Implementation of CAD system for Cancer Detection using SVM based Classification

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Abstract - Breast cancer is one of the most frequently occurring diseases which cause death among women. Masses present in mammogram of breast, primarily indicates breast cancer and it is important to classify them as benign or malignant. Benign and malignant masses differ in geometry and texture characteristics. However, not every geometry and texture feature that is extracted contributes to the improvement of classification accuracy; thus, to select the best features from a set is important. Selecting best feature set again depends on the classification algorithm to be used. Different classification algorithm gives different performance on number of selected features. SVM can generally achieve better performance on using all extracted features. Proposed new system will examine the CAD system for mass classification. It will integrate a support vector machine (SVM).

Key Words: Mammogram, ROI extraction, Segmentation, Features extraction, Feature Selection, Classification

1. INTRODUCTION

Breast cancer is the most common malignancy of women and is the second most common and leading cause of cancer deaths among them. At present, there are no effective ways to prevent breast cancer, because its cause is not yet fully known. Early detection is an effective way to diagnose and manage breast cancer which can give a better chance of full recovery. Therefore, early detection of breast cancer can play an important role in reducing the associated morbidity and mortality rates.

The manual analysis of breast lesions from mammograms presents large interpretation variability amongst radiologists. This variability can be reduced with the aid of Computer Aided Diagnosis (CAD) Systems that can act as a second reader in the analysis of breast lesions. However, for a CAD system to be useful in clinical setting, it must effectively classify as benign or malignant.

Detection, segmentation and classification of breast lesions are the main three steps involved in fully automated CAD systems that can work in the analysis of mammograms. Building a CAD system is difficult because mammograms are marred by low signal to noise ratio for the visualization of breast lesions. In addition, breast lesions present a large variation in terms of shape, size and appearance. A large number of methods have been applied for building automated CAD systems for both types of lesions, namely mass and micro-calciﬁcation, but in this work we focus only on the analysis of masses. The major drawback of current approaches is that they generate a large number of false positives and miss a fair amount of true positive regions during the mass detection stage. Furthermore, mass segmentation is generally based on active contour models and graph-based approaches that rarely capture the large shape and appearance variations of breast masses. Finally, mass classification is generally implemented using sub-optimal hand-crafted features and machine learning classiﬁers such as support vector machines (SVM), linear discriminant analysis (LDA), artificial neural net (ANN), etc. One major limitation of the majority of existing CAD systems is that most of them require manual intervention to obtain mass candidates for segmentation and classiﬁcation.

This paper presents a new approach based on recently developed machine learning models to develop a fully automated CAD system for automated detection, segmentation and classiﬁcation of masses from mammograms. Our proposed solution to the mass detection problem consists of four stages: 1) Segmentation of the Region of Interest (ROI) of the original input mammographic image. 2) Texture and Geometric Features Extraction from the segmented image 3) Feature Selection from the total number of extracted image 4) Classiﬁcation of the mass (tumor) as benign or malignant.

2. LITERATURE SURVEY

Computer-aided diagnosis (CAD) (Nikhil et al 2008; Derya 2009; Leonardo 2009; Lee et al 2000) [1] techniques and systems support radiologists in the role of a second reader and prompting the radiologist to review areas in a mammogram considered. Therefore the development of a successful CAD scheme is necessary to investigate the usefulness for radiologists in their diagnosis to quantify the computer output for radiologists and how to maximize the effect of the computer output on their diagnosis.

Mammography detects breast cancer through the identification of abnormalities such as masses and/or micro-calciﬁcations. However, due to the complexity of breast structure, low disease prevalence, and radiologist fatigue, abnormalities are often ignored. As reported in [2], about 10%-25% of abnormal cases shown in mammography have been wrongly ignored by radiologists. One way to reduce the
false negative rate is to use double reading, which can improve the detection rate by 5%–15% [3].

However, double reading is too expensive and time consuming. Thus, computer-aided cancer detection technology attracts researchers’ and radiologists’ attention. The adoption of a computer-aided detection (CAD) system could reduce the experts’ workload and can improve the early cancer detection rate [4], [5]. A recent report [4] demonstrates that the adoption of CAD improved the detection of breast cancer by 7.62%. Although the adoption of CAD systems has attracted great attention [5], further effort must be put on the investigation of related key technologies. These key technologies include calcification detection, mass detection, architectural distortion detection, and mass classification (malignant or benign).

In [6], the mass classification using gradients was investigated. In the proposed method, a mass boundary was segmented into concave and convex parts, and was polygonized. The features that quantify the extent of the spiculated nature of the boundary and the degree of narrowness of the spicules were extracted for mass classification.

In [7]–[8], morphological features with manually delineated boundaries were adopted for mass classification. Pohlman et al. [8] used an adaptive region growing technique to segment masses. Six features measuring mass shape and boundary roughness were extracted and used for classification. In [9], the mass classification based on the automated segmentation of masses was investigated.

3. PROPOSED SYSTEM

The Proposed CAD system has the following modules:

1. Input Images
2. Noise Reduction and ROI Extraction
3. Segmentation
4. Features Extraction
5. Feature Selection
6. Classification

The below shown is the architecture of the proposed CAD system.

3.1 Input Images Module

In this module we will take input images from Mammographic Image Analysis Society (MIAS), Digital Database for Screening Mammography (DDSM) and UCI Machine Learning Repository. Figure 4.2.1 shows the original mammographic image from the database.

![Figure 2: Original mammogram](image-url)

3.2 Noise Reduction and ROI Extraction

The images which are taken from the standard mammographic databases contain many noises like background noise, pectoral tissue, etc. To get the precise results this noise must be removed by preprocessing which is done by cropping the unrequired area from the image. Then only the Region of Interest is also cropped. Its contrast is improved.
3.3 Segmentation

Once the required ROI is ready, it is then used for segmentation to get the segmented image. Result of image segmentation is a set of segments that collectively cover the entire image, or a set of contours extracted from the image. Fuzzy C-Means Clustering Algorithm is used for segmentation in proposed system. A clustering algorithm organizes items into groups based on similarity criteria. Here the data points can belong to more than one cluster, and associated with each of the points are membership grades which indicate the degree to which the data points belong to the different clusters. It tries to put each of the data points to one of the clusters. It does not decide the absolute membership of a data point to a given cluster; instead, it calculates the likelihood (i.e., the degree of membership). Each item may belong to more than one group (hence the word fuzzy), where the degree of membership for each item is given by a probability distribution over the clusters.

The FCM clustering is obtained by minimizing an objective function shown in below equation:

\[ J = \sum_{i=1}^{n} \sum_{k=1}^{c} \mu_{ik}^{m} |p_i - v_k|^2 \]

**Fuzzy C-Means Algorithm**

Step 1: Set the number of clusters, the fuzzy parameter (a constant > 1), and the stopping condition

Step 2: Initialize the fuzzy partition matrix

Step 3: Set the loop counter \( k = 0 \)

Step 4: Determine the cluster centroids, calculate the objective value \( J \)

Step 5: For each pixel, for each cluster, compute the membership values in the matrix

Step 6: If the value of \( J \) between consecutive iterations is less than the stopping condition, then stop; otherwise, set \( k = k + 1 \) and go to step 4

3.4 Features Extraction

After segmenting the mass from the ROI, we computed a set of features that is related to the geometry and texture of the boundary. A typical benign mass has a round, smooth, and well-circumscribed boundary, whereas the boundary of a malignant tumor is usually speculated, rough, and blurry.

**Geometry Features:** The geometry features represent the shape of the contour of a mass. These are calculated from the boundary pixels after the segmentation. The geometry features can be any among compactness, Normalized Distance Moments, Fourier Features, Normalized Radial Length based features and relative gradient orientation based features.

**Texture Features:** In addition to the shape information of a mass contour, the texture information of the region surrounding the mass boundary also contains important information to discriminate the benign and malignant masses. Thus, we also used the texture information for the mass classification. The texture features extracted are based on Gray level Co-occurrence Matrix.

The libraries used to find out the features are as follows.

- de.lmu.ifi.dbs.jfeaturelib.features.Haralick
- de.lmu.ifi.dbs.jfeaturelib.shapeFeatures.Compactness
- de.lmu.ifi.dbs.jfeaturelib.examples.HaralickDemo
- de.lmu.ifi.dbs.jfeaturelib.examples.HaralickDemoTest

The extracted features are stored in attributes.csv file

3.5 Features Selection

In feature extraction module we have extracted total 31 geometry and texture features. We used WEKA library to select the optimum number of features for classification process.

In WEKA, attribute selection searches through all possible combination of attributes in the data to find which subset of attributes works best for prediction. It employs two objects which include an attribute evaluator and search method. Any
combination of these attribute evaluator and search algorithms can be used to determine the score and rank attribute in a data set.

WEKA tries to use a subset of features and train a model using them. Based on the inferences that WEKA draw from the previous model, it decide to add or remove features from your subset. The problem is essentially reduced to a search problem.

The libraries used to select the features of image are as follows:

Weka.associations
Weka. attributeSelection
Weka.classifiers

Then attributes.csv file is provided as input to the WEKA library. WEKA uses subset evaluation algorithm. On execution of algorithm, features are saved which will be further used for classification.

3.6 Classification

The main task of classifier is to categorize the ROI by considering the geometrical and textural features as benign or malignant. In training phase, data set which is labeled as benign and malignant mass are given to classifier as training data set and the classifier is trained. In testing phase, the unknown data (mammographic image) is given to classifier for actual classification.

Support Vector Machine is the algorithm used for classifying the image. The algorithmic steps are described below. LibSVM is being imported.

Support Vector Machine Algorithm

```java
import libsvm.LibSVM;
import net.sf.javaml.classification.Classifier;
import net.sf.javaml.core.Dataset;
import net.sf.javaml.core.Instance;
import net.sf.javaml.tools.data.FileHandler;

Begin
1. class TutorialLibSVM
2. /* Load a data set */
Dataset data = FileHandler.loadDataset(new File("E:\attributes.csv"), 31, "");
3. /* Constract a LibSVM classifier with default settings*/
Classifier svm = new LibSVM();
    svm.buildClassifier(data);
4. /* Counts for correct and wrong predictions*/
int correct = 0, wrong = 0;
```

5. /* Classify all instances and check with the correct class values*/
```java
for (Instance inst : dataForClassification) {
    Object predictedClassValue = svm.classify(inst);
    Object realClassValue = inst.classValue();
    if (predictedClassValue.equals(realClassValue))
        correct++;
    else
        wrong++;
}
```

4. EXPERIMENTAL RESULTS

From the images, 430 ROIs that contains benign masses and 240 ROIs that contains malignant masses were extracted. Totally, 670 ROIs were used as input for training data set; which contained 430 benign training dataset and 240 malignant training dataset. After that segmentation, feature extraction is carried out on provided training dataset; where the segmented images get stored in the Project0 folder. Then, by clicking TEXTURE and GEOMETRIC FEATURES button, all the 31 texture and geometric features get extracted and get stored in attributes.arff file. Here we have selected all the 31 features since inclusion of all 31 features gives us better results for SVM classifier, as per experiment. Below are the snapshots of execution of the classification program.

For testing, we have taken some benign and malignant images from the same datasets. In the testing GUI, ROI of original mammographic image is provided as input. Then its segmented image is popped after clicking SEGMENTATION button. After that its texture and geometric features are calculated by clicking TEXTURE FEATURES and GEOMETRIC FEATURES button and displayed in the textbox. By clicking CLASSIFY button we will get the message in the message box as “The image is benign/ malignant”.

5. RESULT ANALYSIS

The following experiments were conducted on the entire ROI (n = 670) data set, with all of the 31 features obtained without feature selection. The classification performance is measured by the true positive rate (TPR), the true negative rate (TNR), and the accuracy. The true positive number of a classifier is denoted as TP, the false positive number as FP, the true negative number as TN, and the false negative number as FN. The definitions are as follows:

\[
TPR = \frac{TP}{TP + FN}
\]

\[
TNR = \frac{TN}{TN + FP}
\]

\[
ACCURACY = \frac{TP + TN}{TP + FP + TN + FN}
\]
From 430 Benign ROIs, TP = 390, FN= 40, From 240 Malignant ROIs, TN= 207, FP= 33

From the above obtained mathematical values and equations of TP, TN, FN and FP we get

\[
TPR = 0.9069 \\
TNR = 0.8625 \\
ACC = 0.8910
\]

These results have been cross checked on WEKA explorer using Naïve Bayes algorithm to check the efficiency of the proposed SVM method. The comparative classification performance is shown in below table.

Table -1: The comparative classification performance

<table>
<thead>
<tr>
<th>Accuracy</th>
<th>Naïve Bayes (WEKA)</th>
<th>SVM (JAVA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TPR</td>
<td>0.9069 (390/430)</td>
<td>0.9069 (390/430)</td>
</tr>
<tr>
<td>TNR</td>
<td>0.875 (210/240)</td>
<td>0.8625 (207/240)</td>
</tr>
<tr>
<td>ACC</td>
<td>0.8955 (600/670)</td>
<td>0.8910 (597/670)</td>
</tr>
</tbody>
</table>

From the above observation table the accuracy given by Naïve Bayes algorithm on WEKA explorer is 89.55% while that shown by SVM on JDK is 89.10%.

6. CONCLUSION

Breast cancer is the most common form of cancer among women and is the second leading of death after lung cancer. Early detection and diagnosis of breast cancer increase the treatment options and a cure is more likely. In this thesis, a breast mass classification system is implemented under the JDK environment for benign-malignant mass classification of previously segmented mass candidates. Also, the inclusion of all 31 features for classification stage has a positive effect on the results of classification. The proposed work is carried out on JDK environment using JAVA programming language. Test cases are cross checked on WEKA explorer which gave significant results. Standard datasets like DDSM, MIAS and UCI Machine learning repository are used. A Real Test case from Siddhivinayak Cancer Hospital is being tested which gave positive results. Comparative study shows SVM efficiently gives good accuracy.

REFERENCES


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