Fertilizers Recommendation System for Disease Prediction

PROJECT REPORT

Submitted by

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TABLE OF CONTENT

S.NO	CONTENT	PA
		GE
		NO
1	INTRODUCTION	NO
1	INTRODUCTION	01
2	LITERATURE SURVEY	01
3	THEORETICAL ANALYSIS	02
4		
	EXPERIMENTAL INVESTIGATIONS	03
5	FLOW CHART	05
6	RESULTS	07
_	OT ALEDY ALE	10
7	OUTPUT	10
8	ADVANTAGES AND DISADVANTAGES	14
9	APPLICATION	14
	CONCLUCION	4 50
10	CONCLUSION	15

11	FUTURE SCOPE	15
12	BIBILOGRAPHY	16
	APPENDIX	
	A) PYTHON CODE	

INTRODUCTION

Overview In this project, two datasets name fruit dataset and vegetable dataset are collected. The collected datasets are trained and tested with deep learning neural network named Convolutional Neural Networks (CNN). First, the fruit dataset is trained and then tested with CNN. It has 6 classes and all the classes are trained and tested. Second, the vegetable dataset is trained and tested. The software used for training and testing of datasets is Python. All the Python codes are first written in Jupyter notebook supplied along with Anaconda Python and then the codes are tested in IBM cloud. Finally, a web-based framework is designed with help Flask a Python library. There are 2 html files are created in templates folder along with their associated files in static folder. The Python program 'app.py' used to interface with these two web pages is written in Spyder-Anaconda python and tested.

Purpose this project is used to test the fruits and vegetables samples and identify the different diseases. Also, this project recommends fertilizers for predicted diseases.

2.LITERATURE SURVEY

Existing problem indhu proposed a method for leaf disease detection and suggest fertilizers to cure leaf diseases. But the method involves less number of train and test sets which results in poor accuracy, deepika proposed a simple prediction method for soil-based fertilizer recommendation system for predicted crop diseases. This method gives less accuracy and prediction. Pradnya dahake proposed an IOT based system for leaf disease detection and fertilizer recommendation which is based on Machine Learning techniques yields less 80 percentage accuracies.

Proposed solution In this project work, a deep learning based neural network is used to train the collected datasets and test the same. The deep learning based neural network is CNN which gives more than 90% classification accuracies. By increasing

The more number of dense layers and by modifying hyper parameters such as number of epochs, batch size, the accuracy rate can be increased to 95% to 98%.

3. THEORITICAL ANALYSIS

Block diagram

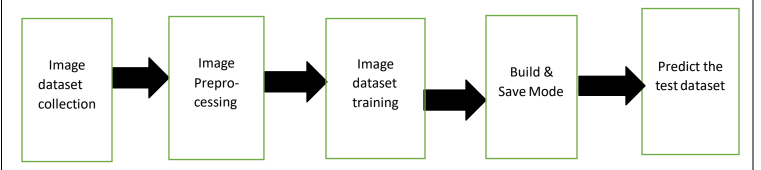


Figure.3.1. Block Diagram of the project

The block diagram of the entire project is shown in Fig.3.1. First step is the image dataset collection followed by image preprocessing. The third step is the training of image datasets with initializing different hyper parameters. Then build the model and save the model file with .h5 format. The final stage is the testing of existing or new datasets using the trained model.

Hardware/Software designing

The software used for training and testing the dataset is Python. The Jupyter notebook (Notebook of IBM cloud also) is used for python programming. The neural network used for training and testing the model is Convolutional Neural Network (CNN).

The CNN has following layers:

• Convolutional layer (32x32 kernal (3x3))

- Max-pool layer (kernel(2x2))
- Flatten layer
- Dense layer (different layers with different size)
- Drop out layer (optional)
- Final output dense layer(size 6x1 for fruit dataset and 9x1 for Vegetable dataset)

In the preprocessing step, images are normalized to 1 and then resized to 128x128. The images are arranged in different batch sizes. Then train set and test set are formed from the collected datasets. In order to do the above steps in Python, the following Python libraries must be imported before starting the process:

- NumPy
- TensorFlow
- Keras
- Matplotlib (optional for data visualization)

The following activation functions used in the CNN training:

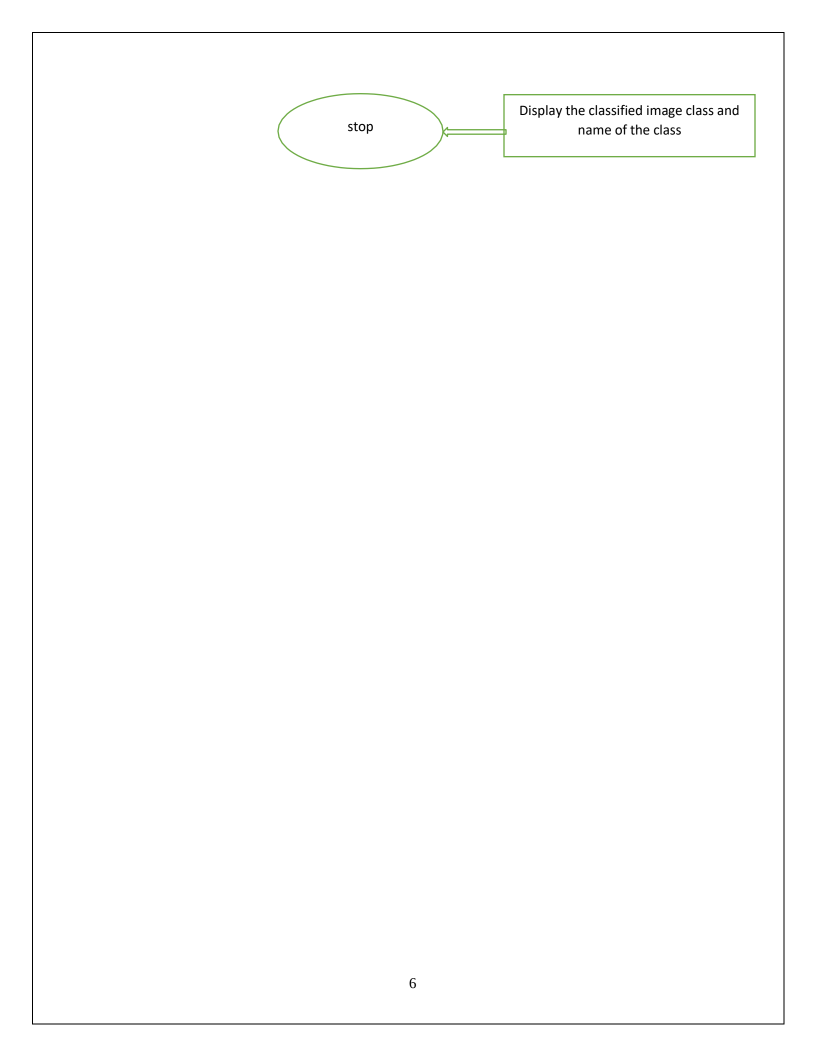
- RELU at the end of convolution layer and Max Pool layer
- SoftMax at the end of output dense layer
- For testing the dataset argmax is used, its an optional

4. EXPERIMENTAL INVESTIGATIONS

Analysis made while working on the solution The batch sizes are varied and tested. For different batch sizes, the CNN gives different accuracies. The batch size

determines the number of iterations per epoch. Another important hyper parameter is the number of epochs. This determines accuracy and it has high influence on accuracy compared to other hyper parameters. The accuracy can be varied from 80% to 90% in vegetable dataset and 95% to 98% in the case of fruit dataset by increasing the number of epochs. The size of test dataset and train dataset also has very high influence on accuracies. The accuracy can be increased by using more number of images in train dataset. The computational time for model building is increased when the size of the train dataset increased and also number of epochs increased. The batch size of train dataset and test dataset also play a vital role in computational time. The Neural Network complexity is increased when more number of convolutional layers increased. If the number of layers increased, better accuracy result will obtain. At the same increasing the number of layers in CNN leads to more training time and also requires more time to build a model. The model .h5 size depends on the size of train dataset and CNN architecture complexity.

5.FLOWCHARTS Start Image Data collection Norm factor=1./255 Batch size=32 for fruit dataset ,16 for vegetable dataset image resize=128*128 Build the model and train the dataset using desired hyperparameters If classified No Accuracy=Accuracy correctly



6.RESULTS

Final findings(output) of the project given below in the form of screenshot: Training and Testing of Fruit dataset

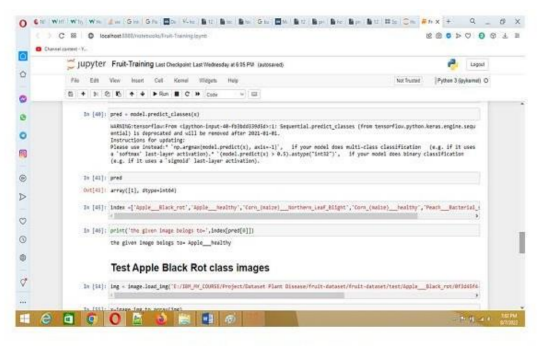


Figure.6.2 Test the Fruit dataset

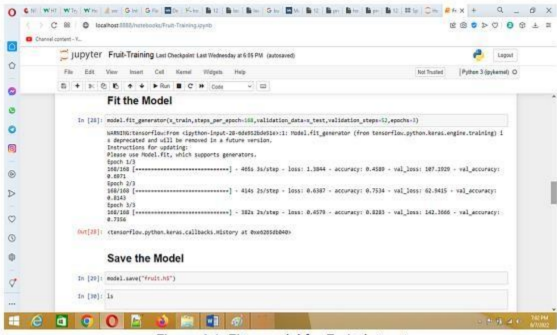


Figure.6.1. Fit a model for Fruit dataset

Train and Test Vegetable dataset

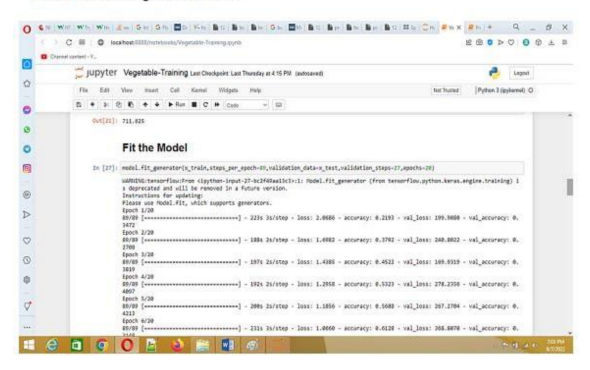


Figure.6.3. Train the Vegetable dataset

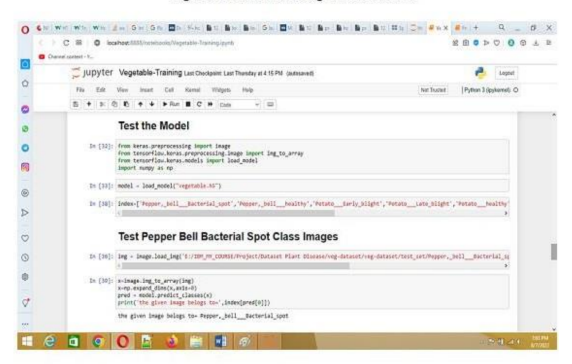


Figure.6.4. Test the Vegetable dataset

Train and Test Vegetable dataset IBM Cloud

Due to CUH limit exceeds, I have downloaded the notebooks and opened in Jupyter notebook

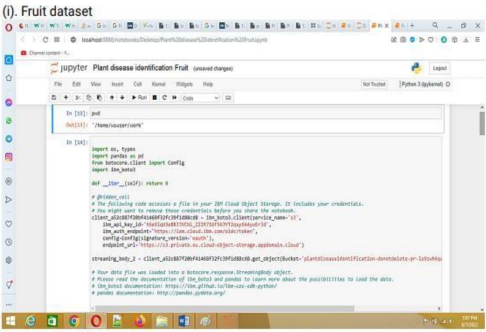


Figure.6.5. Training Fruit Dataset in IBM Cloud

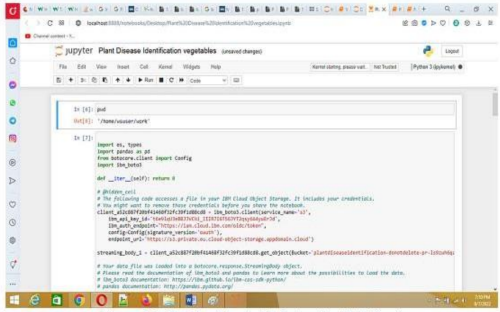
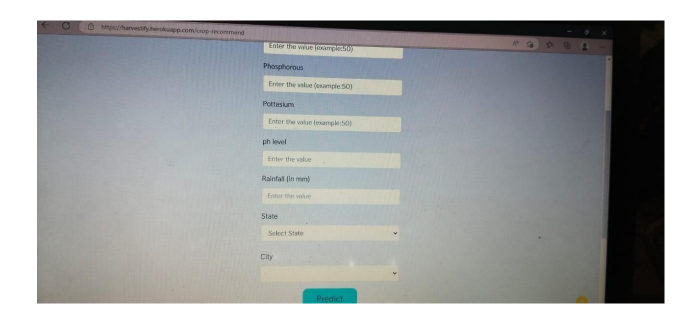
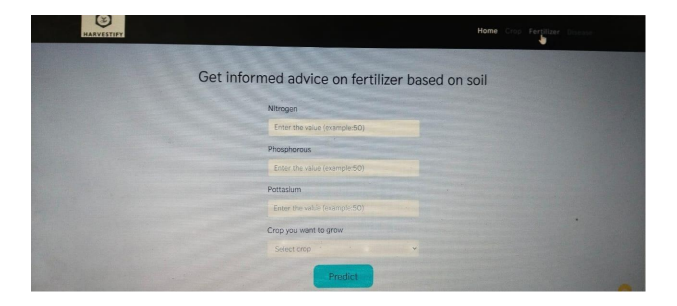


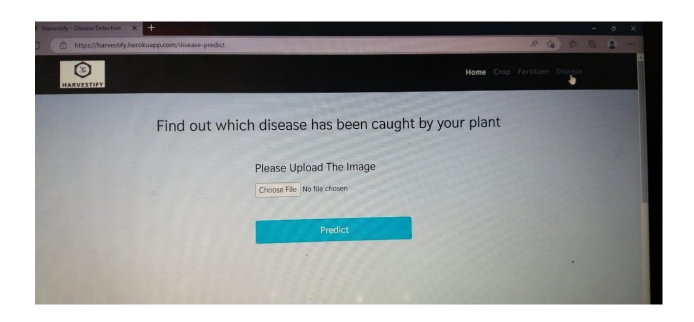
Figure.6.6. Training Vegetable Dataset in IBM Cloud

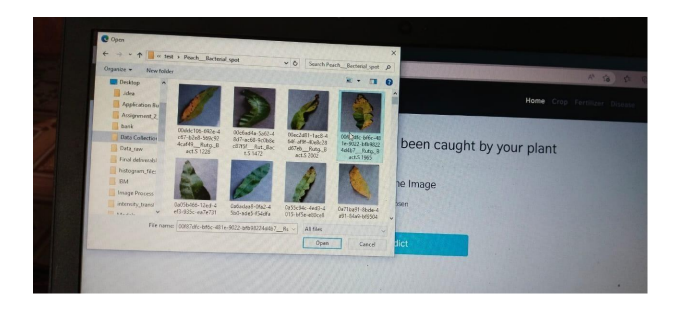
Out put

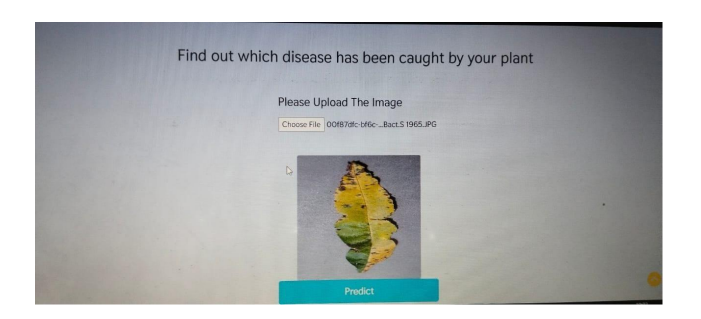












Crop: Peach Disease: Bacterial Spot Cause of disease: 1. The disease is caused by four species of Xanthomonas (X. euvesicatoria, X. gardneri, X. perforans, and X. vesicatoria). In North Carolina, X. perforans is the predominant species associated with bacterial spot on tomato and X. euvesicatoria is the predominant species associated with the disease on pepper. 2. All four bacteria are strictly aerobic, gram-negative rods with a long whip-like flagellum (tail) that allows them to move in water, which allows them to invade wet plant tissue and cause infection.

How to prevent/cure the disease 1. The most effective management strategy is the use of pathogen-free certified seeds and disease-free transplants to prevent the introduction of the pathogen into greenhouses and field production areas. Inspect plants very carefully and reject infected transplants- including your own! 2. In transplant production greenhouses, minimize overwatering and handling of seedlings when they are wet. 3. Trays, benches, tools, and greenhouse structures should be washed and sanitized between seedlings crops. 4. Do not spray, tie, harvest, or handle wet plants as that can spread the disease.

7. ADVANTAGES & DISADVANTAGES

List of advantages

- The proposed model here produces very high accuracy of classification.
- Very large datasets can also be trained and tested.
- Images of very high can be resized within the proposed itself.

List of disadvantages

- For training and testing, the proposed model requires very high computational time.
- The neural network architecture used in this project work has high complexity.

8. APPLICATIONS

- 1. The trained network model used to classify the image patterns with high accuracy.
- 2. The proposed model not only used for plant disease classification but also for other image pattern classification such as animal classification.

3. This project work application involves not only image classification but also for pattern recognition.

9. CONCLUSIONS

The model proposed here involves image classification of fruit datasets and vegetable datasets. The following points are observed during model testing and training:

- The accuracy of classification increased by increasing the number of epochs.
- For different batch sizes, different classification accuracies are obtained.
- The accuracies are increased by increasing more convolution layers.
- The accuracy of classification also increased by varying dense layers.
- Different accuracies are obtained by varying the size of kernel used in the convolution layer output.
- Accuracies are different while varying the size of the train and test datasets.

10. FUTURE SCOPE

The proposed model in this project work can be extended to image recognition. The entire model can be converted to application software using python to exe software. The real time image classification, image recognition and video processing are possible with help OpenCV python library. This project work can be extended for security applications such as figure print recognition, iris recognition and face recognition.

11.BIBILOGRAPHY

- [1]. R Indumathi Leaf Disease Detection and Fertilizer Suggestion", IEEE International Conference on System, Computation, Automation and Networking (ICSCAN), 29-30 March 2019, DOI: 10.1109/ICSCAN.2019.8878781.
- [2]. P. Pandi Selvi, P. Poornima, "Soil Based Fertilizer Recommendation System for Crop Disease Prediction System", International Journal of Engineering Trends and Applications (IJETA) Volume 8 Issue 2, Mar-Apr 2021.
- [3]. H Shiva reddy, Ganesh hedge, Prof. DR Chinnaya3, "IoT based Leaf Disease Detection and Fertilizer Recommendation", International Research Journal of Engineering and Technology (IRJET), Volume: 06 Issue: 11, Nov 2019, e-ISSN: 2395-0056.

APPENDIX

A. Source Code (Jupyter notebook python code)

fruit.ipynb (due to limited page size the code vegetable.ipynb uploaded in github)

#!/usr/bin/env python

coding: utf-8

In[1]: pwd

In[2]: cd E:/IBM_MY_COURSE/Project/Dataset Plant Disease/fruit-dataset/fruit-dataset

Apply ImageDataGenerator functionality to Train and Test set

Preprocessing # In[3]: from keras.preprocessing.image import

ImageDataGenerator train_datagen =

```
ImageDataGenerator(rescale=1./255,shear_range=0.2,zoom_range=0.2,horizonta
1_fli p=True) test_datagen = ImageDataGenerator(rescale=1) # In[4]: pwd
#
                      In[5]:
                                                 x train
train_datagen.flow_from_directory('E:/IBM_MY_COURSE/Project/Dataset
Plant
                                                                 Disease/fruit-
dataset/fruitdataset/train',target_size=(128,128),batch_size=32,class_mode='cate
gorical')
#In[6]:
x_test=test_datagen.flow_from_directory('E:/IBM_MY_COURSE/Project/Datas
        Plant
                   Disease/fruit-dataset/fruit-dataset/test',target size=(128,128),
et
batch_size=32,class_mode='categorical') # # Import the models
#
    In[7]:
                      tensorflow.keras.models
              from
                                                 import
                                                           Sequential
                                                                         from
tensorflow.keras.layers import Dense,Convolution2D,MaxPool2D,Flatten
## Initializing the models 10
# In[8]: model=Sequential()
## Add CNN Layers
#In[9]:
model.add(Convolution2D(32,(3,3),input_shape=(128,128,3),activation='relu'))
# In[10]: x_train.class_indices
## Add Pooling layer
# In[11]: model.add(MaxPool2D(pool_size=(2,2)))
# # Add Flatten layer # In[12]: model.add(Flatten())
## Add Dense Layer
```

```
# In[21]: model.add(Dense(40, kernel_initializer='uniform',activation='relu'))
model.add(Dense(20, kernel_initializer='random_uniform',activation='relu'))
# # Add Output Layer # In[24]: model.add(Dense(6,activation='softmax',
kernel_initializer='random_uniform'))
##Compile the model #
                                    In[25]:
model.compile(loss='categorical_crossentropy',optimizer='adam',metrics=['accur
acy']) # In[26]: len(x_train)
# In[27]: 5384/32
## Fit the Model
#In[28]:
model.fit_generator(x_train,steps_per_epoch=168,validation_data=x_test,validat
ion_st eps=52,epochs=3)
## Save the Model
# In[29]: model.save("fruit.h5")
# In[30]: ls
## Test the Model
#
      In[32]:
                  from
                           keras.preprocessing
                                                   import
                                                              image
                                                                         from
                                                                         from
tensorflow.keras.preprocessing.image
                                          import
                                                      img_to_array
tensorflow.keras.models import load_model import numpy as np
# In[33]: model = load_model("fruit.h5")
## Test Apple_Healthy Class images
```

```
# In[37]: img = image.load_img('E:/IBM_MY_COURSE/Project/Dataset Plant
Disease/fruitdataset/fruit-dataset/test/Apple healthy/00fca0da-2db3-481b-
b98a9b67bb7b105c___RS_HL 7708.JPG',target_size=(128,128)) 11
# In[39]: x=image.img_to_array(img) x=np.expand_dims(x,axis=0)
# In[40]: pred = model.predict_classes(x)
# In[41]: pred
#In[45]:index
=['Apple Black rot', 'Apple healthy', 'Corn (maize) Northern Leaf Blig
ht','Corn_( maize) healthy','Peach Bacterial_spot','Peach healthy']
# In[46]: print('the given image belogs to=',index[pred[0]])
#
       Test
              Apple
                      Black
                               Rot
                                     class
                                            images \# In[54]:
                                                                  img
image.load_img('E:/IBM_MY_COURSE/Project/Dataset
                                                                      Plant
Disease/fruitdataset/fruit-dataset/test/Apple Black_rot/0f3d45f4-e121-42cd-
a5b6- be2f866a0574___JR_FrgE.S 2870.JPG',target_size=(128,128))
# In[55]: x=image.img_to_array(img) x=np.expand_dims(x,axis=0) pred =
model.predict_classes(x) print('the given image belogs to=',index[pred[0]])
## Test Corn Northern leaf Blight class images
# In[56]: img = image.load_img('E:/IBM_MY_COURSE/Project/Dataset Plant
Disease/fruitdataset/fruit-
dataset/test/Corn_(maize) Northern_Leaf_Blight/00a14441-7a62- 4034-bc40-
b196aeab2785___RS_NLB 3932.JPG',target_size=(128,128))
# In[57]: x=image.img_to_array(img) x=np.expand_dims(x,axis=0) pred =
model.predict_classes(x) print('the given image belogs to=',index[pred[0]])
```

```
##
    Test Corn
                Healthy class
                                   images
                                                  # In[58]:
                                                                  img
image.load_img('E:/IBM_MY_COURSE/Project/Dataset
                                                                   Plant
Disease/fruitdataset/fruit-dataset/test/Corn_(maize) healthy/0a68ef5a-027c-
41ae-b227-159dae77d3dd____R.S_HL 7969 copy.jpg',target_size=(128,128))
# In[59]: x=image.img_to_array(img) x=np.expand_dims(x,axis=0) pred =
model.predict_classes(x) print('the given image belogs to=',index[pred[0]]) # #
Test
       Peach
                Bacterial
                                   class
                                          images
                                                         In[60]:
                           spot
                                                                  img
image.load_img('E:/IBM_MY_COURSE/Project/Dataset
                                                                      Plant
Disease/fruitdataset/fruit-dataset/test/Peach Bacterial spot/00ddc106-692e-
4c67-b2e8- 569c924caf49 Rutg. Bact.S 1228.JPG',target_size=(128,128)) 12
# In[61]: x=image.img_to_array(img) x=np.expand_dims(x,axis=0) pred =
model.predict_classes(x) print('the given image belogs to=',index[pred[0]])
# # Test Peach Healthy class images
# In[62]: img = image.load_img('E:/IBM_MY_COURSE/Project/Dataset Plant
Disease/fruitdataset/fruit-dataset/test/Peach___healthy/1a07ce54-f4fd-41cf-
b088- 144f6bf71859 Rutg._HL 3543.JPG',target_size=(128,128))
# In[63]: x=image.img_to_array(img) x=np.expand_dims(x,axis=0) pred =
model.predict_classes(x) print('the given image belogs to=',index[pred[0]])
```