ORIGINAL RESEARCH

An automated mammogram classification system using modified support vector machine

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¹Computer Science Department, Landmark University, Omu-Aran, Kwara State, Nigeria; ²Department of Computer and Information Sciences, Covenant University, Ota, Ogun State, Nigeria **Purpose:** Breast cancer remains a serious public health problem that results in the loss of lives among women. However, early detection of its signs increases treatment options and the likelihood of cure. Although mammography has been established to be a proven technique of examining symptoms of cancer in mammograms, the manual observation by radiologists is demanding and often prone to diagnostic errors. Therefore, computer aided diagnosis (CADx) systems could be a viable alternative that could facilitate and ease cancer diagnosis process; hence this study.

Methodology: The inputs to the proposed model are raw mammograms downloaded from the Mammographic Image Analysis Society database. Prior to the classification, the raw mammograms were preprocessed. Then, gray level co-occurrence matrix was used to extract fifteen textural features from the mammograms at four different angular directions: $\theta = \{0^\circ, 45^\circ, 90^\circ, 135^\circ\}$, and two distances: $D = \{1,2\}$. Afterwards, a two-stage support vector machine was used to classify the mammograms as normal, benign and malignant.

Results: All of the 37 normal images used as test data were classified as normal (no false positive) and all 41 abnormal images were correctly classified to be abnormal (no false negative), meaning that the sensitivity and specificity of the model in detecting abnormality is 100%. After the detection of abnormality, the system further classified the abnormality on the mammograms to be either "benign" or "malignant". Out of 23 benign images, 21 were truly classified as benign. Also, out of 18 malignant images, 17 were truly classified to be malignant. From these findings, the sensitivity, specificity, positive predictive value, and negative predictive value of the system are 94.4%, 91.3%, 89.5%, and 95.5%, respectively. **Conclusion:** This article has further affirmed the prowess of automated CADx systems as a viable tool that could facilitate breast cancer diagnosis by radiologists.

Keywords: cancer diagnosis, CADx systems, radiologists, diagnostic errors, GLCM

Introduction

Breast cancer is a deadly disease that plagued women population in developing countries.^{1,2} However, early detection of the symptoms of breast cancer could facilitate its treatment and it has been established that mammography is the best technique for this task.^{3,4} This technique involves the examination of X-ray images of the breast for symptoms such as architectural distortion, calcifications, masses, etc.

Although mammography has been established to be a viable technique in the diagnosis of breast cancer, its interpretation by Radiologists' are most times in question as different Radiologists may come up with different interpretations.^{5,6} Azar⁷ argued that interpretation of mammogram is a cognitive skill which

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Radiologists develop over time. Nevertheless, digital image processing and machine learning approaches can proffer a solution to many of the inherent problems encounter by Radiologists when interpreting mammogram manually.

Mammogram processing (enhancement and segmentation) enhances visual interpretation. Also, intelligent classifier aids expert in detection and classification of mammograms.⁸

Methodology

The detailed steps employed in the classification task are documented in this section. It involves mammogram acquisition, mammogram processing, feature extraction and the classification models, all of which were implemented in MATLAB 2015.

Mammogram acquisition

Raw mammograms were downloaded from MIAS database.⁹ Out of 322 mammograms available in the database, 126 normal, 60 benign and 48 malignant mammograms were carefully selected. Of these, 37 normal, 23 benign, 18 malignant mammograms were used as test data while the remaining 156 mammograms were used for training the system.

Mammogram processing

Mammography uses low dose x-ray for imaging which produces a low qualitymammograms.¹⁰ In addition, Kayode et al¹¹ ascertained that image acquisition comes with lots of inherent problems which include scratches, dust artifact, scanner induced artifacts and excessive background noise which can further undermine the reliability of X-ray images, therefore, it is expedient to pre-process digital images so as to improve their quality.

Furthermore, image enhancement includes removing noise and adjusting image contrast so that the identification of key features becomes easier. In this article, image enhancement was achieved using Contrast Limited Adaptive Histogram Equalization (CLAHE). Also, segmentation was done in order to separate the needed region also known as the region of interest (ROI) from the entire breast tissue. Otsu threshold algorithm was implemented in MATLAB to extract ROI from the entire mammogram image.

Extraction of features from mammogram's ROI

Radiologists analyze ROI found on mammograms by examining morphological features, such as size, margin and shape of the ROI; this helps them to differentiate between normal and abnormal mammograms and also to differentiate between an abnormal benign mammogram from an abnormal malignant mammogram. However, it should be noted that decisions concerning these morphological features are cognitive and greatly depend on Radiologists' experience and opinion; therefore, it is essential to use computers to facilitate the extraction of GLCM features related to image texture, coarseness and heterogeneity, which are not necessarily seen by unaided eye.

GLCM is the most common statistical method used to compute textural features from gray level images such as mammograms. GLCM features consider the spatial relationship between the pixel of interest and its neighbouring pixels thereby providing us with textural features. Each element (m, n) in the resultant GLCM is simply the sum of the number of times that the pixel with value m occurred in the specified angular direction and distance to a pixel with value n in the input image.¹²

In this article, the thirteen GLCM features, proposed and by Haralick et al¹³ with two other features called cluster prominence and cluster shade, proposed by Tsatsoulis¹⁴ were extracted from mammograms. These GLCM features have been reported to adequately convey information about the textural characteristics of an image. The features are listed in Table 1. The detailed discussion about the features is documented in^{13,14} respectively.

Unlike in the existing works,^{15–18} where a single GLCM feature was created for each image using horizontal direction, $\theta = 0^0$ and distance, D = 1, this study is of the opinion that a single GLCM is inadequate to characterize the textural features of an image, therefore, in addition to the horizontal direction $\theta = 0^0$ and distance D = 1, two diagonals $\theta = \{45^0, 135^0\}$ and the vertical direction $\theta = 90^0$ at distance d = 2 at distance D = 2 which correspond to multiple GLCMs features at four directions and two distances 1 and 2 were computed. The following pseudocode was used to extract the features:

Objective:

1. To Extract GLCM from mammogram images Input:

- 2. ROI of the enhanced image I
- 3. The direction to be used for creating a co-occurrence matrix

Output:

4. The GLCM extracted features vector from the input image I

$$F = \{f_1, f_2, \dots, f_{15}\}$$

Process:

- 5. Begin:
- 6. for θ = {0⁰, 45⁰, 90⁰, 135⁰} do
 7. create *CoocmatrixI*(θ)//*GLCMs at angular directions θ*
- 8. end loop
- 9. for distances $D = \{1, 2\}$ do
- 10. SumCooc = 0 //sum of co-occurrence matrices
- 11. for each angle in θ do
- 12. $SumCooc = SumCooc + CoocmartixI(\theta)$
- 13. end loop
- 14. end loop
- 15. for i=1 to 15 do
- 16. $S_i(D) = compute features(SumCooc, F(i))$
- 17. end loop
- 18. for i=1 to 15
- 19. $AveSumCooc(i) = mean(S_i)$
- 20. features = AveSumCooc(i)
- 21. end loop features
- 22. Return
- 23. End

Classification of mammograms

SVM was used for classification purposes. In MATLAB programming tool, SVM functions support only two classes based on its fundamental principle as a binary classifier. Due to three attributes of interest: normal, benign and malignant, SVM was trained twice, first to classify mammograms to either normal or abnormal, and second, to classify abnormal mammograms to benign or malignant. By this two-stage approach the three attributes of mammograms will be accommodated as illustrated in Figure 1.

Table I	Textural	features	extracted	from	the	dataset
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S/N	Features	Mathematical Expression
١.	*IMC	$f_1 = \frac{HXY - HXY1}{\max(HX, HY)}$
2.	Contrast	$f_2 = \sum_{m,n=1}^{N_g} m - n ^2 P(m, n)$
3.	Correlation	$f_3 = \sum_{m,n=1}^{N_g} \frac{P(m,n) - \mu_x \mu_y}{\sigma_x \sigma_y}$
4.	Cluster Prominence	$f_4 = \sum_{m,n=1}^{N_g} (m - \mu_x + n - \mu_y)^4 P(m,n)$
5.	Cluster Shade	$f_5 = \sum_{m,n=1}^{N_g} (m - \mu_x + n - \mu_y)^3 P(m,n)$
6.	Dissimilarity	$f_{6} = \sum_{m,n=1}^{N_{g}} m - n P(m, n)$
7.	Energy	$f_7 = \sum_{m,n=1}^{N_g} (P(m,n))^2$
8.	Entropy	$f_8 = -\sum_{m,n=1}^{N_g} P(m,n) \log(P(m,n))$
9.	*IMC ₂	$f_9 = \sqrt{(1 - \exp(-2(HXY2 - HXY))))}$
10.	Difference Variance	$f_{10} = \sum_{l=0}^{N_g - 1} l^2 P_{x-y}(l)$
11.	Variance	$f_{11} = \sum_{m,n=1}^{N_{\rm g}} (1-\mu)^2 P(m,n)$
12.	Sum Average	$f_{12} = \sum_{l=2}^{2N_g} lP_{x+y}(l)$
13.	Sum Variance	$f_{13} = \sum_{l=2}^{2N_{\rm g}} (l - f_8)^2 P_{x+y}(l)$
14.	Difference Entropy	$f_{14} = -\sum_{l=0}^{N_{\rm g}-1} P_{x-y}(l) \log(P_{x-y}(l))$
15.	Homogeneity	$f_{15} = \sum_{m,n=1}^{N_{\rm g}} \frac{1}{1 + (m-n)^2} P(m,n)$

Note: $^{\ast}\text{IMC}_1$ and IMC_2 are information measure sof correlation 1 and 2, respectively.

Experimental results

Figure 2 illustrates the Graphic User Interface (GUI) designed for the preprocessing stage. Raw mammograms were fed into the system one after the other then CLAHE was employed for the preprocessing. Afterwards, the suspicious region otherwise known as ROI was extracted from the enhanced image; this is illustrated in Figure 3.

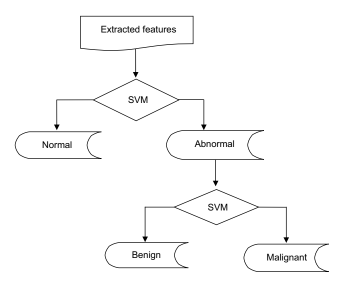


Figure I Two-level classification approach using support vector machine (SVM).

Fifteen GLCM features presented in Table 1 were extracted from the ROI as illustrated with Figures 4 and 5 respectively. The averages of the features at the two distances were calculated as shown in Figure 6 while the overall averages of each of the features which serve as input to SVM classifier, were also calculated as shown in Figure 7.

The first-stage classification entails categorizing the mammograms as either normal or abnormal as illustrated in Figure 8; this is termed the first-stage classification. If the image is normal the algorithm stops. However, if any abnormality is detected, the algorithm further classifies the abnormality into benign or malignant (see Figure 9); this is second-stage classification.

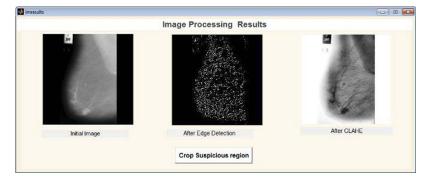


Figure 2 Mammogram enhancement using Contrast Limited Adaptive Histogram Equalization (CLAHE).

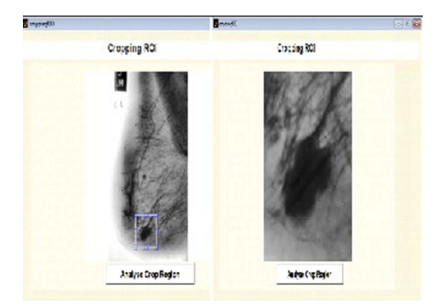


Figure 3 Region of interest (ROI) extraction.

GLCM Feature (a	at 0 degree,D=1)	GLCM Feature (a	at 45 degree,D=1)	GLCM Feature (a	t 90 degree,D=1)	GLCM Feature (a	at 135 degree, D=
IMC1	12.7249	IMC1	12.6761	IMC1	12.7068	IMC1	12.646
Contrast	0.0786677	Contrast	0.0890308	Contrast	0.0951937	Contrast	0.148259
Correlation	0.96745	Correlation	0.963175	Correlation	0.9605	Correlation	0.93867
Cluster Prominence	49.7884	Cluster Prominence	49.5632	Cluster Prominence	49.1133	Cluster Prominence	47.757
Cluster Shade	-4.43132	Cluster Shade	-4.41593	Cluster Shade	-4.42713	Cluster Shade	-4.33945
Dissimilarity	0.0784192	Dissimilarity	0.0885299	Dissimilarity	0.094697	Dissimilarity	0.147759
Energy	0.246767	Energy	0.24136	Energy	0.240294	Energy	0.217814
Entropy	1.71568	Entropy	1.74491	Entropy	1.76052	Entropy	1.89181
IMC2	0.960832	IMC2	0.955819	IMC2	0.952734	IMC2	0.926204
Difference Variance	0.960815	Difference Variance	0.955785	Difference Variance	0.952701	Difference Variance	0.926171
Variance	0.405766	Variance	0.400013	Variance	0.401205	Variance	0.381605
Sum average	12.6714	Sum average	12.6288	Sum average	12.6493	Sum average	12.6017
Sum variance	6.79886	Sum variance	6.78588	Sum variance	6.79688	Sum variance	6.78575
Difference entropy	31.1553	Difference entropy	30.7952	Difference entropy	30.7717	Difference entropy	29.6638
Homogeneity	0.99879	Homogeneity	0.998631	Homogeneity	0.998536	Homogeneity	0.99772

Figure 4 Gray level co-occurrence matrix (GLCM) features computed at D=1.

GLCM Feature (a	at 0 degree,D=2)	GLCM Feature (a	t 45 degree,D=2)	GLCM Feature (a	t 90 degree,D=2)	GLCM Feature (a	t 135 degree, D=2)
IMC1	12.6541	IMC1	12.5589	MC1	12.6174	MC1	12.496
Contrast	0.151817	Contrast	0.164758	Contrast	0.18452	Contrast	0.286959
Correlation	0.937396	Correlation	0.932098	Correlation	0.923466	Correlation	0.881708
Cluster Prominence	48.0463	Cluster Prominence	47.7896	Cluster Prominence	46.7058	Cluster Prominence	44.1581
Cluster Shade	-4.36123	Cluster Shade	-4.31688	Cluster Shade	-4.36368	Cluster Shade	-4.16408
Dissimilarity	0.15094	Dissimilarity	0.16285	Dissimilarity	0.181392	Dissimilarity	0.275763
Energy	0.215584	Energy	0.209718	Energy	0.205798	Energy	0.174695
Entropy	1.90033	Entropy	1.92617	Entropy	1.96193	Entropy	2.13043
IMC2	0.924661	IMC2	0.91883	MC2	0.909763	MC2	0.863921
Difference Variance	0.924618	Difference Variance	0.918766	Difference Variance	0.909617	Difference Variance	0.863238
Variance	0.378509	Variance	0.370611	Variance	0.370855	Variance	0.338104
Sum average	12.65	Sum average	12.5604	Sum average	12.6034	Sum average	12.5068
Sum variance	6.78766	Sum variance	6.7612	Sum variance	6.78357	Sum variance	6.76075
Difference entropy	29.6314	Difference entropy	29.1936	Difference entropy	29.1724	Difference entropy	27.9935
Homogeneity	0.997666	Homogeneity	0.99747	Homogeneity	0.997167	Homogeneity	0.995602

Figure 5 Gray level co-occurrence matrix (GLCM) features computed at D=2.

veightedf			
Weighted GLCM	Feature (at D=1)	Weighted GLCM	Feature (at D=2)
IMC1	15.8777	IMC1	15.8108
Contrast	0.0866523	Contrast	0.159393
Correlation	0.955338	Correlation	0.917936
Cluster Prominence	50.8319	Cluster Prominence	49.1425
Cluster Shade	-5.57082	Cluster Shade	-5.5441
Dissimilarity	0.0864569	Dissimilarity	0.157347
Energy	0.291386	Energy	0.257724
Entropy	1.64448	Entropy	1.81461
IMC2	0.956804	IMC2	0.921645
Difference Variance	0.956791	Difference Variance	0.921531
Variance	0.478462	Variance	0.450529
Sum average	15.8222	Sum average	15.8126
Sum variance	7.73339	Sum variance	7.72534
Difference entropy	41.6018	Difference entropy	39.9965
Homogeneity	0.998667	Homogeneity	0.997551
		Average Weighted 0	

Figure 6 Weighted gray level co-occurrence matrix (GLCM) features computed at distances 1 and 2.

System testing and performance evaluation

The model was tested to determine if the learning algorithm really performed its recognition task of differentiating normal and abnormal image on one hand and malignant and benign on the other hand.

The aim of this step is to introduce the model to the 78 unseen mammograms after it has been trained with a training dataset so as to test how well it performed.

average		
Weighted Averag	ge GLCM Feature	
IMC1	15.8385	
Contrast	0.14112	
Correlation	0.927386	
Cluster Prominence	49.6756	
Cluster Shade	-5.53218	
Dissimilarity	0.139902	
Energy	0.26517	
Entropy	1.77843	
IMC2	0.930237	
Difference Variance	0.930171	
Variance	0.456663	
Sum average	15.8164	
Sum variance	7.72989	
Difference entropy	40.3494	Select Features for Classification
Homogeneity	0.997831	Classify

Figure 7 Overall average gray level co-occurrence matrix (GLCM) features computed.

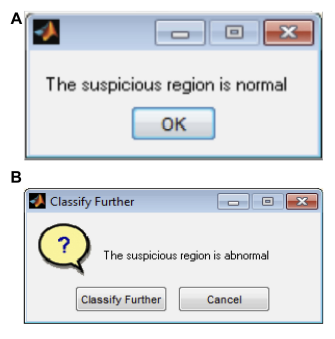


Figure 8 First-stage of classification: (A) normal mammogram classification; (B) abnormal mammogram classification.

First-stage classification

Using the expert's classification provided alongside the dataset as the actual class, in the first-stage classification, all the 37 normal images were classified as normal (no false positive) and all 41 abnormal images were correctly classified to be abnormal (no false negative) meaning that the model can accurately detect abnormality, that is, each of the sensitivity, specificity and accuracy of the model in detecting abnormality is 100%

Second-stage classification

After the detection of abnormality, the system further classified the abnormality on the mammograms to be either "Benign" or "Malignant". This is the second-stage classification. Out of 23 benign images, 21 were classified as truly benign while 2 are misclassified to be malignant. Also, out of 18 malignant images, 17 were classified to be malignant while an image was misclassified to be benign. These values are entered into a confusion matrix as shown in Table 2.

Results from proposed technique and existing works

Table 3 presents the results of existing works who also classified MIAS database mammograms using SVM. These are compared with the results obtained from the proposed technique. Results obtained from existing works have classified mammograms in MIAS database to either normal or abnormal

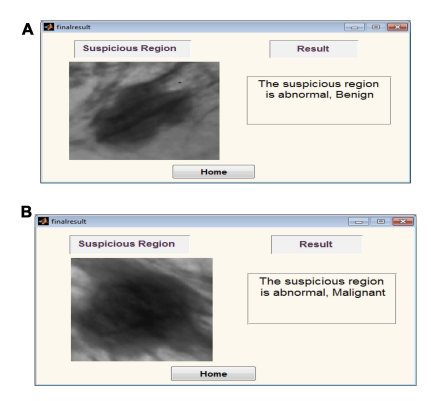


Figure 9 Second-stage classification: (A) benign; (B) malignant.

Expert's classification	System cla	ssification	Total
	Benign	Malignant	
Benign	TN=21	FP=2	23
Malignant	FN=I	TP=17	18
Total	22	19	

Table 2 The system's confusion matrix

Abbreviations: TN = True Negative; FN = False Negative; FP = False Positive and TP = True Positive.

mammograms or benign or malignant mammograms. However, this work reported an automated MIAS database mammograms classification in which mammogram are classified first to abnormal and normal, after which the abnormal mammograms were further classified into benign or malignant mammograms.

Results and discussion

For the first stage classification, the sensitivity, specificity, PPV and NPV of the model are all 100%. This means that the system is 100% accurate of differentiating abnormal mammograms from normal ones.

Using the information on the confusion matrix labeled Table 2, the second level sensitivity, specificity; PPV and NPV of the model were calculated thus:

Sensitivity
$$= \frac{TP}{TP + FN} * (100\%)$$

 $= \frac{17}{18} * (100\%)$
 $= 94.4\%$ (1)

Table 3 Results from proposed technique and existing works

Specificity =
$$\frac{TN}{TN + FP} * (100\%)$$

= $\frac{21}{23} * (100\%)$
= 91.3% (2)

$$PPV = \frac{TP}{TP + FP} * (100\%)$$

= $\frac{17}{19} * (100\%)$
= 89.5% (3)

NPV =
$$\frac{TN}{TN + FN} * (100\%)$$

= $\frac{21}{22} * (100\%)$ (4)
= 95.5%

From the performance metrics, the probability that the system would detect a malignancy among patients that have cancer is 0.944, that is, sensitivity =94.4%; the probability that the system would classify a patient's mammogram image as benign among patients that do not have cancer is 0.913 (specificity =91.3%). Also, the probability that a mammogram would be actually malignant when it is classified to be cancerous by the system is 0.895 (PPV =89.5%) and the probability that the mammogram would be benign when the system says it is not malignant is 0.955, that is, NPV =95.5%.

Conclusion

The significant variability that occurs when interpreting the same mammogram independently by different radiologists

S/N	Authors	Classification category	Sensitivity (%)	Specificity (%)	Accuracy (%)
١.	Lothe et al ¹⁹	NA	92.30	62.50	86.84
2.	Domínguez and Nandi ²⁰	BM	55	85	PPV 0.71
					NPV 0.75
3.	Rejani and Selvi ²¹	NA	88.75	-	-
4.	Moayedi ²²	NA	95.8	99.0	96.60%
5.	Dheeba and Tamil ²³	NA	-	-	86.1%
6.	Kavitha and Thyagharajan ²⁴	BM	100	96	98.00
7.	Zhang et al ²⁵	BM	94.85%	78.20	-
8.	Kamra et al ²⁶	BM	71.43	97.22	93.02
9.	Rouhie et al ²⁷	BM	85.41	91.89	88.65
10.	Damasceno et al ²⁸	BM	99.41	99.84	99.73
11.	Kaur et al ²⁹	NA	-	-	96.90
	Proposed method	NA (BM)	(100) 94.4	(100) 91.3	(100) 92.68

Abbreviations: NA, MIAS classification into normal and abnormal mammograms; BM, MIAS classification into benign and malignant Mammograms; MIAS, Mammographic Image Analysis Society; NPV, negative predictive value; PPV, positive predictive value. leads to diagnostic errors. It is therefore imperative to find an improved method to aid the detection and classification of ROI on mammograms. This article has discussed an automated mammogram classification system that uses a modified SVM classification technique. Performance evaluation results obtained suggested that the system could be used as radiologists' potential tool for supporting decision making in mammogram interpretation. This could in turn help Radiologists to make an accurate and timely decision, thereby increasing the efficiency of their diagnostic skills.

Data availability

Raw mammograms used in this work are publicly available at mini-MIAS: http://peipa.essex.ac.uk/pix/mias/.

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Disclosure

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