is having skin disease or not. The location of the disease that is present in an image and improved performance by CNN model to focus on particular subsections of the images.

4. Requirement Analysis

4.1 Functional requirements

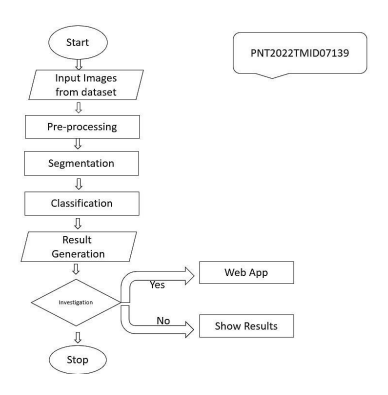
Image Acquisition, Pre-processing Steps such as Colour gradient generator on an image, Cropping and isolating region of interest and Thresholding and Clustering on image, Visual feature extraction, System Training YOLO Model for Skin disease classification with deep learning and CNN, Separate access of application for admin, Diagnosis of Skin disease and Data retrieval and Data manipulation.

4.2 Non-Functional requirements

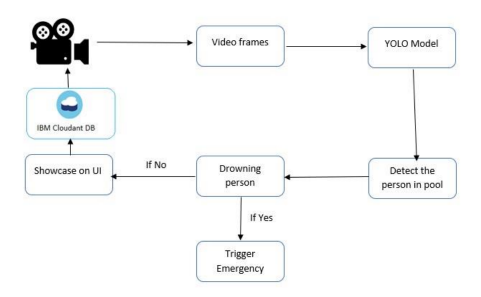
Software Quality Attributes, Prediction, Accuracy.

5. Project Design

5.1 Data Flow Diagram



5.2 Solution and Technical Architecture



5.3 User Stories

Functional Requirement (Epic)	User Story Number	User Story / Task	Story Points	Priority
Prerequisites	USN-1	Install Python IDE, Python packages, Microsoft Visual Object Tagging Tool, Yolo Structure	3	High
Data Collection	USN-2	Dataset should be collected from google or using a Chrome extension such as Fatkun Batch Downloader	3	High
Annotate Images	USN-3	Create A Project in VOTT (Microsoft's Visual Object Tagging Tool)	2	Medium
Training YOLO	USN-4	train our model using YOLO weights	2	Medium
	USN-5	To Download and Convert Pre-Trained Weights	3	High
	USN-6	To Train YOLOv3 Detector	3	High
Cloudant DB	USN-7	Register & Login to IBM Cloud	3	High
	USN-8	Create Service Instant and Credentials	2	Medium
	USN-9	Launch DB and Create database	3	High
Development Phase	USN-10	To build a web application	3	High
	USN-11	Building HTML pages with python code	2	Medium
	USN-12	To run the application	3	High
Testing Phase	USN-13	As a user login to dashboard	2	Medium
	USN-14	As a user import the images with skin diseases to the software application	2	Medium
	USN-15	YOLO processes the image and give the necessary details	3	High

6. Project Planning and Scheduling

6.1 Sprint Planning and Estimation

Sprint	Functional Requirement (Epic)	User Story Number	User Story / Task	Story Points	Priority	Team Members
Sprint-1	Prerequisites	USN-1	Install Python IDE, Python packages, Microsoft Visual Object Tagging Tool, Yolo Structure	3	High	Hemapriya.B Dhivya dharshini.S Dhivya dharshini.J Annapoorani.S
Sprint-1	Data Collection	USN-2	Dataset should be collected from google or using a Chrome extension such as Fatkun Batch Downloader	3	High	Hemapriya.B Dhivya dharshini.S Dhivya dharshini.J Annapoorani.S
Sprint-1	Annotate Images	USN-3	Create A Project in VOTT (Microsoft's Visual Object Tagging Tool)	2	Medium	Hemapriya.B Dhivya dharshini.S Dhivya dharshini.J Annapoorani.S
Sprint-2	Training YOLO	USN-4	train our model using YOLO weights	2	Medium	Hemapriya.B Dhivya dharshini.S Dhivya dharshini.J Annapoorani.S
Sprint-2		USN-5	To Download and Convert PreTrained Weights	3	High	Hemapriya.B Dhivya dharshini.S Dhivya dharshini.J Annapoorani.S
Sprint-2		USN-6	To Train YOLOv3 Detector	3	High	Hemapriya.B Dhivya dharshini.S Dhivya dharshini.J Annapoorani.S
Sprint-3	Cloudant DB	USN-7	Register & Login to IBM Cloud	3	High	Hemapriya.B Dhivya dharshini.S Dhivya dharshini.J Annapoorani.S
Sprint-3		USN-8	Create Service Instant and Credentials	2	Medium	Hemapriya.B Dhivya dharshini.S Dhivya dharshini.J Annapoorani.S

Sprint-3		USN-9	Launch DB and Create database	3	High	Hemapriya.B Dhivya dharshini.S Dhivya dharshini.J Annapoorani.S
Sprint-3	Development Phase	USN-10	To build a web application	3	High	Hemapriya.B Dhivya dharshini.S Dhivya dharshini.J Annapoorani.S
Sprint-3		USN-11	Building HTML pages with python code	2	Medium	Hemapriya.B Dhivya dharshini.S Dhivya dharshini.J Annapoorani.S
Sprint-3		USN-12	To run the application	3	High	Hemapriya.B Dhivya dharshini.S
Sprint-4	Testing Phase	USN-13	As a user login to dashboard	2	Medium	Hemapriya.B Dhivya dharshini.S
Sprint-4		USN-14	As a user import the images with skin diseases to the software application	2	Medium	Hemapriya.B Dhivya dharshini.J
Sprint-4		USN-15	YOLO processes the image and give the necessary details	3	High	Hemapriya.B

6.2 Sprint Delivery Schedule

Sprint	Total Story Points	Duration	Sprint Start Date	Sprint End Date (Planned)
Sprint-1	20	6 Days	24 Oct 2022	29 Oct 2022
Sprint-2	20	6 Days	31 Oct 2022	05 Nov 2022
Sprint-3	20	6 Days	07 Nov 2022	12 Nov 2022
Sprint-4	20	6 Days	14 Nov 2022	19 Nov 2022

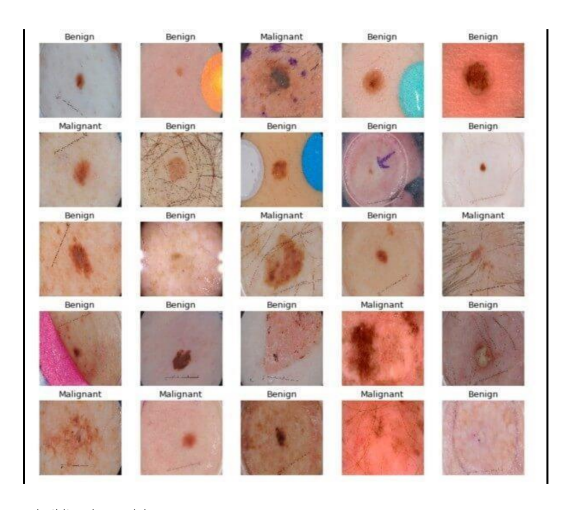
7. Coding and Solutioning

pip3 install tensorflow tensorflow_hub matplotlib seaborn numpy pandas sklearn imblearn

```
import tensorflow as tf import
tensorflow hub as hub import
matplotlib.pyplot as plt import
numpy as np import pandas as
pd import seaborn as sns
from tensorflow.keras.utils import get file
from sklearn.metrics import roc curve, auc, confusion matrix
from imblearn.metrics import sensitivity_score, specificity_score
import os import
glob import zipfile
import random
# to get consistent results after multiple runs
tf.random.set_seed(7) np.random.seed(7)
random.seed(7)
# 0 for benign, 1 for malignant
class_names = ["benign", "malignant"]
Preparing the Dataset
def download_and_extract_dataset():
 # dataset from https://github.com/udacity/dermatologist-ai
 # 5.3GB
 train url = "https://s3-us-west-1.amazonaws.com/udacity-dlnfd/datasets/skin-cancer/train.zip"
 #824.5MB
 valid url = "https://s3-us-west-1.amazonaws.com/udacity-dlnfd/datasets/skin-cancer/valid.zip"
 # 5.1GB
                               = "https://s3-us-west-1.amazonaws.com/udacity-dlnfd/datasets/skin-
                  test url
cancer/test.zip"
                  for i, download_link in enumerate([valid_url, train_url, test_url]):
                                                                                          temp_file =
f"temp{i}.zip"
  data_dir = get_file(origin=download_link, fname=os.path.join(os.getcwd(), temp_file))
print("Extracting", download_link) with zipfile.ZipFile(data_dir, "r") as z:
z.extractall("data") # remove the temp file
  os.remove(temp file)
# comment the below line if you already downloaded the dataset download and extract dataset()
# preparing data
# generate CSV metadata file to read img paths and labels from it def
generate_csv(folder, label2int): folder_name =
os.path.basename(folder) labels = list(label2int) # generate CSV
file df = pd.DataFrame(columns=["filepath", "label"]) i = 0 for
label in labels:
                  print("Reading", os.path.join(folder, label, "*"))
for filepath in glob.glob(os.path.join(folder, label, "*")):
      df.loc[i] = [filepath, label2int[label]]
  output file = f"{folder name}.csv"
                                      print("Saving",
output file)
  df.to_csv(output_file)
# generate CSV files for all data portions, labeling nevus and seborrheic keratosis
```

```
# as 0 (benign), and melanoma as 1 (malignant)
# you should replace "data" path to your extracted dataset path # don't replace if you
used download_and_extract_dataset() function generate_csv("data/train", {"nevus":
0, "seborrheic_keratosis": 0, "melanoma": 1}) generate_csv("data/valid", {"nevus": 0,
"seborrheic keratosis": 0, "melanoma": 1) generate csv("data/test", {"nevus": 0,
"seborrheic_keratosis": 0, "melanoma": 1})
# loading data
train metadata filename = "train.csv" valid metadata filename = "valid.csv" # load CSV
files as DataFrames df_train = pd.read_csv(train_metadata_filename) df_valid =
pd.read_csv(valid_metadata_filename) n_training_samples = len(df_train)
n_validation_samples = len(df_valid) print("Number of training samples:",
n_training_samples) print("Number of validation samples:", n_validation_samples)
train_ds = tf.data.Dataset.from_tensor_slices((df_train["filepath"], df_train["label"]))
valid ds = tf.data.Dataset.from tensor slices((df valid["filepath"], df valid["label"]))
Output:
Number of training samples: 2000 Number of validation samples: 150
# preprocess data def
decode img(img):
# convert the compressed string to a 3D uint8 tensor img =
tf.image.decode_jpeg(img, channels=3)
# Use `convert_image_dtype` to convert to floats in the [0,1] range.
img = tf.image.convert_image_dtype(img, tf.float32) # resize the image
to the desired size. return tf.image.resize(img, [299, 299])
def process path(filepath, label): # load the
raw data from the file as a string img =
tf.io.read_file(filepath) img = decode_img(img)
 return img, label
valid ds = valid ds.map(process path) train ds =
train ds.map(process path)
# test ds = test ds for image, label in
train ds.take(1): print("Image
shape:", image.shape) print("Label:",
label.numpy())
Image shape: (299, 299, 3)
Label: 0
# training parameters
batch_size = 64 optimizer =
"rmsprop" def
prepare_for_training(ds,
cache=True, batch size=64,
shuffle buffer size=1000):
if cache: if
```

```
isinstance(cache, str):
= ds.cache(cache) else:
   ds = ds.cache() #
shuffle the dataset
ds = ds.shuffle(buffer_size=shuffle_buffer_size)
# Repeat forever ds =
ds.repeat() # split to
batches ds =
ds.batch(batch_size)
# `prefetch` lets the dataset fetch batches in the background while the model # is
 ds = ds.prefetch(buffer_size=tf.data.experimental.AUTOTUNE) return ds
valid_ds = prepare_for_training(valid_ds, batch_size=batch_size, cache="valid-cached-data") train_ds =
prepare_for_training(train_ds, batch_size=batch_size, cache="train-cached-data") batch =
next(iter(valid_ds))
def show_batch(batch):
plt.figure(figsize=(12,12)) for
n in range(25):
                ax =
plt.subplot(5,5,n+1)
plt.imshow(batch[0][n])
   plt.title(class_names[batch[1][n].numpy()].title()) plt.axis('off')
show_batch(batch)
```



```
# building the model
# InceptionV3 model & pre-trained weights
module_url = "https://tfhub.dev/google/tf2-preview/inception_v3/feature_vector/4"

m = tf.keras.Sequential([
    hub.KerasLayer(module_url, output_shape=[2048], trainable=False),    tf.keras.layers.Dense(1, activation="sigmoid")
])

m.build([None, 299, 299, 3])
m.compile(loss="binary_crossentropy", optimizer=optimizer, metrics=["accuracy"])
m.summary()
```

Model: "sequential"

Layer (type)	Output Shape	Param #
keras_layer (KerasLa	yer) multiple	21802784

```
dense (Dense)
                                       multiple
                                                                       2049
========== Total params:
21,804,833
Trainable params: 2,049
Non-trainable params: 21,802,784
Training the Model
model_name = f"benign-vs-malignant_{batch_size}_{optimizer}"
tensorboard = tf.keras.callbacks.TensorBoard(log_dir=os.path.join("logs", model_name))
# saves model checkpoint whenever we reach better weights
model Checkpoint = tf. keras. callbacks. Model Checkpoint (model\_name + "\_\{val\_loss:.3f\}.h5", save\_best\_only = True, for each of the control of the contro
verbose=1)
history = m.fit(train_ds, validation_data=valid_ds,
steps per epoch=n training samples // batch size,
              validation_steps=n_validation_samples // batch_size, verbose=1, epochs=100,
callbacks=[tensorboard, modelcheckpoint])
Output:
Train for 31 steps, validate for 2 steps
Epoch 1/100
00001: val_loss improved from
                                                                   inf to 0.49703,
                                                                                                            saving
                                                                                                                             model
vsmalignant 64 rmsprop 0.497.h5
31/31 [=========================] - 282s 9s/step - loss: 0.4646 - accuracy: 0.7722 - val_loss:
0.4970 - val_accuracy: 0.8125
<..SNIPED..>
Epoch 27/100
val_loss improved from 0.40253 to 0.38991, saving model to benign-
vsmalignant_64_rmsprop_0.390.h5
val_accuracy: 0.8359
<..SNIPED..>
Epoch 41/100
Epoch 00041: val loss did not improve from 0.38991
31/31 [=====================] - 21s 690ms/step - loss: 0.2829 - accuracy: 0.8790 - val loss: 0.3948 -
val accuracy: 0.8281
Epoch 42/100
Epoch 00042: val_loss did not improve from 0.38991
31/31 [=====================] - 21s 693ms/step - loss: 0.2722 - accuracy: 0.8831 - val_loss: 0.4572 -
val accuracy: 0.8047
```

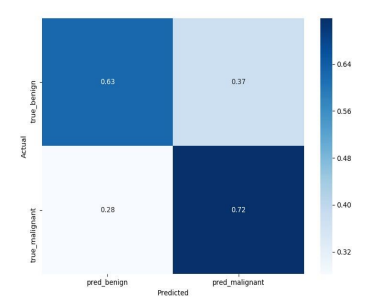
Model Evaluation:

```
# evaluation #
load testing set
test_metadata_filename = "test.csv"
df_test = pd.read_csv(test_metadata_filename)
n_testing_samples = len(df_test) print("Number of testing
samples:", n_testing_samples)
test_ds = tf.data.Dataset.from_tensor_slices((df_test["filepath"], df_test["label"])) def
prepare_for_testing(ds, cache=True, shuffle_buffer_size=1000): if cache: if
isinstance(cache, str):
                          ds = ds.cache(cache)
  else:
   ds = ds.cache() ds =
ds.shuffle(buffer size=shuffle buffer size) return
ds
test_ds = test_ds.map(process_path) test_ds =
prepare_for_testing(test_ds, cache="test-cached-data")
Number of testing samples: 600
# evaluation #
load testing set
test_metadata_filename = "test.csv" df_test =
pd.read_csv(test_metadata_filename)
n_testing_samples = len(df_test)
print("Number of testing samples:", n testing samples) test ds =
tf.data.Dataset.from_tensor_slices((df_test["filepath"], df_test["label"]))
 def prepare for testing(ds, cache=True, shuffle buffer size=1000): if
cache:
  if isinstance(cache, str):
                            ds
= ds.cache(cache)
  else:
   ds = ds.cache()
 ds = ds.shuffle(buffer_size=shuffle_buffer_size) return ds
test_ds = test_ds.map(process_path) test_ds =
prepare_for_testing(test_ds, cache="test-cached-data")
# load the weights with the least loss
m.load_weights("benign-vs-malignant_64_rmsprop_0.390.h5")
print("Evaluating the model...")
loss, accuracy = m.evaluate(X test, y test, verbose=0)
print("Loss:", loss, " Accuracy:", accuracy)
```

Evaluating the model...

```
Loss: 0.4476394319534302 Accuracy: 0.8
def get predictions(threshold=None):
 Returns predictions for binary classification given 'threshold'
 For instance, if threshold is 0.3, then it'll output 1 (malignant) for that sample if the
probability of 1 is 30% or more (instead of 50%)
y_pred = m.predict(X_test) if not
threshold: threshold = 0.5 result =
np.zeros((n testing samples,)) for iin
range(n_testing_samples): # test
melanoma probability if y_pred[i][0] >=
threshold:
   result[i] = 1 # else,
it's 0 (benign)
 return result
threshold = 0.23
# get predictions with 23% threshold
# which means if the model is 23% sure or more that is malignant,
# it's assigned as malignant, otherwise it's benign
y_pred = get_predictions(threshold) def
plot_confusion_matrix(y_test, y_pred): cmn =
confusion_matrix(y_test, y_pred)
# Normalise
 cmn = cmn.astype('float') / cmn.sum(axis=1)[:, np.newaxis]
 # print it print(cmn)
 fig, ax = plt.subplots(figsize=(10,10)) sns.heatmap(cmn,
annot=True, fmt='.2f',
                              xticklabels=[f"pred_{c}" for c in
class_names],
                      yticklabels=[f"true_{c}" for c in
                      cmap="Blues"
class_names],
 plt.ylabel('Actual') plt.xlabel('Predicted')
 # plot the resulting confusion matrix plt.show()
plot_confusion_matrix(y_test, y_pred)
```

Output:

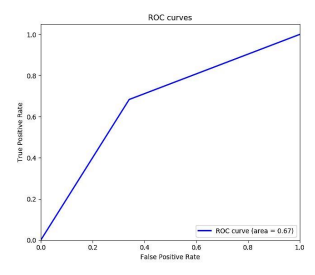


```
sensitivity = sensitivity_score(y_test, y_pred)
specificity = specificity_score(y_test, y_pred)
print("Melanoma Sensitivity:", sensitivity)
print("Melanoma Specificity:", specificity)
```

Melanoma Sensitivity: 0.717948717948718 Melanoma Specificity: 0.6252587991718427

plot_roc_auc(y_test, y_pred)

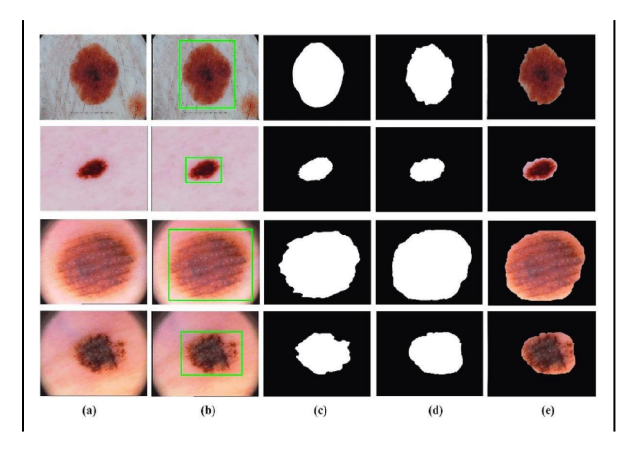
Output:



ROC AUC: 0.671

8. Results

The final results are based on the accuracy results in the form of the melanoma and the non-melanoma skin diseases classifications.



9. Advantages and Disadvantages

9.1 Advantages

Instant Response, improves prediction of Skin Disease, no referral needed, Saves Money and Time, and Confidential Advice.

9.2 Disadvantages

Network Connectivity and Accuracy

10. Conclusion

We have shown that even without a large dataset and high-quality images, it is possible to achieve sufficient accuracy rates. In addition, we have shown that current state-of-the-art CNN models can outperform models created by previous research, through proper data pre-processing, self-supervised learning, transfer learning, and special CNN architecture techniques. Furthermore, with accurate segmentation, we gain knowledge of the location of the disease, which is useful in the pre-processing of data used in classification, as it allows the CNN model to focus on the area of interest. Lastly, unlike previous studies, our method provides a solution to classify multiple diseases within a single image. With higher quality and a larger quantity of data, it will be viable to use state-of-the-art models to enable the use of CAD in the field of dermatology.

11. Future Scope

This implementation of the Structural Co-Occurrence matrices for feature extraction in the skin diseases classification and the pre-processing techniques are handled by using the Median filter, this filter helps to remove the salt and pepper noise in the image processing; thus, it enhances the quality of the images, and normally, the skin diseases are considered as the risk factor in all over the world. Our proposed approach provides 97% of the

classification of t	the accuracy results while another existing model such as FFT + SCM gives 80%, SVM + SCM give	ς
83%, KNN + SCM	gives 85%, and SCM + CNN gives 82%. Future work is dependent on the increased support vectacy in classifying skin illnesses, and SCM is used to manage the feature extraction technique.	
12. Appendi	i x	
GitHub Link:		
https://github.co	om/IBM-EPBL/IBM-Project-38343-1660378786	