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Classification of microcalcification clusters in digital mammograms using a stack generalization based classifier

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Abstract: This paper presents a machine learning based approach for the discrimination of malignant and benign microcalcification (MC) clusters in digital mammograms. A series of morphological operations were carried out to facilitate the feature extraction from segmented microcalcification. A combination of morphological, texture, and distribution features from individual MC components and MC clusters were extracted and a correlation-based feature selection technique was used. The clinical relevance of the selected features is discussed. The proposed method was evaluated using three different databases: OPTIMAM, DDSM, and MIAS. The best classification accuracy (95.00 ± 0.57%) was achieved for OPTIMAM using a stack generalization classifier with 10-fold cross validation obtaining an \( A_z \) value equal to 0.97 ± 0.01.

Keywords: digital mammogram; microcalcification; stack generalization; classification; morphological features.

Note 1.

1. Introduction

Breast cancer is one of the leading causes of cancer death in women [1],[2]. The mortality rate of breast cancer can be reduced by early detection and by using Computer Aided Diagnostic (CADx) systems [3]. Microcalcification (MC) clusters are an important early sign of breast cancer [4]. MC clusters appear as small localized granular points of high brightness within soft breast tissue [5] and it can be difficult to distinguish MC clusters from normal breast tissue because of their subtle appearance and ambiguous margins [6],[7]. Approximately 50% of early diagnosed cases indicated the existence of MC clusters, revealing up to 90% of ductal carcinoma in situ [8]. A typical example of a benign (non-cancerous) and a malignant (cancerous) MC cluster is shown in Fig. 1.

Double reading can improve sensitivity, but a lack of experienced radiologists can be a challenge [9]. CADx can assist radiologists in detecting abnormalities in an efficient way [10],[11] and systems have been developed to provide a second opinion for diagnosis [12]. Previous studies have developed computerized methods to aid the diagnosis of MC clusters. Singh et al. [13] proposed a MC cluster classification technique based on morphology: including size of the calcifications and number of calcifications in a cluster. A region of interest (ROI) around the MC cluster was first enhanced using morphological operations, and two types of features: cluster shape and cluster texture were obtained.

1 The submitted paper is an extended version of the 22nd Medical Image Understanding and Analysis (MIUA) conference paper [11]
A new set of shape features generated by recursive subsampling were added to the feature set, which improved the classification accuracy. Akram et al. [14] proposed an improved Fisher Linear Discriminant Analysis (LDA) approach for the linear transformation of segmented micro-calcification data. In the proposed method, a SVM variant was used to classify benign and malignant clusters. Multi-scale graph topological features were used by Chen et al. [15] using a k-nearest-neighbors classifier. The performance of machine learning techniques was investigated by Rampun et al. [16] by examining the probability outputs from classifiers in conjunction with the classification accuracy and area under the receiver operator curve ($A_z$) to indicate the reliability of CADx.

Bekker et al. [17] proposed a two-phase classification scheme. The method was based on combining decisions from multiple views (craniocaudal (CC) view and mediolateral oblique (MLO) view), implemented by a logistic regression classifier, followed by a stochastic combination of the two view-level (CC and MLO) indications into a final benign or malignant decision. Shachor et al. [18] examined data fusion methods for multi-view MC cluster classification. This data fusion concept was implemented by a special purpose neural network architecture that demonstrated the task of classifying breast microcalcifications as benign or malignant based on CC and MLO mammographic views.

Hu et al. [19] applied a hidden Markov tree model of dual-tree complex wavelet transform (DTCWT-HMT) for microcalcification diagnosis in digital mammograms. DTCWT-HMT was used to capture the correlation between different wavelet coefficients and model the statistical dependencies and non-Gaussian statistics of real signals. The combined features of the DTCWT-HMT and the DTCWT were optimized by a genetic algorithm (GA). An extreme learning machine (ELM) was used as the classifier to diagnose the benign and malignant MC clusters.

A feature selection method was introduced by Diamant et al. [20] based on a mutual information (MI) criterion for automatic classification of MC clusters. The MI based feature selection method was explored for various texture features. Wang et al. [21] used a semi-automated segmentation method to characterize all MCs, and constructed a classifier model to assess the accuracies for microcalcifications and breast masses, either in isolation or combination, for classifying breast lesions. Sert et al. [22], however, used convolutional neural networks along with various preprocessing techniques such as contrast scaling, dilation, cropping etc. to classify microcalcification. Adaptive thresholding and morphological technique was used by Nguyen et al. [23] to segment nuclei for single channel image. A superpixel-based framework was presented for segmentation that used a “hybrid” approach which was intended to integrate the advantage of region-based clustering algorithm and an edge detector with an integrated edge map.

Figure 1. Example MC clusters from the OPTIMAM database: (a) benign MC cluster, (b) malignant MC cluster.
51 features extracted based on size, shape and texture

Figure 2. Overview of the proposed MC cluster classification methodology.

The present work focused on developing a method for discriminating malignant and benign clusters in digital mammograms. Images were first segmented using a wavelet-based method in conjunction with a bi-cubic interpolation technique and a series of morphological operations. A combination of morphological, texture, and distribution features from individual MC components and the MC cluster were extracted and MC clusters were classified with a stack generalization-based classifier. An ensemble classifier was also used to classify MC clusters from digital and digitized mammograms. The most important features were selected and used to classify the MC cluster as benign or malignant. An overview of our proposed approach is presented in Fig. 2.

2. Materials and Methods

2.1. Image databases

We have used the digital mammograms from the OPTIMAM Mammography Image Database [24] which is currently an ongoing project at the Medical Physics Department of the Royal Surrey County Hospital, which contains NHS Breast Screening Programme (NHSBSP) images from different centres across the United Kingdom with an aim to develop a large repository of breast images for research purposes. The database contains 3D and 2D unprocessed and processed breast images, associated annotations and where applicable expert-determined ground truths which describe features of abnormalities like microcalcification, mass, architectural distortions, etc. The images were categorized by radiologists in three clinical categories: normal, benign, and malignant. Core biopsies were also performed where applicable and associated with the opinion provided by the radiologists. In our experiment, patient-based case selection was performed on the digital mammograms, and a total number of 286 cases (136 benign and 150 malignant) were selected, which only contained microcalcification clusters that had associated core biopsy scores. The histological and radiographic scores were not considered for patient-based case selection, as very few images that contained microcalcification clusters were provided with such scores, which was an obstacle to create a balanced database. These mammograms were acquired using a Hologic Selenia mammography unit, with a resolution of 70 microns per pixel and a depth of 12 bits.
Figure 3. Example enhancement of MC clusters: digital mammogram from the OPTIMAM database (top row: 1_1076_463) and digitized mammogram from the DDSM database (bottom row: B_3049_1.RIGHT_MLO): (a) MC patch cropped from the original mammogram (without image enhancement), (b) MC patch cropped after enhancement.

The evaluation also used the digitized mammograms from two different publicly available benchmark databases: the Mammography Image Analysis Society (MIAS) [25], and the Digital Database for Screening Mammography (DDSM) [26]. The DDSM database contains cranial-caudal (CC) and mediolateral oblique (MLO) views of left and right breasts of each patient. The images containing suspicious area have pixel-level "ground truth" information of the abnormality, and a malignancy assessment on a five-point scale according to the American College of Radiology (ACR) Breast Imaging Reporting and Data System (BIRADS) [27]. 280 digitized mammograms containing MC clusters (148 benign and 132 malignant), were used. The MC clusters colocated with masses were not considered, as the existence of mass could mislead the classification results whilst considering the neighborhood of MCs to extract relevant features. The cases were selected at a patient level, and only MLO views were used. The mammograms in the DDSM database were digitised by one of four different scanners: DBA M2100 ImageClear (42 microns per pixel, 16 bits), Howtek 960 (43.5 microns per pixel, 12 bits), Lumisys 200 Laser (50 microns per pixel, 12 bits), and Howtek MultiRad850 (43.5 microns per pixel, 12 bits). For our experiment, only the mammograms obtained using Lumisys 200 Laser scanners were considered to keep inline with the pixel size of another digitised database (MIAS) [25] used for the development and evaluation of the proposed system. The MIAS database [25] contains 322 images among which 24 cases (12 benign and 12 malignant) contain microcalcification clusters. The mammograms in the MIAS
were digitised to 50 microns per pixel. The truth-marking of the locations of the abnormalities were
delineated by an expert radiologist.

2.2. Preprocessing and segmentation

Enhancement was necessary as MC clusters are usually very small, and sometimes can be
situated in dense breast tissue with very low visibility. This phenomenon makes the segmentation
and classification task more difficult and challenging [11]. To overcome this problem, a wavelet-based
algorithm was applied to enhance the mammograms, and the contrast between the MC cluster and
surrounding background tissues was increased, see section 2.2.1. Such contrast enhancement facilitated
the subsequent MC cluster segmentation described in section 2.2.2. Features of MC clusters were
extracted from the segmented image and were used to classify the clusters as benign or malignant.

2.2.1. Mammogram enhancement and patch extraction

A dynamic wavelet-based algorithm [28] was applied to enhance the mammograms. The Discrete
Wavelet Transform (DWT)-based method was used because of its low computational complexity
and special transformed domain properties [29]. The process of mammogram enhancement was
divided into three parts which included decomposition, sharpness estimation and filtering. The image
was first decomposed into individual sub-bands using a multi-level separable DWT [30], [31]. The
log-energies of the vertical, horizontal, and diagonal sub-bands at each decomposition level were

![Figure 4](image_url)

**Figure 4.** (a) Three-dimensional intensity representation of a 158 × 189 pixel area of a digital
mammogram, (b) Calculated object background intensity of the same area, (c) The difference image
between the original image (4a) and the background image (4b).
Figure 5. (a) Enhanced image patch (1_1076_463) from the OPTIMAM database, (b) binary image containing 5% of the highest positive intensity values from the difference image, (c) eliminating single pixels and perform erosion on (b), (d) Image A: pixels having higher value than the specified threshold mentioned in section 2.2.2 are added to (c), (e) contrast enhancement filter applied to the bi-cubic interpolated image of (a), (f) Image B: five percent of the pixels having the highest intensity will be selected from the filtered image, (g) Image C: Logical summation of (d) and (f).

calculated followed by measuring the total log-energy (TLE) of each level. Subsequently, by combining the TLE of each decomposition level [28] the Scalar Sharpness Index (SSI) was calculated. The SSI was later used to estimate the overall sharpness of the images. Higher values of SSI were considered as an indicator of higher sharpness of the image. More details on the wavelet-based enhancement algorithm have been described by Misra et al. [28], where the enhancement approach was applied to satellite images. To enhance the mammograms, the number of sub-bands and the image decomposition level were chosen as 3, as we aimed to obtain the horizontal, vertical and diagonal details from the mammograms. Each sub-band was assigned a predefined weight (0.10) to enhance the diagonal higher spatial frequency. The weight was set to 0.10, as an increase in weight above 0.8 did not provide further increase in enhancement and a weight less than 0.8 provided decay in enhancement. The region containing the MC cluster was cropped, see Fig. 3b, from the enhanced mammogram using the provided annotations. The effect of the enhancement algorithm is shown in Fig. 3b, where it can be noted that the appearance of MC clusters is enhanced for both digital (OPTIMAM) and digitized (DDSM) mammograms from a qualitative point of view.
Figure 6. (a) Elimination of blobs containing one or two pixels from the probability image generated in section 2.2.2 (see Fig. 5g), (b) final probability image, for example case: \(1_{1076}^{463}\), after discarding all blobs from 1 cm\(^2\) pixel block whilst objects inside the block were less than 3. In this example, all the 1 cm\(^2\) pixel blocks contained more than 3 blobs so no object elimination was done.

2.2.2. Probability image generation for MC cluster

A combination of image interpolation, morphological operations, and edge-preserving filtering were applied to generate the probability image of the MC clusters. The enhanced cropped region of interest (ROI), containing the MC cluster, was considered as a three-dimensional plot with the z-axis representing the intensity of each pixel (see Fig. 4a). The whole image was first divided into 30 × 30 sub-regions. The size of sub-regions was set to 30 × 30 to maintain a trade-off between over-segmentation and under-segmentation of the MC clusters. Choosing sub-regions bigger than 30 × 30 would result in over-segmentation in low contrast images where the disparity between the MC cluster and their background is very low. Choosing a size less than 30 × 30 would cause under-segmentation.

Bi-cubic interpolation [32] was applied to each sub-region to obtain pixel intensities of the background tissue: see Fig. 4b. The resulting image, Fig. 4b, was subtracted from the original image, Fig. 4a, to obtain the difference between the original and local background pixel values, Fig. 4c. In Fig. 4b and Fig. 4c, high picks indicate higher pixel intensities and sharp edges in the image. From this difference image (Fig. 4c), the pixels with positive values were identified and a percentage of these (5%) with the highest values were selected to generate a binary image: see Fig. 5b. The reason for selecting the 5% highest pixel values was to avoid under-segmentation. The highest positive pixel values considered as MC clusters were characterized by higher intensity compared to their local background tissue. Single pixels were removed from the generated binary image, and an erosion operation was performed to eliminate false positive pixels, see Fig. 5c. To perform the erosion operation, a square structuring element of size 3 × 3 was used with all values set to one to retain the original morphology of the segmented MC cluster. The lowest value among the 5% selected pixels was specified as a threshold. If the number of the existing pixels, in Fig. 5c, was lower than 10% of the total number of pixels in the cropped image patch, the pixels with intensity higher than half of the previously specified threshold were included in the binary image: see Fig. 5d. Considering the 10% of the total pixels in the cropped image patch will maintain a trade-off between over-segmentation and under-segmentation.

By doing so, enough number of pixels were generated for the binary image (A): see Fig. 5d(d). The above procedure was performed to avoid under-segmentation when the mammogram exhibited very low contrast, which was usually due to erroneous exposure conditions.

Subsequently, a contrast enhancement filter, having a 9 × 9 kernel with its central pixel element equal to 80, was applied to the bi-cubic interpolated image [32]: see Fig. 5e. Five percent of the pixels having the highest intensity were selected from the filtered image, producing another binary image (B), see Fig. 5f. Finally, logical summation (AND) of the two binary images A and B (Fig. 5e and Fig. 5f)
was performed to keep pixels that have high intensity values in comparison with the background intensity of their local neighbourhood tissues: see Fig. 5g.

2.2.3. Specifying MC cluster

The clinical definition of the MC cluster was used for the reduction of false positives from the probability image generated in section 2.2.2. According to the medical definition of clustered MC, more than 3 MCs should reside in a 1 cm$^2$ area [33], which is equivalent to 200 × 200 pixels in the digitized data (DDSM and MIAS) with a pixel size equal to 50 µm, and 143 × 143 pixels in the digital data (OPTIMAM) with a pixel size equal to 70 µm. This results 143 × 143 pixel equivalent to 1 cm$^2$ block area for OPTIMAM, and 200 × 200 pixel equivalent to 1 cm$^2$ block area for DDSM and MIAS.

From the probability image generated in section 2.2.2 (see Fig. 5g) regions containing one or two pixels were removed, as they were considered artifacts [35], and an erosion operation with a 2 × 2 unit element kernel was performed: see Fig. 6a. Here, a 2 × 2 unit element kernel was used for the erosion operation, as a bigger kernel size generated under-segmented images and a smaller kernel had barely any effect. Removal of individual objects with a morphological erosion operation was necessary, because the diagnostic information was based on the existence of a group of MCs [33]. Subsequently, neighbouring pixels with eight connectivity were grouped together [11] and considering the clinical definition of MC cluster formation, the binary image having only 8-connected component was divided into 1 cm$^2$ block areas. This results 143 × 143 pixel equivalent to 1 cm$^2$ block area for OPTIMAM, and 200 × 200 pixel equivalent to 1 cm$^2$ block area for DDSM and MIAS.

Elimination of all the elements inside each 1 cm$^2$ block area were done, where the minimum number of objects inside a block was less than 3 [33], all the elements were removed: see Fig. 6b. In Fig. 6b, no object elimination was done inside any block since all the 1 cm$^2$ blocks contained more than 3 objects; a sample case is shown in Fig. 7, which represents how the images were divided into 1 cm$^2$ block areas, and the elements inside each block were eliminated, where the minimum number of objects inside the block was less than 3 [33]. For better visual understanding, the MC clusters were highlighted in yellow (see Fig. 7c and Fig. 7d) and green (see Fig. 7e and Fig. 7f). Image C was generated for the sample image patch (10_35_242) from the OPTIMAM database (Fig. 7b). All single pixels were eliminated to remove a fraction of false positive MC objects (Fig. 7c). The image was then divided into 1 cm$^2$ pixel blocks, see Fig. 7d. The blocks containing less than 3 MCs, marked by a rectangle, were removed, see Fig. 7e. All the blocks were stitched together to generate the final segmented image (Fig. 7e).

The whole MC cluster may not be covered by the proposed approach. The block area has to be slid to different locations of the patch image to build-up a complete MC cluster network. For the sliding window approach, we would have to come up with a methodology to harmonize the changes in MC clusters between windows and how this representation is affecting the classification. In addition, the sliding window approach would be time consuming, and is an interesting research question to address in future.

3. Segmentation evaluation

The evaluation was carried out using the Dice similarity metric [37] [38], and is in line with our previous work [11]. The reference masks (see Fig. 8b), were generated from the radiologist's annotation outline: see Fig. 8a. Subsequently, individual MCs that reside inside the radiologist's annotation were considered to generate convex hull. This convex hull (see Fig. 8f) and the reference mask (see Fig. 8b), were used to calculate the Dice similarity score (see (Fig. 8g - 8i)). The Dice similarity metric for DDSM and MIAS is presented in Fig. 9.

From Fig. 9, it is clear that the segmentation technique based on the morphological approach works better than the area-rank based segmentation method proposed in [11]. Also, it is to be noted that, the segmentation results generated by applying the method of Oliver et al. [39] gives almost the same similarity score as gained by our proposed morphological operation-based segmentation method,
Figure 7. (a) Enhanced image patch (10_35_242) from the OPTIMAM database, (b) Image C: logical summation of two binary images A and B for image patch (10_35_242), (c) eliminating single pixel from (b) (all MCs are highlighted for better visual understanding), (d) dividing (c) into 1 cm$^2$ pixel blocks: the blocks containing fewer than 3 MCs are marked by a rectangle, the last row and the last column of image blocks were not 1 cm$^2$ pixel block as they were adjusted according to the patch image size. (e) elimination of all MCs inside each 1 cm$^2$ pixel block that contained fewer than 3 MCs (marked by a rectangle), (f) all blocks in (e) are stitched together to produce the final segmented image.

though the similarity score for our proposed approach is slightly higher than with Oliver’s method [39].

4. Classification module construction

To classify MC clusters into benign or malignant, a series of classification algorithms were explored to create an ensemble learner instead of using only one classification method. A set of nine different machine learning algorithms were used, which included k-nearest neighbour (kNN) classification [36], a multilayer perception (MLP) classifier [40], a classification tree [41], random forest [42], support vector machines: using four different kernels (Gaussian RBF, sigmoid, linear, and polynomial) [43], and a Naive Bayes network [44]. All the classifiers individually provide a binary decision by classifying the images as benign or malignant. Each classification algorithm was separately applied to the images and the number of malignancy predictions (votes for malignancy) were counted. Afterwards, the total number of malignancy prediction was divided by the total votes. For example, if eight of the nine classifiers classified a case as malignant, then the final estimation of the ensemble classifier for malignancy will be 89%. The advantage of employing an ensemble classifier was to aggregate a set of models to provide more robust classification results rather than using the opinion from a single classification model. The predictions from individual classifiers were combined using majority voting, and as such the possibility of over-fitting of any particular classifier was avoided. The individual classification results from different classification algorithms will be presented and discussed in section 6.
Figure 8. (a) Annotation by radiologist (B_3121_1.RIGHT_MLO), (b) reference MC cluster mask generated from (a), (c) border extraction from reference MC mask and overlaid on segmented image generated using morphological segmentation approach, (d) MC resides inside the border annotated by expert radiologist, (e) convex hull outline using the border points of segmented blobs residing inside annotation outline, (f) mask generation from convex hull border of segmented image, (g) Dice similarity score (based on morphological segmentation approach) = 0.85599; White region= True positive, Green region= False positive, Magenta region= False negative, (h) Dice similarity score (based on Oliver’s \[39\] segmentation approach) = 0.76514, (i) Dice similarity score (based on area ranking segmentation approach) = 0.5494.

A stacked generalization [46] approach was also applied to create a classifier for classifying the MC clusters. In this approach, the above-mentioned nine different learning algorithms were considered as base classifiers, and the Naive Bayes classifier [44] was used as the meta-classifier (combiner), as previous experiment [45] confirmed that the Naive Bayes classifier as a combiner performed better than majority voting. In a stacked generalization approach, the meta-learner was used instead of averaging to combine predictions of the base classifiers. Predictions of the base classifiers were used as input for the meta-classifier. The meta-classifier attempted to learn the relationships between predictions and the final decision. The meta-classifier also corrected some mistakes of the base classifiers [46].

The aim of this research was to investigate the merit of using a conventional stack generalization approach to classify MC cluster in mammogram. Using modern methods such as auto-encoders or
5. Feature extraction and feature selection

It is crucial to extract and select appropriate features that can classify MC clusters into their clinical categories. MC clusters can be assessed based on specific properties such as: size, shape, number, distribution, etc. [32]. A set of 51 features [51] were computed from the segmented blobs, see section 2.2.3, for extracting the statistical and morphological properties of the MC clusters, which form the feature space. All the computed features characterize either an individual MC or an MC cluster. These features were grouped into three categories: shape, size and texture, see Table 3. Since the number of computed features were large and their discriminating power varied, see Table 3, a feature selection approach was used to obtain the most salient features. More details on the performance of individual features to classify MC clusters are discussed in section 6.

Feature selection was done by employing the CfsSubsetEval [52] attribute evaluator and the BestFirst search method [53] in Weka [53]. CfsSubsetEval [52] evaluated the significance of a subset of features by approximating the individual predictive ability of each feature and the redundancy between them: this meant that features that were highly correlated with the class whilst having low inter-correlation were more likely to be selected [53]. On the other hand, BestFirst [53] searched the feature space subsets by greedy hill-climbing augmented with a backtracking facility [53], which could start from any point and search forwards and backwards, by considering all possible single feature vector additions and deletions [55]. The selected features from unenhanced images were put into a group (α). Subsequently, the same 51 features were extracted from the segmented images that were generated from the enhanced mammograms. The most significant features from the enhanced images were gathered into another group (β), using the same feature selection technique. The common features from group α and group β formed a new feature space.

To ensure the robustness of the feature selection and avoid bias, all the data was divided using 10-fold cross-validation scheme and 9-fold cross-validation scheme respectively. Important features were extracted using the images residing in each fold which showed the same features extracted consistently. When the images were split into different number of groups by changing the fold-number higher and lower than 10, we constantly obtain the same set of features extracted. A similar approach was applied to measure the robustness of the feature selection in a previous publication [51].

The feature extraction and selection technique, as previously mentioned, was applied separately on the digitized and digital databases to investigate whether the provided features from the digital database outperformed those extracted from the digitized database in classifying MC clusters. Table 1
represents the 4 most important features extracted and selected using Digitized database (DDSM), and Table 2 represents the 2 most important features extracted and selected using the Digital database (OPTIMAM) with the associated clinical interpretations.

Table 1. Clinical description of the selected features using the DDSM database for classification of MC clusters.

<table>
<thead>
<tr>
<th>MC cluster classification features</th>
<th>Radiologists characterization features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Summation of the mean of individual MC intensity</td>
<td>Density of MC cluster</td>
</tr>
<tr>
<td>Variance of the standard deviation of the distances from cluster centroids</td>
<td>MC distribution</td>
</tr>
<tr>
<td>MC cluster convex hull area</td>
<td>Cluster size</td>
</tr>
<tr>
<td>Mean of MC perimeter</td>
<td>Individual MC size</td>
</tr>
</tbody>
</table>

Table 2. Clinical description of the selected features using the OPTIMAM database for classification of MC clusters.

<table>
<thead>
<tr>
<th>MC cluster classification features</th>
<th>Radiologists characterization features</th>
</tr>
</thead>
<tbody>
<tr>
<td>MC cluster area</td>
<td>Cluster size</td>
</tr>
<tr>
<td>Size of individual MC</td>
<td>Individual MC size</td>
</tr>
</tbody>
</table>

The in-depth details on the impact of our feature selection approach are described in section 6. Here, all the images were segmented maintaining the clinical grounding of the distribution of the MC cluster which indicate that an area of 1 cm² contains no fewer than 3 MCs [33]. The spatial resolution of mammography is normally ranging from 40—100 µm per pixel, which enables detection of MC clusters at an early stage[15]. The aforementioned feature extraction and selection method was also employed on the segmented images from the digital and digitized databases by randomly considering a 100 × 100 pixel area as 1 cm², to investigate if this had an impact on the MC cluster classification. The results are presented in Table 6 in section 6.

To evaluate the reliability of the feature selection approach, images from the digital and digitized databases were separately divided into ten folds. The process of feature selection was performed on each fold which indicated the same selection of features. Detailed evaluation of the feature selection for MC cluster classification is provided in section 6.

6. Result analysis

To investigate the influence of shape, size, and texture aspects, each individual feature type was separately used for the classification using ensemble learning, see Table 3. The experiment was separately applied on the individual databases, where the features cognate with size provided the highest \( A_z \) values over the shape and texture features for both digital and digitised databases with no feature selection. Whilst only considering the size features, the highest \( A_z \) value (0.87 ± 0.01) was gained for the digital database (OPTIMAM). With feature selection, as described in section 5, the value of \( A_z \) was 0.83 ± 0.01 for OPTIMAM, 0.72 ± 0.01 for DDSM, and 0.68 ± 0.02 for MIAS. The most important size features were related to the area covered by individual MC, eccentricity of individual MCs, eccentricity of MC cluster, MCs distances covered from MC cluster centroid, perimeter of MC cluster, and elongation of MC cluster.
We used 10-fold cross-validation with different seed values. The seed values initialize
randomization of data in each fold. For example, if the value is set to 3, it means that the data
was shuffled among the folds 3 times. Saving the seed value or setting it to the same number each
time guarantees that the algorithm will come up with the same results- identical for each run. In
this experiment, the seed number was set to 1 for the first run and its value was increased by 1 with
each run. Hence, for 10 runs, the maximum seed value was set to 10. In 10-fold cross-validation, the
original sample is randomly partitioned into 10 equal size sub samples (folds). Of the 10 sub samples,
a single sub-sample is retained for testing the model, and the remaining (10-1) sub-samples were
used as training data. The cross-validation process is then repeated 10 times, with each of the folds
used exactly once as the test data. The 10 results from the folds were averaged to produce a single
estimation. The advantage of this method is that all observations are used for both training and testing,
and each observation is used for testing exactly once.

Note that the feature selection was only performed on the training data and therefore it is not
expected that overfitting will happen. By using stratified 10-fold cross-validation we avoided the
risk of over-training. When using the ensemble learning and stack generalization approach, the
hyper-parameters were kept as the default parameters set in Weka, since the advantage of using
default parameters is that we eliminate the risk of introducing optimistic bias by tuning the parameter
to maximize performance \[54\]. The segmentation and feature extraction was implemented using
MATLAB Version: 9.3.0.713579 (R2017b) on Windows 10. The features extracted from the images were
converted from ".mat" format to ".arff" format to facilitate data structures as input for WEKA.

Table 3. A\textsubscript{2} estimation for the classification of MC clusters while applying 10-fold CV using ensemble
learning on segmented image using a block size based on clinical rules.

<table>
<thead>
<tr>
<th>Feature selection</th>
<th>Feature Category</th>
<th>No. of feature</th>
<th>Total feature No.</th>
<th>(A_2) (AUC) DDSTEM MIAS</th>
</tr>
</thead>
<tbody>
<tr>
<td>No</td>
<td>Size</td>
<td>17</td>
<td>51</td>
<td>0.87 ± 0.01</td>
</tr>
<tr>
<td></td>
<td>Shape</td>
<td>17</td>
<td></td>
<td>0.70 ± 0.02</td>
</tr>
<tr>
<td></td>
<td>Texture</td>
<td>17</td>
<td></td>
<td>0.77 ± 0.01</td>
</tr>
<tr>
<td>Yes</td>
<td>Size</td>
<td>7</td>
<td>12</td>
<td>0.83 ± 0.01</td>
</tr>
<tr>
<td></td>
<td>Shape</td>
<td>4</td>
<td></td>
<td>0.71 ± 0.01</td>
</tr>
<tr>
<td></td>
<td>Texture</td>
<td>5</td>
<td></td>
<td>0.78 ± 0.02</td>
</tr>
</tbody>
</table>

![Figure 10](image1.png)

Figure 10. The accuracy of microcalcification cluster classification by individual classifiers: (a)
classification accuracy for K-nearest neighbour, Multilayer perception, J48 decision tree, Random
forests, Naive bayes, (b) classification accuracy by SVM using four different kernels: Gaussian RBF,
Sigmoid, Linear, and Polynomial.
All nine classifiers, described in section 4, were tested individually to assess their performance with results shown in Fig. 10. SVM provided very low classification accuracy compared to the other classifiers, which is caused by low bias and high variance [56]. Another point to note is the SVM trained classifier used the trained data partly to estimate the margin, the support vectors, whereas other function classifiers consider the training set to define the decision function, making them more generalizable. When SVM was discarded from the classifier stack the overall classification preformance decreased [11], while including SVM resulted in improved classification accuracy (around 90% for the DDSM database) [11] indicating the positive influence of SVM on ensemble learning, where a majority voting scheme was applied for improved generalization and to gain more flexibility to maintain strong prediction performance by averaging out classifiers individual mistakes and thus reducing the risk of over-fitting.

For the k-nearest neighbour (kNN) classifier, Fig. 10a, the value of k was set to 5 based on cross-validation. The classification accuracy for digitized and digital mammograms was 93.77% and 81.37%, respectively. Lower value of k caused a decrease in classification accuracy and values higher than 5 provided the same accuracy as for k=5. For a multilayer perceptron (MLP), the number of attributes were summed up with the number of classes and the result was divided by 2 to set the number of hidden layers whilst using the learning rate 0.3 and setting the validation threshold as 20 to terminate the validation testing. Such parameter settings were chosen because it provided the best classification accuracy for digital mammograms (around 84%), but the classifier showed poorer performance for digitized mammograms (around 73% classification accuracy). It is to be noted that, the accuracy was increased to above 92% for both digital and digitized mammograms whilst using a classification tree, i.e. C4.5 (J48). Here, the confidence value was chosen to be 0.25 for pruning and the number of folds was set to 3; in order to determine the amount of data for reduced-error pruning and producing a decision tree. While applying a random forest, the accuracy for digitized mammograms was 84%, but the accuracy for digital mammograms was above 90%. The Naive Bayes classifier provided an increase in classification accuracy for digital mammograms (around 92%), but the accuracy decreased to around 76% for the digital database.

All the classifiers were used to create an ensemble learner (see section 4). The ensemble learner was applied to images from the three different databases: OPTIMAM, DDSM, and MIAS. The performance of the ensemble learner is presented in Table 4. 10-fold cross-validation (10-FCV) scheme and leave-one-out cross-validation (LOOCV) approach were used. For 10-FCV, the images were splitted into 10 folds ensuring that each fold has the same proportion of observations with a given categorical value. In our experiment, each fold contains roughly the same proportions of the two types of class labels (benign and malignant). 10-FCV allows to use different training and testing data which will avoid the over fitting and give better generalization ability. On the other hand, for LOOCV, each observation was held out with training based on the remaining samples.

Two evaluation metrics were used. The first evaluation metric was the overall classification accuracy (CA), which was defined as the percentage of correctly classified MC clusters. The receiver operating characteristic (ROC) curve analysis was used as the second evaluation metric, plotting the true positive rate (TPR) against the false positive rate (FPR) which illustrated a whole range of possible operating characteristics for the classifier model. The ROC analysis was used to assess the predictive ability of the ensemble learner by using the area under the ROC curve denoted by A2 (also know as the AUC) [57] (see Fig. 11). A2 is equivalent to the Wilcoxon signed-ranks test, which is a nonparametric alternative to the paired t-test [58]. All the classification and evaluation aspects were implemented using the Weka [59] data mining suite.

When using 10-FCV, in Table 4, the ensemble learner performed better using only 2 important features which were extracted and selected from the digital database (OPTIMAM) showing an accuracy equal to 89.80 $\pm$ 1.98%. The feature selection was performed using the proposed method described in section 5. The most important 2 features were related to the MC cluster area and size of individual MC. Increase in accuracy was also noticed while using the same 2 important features to classify MC cluster
Table 4. Classification accuracy using LOOCV and 10-fold CV applying all 51 and the 2 most salient features from digital mammogram, and 4 most salient features from the digitized mammogram using ensemble learning. The images were segmented following the clinical grounding of cluster distribution.

<table>
<thead>
<tr>
<th>Database name</th>
<th>Feature number</th>
<th>LOOCV</th>
<th>10-FCV</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>CA</td>
<td>A_2 (AUC)</td>
</tr>
<tr>
<td>OPTIMAM</td>
<td>51</td>
<td>86.49%</td>
<td>0.85</td>
</tr>
<tr>
<td>(286)</td>
<td>4</td>
<td>85.71%</td>
<td>0.84</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>91.12%</td>
<td>0.91</td>
</tr>
<tr>
<td>DDSM</td>
<td>51</td>
<td>73.98%</td>
<td>0.73</td>
</tr>
<tr>
<td>(280)</td>
<td>4</td>
<td>80.66%</td>
<td>0.80</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>88.48%</td>
<td>0.88</td>
</tr>
<tr>
<td>MIAS</td>
<td>51</td>
<td>82.35%</td>
<td>0.79</td>
</tr>
<tr>
<td>(24)</td>
<td>4</td>
<td>100.00%</td>
<td>1.00</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>100.00%</td>
<td>1.00</td>
</tr>
</tbody>
</table>

Figure 11. ROC curves for a stack generalization classifier for the OPTIMAM digital database: (a) 2 features after feature selection (AUC = 0.97), and (b) 51 features (AUC = 0.96).

for the digitized mammograms (85.24 ± 2.52% for DDSM, and 100.00 ± 0.00% for MIAS) compared to Table 3. When considering only the selected two important features, it is investigated that the classification accuracy is lower for the digitized database (DDSM) then the accuracy achieved for the digital database (OPTIMAM). A possible reason for such decrease in accuracy for the digitized mammograms is due to the decreased image quality compared to the digital mammograms, which affected the accuracy of the MC segmentation [60]. As the digital mammograms were higher quality, more accurate segmentation was obtained which potentially influenced appropriate feature extraction and classification results [61]. The accuracy was also high for the same selected features when using the LOOCV scheme: 91.12% for OPTIMUM, 88.48% for DDSM, and 100% for MIAS. Such limitations of digitized mammograms were more pronounced when using 4 important features, extracted and selected from the digitized database (DDSM) using method explained in section 5, and showed decreased accuracy when compared with the selected 2 features from the digital database (OPTIMAM): Table 4.

The stacked generalization approach [46] was applied to create an additional classifier, described in section 4. The outputs of the nine different learning algorithms were collated to model a new dataset. The Naive Bayes classifier [44] was used as the meta-classifier to provide the final classification results [62]. The meta learner was used instead of averaging to combine the predictions of the base classifiers, which provided classification accuracy of 95.75% for the digital (OPTIMAM) database, and classification accuracy of 95.17% for digitized database (DDSM) when applying only 2 important features that were extracted and selected from the digital database (OPTIMAM) whilst using LOOCV.
Table 5. Classification accuracy using LOOCV and 10-fold CV applying all 51 and the 2 most salient features from digital mammogram, and 4 most salient features from the digitized mammogram using stacked generalization. The images were segmented following the clinical grounding of cluster distribution. Naive Bayes was used as the meta-classifier.

<table>
<thead>
<tr>
<th>Database name</th>
<th>Feature number</th>
<th>LOOCV</th>
<th>10-fold CV</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>CA</td>
<td>A_z (AUC)</td>
</tr>
<tr>
<td>OPTIMAM (286)</td>
<td>51</td>
<td>91.89%</td>
<td>0.97</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>92.66%</td>
<td>0.98</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>95.75%</td>
<td>0.97</td>
</tr>
<tr>
<td>DDSM (280)</td>
<td>51</td>
<td>89.96%</td>
<td>0.95</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>92.19%</td>
<td>0.96</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>95.17%</td>
<td>0.98</td>
</tr>
<tr>
<td>MIAS (24)</td>
<td>51</td>
<td>100%</td>
<td>1.00</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>100%</td>
<td>1.00</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>100%</td>
<td>1.00</td>
</tr>
</tbody>
</table>

Comparing Table 4 and Table 5 signifies that the ensemble learner performs poorly providing a decrease in the classification accuracy in all considered cases. This strongly supports the statement that the digital mammograms were higher quality, and more accurate segmentation was obtained which potentially regulate appropriate feature extraction and classification results [61]. It is worthy noting that even though 100% classification accuracy was obtained for the MIAS dataset, the number of sample in MIAS is very small (24 women: 12 benign, and 12 malignant) to draw a significant conclusion in terms of classifying MC cluster, as it has smaller variability then the larger database like DDSM.

The results presented in Table 3, Table 4, and Table 5 are based on the images segmented maintaining the clinical grounding of the distribution of the MC cluster which indicates that an area of 1 cm² contains no less than 3 MCs [33]. Since the spatial resolution of mammography was 40—100 μm per pixel which enabled the detection of MC clusters at an early stage [15]—the feature extraction and selection method presented in section 5 was employed on the segmented images from the digital and digitized databases that treated a 100 × 100 pixel block equivalent to a 1 cm² area. This was done to investigate if such size selection had an impact on the MC cluster classification. The 100 × 100 pixel block is 50% of block size (200 × 200) that was maintained to segment the digitized database (DDSM and MIAS) and 70% of block size (143 × 143) that was maintained to segment the digital database (OPTIMAM). In Table 6, both 51 features, and the selected 4 most important features extracted from the digitized mammogram (DDSM) were used for MC cluster classification using LOOCV and 10-fold CV scheme. Here, the images were segmented, using the approach mentioned in section 2.2.2, without following the clinical grounding of cluster distribution by selecting the block size 100 × 100 to investigate if it had any effects on the MC cluster classification.

The selected 4 most important features provided higher classification accuracy while applying LOOCV and 10-fold CV for the OPTIMAM database (95.77% for LOOCV, and 94.94 ± 0.90% for 10-fold CV), the DDSM databases (93.91% for LOOCV, and 93.98 ± 0.87% for 10-fold CV), and the MIAS database (100% for LOOCV, and 100.00 ± 0.00% for 10-fold CV). Observing that the MIAS provided 100% classification accuracy with the 4 most important features it had a very limited number of samples to draw significant conclusions. The increase in accuracy for the OPTIMAM database
Table 6. Classification accuracy using LOOCV and 10-fold CV applying all 51 and the 4 most salient features from digitized mammogram using stacked generalization. The images were segmented without following the clinical grounding of cluster distribution. Naive Bayes was used as the meta-classifier.

<table>
<thead>
<tr>
<th>Database name</th>
<th>Feature number</th>
<th>LOOCV</th>
<th>10-FCV</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>CA</td>
<td>A_c (AUC)</td>
</tr>
<tr>
<td>OPTIMAM (286)</td>
<td>51</td>
<td>93.66%</td>
<td>0.97</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>95.77%</td>
<td>0.98</td>
</tr>
<tr>
<td>DDSM (280)</td>
<td>51</td>
<td>90.68%</td>
<td>0.96</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>93.91%</td>
<td>0.97</td>
</tr>
<tr>
<td>MIAS (24)</td>
<td>51</td>
<td>100%</td>
<td>1.00</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>100%</td>
<td>1.00</td>
</tr>
</tbody>
</table>

with the 4 most important features over the 51 features derived from the digitized database (DDSM) warrant that the selected features from the digitized database (DDSM) have influence in classifying MC clusters in the digital mammograms (OPTIMAM). This also demonstrated that the feature selection approach proposed in section 5 is robust.

It is noteworthy that, whilst using 10-fold CV, the classification accuracy 94.94 ± 0.90% for the OPTIMAM database using the 4 most important features in Table 6, and the classification accuracy 95.75 ± 0.57% for the same database using the 2 most important features in Table 5 appears to be similar. The same applied when comparing the classification accuracy for the DDSM database. With 10-fold CV and the 4 most important features, Table 6, the DDSM database achieve 93.98 ± 0.87% classification accuracy, since with the 2 most important features and 10-fold CV, in Table 5, the DDSM database obtain an accuracy of 94.90 ± 0.72%. The precision was calculated using an unpaired t-test for the aforementioned circumstances and a p value of p>0.05 was obtained in all cases. This exhibits that similar classification accuracy can be achieved for classifying MC cluster using more number of features (4 most important features) when the feature extraction and selection is performed on digitized database, whereas less features (2 most important features) can be used to obtain similar classification accuracy (around 95%) when the feature extraction and selection is performed on digital database, and the MC are segmented complying the clinical groundings concerning the cluster distribution.

7. Discussion

The proposed method for MC cluster classification was compared with other relevant publications, see Table 7. Akram et al. [12] proposed a tree-based representations for MC clusters, where scale-invariant topological features of MC were extracted showing 91% accuracy for cluster classification. Though high accuracy was achieved, the performance for MC cluster classification on digital mammogram was not reported in this study. In another study by Akram et al. [14], 96% classification accuracy was achieved using digitized mammograms with an improved Fisher Linear Discriminant Analysis (LDA) approach combined with a Support Vector Machine (SVM) variant.

The properties of MC clusters were presented by mereotopological barcodes by Strange et al. [60], where the discrete mereotopological relations between the individual MCs over a range of scales were presented in the form of a mereotopological barcode. The classification accuracy on digitized mammograms reported by Strange et al. [60] was 95% and 80% for the MIAS and DDSM datasets, respectively.

Chen et al. [15] used multi-scale graph topological features and classified MC clusters using k-nearest-neighbours-based classifiers. Their approach obtained 96% accuracy for digital mammograms. Though the accuracy for digital mammogram was high, the number of cases in the digital mammogram database was very low (25 cases), which provided less variability of MC distribution in the sample cases. It is also noteworthy that the digital images were manually annotated...
Table 7. A qualitative comparison of our results with respect to related work.

<table>
<thead>
<tr>
<th>Method</th>
<th>Databases</th>
<th>Cases</th>
<th>Features</th>
<th>Classifier</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Akram et al. [12]</td>
<td>DDSM</td>
<td>288</td>
<td>Tree-based modeling</td>
<td>tree-structure height</td>
<td>CA = 91%</td>
</tr>
<tr>
<td>Akram et al. [14]</td>
<td>DDSM</td>
<td>288</td>
<td>Scalable-LDA</td>
<td>SVM</td>
<td>CA = 96%</td>
</tr>
<tr>
<td>Strange et al. [60]</td>
<td>DDSM</td>
<td>150</td>
<td>Cluster</td>
<td>barcodes</td>
<td>CA = 95%, A2 = 0.82</td>
</tr>
<tr>
<td>Strange et al. [60]</td>
<td>MIAS</td>
<td>20</td>
<td>Cluster</td>
<td>barcodes</td>
<td>CA = 80%, A2 = 0.80</td>
</tr>
<tr>
<td>Chen et al. [15]</td>
<td>MIAS I (Manual Annotation)</td>
<td>20</td>
<td>Topology</td>
<td>kNN/FNN</td>
<td>CA = 95%, A2 = 0.96</td>
</tr>
<tr>
<td>Chen et al. [15]</td>
<td>Digital</td>
<td>25</td>
<td>Topology</td>
<td>kNN/FNN</td>
<td>CA = 96%, A2 = 0.96</td>
</tr>
<tr>
<td>Chen et al. [15]</td>
<td>DDSM (LOOCV)</td>
<td>300</td>
<td>Topology</td>
<td>kNN</td>
<td>CA = 95%, A2 = 0.96</td>
</tr>
<tr>
<td>Chen et al. [15]</td>
<td>DDSM (10-fold CV)</td>
<td>300</td>
<td>Topology</td>
<td>kNN</td>
<td>CA = 85.2 ± 0.05%</td>
</tr>
<tr>
<td>Alam et al. [11]</td>
<td>MIAS (LOOCV)</td>
<td>24</td>
<td>Morphology, Texture &amp; Cluster</td>
<td>Ensemble classifier</td>
<td>CA = 100%, A2 = 1</td>
</tr>
<tr>
<td>Alam et al. [11]</td>
<td>MIAS (10-fold CV)</td>
<td>24</td>
<td>Morphology, Texture &amp; Cluster</td>
<td>Ensemble classifier</td>
<td>CA = 100 ± 0.00%, A2 = 1.00 ± 0.00</td>
</tr>
<tr>
<td>Alam et al. [11]</td>
<td>DDSM (LOOCV)</td>
<td>280</td>
<td>Morphology, Texture &amp; Cluster</td>
<td>Ensemble classifier</td>
<td>CA = 91.39%, A2 = 0.91</td>
</tr>
<tr>
<td>Alam et al. [11]</td>
<td>DDSM (10-fold CV)</td>
<td>280</td>
<td>Morphology, Texture &amp; Cluster</td>
<td>Ensemble classifier</td>
<td>CA = 90.02 ± 1.42%, A2 = 0.89 ± 0.02</td>
</tr>
<tr>
<td>Ours</td>
<td>OPTIMAM (10-fold CV)</td>
<td>286</td>
<td>Morphology, Texture &amp; Cluster</td>
<td>Ensemble classifier (Extended)</td>
<td>CA = 90.97 ± 0.83%, A2 = 0.91 ± 0.01</td>
</tr>
<tr>
<td>Ours</td>
<td>OPTIMAM (10-fold CV)</td>
<td>286</td>
<td>Morphology, Texture &amp; Cluster</td>
<td>Stack generalization (meta-classifier: Naive Bayes)</td>
<td>CA = 89.84 ± 1.69%, A2 = 0.96 ± 0.00</td>
</tr>
<tr>
<td>Ours</td>
<td>OPTIMAM (10-fold CV)</td>
<td>286</td>
<td>Morphology, Texture &amp; Cluster (selected features)</td>
<td>Stack generalization (meta-classifier: Naive Bayes)</td>
<td>CA = 95.75 ± 0.57%, A2 = 0.97 ± 0.01</td>
</tr>
<tr>
<td>Ours</td>
<td>OPTIMAM (10-fold CV)</td>
<td>286</td>
<td>Morphology, Texture &amp; Cluster (selected features)</td>
<td>Stack generalization (meta-classifier: Adapting Boosting)</td>
<td>CA = 96.72 ± 0.46%, A2 = 0.98 ± 0.00</td>
</tr>
</tbody>
</table>

in Chen et al. [15], where delicate lines around small microcalcifications were outlined by an expert radiologist. Such delicate annotation with no false positives might result in higher classification accuracy. In Chen et al. [15], high accuracy, around 95%, was also achieved for a digitised database (MIAS) whilst again considering only a very small number of cases providing limited variation of MC clusters. Conversely, while using a large image database (DDSM), the classification accuracy reduced to 86% for a LOOCV approach and 85.20 ± 0.05% for 10-fold CV. It is worth mentioning that only topological features were taken into account to classify MC clusters, rather than concentrating on the morphological and statistical features of the MC clusters.

In our previous study [11], we acquired high classification accuracy (100%) for the MIAS database (24 cases) using LOOCV and 10-fold CV with an ensemble classifier. For DDSM, the accuracy was
91% (for LOOCV) and 90.02 ± 1.42% (for 10-fold CV). The images used in Alam et al. [11] did not maintain the clinical grounding while segmenting the MC cluster using block processing approach. Also, the experiment was not evaluated on digital mammograms. Promising results were achieved by our developed approach using the images from the digital and digitised databases (OPTIMAM, DDSM, and MIAS). For brevity, we have only shown the results for the OPTIMAM database in Table 7. The comparison of MC classification accuracy for the OPTIMAM database with respect to the DDSM and MIAS databases is represented in Table 3, Table 4, Table 5, and Table 6 in section 6. Whilst using an ensemble classifier for the OPTIMAM database, 87.11 ± 1.38% classification accuracy was achieved, see Table 4. For the DDSM database, the accuracy achieved was 76.28 ± 1.25% for 10-fold CV, which was lower than for the OPTIMAM database. The stack generalization approach, described in section 4, was applied, which provided 89.85 ± 1.69% accuracy without feature selection, and 95.75 ± 0.57% accuracy with feature selection for the OPTIMAM database, see Table 5. To perform quantitative evaluation for the stack generalization classifier, the receiver operating characteristic (ROC) curves for 2 features (Table 5) and 51 features are represented in Fig. 11. Using ROC analysis, we achieved an area under the ROC of $A_z = 0.97$ when using 2 features, whereas for 51 features the value of $A_z$ was 0.96. $A_z$ is equivalent to the Wilcoxon signed-ranks test and a statistical measure, which is a non-parametric alternative to the paired t-test [49], [50]. Additional details on feature selection will be described in section 5. A detailed discussion of the results can be found in section 6.

In addition to this, Table 5 and Table 6 in section 6 reveals that the stack generalization scheme outperformed the ensemble learning approach to classify MC clusters for both the digital and digitized mammograms using LOOCV and 10-fold CV approaches.

Apart from the classifiers that are described in section 4, additional classification algorithms ([63], [64], [65]) were added to construct an extended ensemble learner which provided better classification accuracy 90.97 ± 0.83% for the OPTIMAM database (See Table 7) compared to the accuracy (87.11 ± 1.38%) obtained by the ensemble learner initially used in Table 4 using 10-fold CV.

In Table 7, 95.95 ± 0.57% accuracy was achieved with stack generalization with meta-classifier as Naive Bayes [44]. This accuracy was increased to 96.72 ± 0.46% when using Adaptive boosting [66] as meta-classifier. The Adaptive boosting improved the performance accuracy as it produced a combined classifier whose variance is lower than the variances produced by the weak base learner [67].

It should be noted that most publications in Table 7 used smaller datasets, hence Table 7 represents a qualitative comparison. Table 7 shows how different classifiers classify MC clusters using different types of features. The methods were tested with different settings and data splitting. It is also important to note that the segmented images used in other classification approaches were based on the method proposed by Oliver et al. [39], whereas our proposed classification approach was based on the images segmented using the method proposed by Alam et al. [11], which is why the number of images from the same database in different experiments varies since the under-segmented images generated from the method proposed by Alam et al. [11] were discarded in our experiments. One significant drawback of the developed method was that it performed badly for cases where the MC clusters have no well-defined structure or very few MC were segmented in the cluster region. An extreme situation occurred when only a single MC was identified from the cluster by the segmentation approach explained in Section 2: this influenced the failure to discriminate malignant from benign based on individual MCs morphological feature and texture patterns. However, the experimental results demonstrated the robustness and effectiveness of the developed method when combined with automatic MC detection and feature selection.

8. Conclusion

We have presented a method for discriminating malignant and benign clusters in digital and digitized mammograms. Images from digital and digitized databases were first segmented using a wavelet based method incorporating bi-cubic interpolation and a series of morphological operations were carried out in order to facilitate the feature extraction and classification task from MC segmented
images. A combination of morphological, texture, and distribution features from individual MC components and the whole MC clusters were extracted from mammograms. The most important features were selected and used to classify the MC cluster as benign or malignant. Clinical relevance of the selected features is discussed. ROC curve analysis was used to describe the cluster classification results. The feature extraction and selection were individually done using the digitized and digital mammograms, and afterwards those features were used to classify clusters in the digital database. The proposed method was evaluated using three different databases: OPTIMAM, DDSM, and MIAS. Two different classifiers- ensemble learner and stack generalization, were applied to evaluate the classification result. The best classification accuracy (96.72 ± 0.46%) for the digital database was achieved by using a stack generalization classification with 10-fold CV obtaining an $A_z$ value equal to 0.98 ± 0.00.

**Author Contributions:** N.A. developed the methodology and performed the implementation. E.R.E.D helped with the annotation of the MC cluster in the patch image. R.Z. supervised the investigation, results evaluation, and made substantial contributions to conception and design of this work. All authors discussed the results and contributed to the original draft preparation and editing.

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**Conflicts of Interest:** The authors declare no conflict of interest.

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