Brain Scan Tumor Classification

Project Examples

Contents

Overview	2
Using the Web Application: Do I Have A Tumor?	2
Run Web Application on Localhost	Ę
Accessing the Model	6

Overview

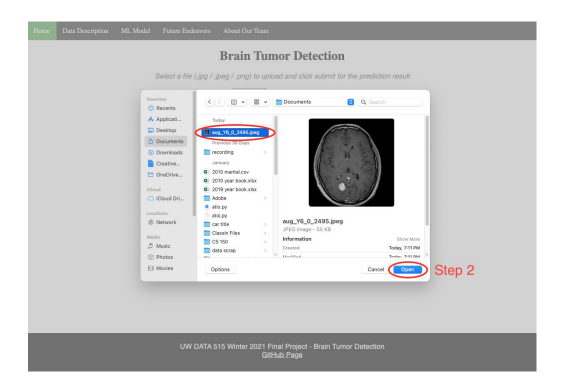
This document provides users with in-depth examples of how to utilize the published web application, how to run the web application on your own local host, and how to access our model directly.

Using the Web Application: Do I Have A Tumor?

The purpose of this project is to decide if a brain is tumorous or not based on an MRI scan input. To utilize the web application you must have an MRI scan image ready (.jpg, jpeg, .png)

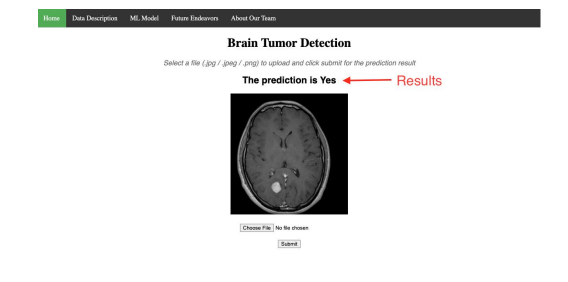
- 1. Navigate to http://doihaveatumor.com/
- 2. Click the 'Choose File' button, and select the 2D brain scan (.jpg, .jpeg, .png supported) from your file explorer





3. Click the 'Submit' button, and wait for the results to display at the top of the page





UW DATA 515 Winter 2021 Final Project - Brain Tumor Detection GitHub Page

 Use the top navigation bar to explore more about the model and our development team

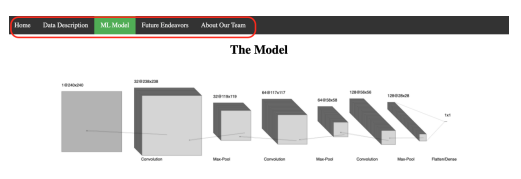


Fig. Illustration of Neural Network Architecture (produced using $\underline{\text{NN-SVG}})$

Our tumor detection model shown above is a 2D CNN with nine hidden layers, and was built using both tensorflow and keras. The input images (data set is described in the "Data Description" tab) were first reshaped into standardized 240x240x1 grayscale arrays using the opency Computer Vision module, they were then normalized through positive global standardization. After being trained on the data, the model predicts if a brain image has a tumor (prediction of 1) or does not (prediction of 0). Below are a few of our model's training parameters and metrics:

Initial Learning Rate: 0.01 Number of Total Training Epochs: 100 Final Validation Set Accuracy: .902

Run Web Application on Localhost

1. In the command line interface run the following command:

git clone https://github.com/aaliyahfiala42/DATA515-Brain-Scan-Classification.git

```
[-bash-3.2$ git clone https://github.com/aaliyahfiala42/DATA515-Brain-Scan-Classification.git Cloning into 'DATA515-Brain-Scan-Classification'... remote: Enumerating objects: 54, done. remote: Counting objects: 100% (54/54), done. remote: Compressing objects: 100% (38/38), done. remote: Total 813 (delta 11), reused 36 (delta 7), pack-reused 759 Receiving objects: 100% (813/813), 121.01 MiB | 19.25 MiB/s, done. Resolving deltas: 100% (398/398), done.
```

2. Install all required packages by navigating to the root directory (using cd

DATA515-Brain-Scan-Classification) and run:

python setup.py install

3. To run the application, navigate to the brain_scan folder (using cd brain_scan)

and run:

python application.py

4. Copy the local host url provided (the http address in the red box) and past the url to a browser of your choice.

```
* Serving Flask app "application" (lazy loading)

* Environment: production

WARNING: This is a development server. Do not use it in a production deployment.

Use a production WSGI server instead.

* Debug mode: off

* Running on http://127.0.0.1:5000/ (Press CTRL+C to quit)
```

Accessing the Model

The model training is provided as a Jupyter Notebook located in the "notebooks" folder under the root directory. To access the model, you must have <u>Jupyter Notebook</u> installed on your local machine, or utilize software such as the <u>Anaconda Navigator</u>, and follow the instruction below using the Anaconda CMD.exe prompt.

1. In the command line interface run the following command:

git clone https://github.com/aaliyahfiala42/DATA515-Brain-Scan-Classification.git

```
[-bash-3.2$ git clone https://github.com/aaliyahfiala42/DATA515-Brain-Scan-Classification.git Cloning into 'DATA515-Brain-Scan-Classification'... remote: Enumerating objects: 54, done. remote: Counting objects: 100% (54/54), done. remote: Compressing objects: 100% (38/38), done. remote: Total 813 (delta 11), reused 36 (delta 7), pack-reused 759 Receiving objects: 100% (813/813), 121.01 MiB | 19.25 MiB/s, done. Resolving deltas: 100% (398/398), done.
```

2. To run the application, in the command line interface run:

jupyter notebook

3. This will then open the Jupyter Notebook interface. Navigate to the notebooks and open the brain_tumor_classification_FINAL notebook. Now you can view, run, and edit the notebook directly.

