Automatic Tumor Grade Classification From MRI

Kelvin Zhang UID: 304569586 Email: k.zhang661@gmail.com Shalini Dangi UID: 304465145 Email: shalinidangi@ucla.edu

Abstract—In this paper, we discuss methods by which prostate cancer can be detected automatically from MRI images using machine learning techniques. Specifically, we explore the results of testing various ML models on T2-weighted images to classify cancerous regions.

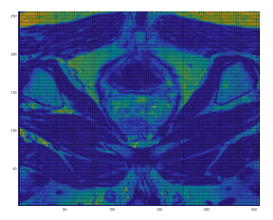
I. Introduction

Prostate cancer (PCa) is one of the most common types of cancer in men, and manifests in the form of malignant growths on the prostate gland. The current standard for PCa diagnosis is transrectal ultrasound biopsy, however manual physician diagnoses is subjective and difficult to standardize and test for errors. MRIs represent a faster, non-invasive alternative and existing research has been directed at automated, software based solutions to prostate cancer detection that are faster, more scalable, and less prone to error than human processing methods. This project will focus on applying machine learning techniques to T2-Weighted MRIs, one of the main parameters in multiparametric MRI analysis, in order to detect the presence and location of PCa growths.

II. METHODS

The data set used for this project consisted of T2-weighted MRI images and corresponding mask images showing cancerous, prostate, and non-prostate regions for 62 different patients, each with varying levels of cancerous tissue (Figure 1). Masks in this data set were manually contoured by a clinician and serve to mark the cancerous pixels shown in the MRI image. In order to provide our models with sufficient classification data, a feature vector was extracted from each pixel in each T2 image. Given a particular pixel, its feature vector consists of the pixel's immediate neighbors within an N-pixel radius (Figure 2). Once extracted, the labeled feature vectors were then used to train supervised learning models provided by SciKit-Learn, which were then tested using k-fold cross validation.

This project explored the following models: random forest classification, AdaBoost, gradient boosting, k-nearest neighbors, decision tree classification, and Multilayer Perceptron (neural network) classification. Additionally, experimental adjustments that we explored in pursuit of optimal performance included feature vector size variation, feature vector pruning, selective pruning of patients, prediction methods after model fitting (binary vs. probabilistic), and threshold variation. Feature vectors were varied by radius N (Figure 2) to determine



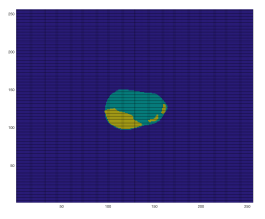


Fig. 1. Example T2-weighted image (top) and corresponding mask (bottom) for patient 1. Mask colors represent 0 - blue, 1 - green, 2 - yellow for non-prostate, prostate, and cancerous prostate regions, respectively. Note that the quality of scans and masks varies greatly among patients.

the vector size yielding the best classification results. The most successful vector size resulting in the highest model evaluation scores was determined to fall within the range of N=4 to N=6. In addition to varying size, feature vectors were also pruned to create a more balanced, relevant data set. This was attempted in three different ways: 1) feature vectors associated with all 0s in the mask (represented a region completely outside of the prostate) were discarded, 2) feature

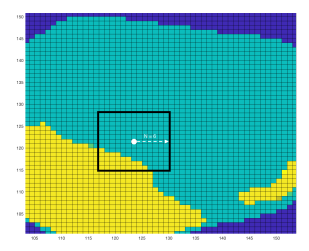


Fig. 2. Illustration of feature vector consisting of all of the pixels within the black box with the center pixel designated by a white circle. The size of the patch is determined by N. (In this example, N=6.) Feature vectors are discarded if the center pixel corresponds to a 0 in the mask (marked by a dark blue color) as this indicates a region outside of the prostate.

vectors associated with at least one 0 in the mask (contained some region outside of the prostate) were discarded, and 3) feature vectors with a center mask value of 0 (centered around a non-prostate region) were discarded. Ultimately, the third pruning attempt was found to be the most successful. Next, after viewing the T2/mask sets for each patient, we discovered large variations in clarity and contrast within the images. As a result, we attempted to selectively prune patients from the set in order to produce a more uniform data set. However, after training models on this narrowed data set, the resulting scores showed little difference from previous runs utilizing all 62 patients. This is perhaps a result of manual pruning without a clearly defined threshold for selection. Thus, this experimental adjustment was discarded. After fitting the model to the test set, we obtained the predicted probability matrix, indicating the probability of a positive/negative prediction according to the model, and used it to plot the receiver operating characteristic (ROC) and precision-recall (PR) curves.

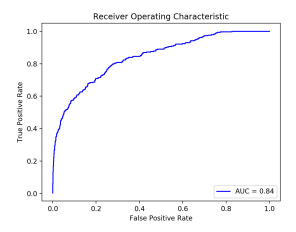
III. EVALUATION

The main metrics we used to evaluate the effectiveness of our models were ROC-AUC and PR-AUC. We selected ROC-AUC since we have an extremely unbalanced data set where there are many more non-cancerous pixels compared to cancerous ones, and being insensitive to class balance would yield reliable measurements regardless of patient or feature vector pruning. Also, since our goal is to be able to compare different learning models, ROC-AUC is useful because it considers all operating points instead of focusing on a particular one.

PR-AUC was used because unlike the ROC curve, it also takes into account the fraction of true positives among positive predictions. This is means that it depends on the frequency of the positive class, in our case cancerous pixels, which could give us a better representation of the differences in our

algorithms given that our positive class is not only much rarer but also much more interesting than the negative case. For our application, the true negative does not offer much value and we would much rather extract more information about the recall of the models, Failing to detect a tumor would be disastrous compared to detecting healthy tissue. The precision recall tradeoff would be essential to selecting the final model for actual application as it should prioritize high recall and the ability to detect all of the cancerous regions, but also demonstrate the compromise of precision so as to not waste expensive and labor-intensive medical resources.

To perform the actual cross validation of our models, we took two difference approaches. We first used leave-one-out (LOOCV) using a single patient as the test subject, but that led to large variance in our diagnostic statistics between patients. We then decided to opt for k-fold cross validation since it mitigates this, in exchange for being slightly pessimistically biased in terms of performance.



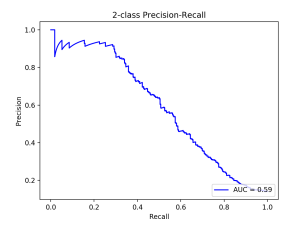


Fig. 3. Receiver Operating Characteristic curve for GBC with n=6 (top), Precision Recall curve for GBC with n=4 (bottom). These were the best performing models for their respective diagnostic statistic.

IV. RESULTS

ROC-AUC varied dramatically across models using LOOCV, with some underperforming near 0.5 and GBC with

N=6 achieving the highest score of 0.8404. Using k-fold validation GBC with N=5 achieved the highest average ROC-AUC of 0.5805.

PR-AUC tended to fluctuate between specific images as expected since it is affected by class balance. We considered precision recall performance by taking the difference between PR-AUC and baseline precision of positive labels/total labels. In our LOOCV testing, this was maximized by GBC at N=4 with a score of 0.589 and an improvement over baseline of 0.411, while k-fold validation was maximized for GBC at N=4 with a score of 0.413 and an improvement over baseline of 0.241.

In addition to diagnostic statistics, we were also interested in finding the ideal threshold for this application and order to choose a particular model. In Figure 3, we can see the best performing ROC and PR curves. Purely looking at the ROC curve, averaged over N=6, we found that the mode threshold value that maximized the difference between true positive rate and false positive rate was 0.2. However, as we discussed previously, this is likely not the best metric for our use case. Based on the importance of finding all of the positive cases, we should be prioritizing recall heavily, and from the PR curve we can see that this can be obtained for a relatively linear trade off in precision. Our ROC curve also encourages this as recall increases steeply with respect to FPR, thus allowing us to achieve very high recall with small concessions.

V. DISCUSSION

Ultimately, we observed the best ROC-AUC and PR-AUC performance from the SciKit-Learn Gradient Boosting Classifier (Figure 3). It had the overall best scores for both ROC-AUC and PR-AUC for both types of cross validation. Given more time to explore this project, extensions that might result in better ROC-AUC and PR-AUC scores might involve normalizing pixel values to adjust for discrepancies in quality among patient data, more intelligently pruning outliers, and testing more deep learning models.

The next logical step would be to use our predictions to create fully segmented reconstructions of the prostate cancer, which would then allow for on the fly Gleason classification and other medical diagnoses. Although we were unable to generalize strong results through all of the models we tested, based on the data we gathered, we hypothesize that with additional fine-tuning of the individual estimators, careful curation of the data, and further experimentation including the extensions listed previously, machine learning models have the potential to offer a very powerful approach to this medical imaging problem.

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