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Breast mass classification in sonography with transfer learning using a deep convolutional neural network and color conversion

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Abstract

Purpose: We propose a deep learning based approach to breast mass classification in sonography and compare it with the assessment of four experienced radiologists employing BI-RADS 4th edition lexicon and assessment protocol.

Methods: Several transfer learning techniques are employed to develop classifiers based on a set of 882 ultrasound images of breast masses. Additionally, we introduce the concept of a matching layer. The aim of this layer is to rescale pixel intensities of the gray scale ultrasound images and convert those images to red, green, blue (RGB) to more efficiently utilize the discriminative power of the convolutional neural network pre-trained on the ImageNet dataset. We present how this

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The authors have no conflicts to disclosure.

conversion can be determined during fine-tuning using back-propagation. Next, we compare the performance of the transfer learning techniques with and without the color conversion. To show the usefulness of our approach, we additionally evaluate it using two publicly available datasets.

Results: Color conversion increased the areas under the receiver operating curve for each transfer learning method. For the better-performing approach utilizing the fine-tuning and the matching layer, the area under the curve was equal to 0.936 on a test set of 150 cases. The areas under the curves for the radiologists reading the same set of cases ranged from 0.806 to 0.882. In the case of the two separate datasets, utilizing the proposed approach we achieved areas under the curve of around 0.890.

Conclusions: The concept of the matching layer is generalizable and can be used to improve the overall performance of the transfer learning techniques using deep convolutional neural networks. When fully developed as a clinical tool, the methods proposed in this paper have the potential to help radiologists with breast mass classification in ultrasound.

Keywords

BI-RADS; breast mass classification; convolutional neural networks; transfer learning; ultrasound imaging

1. Introduction

Breast cancer is one of the most common cancer in American women¹. Ultrasound (US) imaging is the most common adjunct imaging modality used to evaluate mammographic findings, palpable masses and to guide biopsy. Breast US is also the primary imaging modality in the evaluation of breast complaints in women under the age of 30. In comparison to other imaging modalities, US is relatively low cost, readily available and can accurately differentiate cysts vs. masses. However, accurate diagnosis with US requires experienced and trained radiologists to evaluate cystic and solid breast masses. To support the radiologists and standardize the reporting process the Breast Imaging Reporting and Data System (BI-RADS) was developed by the American College of Radiology (ACR)². BI-RADS lexicon characterizes US mass features based on shape, margins, orientation, echo patterns and posterior acoustic features. BI-RADS also provides assessment categories and recommendations as well as guidance on reporting. Although BI-RADS standardizes the reporting, the assessment of mass features is still subjective and depends on radiologist's experience and training. There is increased interest in potential use of US for screening, particularly in women with dense breast tissue. Screening breast US can detect additional mammographically occult cancers but it has a low positive predictive value of about 8% leading to increased number of unnecessary biopsies and higher rate of short-term follow $up^{3,4}$. To further help the radiologists correctly and objectively assess breast masses, various computer-aided diagnosis (CADx) systems have been proposed^{5,7}. These systems process US images to provide as output the probability that the examined masses are malignant.

CADx pipeline commonly includes four steps: image preprocessing, mass segmentation, feature extraction and classification. The performance of a CADx system is related to applied features that are usually engineered by employing expert knowledge. The usefulness

of the hand-selected features is reported to be that morphological features are the most effective for breast mass classification⁷. More angular and indistinct margins are expected in the case of malignant masses so the aim of the morphological features is to assess mass shape and margin. Various morphological features were inspired by the BI-RADS descriptors and aim to computerize BI-RADS related features. The morphological features were successfully employed to discriminate breast masses in several studies^{6–16}. However, efficiency of morphological features may depend on image preprocessing, US scanner, the specific view of the mass and applied segmentation algorithm^{17,18}.

Deep learning methods utilizing convolutional neural networks (CNNs) are gaining momentum in medical image analysis. CNNs for classification process an input image using different network layers to provide as output the probability that the examined image contains particular pathology. Due to limited medical datasets it is usually more efficient to use transfer learning and adjust a pre-trained deep model to address the classification problem of interest. Transfer learning methods were employed for breast mass classification and segmentation in several studies^{19–25}. Additionally, deep learning was used to detect breast lesions²⁶ and differentiate breast masses with shear-wave elastography²⁷. The better performing pre-trained deep learning models have been developed using RGB color images^{28–31}. However, medical images, including US images, are commonly gray scale, which raises question about how to efficiently utilize the discriminative power of a pre-trained model. In order to use a pre-trained model, the most widely used approach is to duplicate the gray scale intensities across all color channels^{19-22,25}. Another approach is to modify the convolutional layers of a pre-trained model, for example it is possible to convert the RGB images originally employed to develop the model to gray scale and use them to modify the model²¹. The second approach, however, may not lead to better classification performance²¹. Modification of the first layers influences the deeper layers in a network and does not necessarily improve the overall performance.

In this work we propose how to more efficiently utilize the color dependent representational capacity of a deep CNN to improve breast mass classification in US images. Instead of using duplicated gray scale images as input or modifying first convolutional layers, we introduce the concept of a matching layer (ML). This additional layer is added before the original input layer of the pre-trained CNN to convert gray scale US images to RGB. We show that this transformation can be efficiently learned during fine-tuning using the back-propagation algorithm. Next, we show that our approach leads to improved performance. Additionally, the usefulness of our approach to classification is depicted using two publicly available datasets of breast mass US images. To show the potential clinical value of the employed methods, our best performing classifier is compared with the performance of four experienced radiologists utilizing BI-RADS lexicon categories for overall breast mass assessment.

This manuscript is organized in the following way. First, we describe the datasets employed in this study. Then, transfer learning methods utilizing deep CNNs are detailed and we describe how these methods were applied to address the problem of the breast mass classification. Next, we describe how the gray scale US images were converted to RGB

via the ML. Results are presented and we discuss the advantages and the disadvantages of the applied methods.

2. Materials and Methods

2.1. Datasets

The main dataset employed in this study contained 882 US images of unique breast masses, one mass per patient, consisting of 678 benign and 204 malignant lesions. The dataset was divided into training, validation and test sets. The training set contained images of 582 breast masses (23% malignant) while the validation and the test sets both contained 150 masses each (23% malignant). In the case of the benign masses, the three sets contained similar number of fibroadenomas (42%), simple (26%) and complicated cysts (9%) as determined by one radiologist in the study. From a total of 14 malignant histological findings the following were included: 29% invasive ductal carcinoma, 21% invasive lobular carcinoma, 21% intraductal carcinoma, 11% ductal carcinoma in situ (all types), 10% invasive and in situ carcinoma, 20% other malignancies. The proportion of the five dominant findings was maintained in each dataset (training, validation and test). The distribution of mass types closely corresponded to the 5-year average mix of cases (<2% differences) at the Moores Cancer Center, University of California, San Diego. DICOM B-mode images (8 bit) were retrieved retrospectively in chronological order from institutional archive under approval by the Institutional Review Board and privacy compliance. Following BI-RADS criteria, cases were included if a breast mass was identified in at least two views sonographically, with only one image used. Biopsy was performed in 65% and the remainder had benign clinical follow-up of at least two years. Exams with no mass present (BI-RADS 1), inconclusive pathology results, significant artifacts or known cancers were excluded, but none were specifically included or excluded for race, ethnic background or health status. Self-reported racial/ethnic descriptions were White (69%), Asian/Pacific Islander (12%), Hispanic (7%), Black (5%), Native American/Eskimo (<1%), Other (3%), and not reported (4%). Age ranged from 18–90 years (mean 51±15). Mass size ranged from 2.5 to 98 mm² (mean 12.8±9.3 mm²). Sonography was performed at an ACR accredited center following standard clinical protocol with one of three scanners: Siemens Acuson (59%), GE L9 (21%) and ATL-HDI (20%). The BI-RADS category was assigned to each mass in the test set independently by four senior sub-specialty radiologists, reviewing the cases in random order in two sessions using a standard hard-copy BI-RADS classification form that includes final assessment category, descriptors and recommendations for follow-up interval or biopsy. The radiologists were not aware of the confirmed findings for any case.

To show the usefulness of the methods proposed in this paper, we also employed two publicly available breast mass datasets^{26,32}. The first one, named UDIAT, consists of 163 B-mode images of breast masses (53 malignant and 110 benign) collected using Siemens ACUSON scanner from the UDIAT Diagnostic Centre of the Parc Tauli Corporation, Sabadell (Spain). This dataset was used by the authors to develop deep learning based algorithms for the breast mass detection²⁶ and segmentation²². The second dataset, named OASBUD (Open Access Series of Breast Ultrasonic Data), consists of raw ultrasonic echoes (before B-mode image reconstruction) acquired from 52 malignant and 48 benign masses

with the Ultrasonix SonixTouch Research scanner from patients of the Oncology Institute in Warsaw (Poland). For each mass two perpendicular scans were recorded. The OASBUD was originally used to assess the statistical properties of backscattered ultrasound echoes in breast tissue^{33,34} and to differentiate breast masses using transfer learning with CNNs²⁰. Detailed descriptions of both datasets can be found in the original papers^{26,32}.

2.2. Transfer learning

In this study we used the VGG19 neural network publicly available in TensorFlow.^{29,35} The CNN was pre-trained on the ImageNet dataset that contains over 1.2 million RGB images corresponding to 1000 classes²⁸. The model includes five large blocks of sequentially stacked convolutional layers followed by a block of fully connected (FC) layers. Convolutional layers extract different information from images. The first layers include edge and blob detectors while the deeper layers include ImageNet class-related features. This CNN was reported to be useful for various medical image analysis tasks³⁶, including breast mass classification^{19,20}, so it was selected for this study to enable comparison to other's results.

We employed two approaches to neural transfer learning³⁷. The first utilized the pre-trained model as a fixed feature extractor. In this case, the model architecture was not modified. The aim of the second approach was to fine-tune the CNN using the new dataset, in our case breast sonograms. In order to perform fine-tuning the CNN architecture is usually modified, the last layers of the network are replaced with custom FC layers. Next, the back-propagation algorithm is used to adjust the model to the new classification problem.

The main dataset was augmented in order to improve training and provide more diverse images to the network. First, each US image of a breast mass was median filtered and cropped with a fixed exterior margin of 30 pixels using the region of interest (ROI) provided by the radiologist and resized to the default VGG19 image size of 224*244. Additionally, the images were flipped and shifted by 15 pixels horizontally. Image shift was applied before cropping. Due to the augmentation, the number of images in each set increased six times. We decided not to perform image rotation or shift in longitudinal direction, as this would alter some of the known attributes of breast masses such as posterior shadowing and enhancement, potentially decreasing classification performance³⁸. Example images are presented in Fig. 1. The B-mode images from the UDIAT dataset were pre-processed and augmented in a similar way as in the case of our dataset. Raw ultrasonic echoes from the original paper³². Echo amplitude was computed with the Hilbert transform and logarithmically compressed to reconstruct the B-mode image. Next, the data were pre-processed and augmented as in the case of the other datasets.

2.2.1. Neural feature extraction—We implemented two efficient neural feature extraction methods employed in a recent paper¹⁹. First, features for classification were extracted from each of five max pooling (MP) layers of the original VGG19 model³⁷. Next, features corresponding to each block were averaged along spatial dimensions and normalized using \hat{P} norm. Features vectors corresponding to each block were then combined

to form the final MP feature vector. Second, we additionally used the first FC layer to extract features. For both methods all zero variance features in respect to the training set were discarded. For classification we employed the Support Vector Machine (SVM) algorithm³⁹. In our study we applied two different approaches to model development. The main dataset was divided into training, validation and test sets. Training and validation sets were used to determine the best performing hyper-parameters and the optimal kernel of the SVM classifier via the grid search algorithm. For the C and γ parameters of the SVM algorithm the grid included parameters in range of [0.00001, 0.0001, ..., 1, 10] and [1, 5, ... 100]. The parameter grid also included the kernels, namely the linear and radial basis function kernels. Next, the best performing model was evaluated on the test set. However, the UDIAT and OASBUD datasets are too small to divide them into three separate sets in a convenient way. Therefore, we applied 10-fold cross-validation (case based) to evaluate the classification. Within each fold, additional 5-fold cross-validation was used to find optimal hyper-parameters of the SVM classifiers. To address the problem of class imbalance, we employed class weights inversely proportional to class frequencies in the training set. The SVM classifiers were developed and evaluated using the augmented datasets. To determine a posteriori probability that a mass in the test set is malignant, we averaged the probabilities calculated for each image of this particular mass. Next, to assess the classification performance we determined the receiver operating characteristic (ROC) and calculated the area under the ROC curve (AUC). Sensitivity, specificity and accuracy of the better performing algorithms were calculated using the ROC curve for the point on the curve that was the closest to $(0, 1)^{40}$. Welch's t-test at significance level of 0.001 was used to determine whether there is a difference in AUC values. All calculations were performed in Python.

2.2.2. Fine-tuning—The architecture of the original VGG19 model was modified in order to perform fine-tuning. Unfortunately, the UDIAT and OASBUD datasets were too small to efficiently fine-tune the VGG19 model, therefore fine-tuning was employed only in the case of the main dataset. The way to modify the architecture was determined using the validation set. The original FC layers, developed for the ImageNet classes, were replaced with a FC layer with 4096 units followed by a FC layer with 256 units and a single unit employing sigmoid activation function suitable for binary classification (in this case, benign or malignant). For the first two FC layers we used a rectifier activation functions. Initial weights of the layers were set using the Xavier uniform initializer. With the use of the validation dataset, we found that the highest performance can be obtained if the first four convolutional blocks are frozen and only the fifth block and the fully connected layers are fine-tuned. We also found that the fine-tuning of the first convolutional block does not improve the classification performance. To fine-tune the VGG19 neural network we used the mini-batch stochastic gradient descent with Nesterov update. The learning rate was initially set to 0.001 and was decreased by 0.00001 per epoch up to 0.00001. The momentum and the batch size were set to 0.9 and 40, respectively. The binary cross-entropy loss was employed with weights inversely proportional to class frequencies in the training set. To reduce over-fitting, we applied dropout with 80% dropout probability to the first fully connected layer. The experiments were performed on a computer equipped with a GeForce GTX 1080 Ti graphics card. The AUC value on the validation set was monitored during the

training. As in the case of the SVM algorithm, we selected the model that maximized the AUC value on the validation set.

2.3. Matching layer

Recently proposed CNN based breast mass classification approaches employed gray scale US images as input to the pre-trained models^{19–22}. In this paper, we propose to adjust the gray scale US images to the pre-trained model instead of duplicating gray scale images across the channels or modifying the first convolutional layer of the CNN. By performing this transformation, we aim to utilize more efficiently the representational capacity of the deep model. For this task we use a ML that transforms the input gray scale images to RGB images via a linear transformation:

$$I_{out} = \overrightarrow{a} I_{in} + \overrightarrow{b}$$

where I_{in} is the gray scale image, I_{out} is the output RGB image, \vec{a} and \vec{b} are the transformation parameters that shall be determined during training. This transformation is a 1D convolution with a bias term. 1D convolutions were employed, for example, by the authors of the GoogleLeNet CNN to reduce the dimensionality of the input data³¹. In our case, we use a 1D convolution layer to artificially increase the dimensions of the input images and ideally perform color conversion from gray level images to RGB. Fig. 2 depicts the modified VGG19 architecture that includes the ML layer in the front. In this study, we determined the parameters of the ML layer during fine-tuning using the back-propagation algorithm in a way that minimizes the loss function.

3. Results

First, we used the gray scale images to perform classification of the B-mode images from the main dataset following the standard approach^{19,20,22,25}. For each US image, the gray level intensities were copied to RGB channels and the VGG19 CNN was tuned. The highest AUC value on the validation set, equal to 0.921, was obtained after 16th epoch. The corresponding AUC value on the test set was equal to 0.895. Next, we extracted the MP features and the FC features using the original VGG19 model and used those features to train the SVM classifiers. The validation set was employed to select the best hyper-parameters. For both feature sets we obtained similar AUC values equal to 0.849 on the test set. The classification performance is depicted in Table I and the ROC curves are shown in Fig. 3.

Next, we fine-tuned the VGG19 CNN combined with the ML using the back-propagation algorithm. The highest AUC value on the validation set, equal to 0.961, was obtained after 7th epoch and corresponded to AUC value of 0.936 on the test set, see Table II and Fig. 4. To visualize how the ML works, we converted two gray scale US images into RGB images with results depicted in Fig. 5. Following the conversion, the image was dominated by light blue and yellow colors.

The converted RGB US images were utilized to extract the MP features and the FC features using the original VGG19 model (not fine-tuned). Again, the validation set was employed

to find the best performing hyper-parameters for the SVM classifiers. For the RGB images we obtained higher AUC values on the validation and the test set, equal to 0.889 and 0.873 for the MP features and the FC features, respectively. Color conversion improved the classification performance. In comparison to gray US images, the AUC values significantly increased by around 0.04 (*p*-values<0.001), see Table I. The ROC curves calculated for the classifiers developed using the ML layer are depicted in Fig. 4.

In the next step, we extracted the MP features and the FC features from the VGG19 model using the B-mode images from the UDIAT and OASBUD datasets. For each US image, the gray level intensities were copied to RGB channels. In the case of the UDIAT dataset, we obtained AUC values of 0.858 and 0.849 for the MP and the FC features, respectively. For the OASBUD dataset the corresponding AUC values for the MP and the FC features were equal to 0.819 and 0.791, respectively. Unfortunately, due to small sizes of these datasets we were not able to perform fine-tuning in an efficient way, so we employed the ML layer developed using the main dataset. As in the case of our dataset, all B-mode images from the UDIAT and OASBUD datasets were converted to RGB using the ML and utilized to extract the FC and the ML features. The same cross-validation folds were used to evaluate the performance. Results showed that due to the color conversion the classification performance increased. For the UDIAT dataset, the AUC values for the MP and the FC features increased to 0.873 and 0.893, respectively. In the case of the OASBUD, the AUC values were equal to 0.831 and 0.881 for the MP and the FC features, respectively. The ML improved significantly the AUC values (*p*-values<0.001) in the case of the FC features (both datasets). However, for the classifiers trained using the MP features the improvement was too small to provide statistically significant difference. Results are depicted in Table II. Fig. 6 shows how the ML converts the B-images from the UDIAT and OASBUD datasets to RGB.

Four radiologists participated in our study. Table III shows the distribution of BI-RADS categories for the masses in the test set. The Fleiss' kappa was equal to 0.41 indicating moderate agreement of the radiologists in BI-RADS category final assessment. Table IV presents the classification performance of the radiologists employing the BI-RADS categories. The AUC values ranged between 0.806 and 0.882 with mean of 0.849. The AUC value for the better performing CNN with ML was significantly higher than the highest AUC value for the radiologists, 0.936 vs. 0.882 (p-value<0.001). Accuracy, sensitivity and specificity were determined for the BI-RADS category 3 (probably benign) used as the benign cut-off. In this case, the sensitivity of the radiologists was excellent (mean (0.992), but the specificity was lower (mean (0.412)), indicating an expected bias towards not missing a positive mass. To additionally compare the assessment of the radiologists with our better performing method, we employed the following procedure. First, majority voting was applied to assign a single BI-RADS category to each breast mass. Ties were handled by assigning the higher BI-RADS category. Second, we investigated the relation between the output of the network (a posteriori probability of malignancy) and the BI-RADS category (after voting) assigned to each mass. Fig. 6 shows that the probability of malignancy increases with the BI-RADS category, as expected. To confirm this observation, the Mann-Whitney statistical tests with Bonferroni correction were applied. Results show that four groups are statistically different (*p*-value<0.05), expect for the BI-RADS categories 2 and

3 (*p*-value=0.22). These two groups contained only benign lesions for which the network assigned small probabilities of malignancy.

To illustrate the examples that were difficult for our better performing model to classify (fine-tuned VGG19 model with the ML), we extracted from the test set the images of malignant and benign breast masses that were assessed with the highest and the lowest confidence level for each class, these are shown in Fig. 8. For example, Fig. 8b) shows a benign mass that was assessed with the highest confidence of being malignant by the classifier. For this mass, the radiologists assigned BI-RADS categories 4, 4, 4 and 5, which indicates that according to the radiologists this mass was suspicious for malignancy. Fig. 8d) depicts a malignant mass that was assessed with the lowest confidence of being malignant by the classifier. All radiologists assigned BI-RADS category 5 to this mass. The oval shape, moderate posterior enhancement and fairly uniform anechoic pattern of this mass might be the reason why the model performed worse in this case.

4. Discussion

Our study demonstrates potential usefulness of a deep CNN based approach for breast mass classification in US images on three different datasets. In the case of the main dataset, we evaluated three transfer learning methods and achieved good results with each of them. The highest AUC value of 0.936 was achieved for the fine-tuned VGG19 model combined with the ML. Using the MP and the FC features we obtained AUCs in range from 0.849 to 0.889. In the case of the other datasets we obtained the AUC values in range from 0.791 to 0.893. Although we were not able to fine-tune the model using the smaller datasets, we still obtained similar classification performance using the ML and the FC features as in the case of the main dataset. For the UDIAT dataset the performance was almost the same, while for the OASBUD dataset the AUC values were lower by about 0.03 (without the ML). All these results are comparable with the results reported in previous papers^{19,20,25}, where the AUC values of around 0.85 were obtained. In one of the studies the GoogleLeNet was fine-tuned to classify breast masses in US images²¹. The authors achieved high AUC value of 0.96. However, the authors used a large set of over 7000 US images to develop the model. In our case, we used a set of 882 breast mass images to develop the model, which may explain the difference in AUC values. Supposedly, with a larger dataset it is possible to fine-tune the model in a more efficient way.

In our study fine-tuning proved to be more efficient than the SVM algorithm utilizing directly extracted CNN features; we obtained higher AUC values by around 0.04 for the main dataset. The better performance may be partially explained by the fact that we fine-tuned the last convolutional block of the VGG19 model. The usefulness of the convolutional blocks of the VGG19 CNN for breast mass classification was evaluated separately in one of the previous papers²⁰. The performance of the last 5th convolutional block was lower than in the case of the 4th block, which suggests that the 5th block is specifically related to recognition of the ImageNet objects. The approach employing fine-tuning, however, is more challenging and troublesome to develop. It requires to replace the FC layers of the original CNN with custom layers and determining which layers of the original network should be trainable during the fine tuning. Moreover, the learning rate and other hyper-parameters

have to be correctly selected to yield good classification performance on the validation set. Iterating over methods of model development is time consuming and may not yield good results at first. Moreover, in the case of a small dataset, the fine-tuning might not be efficient and it is more reasonable to utilize the FC or the MP features.

The concept of the color conversion has been widely employed in the image analysis field⁴¹. Usually a type of color conversion is used before image segmentation 42,43 . In the case of deep learning, color conversion was employed in deep colorization⁴⁴ or neural style transfer⁴⁵. Our study shows that color distribution appears to be an important factor and that it should be taken into account to more efficiently use transfer learning with pre-trained deep models. By utilizing the ML we were able to obtain better classification performance. The first layers of the pre-trained network commonly include color blob detectors. Modifying these layers may not necessarily lead to better performance, because those layers are somehow connected with the deeper layers in the network. With the color conversion it was possible to more efficiently use the pre-trained CNN. This advantage is clearly depicted in the case of the MP and FC features extracted from all datasets. The ML developed using the main dataset proved to work with other datasets as well. This shows the universality of our approach. For example, in the case of the FC features extracted using the OASBUD dataset due to the color conversion the AUC value increased from 0.791 to 0.881. Moreover, the color conversion enabled us to more efficiently fine-tune the VGG19 CNN. In this work we used linear transformations to convert gray scale US images to RGB. Regular image pre-processing required in the case of the VGG19 network could be performed using only the bias term \vec{b} . We decided to additionally employ the scaling parameter \vec{a} for several reasons. First, color inversion in particular channels may improve the performance. Second, the area of mass in US image is commonly hypoechoic or anechoic while the surrounding tissue is significantly brighter. For the network to perform well it might be useful to rescale this relation and for example decrease this difference in brightness levels. The proposed approach, however, is general and not limited to gray scale images or to US, so it can be applied to RGB images as well. Moreover, it is possible to use other transformations including nonlinear ones, which could be useful for processing images from different medical modalities. The advantage of our approach is that the transformation is determined during fine-tuning automatically. In some sense, our approach can be perceived as adding an additional neural network in front of the pre-trained one. The main issue is that the transformation may change the range of intensities to one outside [0, 255] making the transformed image difficult or impossible to visualize.

The CNN based methods developed using the main dataset achieved results comparable to or higher than the AUC obtained by the radiologists. Our best performing approach showed an AUC value that was higher by 0.04 to 0.13 than the range of AUCs calculated for the radiologists. These results demonstrate potential clinical usefulness of the developed classifiers. However, it is important to recognize that diagnostic breast US is commonly used to determine whether and where to perform the biopsy rather than to determine if a lesion is benign or malignant. In order to avoid missing a potential malignancy, radiologists achieve very high sensitivity at the expense of lower specificity.³⁴ The better classifier in this study achieved higher specificity (0.90) but lower sensitivity (0.85) than all of the radiologists.

However, according to the ROC curve in Fig. 4, at a sensitivity of 1.0 corresponding to that of the radiologists, our better performing CNN achieves higher specificity of 0.65 compared to 0.3–0.5. We also showed that the output of the better performing model (*a posteriori* probability of malignancy) is related to the BI-RADS category. This, however, was expected since both measures aim to assess the level of malignancy.

To more thoroughly evaluate the classifier utility in a clinical environment, it will be desirable to employ the CNN as a decision tool with the radiologist "in the loop." For a particular mass, the CNN result may be overridden by the interpreting radiologist in the final assessment. The current results do not describe a complete medical device ready for clinical use but they may provide necessary information to properly design and study such a tool. In our approach the radiologists identify the mass and select an ROI, hence requiring interaction by the radiologist. It remains to be seen if this can be done efficiently or semi-automatically without impacting workflow, and if it is useful for the radiologist with difficult cases where accurate assessment of the type of breast mass is desired. In future, it would be also interesting to investigate whether the CNNs can be used to classify malignant and benign breast mass subtypes.

5. Conclusion

In this study we utilized the VGG19 CNN for breast mass classification and introduced the concept of the matching layer, a layer that is used to convert gray images to RGB. We demonstrated that with the ML it was possible to perform more efficient classification than in the case of duplicating gray level US images across the RGB channels. The concept of the ML is general and can be applied to various problems to enhance CNN based transfer learning techniques. The AUC obtained for our better performing classifier was higher than that of four expert radiologists who utilized BI-RADS lexicon and categories. Even at sensitivity of 1.0 the classifier achieved higher specificity in comparison to the radiologists. In future we are going to perform additional studies to determine the clinical usefulness of the employed methods.

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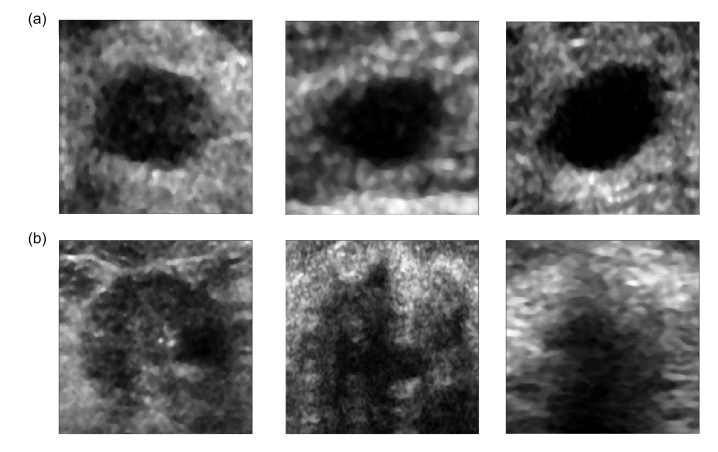
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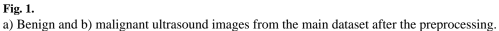
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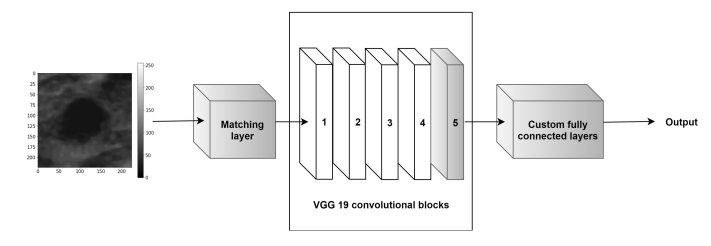
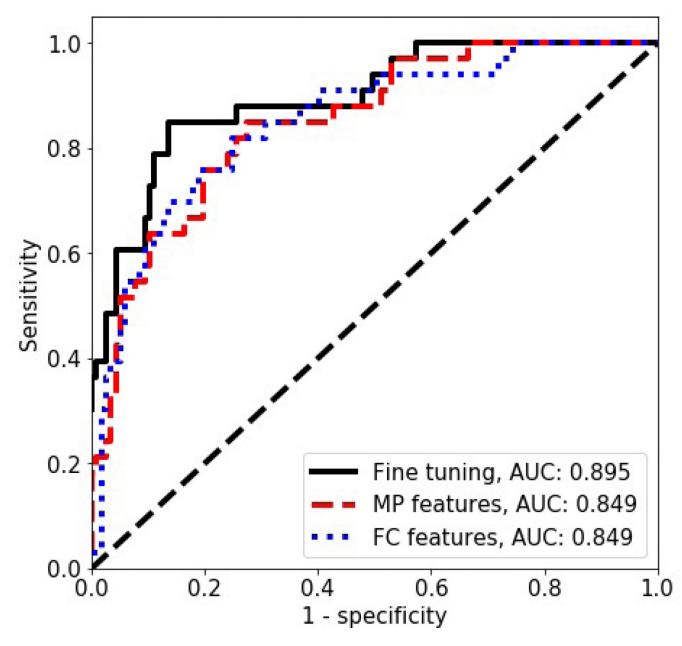


Fig. 2.

The modified architecture of the VGG19 CNN, gray colors indicate the trainable layers. We propose to add a ML in front of the pre-trained model to transform the gray scale US images to RGB to utilize more efficiently the representation capacity of the deep model.

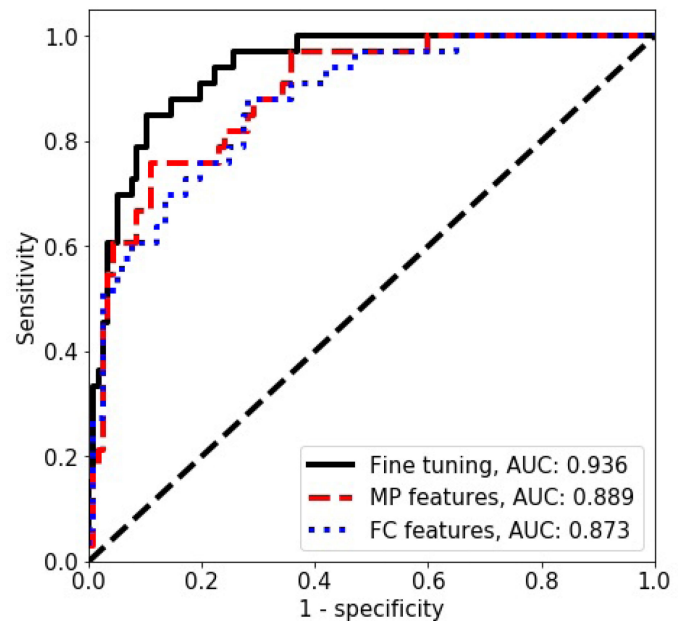
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The ROC curves for the CNN based classification without the ML in the case of the main dataset.

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The ROC curves for the CNN based classification with the ML in the case of the main dataset.

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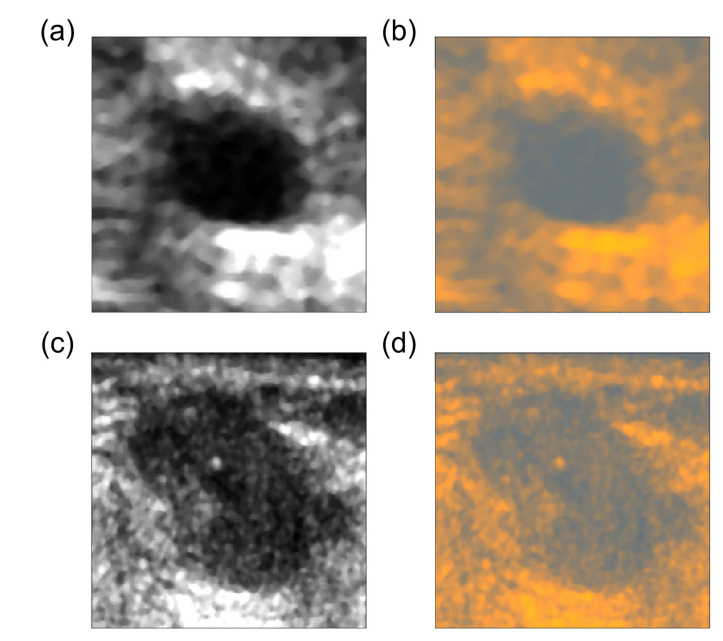


Fig. 5.

Conversion of gray scale US images from our dataset to RGB using the ML, a) a benign lesion image and b) its conversion, c) a malignant lesion image and d) its conversion. By using the color conversion, it was possible to more efficiently use the CNN model pre-trained on RGB images.

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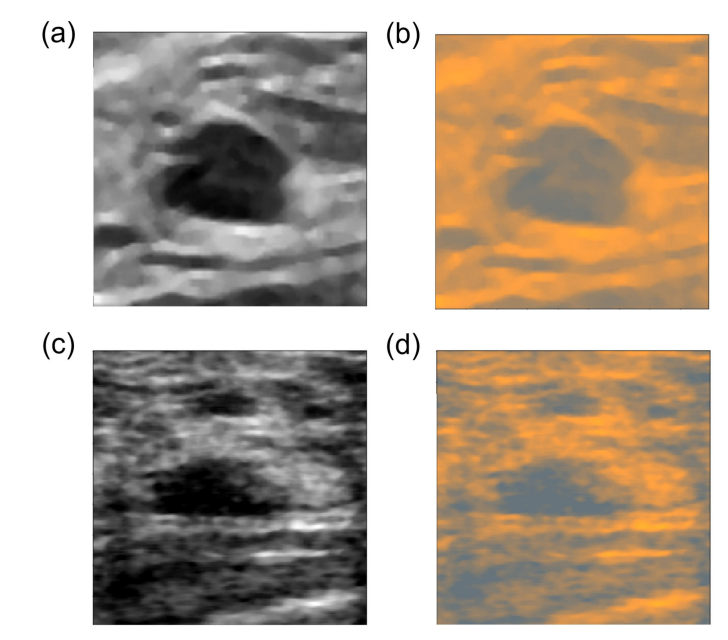
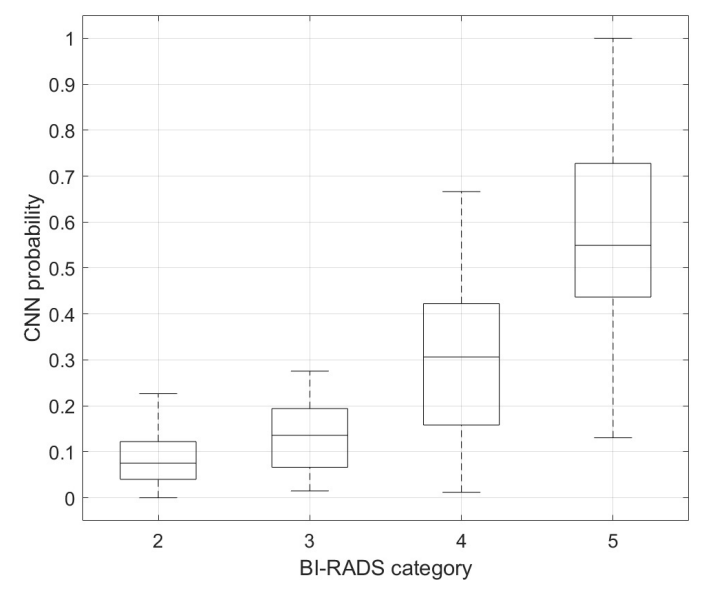


Fig. 6.

a) Image from the UDIAT dataset and b) its conversion to RGB, c) image from the OASBUD and d) its conversion to RGB. By using the color conversion, it was possible to more efficiently use the CNN model pre-trained on RGB images.

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Relation between the output of the CNN with the ML and the average BI-RADS category assigned to each breast mass.

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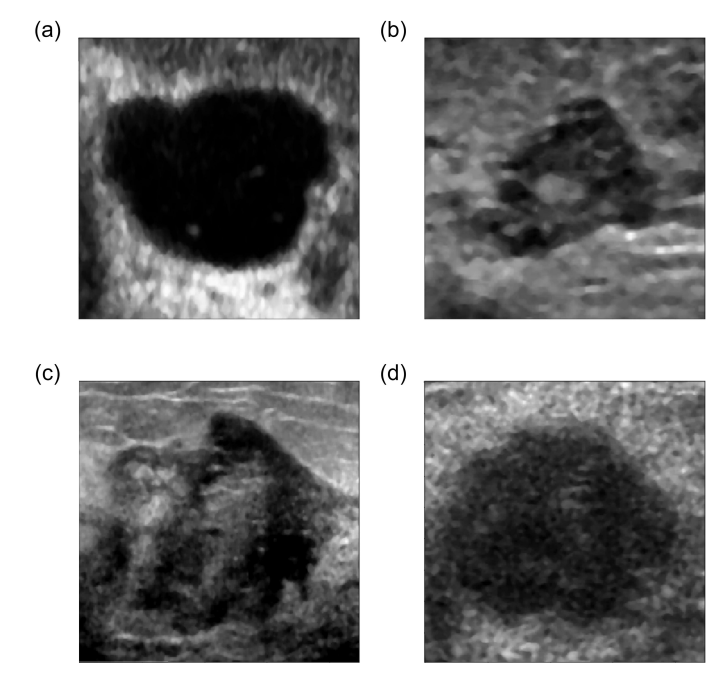


Fig. 8.

Benign masses assessed by the fine-tuned model as malignant with a) the lowest (*a posteriori* probability of 0.98, BI-RADS: 2, 2, 2, 2) and b) the highest confidence level (*a posteriori* probability of 0, BI-RADS 4, 4, 4, 5), and malignant masses assessed as malignant with c) the highest (*a posteriori* probability of 1, BI-RADS: 4, 5, 5, 5) and d) the lowest confidence level (*a posteriori* probability of 0.33, BI-RADS 5, 5, 5, 5).

Table I.

Classification performance of the models developed without the ML for our dataset. The standard deviations of the parameters were calculated using bootstrap.

Method	AUC	Accuracy	Sensitivity	Specificity
Fine tuning	0.895±0.031	0.860 ± 0.024	0.848 ± 0.056	0.863±0.026
MP features	0.849±0.036	0.793±0.036	0.757±0.054	0.803±0.048
FC features	0.849±0.036	0.800 ± 0.038	0.757±0.054	0.812±0.046
Fine tuning, ML	0.936±0.019	$0.887 {\pm} 0.028$	0.848±0.039	0.897±0.035
MP features, ML	0.889±0.029	0.860 ± 0.044	0.757 ± 0.058	0.889±0.062
FC features, ML	0.873±0.036	0.753±0.044	0.879 ± 0.058	0.718±0.053

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Table II.

Classification performance of the models developed with and without the ML in the case of the UDIAT and OASBUD datasets. The standard deviations of the parameters were calculated using bootstrap.

Dataset	Method	AUC	Accuracy	Sensitivity	Specificity
	MP features	0.858±0.029	0.853±0.024	0.796±0.043	0.880 ± 0.027
UDIAT	FC features	0.849±0.031	0.822±0.037	0.759±0.043	0.853±0.053
	MP features, ML	0.873±0.027	0.840±0.023	0.833±0.037	0.844±0.027
	FC features, ML	0.893±0.030	0.840 ± 0.024	0.851±0.042	0.834±0.030
OASBUD	MP features	0.819±0.030	0.760±0.029	0.692 ± 0.057	0.833±0.057
	FC features	0.791±0.035	0.750±0.031	0.750 ± 0.044	0.750±0.050
	MP features, ML	0.831±0.031	0.760±0.031	0.762 ± 0.059	0.750±0.061
	FC features, ML	0.881±0.023	0.830±0.026	0.807±0.039	0.854±0.036

Table III.

Distribution of BI-RADS categories for the test set, Fleiss' kappa was equal to 0.41.

		BI-RADS			
		2	3	4	5
Dadialagiat 1	Benign	44	15	57	1
Radiologist 1	Malignant	0	1	23	9
Dadialagiat 2	Benign	37	13	13 64	3
Radiologist 2	Malignant	0	0	16	17
Dadialagiat 2	Benign	41	11	61	4
Radiologist 3	Malignant	0	0	12	21
D 11 1 1 4	Benign	19	13	60	25
Radiologist 4	Malignant	0	0	3	30

Table IV.

Performance of the Radiologists employing BI-RADS for benign cut-off set to BI-RADS 3.

	AUC	Accuracy	Sensitivity	Specificity
Radiologist 1	0.806 ± 0.028	0.607	0.967	0.504
Radiologist 2	$0.848 {\pm} 0.028$	0.553	1	0.427
Radiologist 3	0.882±0.026	0.567	1	0.444
Radiologist 4	0.860±0.027	0.433	1	0.273
Mean	0.849	0.540	0.992	0.412