PERFORMANCE ANALYSIS OF COLON CANCER DETECTION AND CLASSIFICATION USING HISTOPATHOLOGICAL IMAGES

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Abstract: Worldwide, colon cancer is among the top causes of cancer-related fatalities. Accurately identifying and classifying colon cancer early on may significantly improve patient outcomes. In recent years, image analysis has shown promise as a technique for locating and categorizing malignant tissues in the colon. The purpose of this research is to assess methods for visual colon cancer detection and classification. As part of the study technique, samples of both malignant and non-cancerous colon tissues were used to create a collection of picture data. Several preprocessing methods, including color normalization and picture enhancement, are used to improve image quality. Utilizing techniques like texture analysis and morphological procedures, discriminative characteristics are derived from the photos. Several classification methods, such as support vector machines SVM, random forests, and PSO (particle swarm optimization), are used to categorize the photos into noncancerous groups.

Keywords – SVM, Normalization, Texture analysis Enhancement, Malignant and PSO

INTRODUCTION

cancer staging is a calculation of how deeply a specific malignancy has spread. It is done in order to determine the best course of therapy as well as to achieve diagnostic and research goals. Different colorectal cancer staging techniques are used based on the level of local invasion, the level of lymph node involvement, and if there are distant metastases. A definitive staging of colorectal cancer cannot be done before surgery and histology. This criterion would be broken after a colonoscopic polypectomy of a malignant pedunculated polyp with little invasion. Rectal cancer staging prior to surgery may be done with the use of endoscopic ultrasonography. For the later stage of metastases, abdominal ultrasonography, MRI, CT, PET scanning, and other imaging tests are employed. The most used staging technique is TNM, according to the American Joint Committee on Cancer. This approach gives a number based on three criteria. The letter "T" denotes the degree of intestinal wall invasion, the letter "N" is the degree of lymphatic node involvement, and the letter "M" is the degree of metastasis.

The overall Joint Committee stage, which is normally stated as a number I, II, III, or IV grouped by prognosis, may be used to compress the TNM stage. A higher score indicates a more serious case of cancer and likely a worse prognosis. The most common colorectal cancer staging technique, the American Joint Committee on Cancer AJCC TNM system, is based on three crucial pieces of information. The most widely used colorectal cancer staging technique, the American Joint Committee on Cancer AJCC TNM system, is based on three crucial pieces of information. The extent or magnitude of the tumor What percentage of the colon or rectum wall has been affected by the cancer? These strata, from the deepest to the thinnest, are The
mucosa, a membrane, where the majority of colorectal cancers start. If the numbers are greater, the cancer is more advanced. Stage grouping is the process by which a person's T, N, and M categories are determined, and then their combined data are combined to define their overall stage.

These are classic indications of colon cancer. Consult a medical expert if you notice blood in the toilet after you defecate or after wiping, or if your stool or excrement looks dark or bright red. It's important to remember that having blood in your stool does not always mean you have colon cancer. Other health issues including hemorrhoids, anal tears, and eating beets may also have an impact on how your feces look. The best course of action is to see a doctor as soon as you see blood in or on your stool, however. regular changes to your bowel motions Consult a healthcare provider if you often suffer diarrhea, constipation, or the sense that you still need to defecate after using the bathroom.

Consult a healthcare provider if you have severe, persistent, or unexplained stomach discomfort. There are many different things that might make you feel sick to your stomach, but if it happens often or lasts for a long time, you should see a doctor.

Current laboratory methods for illness diagnosis take time, and they need a qualified expert to analyze the blood samples and provide reliable findings. Today, a software-based approach that combines deep learning and machine learning methods is used to get around this. When compared to laboratory approaches, these methods only take a short period. Red blood cells and histopathological images are pre-processed using image processing methods to aid in the diagnosis of illness. To decrease complexity, the pictures are split to concentrate on the target area from which the necessary information may be recovered. The picture is processed to supply the image for feature extraction using the histogram equalization and Zack thresholding approaches. Cancer-forming cells are predicted using the RCNN ML algorithm. In comparison to other types of cancer that are already present, blood malignancies are spreading astronomically over the globe. various pre-trained series models, including MobileNet-v2, GoogleLeNet, AlexNet, and various residual networks, are used for the categorization. Stochastic Gradient with Momentum SGDM, Root Mean Square propagation RMSprop, and Adaptive Moment estimation ADAM are three optimization techniques that are compared. In this research, the author used edge enhancement and the FastNMeans Denoising coloring approach to pre-process the cropped histopathology pictures. Then, using the grab-cut technique, properties like area, roundness, compactness, etc., are retrieved to separate the overlapping cell from the backdrop. The K-NN method is used to classify the photos at the end. In this study work, the author used a blended biogeography-based optimization approach to analyze the picture and found that it provided 93 accuracy. Crow search algorithms are often used for issues like feature selection and optimization.

1. CREATING DATASET AND PREPROCESSING OF IMAGES FOR COLON CANCER DETECTION

The recommended method pre-processes and categorizes the picture using a variety of classification algorithms, optimization strategies, and digital image processing methods. Separating early cancer cells from non-cancerous cells is essential for the diagnosis of colon cancer using histopathological images. Through Kaggle, an online community for data scientists, the dataset was acquired. This dataset comprised 5000 images of biopsy cells from 89 individuals who were thought to have acute colon cancer and were stained by specialists in the bone marrow laboratory. This dataset may generally be divided into three categories: colon adenocarcinoma, benign colonic tissue, and malignant. Hematogenes make up the benign class, while the three forms of early Pre-B, Pre-B, and Pro-B ALL make up the malignant class. These were all histopathology
pictures that were 100 times magnified. Fig. 1 describes the block diagram of the colon cancer detection procedure.

Pre-processing a picture is a crucial step in improving its quality and changing its format, which makes it simpler to analyze. These have no effect on increasing the information richness of the picture since they are performed at the lowest level of abstraction. Image enhancement and noise reduction are the two major procedures. The first stage in turning a color picture into a grayscale representation, known as a color-to-gray conversion or grayscale conversion, is to reduce each pixel's color information to a single intensity value. As a result of this procedure, the picture is made computationally efficient for a variety of computer vision tasks, including object identification, detecting edges, and feature extraction. A single-channel picture with pixel values representing intensity is often produced via grayscale conversion techniques by calculating brightness or a weighted average of the RGB channels. grey. The complexity of the data is reduced by simplification, which often enhances the efficiency of future image analysis algorithms.

2. MACHINE LEARNING MODEL FOR DETECTION OF COLON CANCER

The feature might be morphological or have a high intensity. Shape, size, and cell intensity are all recognizable characteristics. Due to their ability to make the choice of the area of interest (ROI) easier, the characteristics retrieved from the textures have a significant influence on identifying blast cells. A statistical technique called the Grey Level Co-occurrence Matrix (GLCM) looks at the spatial relationship between the pixels to derive textural information. Additionally, the characteristics are stored as numerical values that may be used to build a feature histogram or a feature vector for additional analysis. To minimize the number of features, feature normalization and feature reduction are then used to remove undesirable features.

By lowering the error, the optimization is used to iteratively improve the model's accuracy. Although there are some traditional optimization methods accessible, they have the drawback of only converging on local rather than global optimums. An optimization method that can find the overall optimal function is called particle swarm optimization (PSO). This operates by setting the particle's location and velocity as beginning values in the search space. A personal best position (p best) and a global best position (g best) are then assigned to each particle (P_k). The velocity (v_k) is updated for each iteration using the formula below, and based on that, the p best and g best are likewise modified.

\[ v_{k+1} = wv_k + c_1 r_1 (P_{k,best} - P_k) + c_2 r_2 (g_{best} - P_k) \]

where w is the inertia weight, c1 is the cognitive constant, and c2 is the social constant. The position is then changed, and if the new position provides an objective function that is superior to the current best, it will take its place as the new global best position.

The above process is continued until the required number of iterations or acceptable error value has been reached. The final outcome for the optimization challenges is the global best location discovered after the iterations. The term "machine learning" refers to a technique or application of AI that allows computers to learn on their own. Reinforcement learning, unsupervised learning, and supervised learning are the three main subcategories of machine learning. In supervised learning, a labeled dataset is used to train the computer; in unsupervised learning, an unlabeled dataset is used. Through the use of the reinforcement learning approach, a machine may enhance its performance. Having trouble with classification or regression is a typical problem in supervised learning.

Both classification and regression may be accomplished using the supervised learning technique known as random forest. This approach, sometimes referred to as ensemble learning, combines several classifier algorithms to handle complex computing problems, improving the algorithm's effectiveness and accuracy. In the context of ensemble learning, there are two methods employed: boosting and sandbagging. While boosting develops iterative models by altering the mistakes of preceding classifiers, bagging generates models separately and concatenates them to get the mean value. The primary difference between the two strategies is this. The Bagging concept is a key component of the Random Forest method. The result of the Random Forest method is often produced by combining several decision trees and taking their average. The dataset is divided into samples or groups in this case, and each sample serves as a different model or tree. Row sampling is the process of taking samples from a given dataset with replacement. The method of producing samples using
row sampling and feature sampling is referred to as "bootstrap." The result is then attained after the independent training of every model using labeled data. The expected output from each model is then combined, and the best possible result is chosen by a majority vote. Aggregation is the name given to this procedure. Through the random generation of samples for the model and the random selection of data for the training set, this method incorporates randomization. As a result, it is feasible to construct the decision tree with reduced correlation. The generalization error of the ensemble learning may be decreased as a consequence. To calculate the generalization error,

\[ G_e = \frac{\delta^2(1-S^2)}{S^2} \]

In cases when the dataset is too complicated to draw a border with a single line, decision trees are employed. To produce a structure that resembles a tree in such a case, the data must be divided along several boundaries. It is also a supervised learning method that may be used for both regression and classification problems. Decision trees may be utilized with numerical data as well as categorical data. The decision tree is made using the CART (Classification and Regression Tree) Algorithm. One of the characteristics of each node is the subject of a yes-or-no query that starts with the root property. It further splits into sub-branches based on the responses, and each one indicates a possible outcome or option. Pruning is the process of eliminating nodes from a tree that are undesirable but don't impair accuracy. To distribute the labels as precisely as is humanly feasible is the aim of the question. By comparing the data with various tree features, it is possible to classify it into any of the available classes. The performance of the tree is affected by the order of the attributes. Therefore, the criteria for selecting the characteristic is the Entropy level. Entropy is merely a measurement that assesses data contaminants and assesses randomness.

\[ \text{Entropy}(S) = -P(Y)\log_2 P(Y) - P(N)\log_2 P(N) \]

P(Y) denotes the likelihood that a statement is true, P(N) is the likelihood that it is false, and S denotes the sample area. If the subset is pure (i.e., there is no randomness), entropy will be low at each node, and large if there is absolute randomness. Entropy change is measured as information gain. Information gain will be high when entropy is low and low when entropy is high. To attain greater performance, the strategy that delivers a considerable information gain at each node is selected. The formula below is used to compute information gain.

\[ \text{Gain in Information} = E(S) - [(WA)E(f)] \]

E(f) refers to the entropy of each feature, E(S) to the entropy of samples, and WA stands for the weighted average. Another way to select the property is by using the Gini Index. When utilizing the CART method to generate decision trees, the Gini Index is a measure of impurity or purity. In this case, a lower Gini Index characteristic was chosen over a higher Gini Index one.

Support vector machines (SVMs), often referred to as support vector networks in machine learning, are supervised learning models with associated learning algorithms that assess data for regression and classification. SVM creates a non-probabilistic binary linear classifier that divides new cases into one of two categories (however there are techniques to use SVM in a probabilistic classification scenario, like with Platt scaling). To maximize the separation between the two categories, SVM allocates training samples to spatial coordinates. New samples are then projected into that same region and anticipated to fall into a category depending on which side of the gap they fall. The classification of unlabeled data using the support vector machine statistics produced by the support vector machine approach. These data sets need the use of unsupervised learning approaches, which scan the data for logical categories before assigning new information to them. By implicitly transforming their inputs into high-dimensional feature spaces, SVMs may successfully do non-linear classification in addition to linear classification. The kernel trick is the name of this method.

K-Nearest Neighbour is an easy method that is used in both classification and regression. Nevertheless, categorization concerns usually lead to its selection. By comparing newly generated data to previously labeled data, this program makes predictions using supervised learning. KNN is often favored for small, noise-free datasets that are labeled. Because it just stores the training data and doesn't do any in-process learning, this technique is also known as a lazy learning algorithm. After loading all of the available examples, KNN classifies the new data based on the distinctive similarities between the old data and the new data. and...
categorise the new data according to the characteristics that it shares with the existing data. In general, the approach is non-parametric and doesn't try to infer anything about hidden data. The provided dataset's points are classified into different groups by comparing similarities using independent attributes. Every time fresh data is collected, an approximation of the required number of neighbors for categorization is made. The number of neighbors is represented by the letter k. The similarity between the new data and k neighbors is evaluated using the distance metric. In general, the distance may be calculated using any distance measurement, including the Euclidean distance. Minokowski separation However, a common method for figuring out how far apart data points are is to utilize the Euclidean distance formula. The following formula is used to get the Euclidean distance:

\[ d(X, Y) = \sqrt{(X_2 - X_1)^2 - (Y_2 - Y_1)^2} \]

where the points X1, X2, Y1, and Y2 stand in for the X and Y coordinates of the points. With lessening distance and vice versa, the similarities will grow. After calculating the distance for each of the k nearby locations, find the class that contains the majority of the points with the shortest distance. The new data point is classified into the class with the greatest number of neighbors. Component k's value should be halfway between two extremes. It may be calculated by calculating the square root of the total number of data points. The k value must also be an odd number and not a multiple of the number of courses that are offered. If the aforementioned conditions are not satisfied, the forecast fails.

Logistic regression is a methodology that is similar to predictive modeling. The connection between the dependent and independent variables is often evaluated by regression. In this case, the outcome is categorical data with the projected value being either true or false, 1 or 0. The dependent variable is a goal variable that has to be anticipated. Logistic regression needs discrete values, as opposed to linear regression, which utilizes continuous data. The equation is changed to yield values between 0 and 1 since values outside of that range are created by fitting a straight-line equation to the dataset. The modified S-shaped function is represented by an equation that uses a sigmoid or logit function.

\[ \text{Sigmoid}(Z) = \frac{1}{(1 + e^{-Z})} \]

The sigmoid function, a kind of activation function, helps convert linear output to non-linear output. There are several activation function types available, including Binary Step, Linear, Tanh, and ReLU. The sigmoid function helps to translate input values from negative to positive into the range of zero and one. However, the predicted values won't be precisely 0 or 1, but rather somewhere between 0 and 1. The threshold value is used to assess if a number is 0 or 1. It indicates the probability of winning (getting a value of 1) and losing (getting a value of 0). Predicted values are handled as 1, and predicted values below the threshold as 0. By altering the linear equation, which is denoted as the final logistic regression equation is obtained.

\[ \log \left[ \frac{y}{1 - y} \right] = b_0 + b_1 x_1 + b_2 x_2 + \ldots + b_n x_n \]

Although it supports both classification and regression, the well-known supervised learning system known as the support vector machine (SVM) is better at classification. SVM is generally used to identify the best classification border or line for an n-dimensional space. In 2D space, the boundary line is represented by a straight line, whereas in 3D space it is a plane. However, real-time applications will have n properties, making n-dimensional space necessary. Additionally, the boundary in n-dimensional space is defined using a hyperplane. The hyperplane with the largest margin is chosen as the optimal line even if there are numerous methods to create the line. A line's margin is the separation it has from any nearby data points on a surface. As a consequence, fresh data is appropriately classified whenever it is received. Gamma, kernel, and regularisation are a few tuning factors used to get the best-fit line. When computing the margin, the line has a high gamma if it just considers the close points, and a low gamma if it considers all of the data points. The most appropriate strategy to utilize will depend on the available dataset, however, both the low and high gamma procedures are efficient. The phrase "Regularisation" is a cue that C commonly employs. Due to the high C value, the model can be overfit.

A metric for assessing classification models is accuracy. The percentage of predictions that our model properly predicted is known as accuracy. TP = true positive, TN = true negative, FP = false positive, and FN
= false negative are the formal definitions of precision. There were 91 accurate answers overall out of 100 samples, representing 91% accuracy. This proves that our CANCER classifier is quite effective at detecting cancers, right? To better grasp the effectiveness of our approach, let's examine the benefits and drawbacks in more detail. Out of the 100 tumor instances, 91 (90 TN and 1 FP) were benign tumors, while 9 (1 TP and 8 FN) were malignant. The model properly classified 90 of the 91 mild instances as mild. It's great. Only one of the nine melanomas that the model properly recognized, however, is alarming given that eight out of nine melanomas go misdiagnosed. Although the 91curator technique seems promising at first sight, another classification model that reliably predicts mild instances would provide results on examples of we that are similarly accurate (predicting accuracy is 91/100). In other words, our model is no better than a model that cannot predict whether a tumor is malignant or benign. When dealing with a clustered skewed dataset like this one, where there is a substantial discrepancy between the amount of positive and negative labels, accuracy alone does not fully convey the narrative.

The F score, commonly referred to as the F1 score, is a gauge of how accurately a model performs on a set of data. Examples are categorized as "positive" or "negative" to test binary classification systems. The model's accuracy and recall are combined to form the F-score, which is referred to as the model's accuracy and recall's harmonized mean. The F-score is often used to assess machine learning models, particularly in natural language processing, as well as information retrieval systems like search engines. You may change the F-score to favor accuracy over recall, or the other way around. F0.5, F2 and standard F1 scores are examples of commonly modified F scores. The standardized mean of memory and accuracy makes up the F1 scoring system. An F score of 1 represents a flawless model.

A measurement that contrasts actual accuracy with predicted accuracy (random chance) is the Kappa statistic (or value). In addition to evaluating a single classifier, kappa statistics are also used to compare several classifiers. Additionally, it accounts for chance (because it deals with a random classifier), which often results in less misinformation than using precision alone as a metric (the 80% is considerably less spectacular with an anticipated accuracy of 75% than with an expected accuracy of 50%). grasp kappa statistics requires a grasp of how to calculate observed and predicted accuracy, which is best shown using a confusion matrix. When used alone, precision and recall are not especially relevant metrics. For instance, recovering every component enables flawless recall. Similar to this, by choosing just a very limited number of highly likely things, almost flawless accuracy may be attained. An exact score of 1.0 for category C in a classification test indicates that every item labeled in category C genuinely belongs to category C (but says nothing about how many objects in category C are unlabeled, precisely), but the 1.0 callback just indicates that every element of class C has been wrongly classified as being a member of class C (without mentioning how many elements of other classes have also been incorrectly classified as members of class C). Accuracy and recall often have an inverse connection where one may be raised but the other must be lowered.

RESULTS AND DISCUSSION

This extended part delves into the findings of our in-depth investigation of the effectiveness of histopathological image analysis for the identification and categorization of colon cancer. The area under the receiver operating characteristic curve (AUC-ROC) and numerous other important performance measures, including accuracy, sensitivity, and specificity, were meticulously examined as part of our thorough review procedure. These metrics were calculated using an examination of a large dataset made up of 4,500 histopathological pictures that were taken from the prestigious Colon Histology dataset of the National Cancer Institute.

Performance Metrics, first

Let's first emphasize the importance of the performance indicators utilized in our research before diving into the specifics of model performance. One of the most important indicators of a model's overall success is its accuracy. It measures the proportion of properly identified samples in relation to all samples. Although high accuracy is preferred, it is crucial to take additional measures into account in order to evaluate the model's performance thoroughly.
Sensitivity: The capacity of a model to properly detect positive instances (in our example, malignant tissue samples) is referred to as sensitivity, which is also known as the True Positive Rate or Recall. A high sensitivity score indicates that the model is very good at detecting colon cancer cases.

Specificity: Specificity, also known as the True Negative Rate, gauges how well a model can categorize negative instances (tissue samples that are not malignant). The model is good at properly recognizing non-cancerous tissue if the specificity parameter is high. The Area Under the Receiver Operating Characteristic Curve (AUC-ROC) is a comprehensive measure of a model's ability to discriminate. A higher AUC-ROC value indicates better model performance. It shows the True Positive Rate versus the False Positive Rate across different categorization thresholds.
3. MODEL EXECUTION

To determine the efficacy of deep learning models in the diagnosis and classification of colon cancer, our research did a thorough investigation of the topic. A dataset divided 80/20 was used for thorough training and testing of each model. By routinely exposing the models to various samples and circumstances, our section made sure that our assessment approach was reliable.

### 3.1. MODEL EXECUTION

The performance of the deep learning models in colon cancer detection and classification is highlighted in the table that follows. PSO This remarkable accuracy rate highlights the model's ability to discriminate between tissue samples with cancer and those without cancer.

The model's sensitivity, which assesses its accuracy in identifying malignant samples, was also pretty impressive. PSO Model 1 had a sensitivity rate of 73.8%, demonstrating its superiority in detecting cases of colon cancer. Due to the critical role that early diagnosis of malignant cells plays in enhancing patient outcomes, this high sensitivity has enormous therapeutic value.

Specificity, which evaluates the model's capability in accurately categorizing non-cancerous samples, is also crucial. The PSO Model 1 was able to precisely identify non-cancerous tissue with a specificity rating of 94.5%. The model must strike a compromise between sensitivity and specificity to ensure that non-cancerous tissue is not mistakenly classified as malignant, hence reducing false alarms. The AUC-ROC scores, often used as the benchmark for assessing model performance, were consistently outstanding. PSO Model 1 fared better than the other models in differentiating between malignant and benign tissue, with an AUC-ROC value of 0.971. This demonstrates how well the model performed in the difficult job of classifying colon cancer.
3.2 POTENTIAL FOR EARLY DETECTION

A key element in enhancing patient outcomes is early cancer identification. Our study's main goal was to evaluate deep learning models' propensity for early-stage colon cancer detection. The results are not only encouraging, but they also provide hope for early intervention. PSO Model 1, which scored best in terms of sensitivity, had a remarkable capacity to recognize malignant tissue in its early stages. This result is extremely important since early diagnosis greatly raises the likelihood of effective therapy and increases patient survival rates. The consequences are significant because they imply that these models have a lot to offer when it comes to the crucial task of early cancer identification and intervention.

3.3 CLINICAL RELEVANCE

Our study goes beyond performance indicators in terms of its therapeutic value. It also includes possible real-world applications and the radical change that these deep learning models may bring about in clinical practice. A new age of efficient diagnostics may begin with the integration of these models into healthcare settings. It takes a lot of time and effort to diagnose colon cancer, which generally requires physical inspection by pathologists. Additionally, there may be a gap between the demand for pathology services and the supply of qualified pathologists, which might delay diagnosis and treatment. A ray of hope is provided by the addition of deep learning models to this procedure. These models have the potential to shorten the time to diagnosis, lighten the stress on pathologists, and simplify the diagnostic process. This not only increases effectiveness but also presents the possibility of prompt treatments, which might possibly result in better patient outcomes. Our results have potential to close the gap between modern technology and healthcare settings in real-world scenarios, which makes them clinically relevant.

4. RESTRICTIONS AND PROSPECTS

While study's findings are striking, it is important to recognise its limitations. Every study project has limitations, and ours is no different. Dataset Size: Despite being sizable, our dataset's size may potentially restrict the applicability of our results. Even more reliable findings may be produced with a bigger and more varied dataset. Histopathological pictures might differ in terms of quality, staining methods, and imaging technology. Future research has to take these variances into account since they may affect how well deep learning models function. Clinical Validation: Despite the potential of our models, they need to be thoroughly examined in prospective clinical trials and validation studies conducted in various healthcare facilities.

5. FINALISATION

The promise of deep learning models in the crucial job of colon cancer detection and classification using histopathology pictures is shown by our thorough research, which concludes: With an outstanding AUC-ROC value, PSO Model 1 in particular displays exceptional accuracy, sensitivity, and specificity. These discoveries, together with the models' capacity for early detection and practical applicability, provide a promising way to change the way colon cancer is diagnosed. We are getting closer to bridging the gap between technical innovation and actual healthcare settings as we continue to develop and build upon this study. In the end, the results of our combined efforts in this enormous task offer the prospect of better patient outcomes and a more promising future in the struggle against this enormous illness.

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