Neural Networks for Neural Tumors

Final Project for ECS 171 - Machine Learning at UC Davis, with Professor Setareh Rafatirad

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Abstract—This report documents the process of creating a machine learning model that classifies brain tumors given MRI brain scan imagery. This project saw the creation of three successful models using the following algorithms: Convolutional Neural Network, Support Vector Machine, and Random Forest. Disclaimer: this project is for educational purposes only and not meant for medical use or research.

Index Terms—Brain tumor classification, Convolutional Neural Network (CNN), MRI brain imagery, Random Forest (RF), Support Vector Machine (SVM).

I. INTRODUCTION

As a team of five young computer science students with no prior machine learning experience, we ambitiously decided to tackle the real-world challenge of brain tumor classification. Through hours and hours of research and trial and error, we learned a lot, and we have a lot to share.

We hope this report serves not only as an informative summary of our findings, but also as motivation for your own projects. We faced many challenges during this project, but in the end created something we are all very proud of, despite our lack of experience.

A. Motivation

As of 2025, there are more than 120 different kinds of brain tumors¹. Population statistics gathered by braintumor.org from 2015 through 2019 led researchers to estimate that in 2023, over 94,000 people would be diagnosed with a primary brain tumor, in America alone².

Luckily, 72% of primary brain tumors are benign. While not completely harmless, benign tumors are much less fatal than malignant (cancerous) tumors. However, that still leaves a concerning 28% of tumors that are malignant. Of those 28%, an estimated 65% of cases are proven to be fatal. Among the malignant tumors, glioblastoma is the most common, with a survival rate of only 6.9%. These numbers are tragic, and at the moment, the best way we know to reduce them is through early detection.

Hundreds of thousands of MRI brain scans are performed every day. With the scans being so complex, it can sometimes be hard for even a trained professional to determine whether or not a patient has a brain tumor, and even harder to tell which kind.

A tool that could detect and identify brain tumors within seconds, and with high accuracy, would not only save a lot of time and money, but lives. As mentioned above, early detection is absolutely key to increasing survival rates. Getting more people in and out of MRI machines and having their scans be automatically processed and analyzed would save countless lives. Not to mention the impact such a tool would have on developing nations who might not have as many specially trained doctors that can readily perform MRI analysis. Luckily in the realworld, hospitals are already adopting and developing these ideas.

B. Approach

A classification machine learning model is a great approach to this problem. Using a convolutional neural network and an image dataset, we can train a model to accurately classify brain tumors. With its complex structure, a CNN can detect nuances and patterns in the scans that are otherwise hard to spot.

This project focuses on three general brain tumor classes: glioma, meningioma, and pituitary. Each class contains many of the 120+ tumor types mentioned before. The model will also have a "no tumor" class. The aim for the model was to make predictions with over 95% accuracy.

II. LITERATURE REVIEW

A. Brain Tumor Classification in MRI Image Using Convolutional Neural Network³

In this article from 2021, authors Khan et al. examine convolutional neural network models for brain tumor classification, reporting classification accuracies ranging from 80% to 98%. The authors highlight two primary challenges in training such models: (1) the scarcity of accurately labeled data, which increases training complexity and cost, and (2) inconsistency in image dimensions, given that CNNs require fixed-size input. Their dataset is partitioned into training, validation, and testing sets, serving model training, hyperparameter tuning, and final evaluation, respectively. Image preprocessing is performed using Canny Edge Detection, an opensource computer vision technique. To address the limited dataset size, the authors employ data augmentation methods such as flipping, rotation, and brightness adjustments to enhance variability. The model utilizes the ReLU activation function and stochastic gradient descent for optimization.

Given our limited computational resources, we plan to utilize a pre-trained model. The eight-layer architecture discussed in the paper provides a useful baseline. The preprocessing techniques mentioned are also great starting points for us, and let us know what we should look out for. Furthermore, we intend to implement data augmentation techniques to improve generalization on our small dataset. The use of ReLU and stochastic gradient descent aligns well with our current understanding, and we aim to assess their impact on our model's performance.

B. A Deep CNN Based Multi-Class Classification of Alzheimer's Disease Using MRI⁴

In this article, Farooq, Anwar, Awais, and Rehman demonstrate their implementation of a four-way Alzheimer's disease classifier. Considering their model also takes MRI imagery as input and has four class target labels, their methodology provides valuable insights. The authors began their model development with data preprocessing and a technique known as skull stripping, which is the process of removing non-brain tissues like the skull or fat from MRI imagery. We could implement skull stripping in our model to improve the signal-to-noise ratio and

reduce input complexity, which could help against overfitting.

Additionally, this article also mentions the utilization of data augmentation to increase the size of the dataset: they simply flipped the images along the horizontal axis, which is valid due to the left and right symmetry of the brain. This method grew their dataset from 355 MRI scans to 38,024. We can implement the same method in our project to increase the sample size, and combat overfitting, making our model more accurate.

C. Brain Tumor Classification Using Convolutional Neural Networks⁵

In this publication, authors Seetha and Raja demonstrate the effectiveness of convolutional neural networks for brain tumor classification. Their proposed CNN-based classification approach achieves an impressive 97.5% accuracy rate while maintaining low computational complexity compared to other methods. Seeing such a high accuracy be achieved is both inspiring and motivating for us. Considering their model also predicts four tumor types, their high accuracy rate sets a benchmark we should aspire to, and their focus on computational efficiency aligns with our resource constraints.

The authors implemented a deeper architecture design using small kernels, with neuron weights intentionally kept small to optimize performance. Their experimental results demonstrated that their CNN model outperformed other state-of-the-art methods in both accuracy and efficiency. Like our approach, they used a dataset containing various brain MRI scans and faced similar challenges with image preprocessing. Their optimization of kernel size and weight parameters offers valuable insights for our model development. Given that our initial prototype showed signs of overfitting (93% training accuracy with 42% testing accuracy), we should consider implementing some of the suggested architectural decisions regarding smaller kernels and weight optimizations, while continuing with our planned transfer learning approach to improve generalization.

III. DATASET DESCRIPTION

The dataset used for this project comes from a group of brilliant students at Savitribai Phule Pune

University in India. It is an open source dataset that can be found for free on Kaggle⁶. The dataset is reviewed and well respected. It consists of 3264 labeled MRI scan images. Like hinted at before, the images are divided into four categories: three different tumor types, and one set of "no tumor" scans. Because each image in our dataset is already labeled, our model's training is considered supervised, as opposed to unsupervised. An unsupervised model would require a much different approach, most likely using different algorithms.

A. Preprocessing

Due to the non-numerical nature of image datasets, our input features come in a different form than they normally would with a numerical dataset. Instead of having numerically represented features such as 'age', 'blood pressure', and so on, our dataset only has images. Technically, each image is a set of numbers (pixel intensities), but the idea remains the same. There is no standard feature selection process, or process to choose which features to train the model on. Instead, things like image preprocessing and model hyperparameters are key to training an accurate model.

As explained under the *Literature Review* section, preprocessing steps might include techniques like image resizing, cropping, normalization, data augmentation (rotating, flipping, zooming), and more. Going even further, the 'skull stripping' technique mentioned by Farooq et al.⁴ can be used to isolate the brain tissue in an MRI scan. This is helpful for training a model, since we do not care about the skull or fat in the scans, it is considered noise. We do not want the model to train on those features of the image.

IV. EXPLORATORY DATA ANALYSIS

As seen in the left histogram of *Figure 1* on the next page, there is a relatively even distribution of data among each class in the training set, at around 825 images. However, the no tumor class has significantly less, with 395 images. We suspect this is because the scans without tumors have much less variance. And because of the reduced variance, it should be much easier for the model to detect "no tumor" scans. We did not think the lower number of images would have a negative effect on the model's

ability to classify images as "no tumor", and we were right. However, if we did see poor results in that classification specifically, we could have tried training the model on 395 images for each class to see if that helped with the overfitting of the other classes. On the other hand, we could have performed data augmentation on the "no tumor" class, bringing its number of images up to the 825 range like the other classes. In hindsight, we did not have to apply either of these techniques.

Shifting our focus toward the testing dataset, again looking at *Figure 1*, but this time at the right histogram, we see that there is a much more uneven distribution among each class. This does not affect our model's performance, since it is not being trained on the testing data. However, we thought the results of the pituitary class could be skewed considering the relatively low number of scans, but it was also not a problem. If we had seen inconsitencies, it could have been useful to test against more pituitary imagery to ensure the model can handle more variations of scans. We could again have performed data augmentation to create more pituitary data. Testing against more data could give us more confidence in our model.

Taking a look at *Figure 2*, again on the next page, we can see that the dataset is comprised of images of many varying sizes. There is a concentration around the 200x200 and 500x500 sizes. With this information, we know we will have to perform some image preprocessing to convert all images into the same size so the model has consistent input.

V. METHODOLOGY

A. Feature Selection

As mentioned in the dataset description, we don't have labels for our input features like a numerical dataset would have. We only have our images, which are technically each a set of pixel intensity values, but it means we can't select certain features to train our model on. However, we can still perform image preprocessing and hyperparameter tuning to increase our model's accuracy.

Looking at *Figure 2* again, we see all of our images have different dimensions. As a preprocessing step, we will resize all images to be 224x224 pixels, the conventional CNN input size. The resizing step is not as simple as it sounds though, we must be

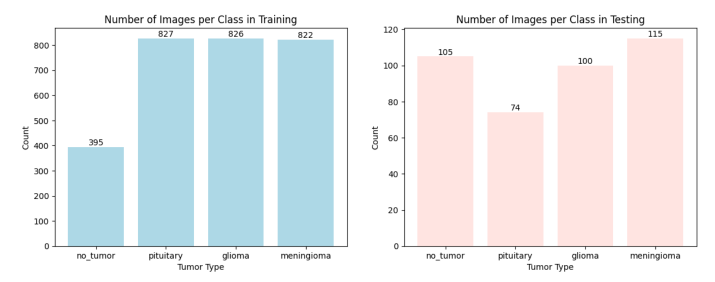


Fig. 1. The dataset has a consistent number of training images across classes, except for "no tumor". This should not be an issue because "no tumor" imagery has much less variance. For testing, the distribution of images is more varied, but this does not affect model performance, since the model is not trained on this data.

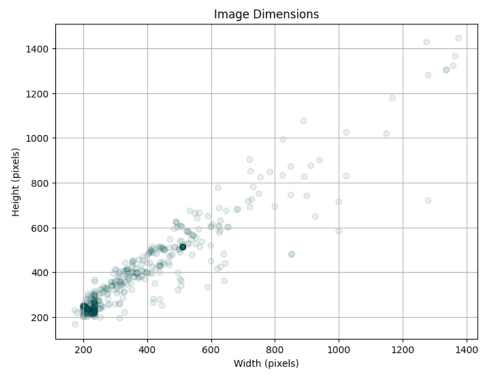


Fig. 2. It is clear to see the images in the dataset are of many varying dimensions. We do see higher densities around the 200x200 and 500x500 dimensions, though. Regardless, such variation means image preprocessing and resizing is necessary before training.

careful not to create unwanted noise in our data. After imagine resizing, we then normalize each pixel's value from its 3-channel RGB form.

The training dataset consists of 2870 MRI scans. If needed, we can perform data augmentation as mentioned previously. We can rotate, flip, and zoom

the images to create 'artificial' data that would hopefully reduce any overfitting and allow for the model to better capture underlying patterns. We also would expect this technique to allow for the model to predict against a larger variety of testing data (i.e. rotated images).

Along with input features, the hyperparameters of a model are crucial to achieving a good fit. The key hyperparameters for the CNN model were: batch size for gradient descent, training epochs, loss function, dropout rate, and a few others. Techniques like grid search can be used for hyperparameter tuning, more on that later.

B. Early Model Development

Our first prototype model was a CNN with four convolutional blocks, followed by fully connected layers using the Keras Sequential API. We trained the model over 20 epochs, with each epoch giving us higher accuracy on the training data, peaking at 93%. However, when performing predictions on our testing data, the model saw a big dip in performance. With an average accuracy of 42%, we knew immediately that our model was heavily overfitting the training data.

To improve upon the first model, we applied k-fold cross validation and tuned hyperparameters such as dropout rate, number of layers, number of neurons, etc. This was the first time we used grid search to tune hyperparameters. It was also at this stage we decided that going forward, we would use transfer learning to reduce the number of learned parameters and implicitly filter irrelevant features. Transfer learning, which is the process of leveraging a pre-trained model to improve your own, can greatly reduce overfitting risks.

VI. EXPERIMENTAL RESULTS

Neural networks like CNNs are very good at detecting features and learning patterns in data. Because of this, we can leverage deep learning to preprocess data, and then use that data to train other models. So far, this report has focused on developing a CNN model with high accuracy by iteratively refining both model architecture and training strategy. While CNNs are the most common algorithm choice for image classification, they are not the only option. Image classification can be done by various

other algorithms, including support vector machine (SVM), and random forest (RF).

While the detailed performance of each model we created will be discussed in the following sections, this section outlines the chronological development and iterative experimentation that lead us to our final models.

A. Baseline CNN Model

As mentioned before, our very first model was built from scratch using TensorFlow's Sequential API. The model consisted of four convolution blocks with subsequent increases in the number of filters (kernels). Each block was followed by a max pooling layer helping reduce the spatial dimensionality. Additionally, each hidden layer of the model was accompanied by a dropout layer to mitigate overfitting. The last hidden layer of the model was a dense layer of 128 neurons using the ReLU activation function. The output layer was four neurons using Softmax as the activation function. Softmax, given its consistent output between 0 and 1, is the standard choice of activation function for the output layer in multi-class classification.

This model was trained over 20 epochs using the Adam optimizer and categorical cross-entropy as the loss function. Again, as mentioned before, the model ended up with a training accuracy of 93% and a testing accuracy of 42%, implying significant overfitting.

B. Transfer Learning with ResNet50

In our next step, we decided to employ transfer learning. We started with the ResNet50 model, a pre-trained model trained on the ImageNet dataset. We leveraged the architecture of ResNet50 as a feature extractor by eliminating its output layer and freezing all other layers. The classification head was built on top of the pre-trained model, which served as a convolutional base. The base was followed by a Global Average Pooling layer to compress spatial dimensions. The classifier was again completed by a fully connected dense layer of 128 neurons with the ReLU activation function and a Softmax output layer. The layer of 128 neurons was accompanied by a dropout rate of 0.3 to help prevent overfitting.

The model was trained over 20 epochs. The transition to transfer learning yielded a significant

improvement in validation accuracy (89% compared to 76% in the first model) while maintaining a reasonable training accuracy of 93%. Motivated by this encouragement, we experimented with selective fine-tuning the model by unfreezing the last 30 layers of the ResNet50 pre-trained model, allowing the weights of those layers to be adjusted during backpropagation. With this adjustment, the model ended up with a training accuracy of 97%, and a testing accuracy of 79%. Although the model was still slightly overfitting, we had made huge improvements. At this stage, we also introduced new training practices such as the learning rate hyperparameter and early stopping, which shaped our training strategy for later experiments.

C. Dataset Reorganization and ResNet101 Upgrade

Motivated by the significant improvement in the previous stage, we had defined that transfer learning would be one of the most critical techniques in developing our CNN model. We decided to try different pre-trained models, from simpler models such as VGG16 and VGG19, to more complex ones in the ResNet and EfficientNet families. Even with either layer-wise fine-tuning or standard fine-tuning techniques applied to those models, we had noticed a common trait: persistent underperformance on the testing test (with an accuracy around 70% to 80%). This discrepancy prompted us to re-evaluate our dataset. Initially, we had used the dataset split provided by Kaggle. We suspected that there might have been an imbalanced presentation across classes or feature distribution in the default split. To resolve this issue, we decided to recombine the training and testing sets with shuffle, then split the entire dataset with a ratio consistent with that of the default split.

With limited available computational resources, we also upgraded our pre-trained model to ResNet101, a more complex model compared to ResNet50. We employed the same structure as in the ResNet50 setup. The model was initially trained over 20 epochs with all ResNet101 layers frozen, yielding a training accuracy of 92% and validation accuracy of 85%. When the fine-tuning technique was applied (with the last 40 layers unfrozen), the model reached 99% training accuracy and 96% validation accuracy after being trained over four additional epochs. The model also performed much

better on the testing data, yielding 97% accuracy with reasonably high F1-scores for each class label. This was our breakthrough.

By employing ResNet101 with layer-wise finetuning and a better dataset split, we ended up with a highly robust CNN model with consistent performance across training, validation, and testing data.

D. Non-deep Learning Models with feature extraction

To develop other classifiers that incorporate algorithms rather than deep learning, we decided to employ Support Vector Machine (SVM) and Random Forest (RF) models. We leveraged the finetuned CNN model as the feature extraction for our selected dataset in order to obtain high accuracy while keeping the training process relatively simple. More specifically, we extracted the output of the GlobalAveragePooling2D layer of the fine-tuned CNN model, which includes 2048-dimensional feature vectors, and that result would serve as the input for each non-deep learning model. In other words, our SVM and RF models were trained on a more semantically meaningful input space compared to raw pixels.

After conducting hyperparameter tuning, we obtained an overall accuracy of 96% and 95% for SVM and RF classifiers respectively. In general, these results demonstrate that classical classifiers, when paired with feature extractors, can deliver highly competitive performance for image classification, with significantly reduced training complexity.

VII. EVALUATION

As previously mentioned, we trained three machine learning models for this project: a CNN, an SVM, and an RF model. In this section, the performances of these models are assessed using standard classification metrics such as accuracy, precision, recall, and F1-score. Accuracy describes the rate of correct predictions. Precision is the measurement of true positives against false positives. Recall measures true positives against false negatives. And F1-score is the harmonic mean of precision and recall.

TABLE I CNN METRICS

Class	Precision	Recall	F1-score	Support
glioma	0.98	0.93	0.95	109
meningioma	0.93	0.97	0.95	105
no tumor	0.97	0.98	0.98	60
pituitary	0.99	0.99	0.99	118

TABLE II SVM METRICS

Class	Precision	Recall	F1-score	Support
glioma	0.98	0.91	0.94	109
meningioma	0.89	0.97	0.93	105
no tumor	0.98	0.97	0.97	60
pituitary	0.98	0.98	0.98	118

TABLE III RF METRICS

Class	Precision	Recall	F1-score	Support
glioma	0.98	0.89	0.93	109
meningioma	0.87	0.98	0.92	105
no tumor	1.00	0.93	0.97	60
pituitary	0.97	0.98	0.98	118

We can see from Tables 1, 2, and 3 that the CNN demonstrated the highest overall accuracy at 97%, followed closely by the SVM at 96%, and the RF at 95%. The F1-scores of each model mirrored this trend, with the CNN achieving an average F1-score of 0.97, the SVM 0.96, and the RF 0.95. These scores indicate that the CNN not only achieves better overall classification performance, but also maintains consistency across classes in terms of precision and recall.

Diving into per-class metrics, again looking at the three tables, we observe that for the label "glioma", all three models had an extremely high precision of 0.98, indicating very few false positives. In terms of recall, the CNN exhibited a score of 0.93, the SVM had a lower recall at 0.91, and the RF had the lowest recall at 0.89. These values highlight that both the SVM and the RF have higher false negative rates compared to the CNN for the "glioma" class, meaning they classify glioma samples as other than glioma.

In the case of the "meningioma" class, while the CNN had the best precision at 0.93, the SVM and

the RF had equal or superior recalls at 0.97 and 0.98 respectively, with the CNN at 0.97. The SVM and the RF had reduced precision scores at 0.89 and 0.87, which is a notable difference from the CNN's aforementioned score. This pattern indicates that the SVM and the RF saw more false positives, but equal or less false negatives. So if the goal was to reduce the number of "meningioma" tumors classified as something else, the RF would be favorable by that 1% difference we see in the recall scores.

In classifying "no tumor", the RF achieved perfect precision of 1.00, which means that every time RF predicted "no tumor", the model was correct. However, its recall at 0.93 was the lowest among the three, pointing to a higher rate of false negatives. The CNN balanced both precision and recall well, at 0.97 and 0.98 respectively, outperforming both alternatives in overall reliability for this class.

For the label "pituitary", all three models performed strongly, with the CNN leading both in precision and recall both at 0.99. The SVM and RF followed closely with both metrics at 0.98 and 0.97-0.98, respectively, indicating that all models are

highly effective in classifying "pituitary" tumors.

Unlike accuracy, which can be misleading if one class dominates the dataset, the F1-score provides a balanced view of the model's ability to correctly identify each class without favoring either false positives or false negatives. This makes it especially useful in medical imaging applications, where both types of errors can have serious implications for patient diagnosis and treatment.

Because the CNN is the best general model of the three, we will take these next few paragraphs to analyze its F1-scores and their meanings. The model's F1-score of 0.95 for glioma tumors indicates that while the model performs well in identifying most glioma cases, there is a notable trade-off between precision and recall. The reduced recall value suggests the model may occasionally misclassify glioma tumors as other types or miss them altogether.

The "meningioma" class also has an F1-score of .95, indicating a very similar story, except this time in reverse. The model has a lower precision compared to recall, meaning the model tends to classify tumors as "meningioma" when in fact they are not. This could be due to visual overlap with other tumor types, particularly gliomas, or due to greater variability in meningioma imaging.

The F1-score of 0.98 for the "no tumor" class highlights the model's strong performance in distinguishing healthy scans from non-healthy ones. The high precision and recall underlying this score indicate that the model rarely misclassifies healthy brains as having a tumor (false negatives) and is also unlikely to miss the presence of a tumor (false positives). This reliability is crucial in medical settings, where a false alarm can lead to unnecessary procedures, and a missed tumor can be life-threatening.

With the highest F1-score of 0.99, the model shows excellent performance in detecting pituitary tumors. This suggests that the "pituitary" class is the most distinct or consistently represented in the dataset, allowing the model to make confident predictions. High scores for both precision and recall indicate that pituitary tumors are both correctly identified and rarely confused with other classes. This robustness may stem from more uniform imaging patterns or stronger contrast features in this tumor type.

VIII. CONCLUSION

The above evaluation highlights the convolutional neural network's strength in handling complex, high-dimensional image data, likely due to its hierarchical feature extraction capability. Its superior balance in both precision and recall in all tumor types supports its suitability for real-world clinical deployment, where false positives and false negatives carry significant risk.

The support vector machine model performed comparably well, especially in the detection of meningioma and pituitary tumors, and may offer a lighter alternative in environments with limited computational resources. However, it showed reduced precision in some classes, which made it slightly less robust overall.

The random forest model, while achieving remarkable precision, particularly for the "no tumor" class, exhibited reduced recall across multiple classes. This suggests a tendency to make fewer false positives, at the expense of missing actual tumor cases, which, as said before, is a very serious and life-threatening issue in medical diagnostics.

Simply because of the extreme high-stakes nature of brain tumor classification, none of these three models we presented should be used in real-world medical diagnosis or research. However, if you had to choose one, the CNN is the clear winner. It was expected from the start given the reputation of CNNs as image classifiers, but it must be said that the results of the SVM and RF models were much better than we expected.

IX. OUR TEAM

Thank you for reading our paper. We had a great time researching, engineering, and writing up this project. The link to our GitHub¹⁰ repository can be found in the references of this report. The repository holds our source code, the dataset, this report, and some more supplementary files. Please feel free to fork the repository and mess around with it as you wish.

A. Demonstration

For those interested in seeing the model in action, a short five minute demonstration of our model can be found on YouTube¹¹. The demo briefly walks

through how to use the model, how the frontend was developed, and how the model is invoked.

B. Roles and Contributions

Joe: As the team leader, Joe took charge of logistics. He organized meetings, defined the roadmap for the project, and kept the team on track. He also led the exploratory data analysis subproject, and was the main author of the final report.

Nam: Throughout the project, Nam was primarily responsible for the development and implementation of the machine learning models. Nam took over the experimental process of optimizing each model, researching architectures and frameworks to successfully develop models with different algorithms. In general, Nam's contributions encompassed every stage of model development, from initial prototyping to completion.

Jonathan: Through literature review and careful research, Jonathan laid the groundwork for the project, finding out which techniques might be useful to us. Jonathan was also responsible for writing the evaluation section of the final report, analyzing the performances of each model by talking about the evaluation metrics such as F1-scores, precision, and recall.

Richard: Richard contributed to the project by assessing the performance of the three machine learning models. Based on these analyses, he provided the rationale for selecting the CNN as the final model. Additionally, he conducted comprehensive literature review to summarize key developments in the field which served as a foundational reference for the project.

Aryan: Aryan participated in group meetings, worked on literature reviews, and helped the group brainstorm ideas and find the dataset. Most importantly, he built and deployed the fullstack applica-

tion, and recorded the demonstration video to go along with it.

C. Project Timeline / Roadmap

- April 19: First team meeting, finding a dataset
- April 25: One-pager project description
- May 01: Complete EDA
- May 02: Initial model (42% accuracy)
- May 06: Mid-Quarter Report
- May 15: Model improvements (80% accuracy)
- May 18: Dataset reorganization (95% accuracy)
- May 20: Began drafting final report
- May 22: SVM and RF models implemented
- May 24: Frontend developed
- June 07: Final report complete

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