

$\begin{array}{c} {\bf Glioma\text{-}meningioma\ tumor\ classification}\\ {\bf on\ MRI\ scans} \end{array}$

Written By:Nikolaos Mouzakitis

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1 Introduction

Brain tumors are among the most complex and dangerous diseases affecting the central nervous system, with over 300,000 new cases diagnosed globally each year. Early and accurate diagnosis is crucial to initiate effective treatment and improve patient survival rates. Magnetic Resonance Imaging (MRI) has become the gold standard for detecting and characterizing brain tumors due to its high resolution and soft-tissue contrast, enabling detailed visualization of tumor morphology and location.

Meningiomas and gliomas are two common types of primary brain tumors with distinct histological origins and clinical behaviors. While meningiomas are typically slow-growing and benign, gliomas are often aggressive and malignant, requiring radically different therapeutic approaches. Manual differentiation by radiologists, however, can be time-consuming and prone to inter-observer variability due to the subtle visual differences in early-stage tumors.

To address these challenges, an automated classification system leveraging machine learning could serve as a decision support assisting tool, reducing diagnostic workload, increasing consistency, and tackling human bias. Such systems not only streamline clinical workflows but also provide a framework for future research into AI assisted diagnostics in neuro/oncology and medicine application domains in general.

In this project, has been attempted to develop and evaluate a machine learning-based system to classify MRI scans into meningioma or glioma categories using a hybrid feature extraction approach. By combining domain-inspired radiomic features (e.g., texture, shape, and intensity statistics) with frequency-domain features derived from Fourier analysis and customengineered features, both spatial and spectral tumor characteristics are captured. Two models—a Random Forest classifier (1,000 trees with feature randomization) and a nonlinear Neural Network (MLP with *100-50 hidden neurons*)— are trained and compared to identify the optimal balance between performance and interpretability for clinical deployment.

2 Related Work

In [4], authors review the usage of AI based radiomics and radiogenomics in glioma, while emphasizing in their roles in diagnosis, treatment response prediction and understanding tumor heterogeneity. Also the challenges in standardizing feature extraction and analysis methodologies are addressed.

Duron et al. [3], in their research proposed a radiomics-based classification model for distinguishing gliomas from meningiomas by utilizing T1weighted MRI scans. The author's approach involved extraction of a vast set of radiomic features which was then followed by the application of machine learning techniques for training and validating a classifier. Results from this particular study demonstrated high diagnostic performance, showing the potential of data-driven radiomics for the support of non invasive tumor characterization. This work can serve as a foundational reference for MRI based tumor classification and can support development of automated systems that may offer assistance in clinical decision-making, similar to the goals of the current project.

Li and co-authors[5], in their work created radiomic models for the prediction of meningioma grade and Ki-67 index, by integrating clinical and radiological features. The models used, demonstrated the potential of radiomics in assessing the biological behavior of meningiomas.

3 Hw and Sw Requirements

The software stack used to implement the classification system consists of Python libraries such as SimpleITK, OpenCV, scikit-learn, pandas, PyRadiomics, matplotlib, seaborn. The code and the MRIs used for this project report are available in the following repository [2].

3.1 Data Details

MRIs were sourced from [1], which contains 7023 MRIs of human brain, divided in 4 categories: *glioma - meningioma - no tumor and pituitary*. For this report's purpose the first two categories are utilized, and selected 1000 MRI scans from both *glioma* and *meningioma* classes.

3.2 Method

Related to the methodology of the classification system, for the preprocessing step all images are loaded using SimpleITK and converted into NumPy arrays and their respective pixel intensities are normalized into the range of grayscale images ([0, 255]). In the next step a 10-pixel masking takes place, in order to reduce the black surrounding areas appearing in every MRI. This border mask is applied and excludes irrelevant regions at the edges of the images. The number of the extracted features can be divided into three subcategories:

- 1) Features acquired from Pyradiomics: by utilization of PyRadiomics first-order statistics (mean, variance, entropy) and texture features (GLCM, GLRLM, GLSZM) are extracted.
- 2) Custom Features: have been designed and implemented in order to intuitively help detecting a tumor alike object on an MRI. (intensity_skewness, intensity_outlier_score, high_intensity_area, max_circularity, top3_circularity_mean, solidity_outlier, abnormal_area_ratio, circular_area_score, asymmetry_score,

asymmetry_outlier, boundary_sharpness_mean, boundary_sharpness_max, boundary_sharpness_outlier).

3) Frequency Domain Features: Energy, entropy, mean, standard deviation and skewness in low, mid, and high-frequency bands using FFT.

The frequency feature extraction module analyzes an image's frequency content by decomposing its Fourier spectrum in three frequency bands: low, mid, and high frequencies. In the first place, the image is converted to grayscale (in the case it was an RGB one), and its 2D Fast Fourier Transform (FFT) is computed. The frequency spectrum is then partitioned into concentric circular bands centered around the zero frequency, with radius scaled adaptively to the image dimensions (20%, 20–60%, and 60–100% of the maximum frequency). For each one of the bands, five statistical features have been extracted: energy (total squared magnitude), entropy (spectral uniformity), mean intensity, standard deviation and skewness (asymmetry of the distribution). These utilization of these features characterize the image's texture and structural patterns This way is handling possible variations of the image sizes due to the adaptive band scaling.

In total, 139 features were examined, which is the combination of the three previous described categories. By conducting the feature extraction process, min-max normalization is performed in all the generated feature values mapping them on the [0, 1] range as we can observe in Figure 1. This have been performed as a necessary step before training machine learning models to help the models generazibility and for models skewness avoidance.

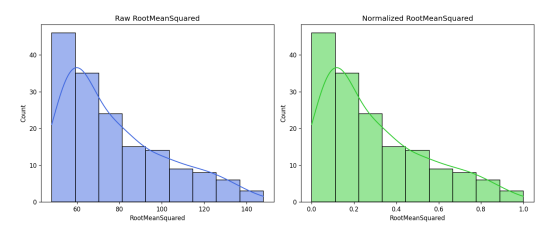


Figure 1: Comparisson of feature distribution prior and after the *min-max* normalization.

4 Evaluation

In the classification stage, two different classifiers have been trained and evaluated, a Random Forest classifier and a MultiLayer Perceptron Neural Network. Classification evaluation examined using different configurations of features in order to be able to compare the contribution of each of the feature sets in the classification accuracy and estimate the degree of their contribution.

The Random Forest model, an ensemble based method known for its robustness and interpretability, was trained using 100 decision trees and a fixed random seed for ensuring reproducibility. For the second classifier, the neural network model was configured having two hidden layers comprising 100 and 50 neurons respectively, trained for up to 500 iterations.

4.1 Classification with texture, shape and statistical features

Classification evaluation with texture, shape and statistical features results are presented below.

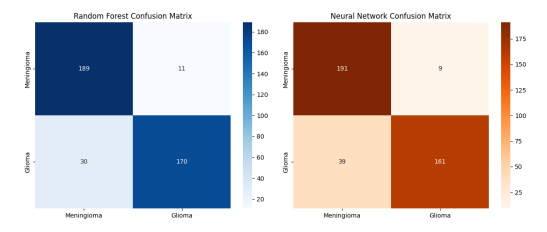


Figure 2: Metric results.

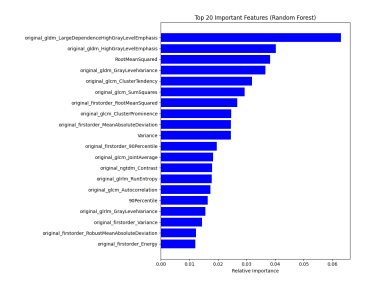


Figure 3: Top20 features of RF.

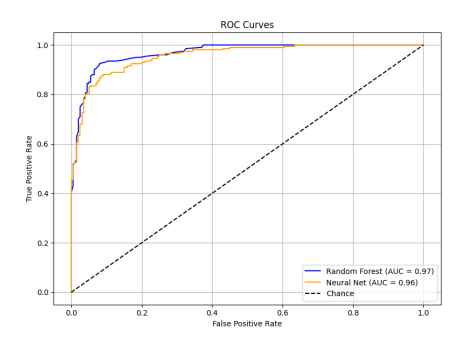


Figure 4: RoC curves for RF and MLP-nn.

Training Rando	m Forest			
Random Forest Accuracy: 0.89 AUC: 0.898				
Classification	Donast.			
Classification	precision	recall	f1-score	support
Θ	0.86	0.94	0.90	200
1	0.94	0.85	0.89	200
accuracy			0.90	400
macro avg	0.90	0.90	0.90	400
weighted avg	0.90	0.90	0.90	400
Training Neura	l Network			
Neural Network Accuracy: 0.88 AUC: 0.880				
Classification	Report:			
	precision	recall	f1-score	support
Θ	0.83	0.95	0.89	200
1	0.95	0.81	0.87	200
accuracy			0.88	400
macro avg	0.89	0.88		400
weighted avg	0.89	0.88	0.88	400

Figure 5: Classification report

4.2 Classification with frequency features

Classification evaluation's results using the extracted frequency features are presented in the next figures.

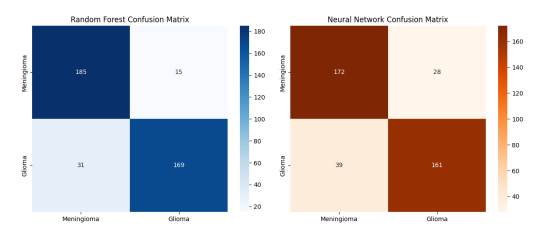


Figure 6: Metric results.

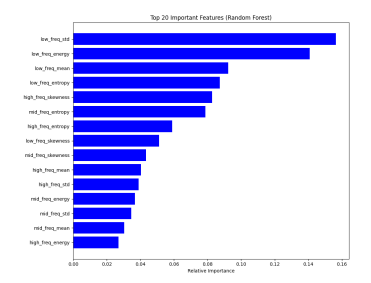


Figure 7: Top20 features of RF.

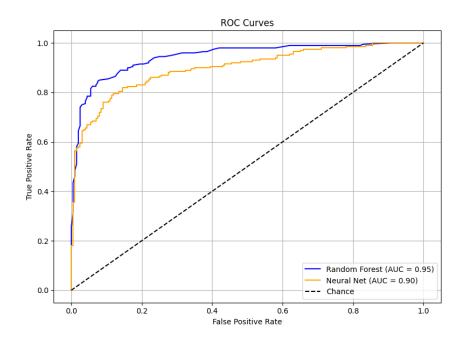


Figure 8: RoC curves for RF and MLP-nn.

Training Random Forest Random Forest Results: Accuracy: 0.885 AUC: 0.885					
Classificatio	n Report: precision	recall	f1-score	support	
0 1	0.86 0.92				
accuracy macro avg weighted avg	0.89 0.89		0.88	400 400 400	
Training Neural Network Neural Network Results: Accuracy: 0.833 AUC: 0.833					
Classificatio	on Report: precision	recall	f1-score	support	
0 1	0.82 0.85		0.84 0.83		
accuracy macro avg weighted avg	0.83 0.83	0.83 0.83			

Figure 9: Classification report

4.3 Classification utilizing all features

Classification results utilizing all the available features extracted are presented in following figures.

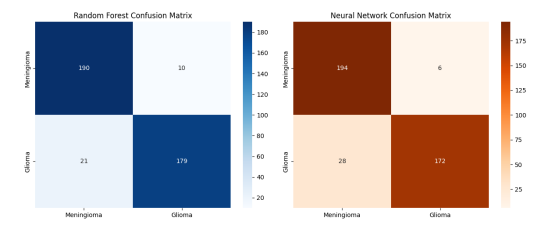


Figure 10: Metric results.

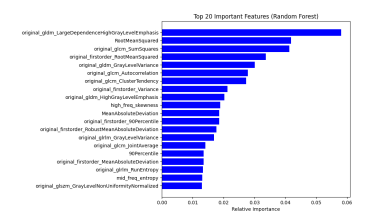


Figure 11: Top20 features of RF.

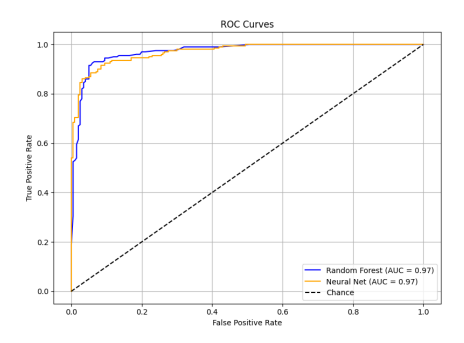


Figure 12: RoC curves for RF and MLP-nn.

Training Random	Forest		j	
Random Forest R Accuracy: 0.922 AUC: 0.922				
Classification				
р	recision	recall	f1-score	support
0 1	0.90 0.95	0.95 0.90		
accuracy			0.92	400
macro avģ	0.92	0.92	0.92	400
weighted avg	0.92	0.92	0.92	400
Training Neural	Network			
Neural Network I Accuracy: 0.915 AUC: 0.915	Results:			
Classification I p	Report: recision	recall	f1-score	support
0 1	0.87 0.97	0.97 0.86	0.92 0.91	
accuracy				400
macro avg	0.92			
weighted avg	0.92	0.92	0.91	400

Figure 13: Classification report

As shown in the follwing plot, both the Random Forest (RF) and MLP neural network (MLP-NN) models exhibit improved classification accuracy as the size of the MRI dataset increases. Initially, with small sample sizes (100–400 images), the models perform similarly, with MLP-NN slightly outperforming RF. An observation to note is the performance spike of Random Forest at 500 samples, reaching 92%, while MLP-NN temporarily dips to 87%. This can indicate that the RF model benefits more from mid-sized datasets, possibly due to its ensemble nature. Conversely, the MLP-NN may have required different hyperparameters or regularization to generalize effectively at that specific size. As the dataset grows further (800–2000 samples), both models recover and show strong accuracy, with MLP-NN gradually surpassing Random Forest, particularly at 2000 samples where it achieves 91.5% vs 92% for RF.

Overall, while RF demonstrates stable performance even with moderate data, MLP-NN shows greater potential at larger scales.

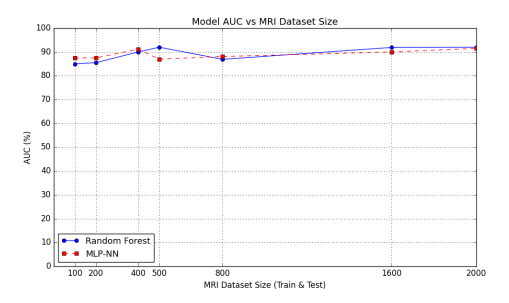


Figure 14: AUC against combined train/test MRI size.

4.4 Evaluation Findings

The results from the three feature configurations demonstrate that combining all available features (texture/shape/statistical and frequency domain features) yields the highest classification performance. Key observations include:

• Feature set comparison:

- All features achieved the best results, underscoring the complementary nature of radiomic, custom-designed, and frequency-domain features.
- Texture/shape/statistical features alone provided competitive performance, suggesting and confirming the strong discriminative power they yield for tumor classification.
- Frequency features were slightly less effective in isolation but contributed meaningfully when integrated with other features.
- Glioma classification improvement: Misclassification rates for glioma dropped significantly (from 31 out of 200 MRIs (15.5%) using texture/shape features only to 21 out of 200 MRIs (10.5%) and from 38 out of 200 MRIs (19%) using texture/shape features only to 28 out of 200 MRIs (14%)) when all features were utilized. This highlights the importance of hybrid feature engineering for capturing tumor heterogeneity.

• Model performance: The Random Forest classifier consistently outperformed the MLP neural network across all configurations, particularly in terms of interpretability (e.g., feature importance rankings) and robustness (AUC metrics).

4.5 Classifier Optimization

At this point, hyperparameter optimization is explored in order to achieve better classification results utilizing the created models. In our workflow, first a Random Forest classifier was utilized with hyperparameter tuning to identify the 30 most important features from the dataset. The hyperparameters such as the number of trees (n_estimators), tree depth (max_depth), minimum samples per split and leaf (min_samples_split, min_samples_leaf), and others are optimized using randomized search with cross-validation.

This ensures that the Random Forest model is tuned for the purpose of capturing the most relevant patterns in the data but also for preventing overfitting. After determining the feature importances, the top 30 features are selected as inputs for the next step. These selected features are then fed into a neural network (MLPClassifier) with fixed hyperparameters (e.g., activation function, solver, learning rate, and number of iterations) for performing the final classification. Using the best subset of features reduces dimensionality and noise, and can improve the neural network's performance and training efficiency, while benefiting from complementary strengths of both models for robust classification.

The models who yielded the best performance, have also been modified from their previous configuration. For the Random Forest, the decision trees we increased by a factor of ten to 1000 decision trees, each using the Gini impurity criterion for spliting nodes and a random subset of 15 features per split to ensure diversity and reduce overfitting. In addition, the Neural Network remained a two layer MLP with 100 neurons in the first hidden layer and 50 in the second, activated by ReLU non-linearity. It uses the L-BFGS optimizer, a memory-efficient gradient based method suitable for smaller datasets.

Results from tuned-modified models in this way are presented below.

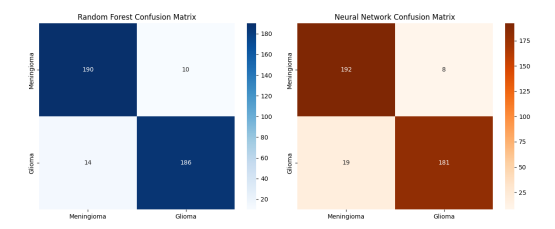


Figure 15: Metric results.

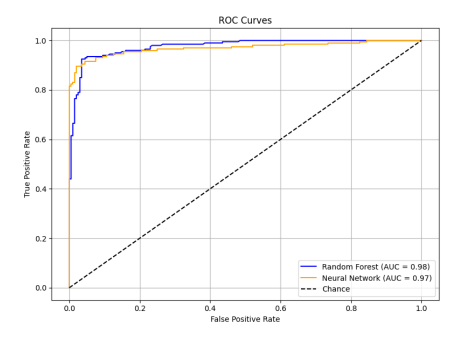


Figure 16: RoC curves for RF and MLP-nn.

After hyperparameter tuning, the Random Forest classifier's accuracy improved from 0.922 to 0.94, indicating a notable gain of approximately 1.8%. This improvement suggests that optimizing parameters like the number of trees, tree depth, and feature selection/reduction helped the model to better capture patterns in the data, resulting in more accurate predictions.

In a similar way, the neural network 's accuracy increased from 0.915 to 0.932, a gain of about 1.7%. This demonstrates that tuning hyperparameters such as the network architecture, learning rate, and using only the top 30

features selected by Random Forest's model allowed better generalization on the test data.

In overall, both models benefited from hyperparameter optimization, with the Random Forest showing a slightly higher accuracy gain. Improvements like these highlight the importance of fine tuning model settings in order to achieve maximization in classification performance, even in cases of starting from already strong baseline models.

As for the model agreement, a comparative analysis plot in Figure 16, shows that both classifiers largely agree and perform well together, with most samples correctly identified by both. Each model also has some unique correct predictions, demonstrating by this their complementary strengths. More strategies could be employed (f.e voting etc) that could improve an overall system's performance. There is a relatively small number of samples misclassified by both models(11/400) samples, highlighting area for further improvements for this project.

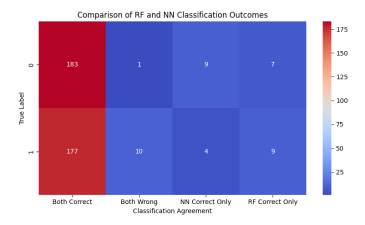


Figure 17: RoC curves for RF and MLP-nn.

5 Histogram Matching enchancement

Analysis of the histograms of the MRIs on the dataset is performed and is presented below. Results of analysing the intensities of the MRIs per category (Figure 17) demonstrate that unnormalized intensity distributions introduce scanner or acquisition protocol-dependent variability that could dominate the signal of interest. Via the application of histogram matching technique image intensities should be aligned on the intensity distributions while preserving the relative contrast relationships essential for the exact tumor classification.

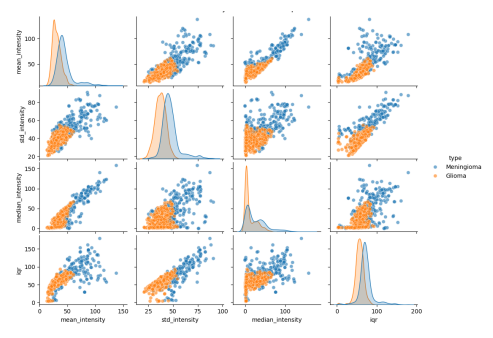


Figure 18: Analysis plot of all the MRI histograms.

Also in Figure 18 we can observe graphical the distribution intensities of the MRIs for both classes.

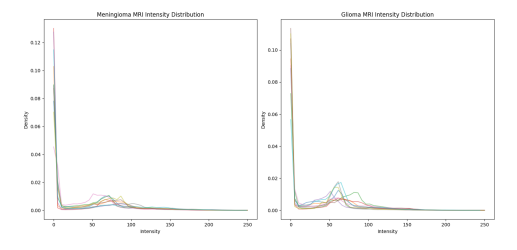


Figure 19: Intensity distributions of MRI histograms.

5.1 Reference image selection

For the process of the selection of an optimal reference image for histogram matching, MRIs have been systematically evaluated from the dataset (sampling 300 candidates by default) and choose the one that maximized class

separability after histogram matching. Each candidate reference image, is tested on a subset of images (20 per class) by performing histogram matching and then calculating a separability score based on the difference in mean intensities between glioma and meningioma classes normalized by their combined standard deviation. The candidate that produces the highest score (indicating by that the best separation between classes after histogram matching) is selected as the final reference image. By following such an approach chooses a reference that maintains discriminative intensity characteristics between the two tumor types while using a single unbiased reference for all images.

The effect of histogram matching on 500 sampled MRIs belonging to meningioma category and 500 sampled MRIs belonging to glioma category is visualized in Figure 19.

```
=== Pre-Histogram Matching Intensity Analysis ===
[Pre-HM] Class 0 Intensity Stats:
    Mean: 44.75 ± 16.10
    Std: 47.41 ± 9.15

[Pre-HM] Class 1 Intensity Stats:
    Mean: 30.61 ± 7.49
    Std: 37.33 ± 5.54

=== Unbiased Histogram Matching ===
[Post-HM] Class 0 (Meningioma) Intensity Stats:
    Mean: 89.41 ± 4.22
    Std: 62.45 ± 3.89

[Post-HM] Class 1 (Glioma) Intensity Stats:
    Mean: 93.91 ± 6.49
    Std: 57.95 ± 6.71
```

Figure 20: Intensity metrics of 1000 MRIs prior and after the application of histogram matching.

Prior to histogram matching, there was a greater difference in intensity statistics (mean and standard deviation) between the two classes. Meningioma MRIs exhibited a higher mean intensity (44.75 \pm 16.10) and standard deviation (47.41 \pm 9.15) compared to glioma MRIs (30.61 \pm 7.49 mean; 37.33 \pm 5.54 std), indicating class-specific variations in brightness and contrast. After performing histogram matching, the intensity distributions of both classes are aligned more closely to a common reference, resulting in similar mean intensities (89.41 \pm 4.22 for healthy, 93.91 \pm 6.49 for tumor) and reduced variation in standard deviation between the classes (62.45 \pm 3.89 vs. 57.95 ± 6.71). This transformation effectively normalized intensity-related differences, allowing downstream radiomic feature extraction and classification models to focus more on structural and textural patterns.

5.2 Evaluation of classification with histogram matching

Utilizing the histogram matching with the previously described reference image selection method, and by extracting all the features, classification has been re-evaluated. Results of using all the available features extracted from histogram matched MRIs are presented in following figures.

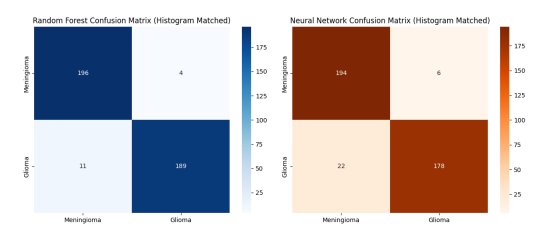


Figure 21: Metric results.

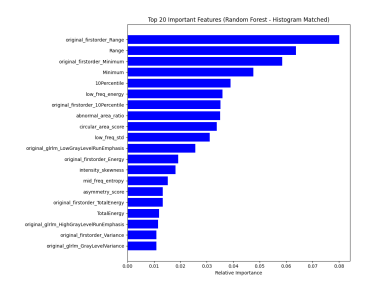


Figure 22: Top20 features of RF.

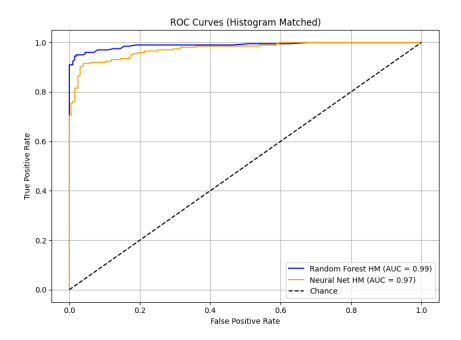


Figure 23: RoC curves for RF and MLP-nn.

Training Random Random Forest R Accuracy: 0.963 AUC: 0.962				images	
Classification p	Report: recision	recall	f1-score	support	
0 1	0.95 0.98	0.98 0.94	0.96 0.96	200 200	
accuracy macro avg weighted avg	0.96 0.96	0.96 0.96			
Training Neural Network on histogram-matched images Neural Network Results (Histogram Matched): Accuracy: 0.930 AUC: 0.930					
Classification Report: precision recall f1-score support					
0 1	0.90 0.97	0.97 0.89		200 200	
accuracy macro avg weighted avg	0.93 0.93	0.93 0.93	0.93 0.93 0.93	400 400 400	

Figure 24: Classification report

Histogram matching with an optimized reference image improved classification accuracy by 4.2% (RF) and 1.5% (MLP-nn), suggesting that the intensity normalization enhances model generalizability. The larger gain was

observed in the Random Forest and by observing the figure refering to the top 20 features contribution for the RF-classification we can inspect that four (20%) of the custom extracted features are included in the list.

6 Conclusion

In this project has been attempted to create an application of an automated system for classifying gliomas and meningiomas using MRI scans, utilizing a combination of radiomic, custom-designed, and frequency-domain features. The Random Forest and Multi-Layer Perceptron classifiers demonstrated strong performance, with the Random Forest achieving slighter superior classification accuracy.

The integration of all feature types yielded the highest classification accuracy, suggesting a complementary nature of texture, shape, statistical, and frequency-domain features. Fine-tuning the models led to improvements, underscoring the importance of parameter selection from all the categories for maximizing performance. In addition, intensity normalization via histogram matching further enhanced model generalizability, reducing scanner dependent variability and improving the outcomes.

References

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