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Application of Artificial Intelligence in Microwave Radiometry (MWR)

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Abstract: Microwave radiometry is being developed more actively in recent years for medical applications. One such application is for diagnosis or monitoring of cancer. Medical radiometry presents a strong alternative to other methods of diagnosis, especially with recent gains in its accuracy. In addition, it is safe to use, non-invasive and has a relative low cost of use. Temperature readings were taken from the mammary glands for the purpose of detecting cancer and evaluating the effectiveness of radiometry. Building a diagnostic system to automate classification of new samples requires an adequate machine learning model. Such models that were explored were random forest, XGBoost, k-nearest neighbors, support vector machines, variants of cascade correlation neural network, deep neural network and convolution neural network. From all these models evaluated, the best performing on the test set was the deep neural network with a significant difference from the rest.

1 INTRODUCTION

Microwave radiometry has seen in recent years increased usage and interest for further development and research within medical applications (Vesnin et al., 2017). This has resulted in a significant improvement of the system’s accuracy in taking internal temperature measurements. One of its main applications is for cancer detection and monitoring, such as breast cancer which will be the focus of this paper.

However, while gaining momentum in its utilization it is still not widely adopted. Main reason for this can be attributed to the fact that it has recently being adopted for medical use and so medical or clinical professionals have not yet received adequate training to interpret the information. However, this leads to the initial issue, if the professionals are unable to use the system then they will be more hesitant to acquire them.

The deadlock can be resolved with the introduction of an automated diagnostic systems which will extract useful information from the readings and offer a diagnostic prediction. For this paper, the focus will be in evaluating how effective such data alone can be used for diagnosis of cancer, using data collected from mammary glands. Additionally, a further scope is to contribute into determining an ideal machine learning algorithm for such a task.

The paper will start off with a brief development history of radiometry within the medical field, what it captures and why it is an attractive system to be used for cancer detection and monitoring. Following, a description of the data set will be provided which includes readings from radiometry of the mammary glands for cancer detection. In addition, any pre-processing that was conducted on the data will be documented. After the setup information has been provided, the description and results of various non-neural network and neural network models will be presented on classifying low or high risk of presence of breast cancer. Finally, the paper will finish off with the conclusions and some possible future work.


2 MICROWAVE RADIOMETRY

Microwave radiometry’s started from the theoretical research of James Clerk Maxwell and experimentally verified by Heinrich Hertz, with development of the first radars in the 1930s (Skolnik, 2018). Later, during WWII Robert invented a kind of radio receiver, was known as “Dicke Radiometric Receiver” or just “Dicke Radiometer”. His radiometer used a switchable resistor, the “Dicke Resistor”, as a technique to allow for noise temperature calibration (Dicke, 1982).

However, only later on, from 1970s, such technology was first applied for medical and clinical usage (Myers et al., 1979; Bolomey et al., 1982; Peronnet et al., 1983; Pichot et al., 1985). But there was no significant investment until the late 1990s, which gain a lot more interest from the medical scientific community (Conceicao et al., 2016). Since then it has found applications for detecting or monitoring breast cancer (Vesnin et al., 2017), thermal denaturation of albumin (Ivanov et al., 2018), carotid artery diseases (Drakopoulou et al., 2018), brown adipose tissue activity (Crandall et al., 2018), rheumatoid arthritis (Pentazos et al., 2018), inflammation levels in joints (Laskari et al., 2018), brain temperature (Rodrigues et al., 2018) and transcapillary water exchange in the lungs (Bondar et al., 2017).

It is able to capture the temperature at the skin or at a depth from the surface, which is particular useful for diagnosing and monitoring treatment progress of cancerous tumors (Vesnin et al., 2017). It achieves this by measuring the electromagnetic radiation emitted by the tissues in the microwave range (Vesnin et al., 2017). The measurement obtained depends on the variation of the properties of the various biological tissues (Semenov, 2009). In turn, these properties are impacted by the level of water found in the tissue, with a significant difference between muscle, which is high in water levels, and fat and bone, which have low levels (Gabriel et al., 1996a; Gabriel et al., 1996b). In addition, both physiological and pathological conditions can alter the levels of dielectric properties of the tissues (Semenov, 2009).

Specifically for cancer, it has been found that tumors emit heat which is connected to their growth rate (Gautherie, 1980). As the tumorous cells grow, they replicate themselves at a much higher rate leading to the release of higher amounts of energy compared to neighboring healthy cells. The tumors’ ability to create new vasculature will determine its maximum volume (Schneider and Miller, 2005). At such a stage, cell growth and cell death rate reach an equilibrium. However, when the growth slows down this will result to near normal temperature readings making cancer detection more difficult for such cases (Vesnin et al., 2017).

It is an attractive complementary technique to other methods of diagnosis of cancer, such as mammography or biopsy (Vesnin et al., 2017). The main reason stated by Vesnin et al. is that advances in the system have allowed it to achieve high sensitivity and specificity in cancer detection. Also, it is a non-ionizing safe method, noninvasive, results are obtained quickly and has a low cost (Vesnin et al., 2017; Semenov, 2009). Hence, this enables it to be used at any frequency, for any age group and by someone during pregnancy or lactation. In addition, it can also occupy a supportive role in decision making for professionals as for it adds information not obtainable from other methods. Such information is the thermal activity of the tissue, the rate which cancerous cells multiply and the level of risk for mutagenesis (Vesnin et al., 2017).

3 DATA SET

3.1 Description

To conduct the evaluation of microwave radiometry for its effectiveness in detecting cancer a data set comprising temperature values from mammary glands was used. The values were recorded using the RTM-01-RES (www.mmwr.co.uk) device from various medical centers (Zenovich et al., 2016). The device captures temperature readings at nine different locations on each gland, one at the nipple (defined as point 0) and the rest equidistant around the nipple (points 1 to 8), plus at the axillary region (point 9). In addition, two more locations where captured at the lower chest (defined as points T1 and T2), as references to normalize ambient temperature variations. For each of these points, the temperature was measured at the skin and at a depth from the skin of 5cm. A graphical representation of the capture points can be seen in Figure 1.

In total, there are 363 pairs of mammary glands of which 77 are classified as healthy or low risk (labeled as class 0) and 286 classified as potentially cancerous or high risk (labeled as class 1). For each sample, to be classified as low risk both glands must be considered as healthy and for it to be considered as high risk then at least one must be of high risk. Individual glands compromise of 319 low risk and 407 as nodal cancer, 119 as diffuse changes with no presence of cancer and 90 as nodal changes with no presence of cancer. All the following experiments had
the data class balanced split into three sets, training, validation and test set, with allocated percentage of 60% (low risk: 46 and high risk:171), 20% (low risk: 15 and high risk:57) and 20% (low risk: 16 and high risk:58) respectively.

Figure 1: Sampling points on each mammary gland (0-8) including the axillary point (9). Points T1 and T2 are used as reference values when normalizing the values against ambient temperature (Zenovich et al., 2016).

3.2 Ambient Temperature Normalization

Having collected the data at multiple locations and at different times the ambient temperature varies for each sample in which the values will not be directly comparable. Typically, the measurements were taken under temperatures ranging from 20 to 27 degrees Celsius. When analyzing temperature values for prediction systems breast size, age and external conditions that can impact the results must be taken into account (Anisimova, 2013; Kobrinskiy, 2008). Hence, a previous research (Losev and Lvshinskiy, 2015) that used the same data set proposed and evaluated a normalization algorithm to overcome this issue. The algorithm was defined as such (Losev and Lvshinskiy, 2015):

For every point \( t_{d,i,j} \) captured, plot their values against one of the control temperature points \( T_{c,d,j} \), where \( i = 0...9, c \in \{1, 2\}, j \in \{\text{skin, depth}\} \) and \( d = 0...n - 1 \), with \( n \) the total number of samples:

1. On the plotted graph between temperature points and one of the reference values, we use linear regression to find \( a \) and \( b \) such that the error is minimized through least square fit method on the function:

\[
 t_{d,i,j} = a \cdot T_{c,d,j} + b \quad (1)
\]

2. Calculate the average value of the temperature point such that:

\[
 T_{\text{avg},c,j} = \frac{1}{n} \sum_{d=0}^{n-1} T_{c,d,j} \quad (2)
\]

3. Update the temperature points:

\[
 t_{d,i,j} = t_{d,i,j} + a \cdot (T_{\text{avg},c,j} - T_{c,d,j}) \quad (3)
\]

4. Replace the control points with the average value found:

\[
 T_{c,d,j} = T_{\text{avg},c,j} \quad (4)
\]

for \( d = 0...n - 1 \)

Losev and Lvshinskiy showed that when applying their proposed algorithm, it improved the specificity and sensitivity of a regression prediction model. There was a strong linear correlation coefficient between all points 0-9 against either of the two reference points as the temperature increased. While they showed comparing against both reference points resulted in improvement in predictions, using reference T2 obtained slightly better performance of overall 4%. Therefore, for the following experiments evaluating various prediction models this normalization algorithm was applied against reference point T2. In turn, this allowed the removal of the two control points as for all samples would have the same value.

3.3 Oversampling

As described in section 3.1, the data set is heavily imbalanced towards the high risk class with a total of 77 against 286 samples. Consequently, this introduces a bias towards the higher proportion samples in which most machine learning algorithms will favor when classifying (Krawczyk, 2016). Some algorithms can handle this imbalance by introducing sample weights giving more importance to the least represented class or by introducing an appropriate metric (He and Ma, 2013). While both of these techniques were used in the experiments where applicable, applying oversampling (He and Ma, 2013) guarantees consistency between the various algorithms.

The techniques explored were random resampling, Synthetic Minority Over-Sampling Technique (SMOTE) with regular, borderline 1, borderline 2 and Support vector Machine (SVM) variations (Chawla et al., 2002; Han et al., 2005) and Adaptive Synthetic (ADASYN) (He et al., 2008). The oversampling techniques were compared using a random forest (Breiman, 2001) from the scikit-learn library (Pedregosa et al., 2011) having set a sample weight
importance to handle imbalance. Additionally, for each case the tree was optimized using the hyperopt (Bergstra et al., 2015) library with the tree of Parzen (TPE) (Bergstra et al., 2011) optimizer and weighted Geometric Mean (G-mean) loss (Kubat and Matwin, 1997; Barandela et al., 2003) as the loss function to minimize on. Using both weight balance and G-Mean loss means that it eliminates the need for oversampling. However, we want to evaluate whether oversampling is equivalent and interchangeable with these techniques and does not negatively impact the results. The results of the various oversampling techniques are summarized in Table 1. The main metric used for comparison is G-mean loss then sensitivity and specificity and lastly accuracy. Improvements against no oversampling based on the loss function is observed for SMOTE with all variations but SVM. However, with borderline 1 variation one can observe a significant improvement especially with specificity without sacrificing significantly the model’s sensitivity. It obtained a G-mean loss value of 0.3268, sensitivity of 0.8621, specificity of 0.5 and accuracy of 0.7838. Hence, for the model evaluations the low risk class of the training set was oversampled using SMOTE borderline 1. Oversampling was used until that of the high risk, that is 286.

4 MODEL EVALUATIONS

4.1 Non-neural Network Models

Non-neural network models are still a vital alternative to neural network ones and can set a good baseline for future models (Wilkins et al., 1996; Lim et al., 2000). Non-neural networks models usually can train their weights with much less time than compared to their counterparts. Also, this leads to requiring fewer computational resources, allowing them to be trained on personal machines. Lastly, they require less hyperparameter tuning and setup time and do not require an architecture to be designed specifically for the problem, making them production-ready sooner. While having all these benefits, the results can also be in par with what is obtained from neural networks but depends on the complexity of the problem at hand. Additionally, the best non-neural network model can act as a base line for comparison of various network architectures.

The models evaluated were Random Forest (RF), XGBoost (Chen and Guestrin, 2016), K-Nearest Neighbors (K-NN) (Cover and Hart, 2006), Support Vector Machine (SVM) with linear kernel and radial basis function (RBF) (Burges, 1998; Cortes and Vapnik, 1995). The algorithms were obtained from the scikit-learn library, apart from XGBoost which was obtained from its own library (Chen and Guestrin, 2016). Each of these models, their optimal hyper-parameters were determined through the usage of the hyperopt library with the TPE optimizer. Additionally, the loss function used to minimize the error on was the weighted G-mean loss.

From the models evaluated, the top performer based on the lowest achieved weighted G-mean loss value is XGBoost with a value of 0.3994. It also obtained a decent sensitivity of 0.7069 but just 0.5 on specificity and a biased accuracy of 0.6622. Following came SVM with linear kernel obtaining a weighted G-mean loss of 0.4241 and closely in third RF with 0.4281. K-NN and SVM with RBF obtained significantly worse results with a weighted G-mean loss of 0.4829 and 0.5687 respectively. The full summary of the results on the test set are shown in Table 2.

4.2 Neural Network Models

Here the neural networks will no longer directly use the weighted G-mean loss function to optimize the parameters on but instead use a categorical cross entropy function (de Boer et al., 2005) to measure the error of the network. The weighted G-mean loss was not used because it is not possible to obtain a differentiable global G-mean loss on batch operations. However, to be able to compare the results to that obtained in the non-neural networks a non-weighted G-mean batch-wise loss function was applied. Also, to be able to obtain a respective weighted loss value from the batch-wise loss function class weight balancing was preferred over oversampling. Additionally, it was used as an early stopping criteria on the validation set. Its implementation is the same as that of a normal G-mean loss function and was executed at the end of each batch, which was set to a size of 50 samples, during training and obtained the average at the end of each epoch.

By using categorical cross entropy, the class labels were transformed to binary values by applying one-hot encoding. Hence, the classes were represented as vectors with the low risk class (0) as (1, 0) and the high risk (1) as (0, 1). Additionally, the loss function assumes that the passed input represents the probability for each encoding to be true. That is, it expects a vector which sums to 1 and each individual value is within [0, 1]. For the network to oblige by this constraint, the output layer’s activation function used was a softmax function (Bishop, 2006) which from a vec-
Table 1: Summary of the results on the test set of a random forest classifier when using oversampling on the least represented class (low risk) in the data set so it becomes balanced.

<table>
<thead>
<tr>
<th>Oversampling</th>
<th>G-Mean Loss</th>
<th>Accuracy</th>
<th>Sensitivity</th>
<th>Specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td>No oversampling</td>
<td>0.3894</td>
<td>0.7702</td>
<td>0.8793</td>
<td>0.375</td>
</tr>
<tr>
<td>Random</td>
<td>0.3994</td>
<td>0.6622</td>
<td>0.7069</td>
<td>0.5</td>
</tr>
<tr>
<td>SMOTE regular</td>
<td>0.3749</td>
<td>0.6622</td>
<td>0.6897</td>
<td>0.5625</td>
</tr>
<tr>
<td>SMOTE borderline1</td>
<td>0.3268</td>
<td>0.7838</td>
<td>0.8621</td>
<td>0.5</td>
</tr>
<tr>
<td>SMOTE borderline2</td>
<td>0.3693</td>
<td>0.7568</td>
<td>0.8448</td>
<td>0.4375</td>
</tr>
<tr>
<td>SMOTE SVM</td>
<td>0.4126</td>
<td>0.7297</td>
<td>0.8276</td>
<td>0.375</td>
</tr>
<tr>
<td>ADASYN</td>
<td>0.401</td>
<td>0.7027</td>
<td>0.7759</td>
<td>0.4375</td>
</tr>
</tbody>
</table>

Table 2: Summary of the results on the test set for the non-neural network models.

<table>
<thead>
<tr>
<th>Model</th>
<th>G-Mean Loss</th>
<th>Accuracy</th>
<th>Sensitivity</th>
<th>Specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF</td>
<td>0.4281</td>
<td>0.7027</td>
<td>0.7831</td>
<td>0.375</td>
</tr>
<tr>
<td>XGBoost</td>
<td>0.3994</td>
<td>0.6622</td>
<td>0.7069</td>
<td>0.5</td>
</tr>
<tr>
<td>K-NN</td>
<td>0.4829</td>
<td>0.527</td>
<td>0.5345</td>
<td>0.5</td>
</tr>
<tr>
<td>SVM Linear Kernel</td>
<td>0.4241</td>
<td>0.6216</td>
<td>0.6551</td>
<td>0.5</td>
</tr>
<tr>
<td>SVM RBF Kernel</td>
<td>0.5687</td>
<td>0.7432</td>
<td>0.7826</td>
<td>0.0625</td>
</tr>
</tbody>
</table>

tor of real number outputs a probability distribution. The optimal hyperparameters were found through the usage of grid search based on the validation results of the G-mean loss. Additionally, the architecture which includes the number of layers and neurons, activation functions, optimizers and regularization methods were determined through experimentation with a variety of combinations. All the networks were implemented using Keras (Chollet et al., 2015) with Tensorflow (Abadi et al., 2016) backend.

4.2.1 Cascade Correlation Neural Network

On this specific data set, the best performing diagnostic model concluded from a variety of models from a previous research (Zenovich et al., 2016) was a Cascade Correlation Neural Network (CCNN) (Fahlman and Lebiere, 1990). Subsequent goal of this paper is to further explore and improve the CCNN. For the evaluation, the previous network will be re-implemented so results are comparable. This model will be distinguished as the base CCNN model. Further, taking advantage of the previously positive results, another two variations are being proposed in this paper and are defined as improved and extended CCNN models, in an attempt to further refine the results.

The CCNN was proposed by Fahlman and Lebiere (1990) as an approach that is not only limited to tuning the parameters of the network but also dynamically determining the optimal architecture, constraint to the number of hidden layers. The network initial consists of a fully connected input and output layers, which their size is defined by the problem. Then the algorithm executes these following steps until convergence:

- All weights of units connected to the output layer are trained until the minimum error is reached
- A pool of candidate units are generated which have as input the output of all previously added layers excluding the output layer’s
- These candidate units are trained such that their output maximizes the correlation coefficient between the residual error of the network
- The candidate that has the maximum correlation is selected to be added to the network. Its input weights are frozen and its output is connected with the output layer

The network continues this iterative process until the addition of a unit does not lead to a smaller error than the previous execution.

The base CCNN model reflects closely the initially proposed algorithm (Fahlman and Lebiere, 1990) with some minor changes over and above those mentioned in section 4.2. The hidden and candidate units used the sigmoid function as their activation function. Additionally, the weights were initialized randomly from a normal distribution which had a mean of 0 and standard deviation of 0.5 and the bias was set to 0. After every loop the weights of the
output layer were reinitialized to avoid being stuck at bad local minimums. Furthermore, the optimization function used was Stochastic Gradient Descent (SGD) (Bottou, 2010). Its learning rate was set to 0.00001 and 0.000005 for the output and hidden candidate layers respectively. Noting that in the previous research (Zenovich et al., 2016) the authors used Simulated Annealing, but SGD was preferred hoping for better generalization. Finally, the candidate pool size was set to 16 and each candidate layer had two units, the same as the output layer.

For the proposed improved model, only the differences from the base one will be noted. The focus of this CCNN model is to utilize more recent techniques to improve performance. Firstly, the weight initialization scheme was changed from random distribution to Xavier (Glorot and Bengio, 2010) sampling from a normal distribution. Also, the optimizer was changed to Adam (Kingma and Ba, 2014) as a further improvement to SGD. Its learning rate was set to 0.00001 and 0.000005 for the output and candidate layers respectively. For both cases, the decay of first-order gradient to 0.9, decay of second-order gradient to 0.99 and a small epsilon of 1e-08. Additionally, the activation functions of the hidden layers were changed to Rectified Linear Units (ReLUs) (Nair and Hinton, 2010). Lastly, for the output layer warm-start weight initialization was added to carry over weights that contributed the most to lowering the loss value.

The extended model, building from the improved model, focused on further expanding the capabilities of dynamically constructing the architecture by also introducing regularization layers to the pool in an attempt to improve generalization. The hidden candidate layer was changed to have the following format and strict order:

- Gaussian noise layer with a mean of 0 and a standard deviation of 0.5
- Dense layer (original unit)
- Batch normalization layer (Ioffe and Szegedy, 2015) with momentum at 0.99, epsilon at 0.00001 and a trainable beta value
- Dropout layer (Srivastava et al., 2014), which randomly drops one of the two units

The candidate pool consisted twice of all possible combinations of the regularization layers, while strictly maintaining the order presented. Thus, the total pool size was maintained to 16 with only two candidate layers being the same in comparison to all 16 in the two previous models.

Based on the G-mean loss value on the test set the best performer from the CCNNs was the improved variation with a value of 0.5417, accuracy of 0.5541, sensitivity of 0.6207 and specificity of 0.375. A marginal difference followed the extended model with G-mean loss of 0.5495 and lastly, with significantly worse results, the base model with a value of 0.5889. The full summary of the results on the test set can be found in Table 3.

On the validation results there is a significant point to note out. The G-mean loss value obtained by the models on the validation set were 0.3512, 0.2677, 0.1578 for the base, improved and extended models respectively. The extended model was able to extract more information from the training set to improve its score on that of the validation. However, having nearly the same score as the top performer, there was at least no loss of information compared to the improved model, but the recognition of patterns that were useful on the validation set were not so for the test set. This is possibly due to the fact that the data set contains considerable number of outliers, as far as the network is concerned. This in turn prevents separation of the data in such a way that each set samples from the possible distribution of the problem, hindering generalization capabilities.

### 4.3 Deep Neural Network

A Deep Neural Network (DNN) was also constructed to compare the performance of the cascade networks. Specifically, it was used to evaluate the results, training speed and memory usage between the models. The design of the DNN was based on the results obtained previously in section 4.2.1. Thus, the network will also focus on using various generalization methods.

The DNN’s hidden layers used ReLu for their activation function and their weights were initialized used Xavier’s method. Also, the optimizer used was Adam with a learning rate of 0.00005, the decay of first-order gradient at 0.9, decay of second-order gradient at 0.999 and an epsilon value at 1e-8. For regularization, Gaussian noise layers, with a standard deviation of 0.2 and mean of 0, and dropout layers, with 20% dropout rate, were included in the model. Additionally, batch normalization layers were added with momentum set to 0.99, epsilon to 0.0001 and a trainable beta value. Lastly, details described in section 4.2 still apply here. The final layout of the network consists of five hidden layers, excluding the input and output layers. The network’s architecture was formed as following:

- Input layer with 40 units
- Batch normalization, Gaussian noise and dense layer with 1000 units
Table 3: Summary of the results on the test set for the neural network models.

<table>
<thead>
<tr>
<th>Model</th>
<th>G-Mean Loss</th>
<th>Accuracy</th>
<th>Sensitivity</th>
<th>Specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base CCNN</td>
<td>0.5889</td>
<td>0.4324</td>
<td>0.4483</td>
<td>0.375</td>
</tr>
<tr>
<td>Improved CCNN</td>
<td>0.5417</td>
<td>0.5541</td>
<td>0.6207</td>
<td>0.3125</td>
</tr>
<tr>
<td>Extended CCNN</td>
<td>0.5495</td>
<td>0.5405</td>
<td>0.6034</td>
<td>0.3125</td>
</tr>
<tr>
<td>Deep Neural Network</td>
<td>0.2843</td>
<td>0.7703</td>
<td>0.8103</td>
<td>0.625</td>
</tr>
<tr>
<td>Convolution Neural Network</td>
<td>0.3637</td>
<td>0.6081</td>
<td>0.5862</td>
<td>0.6875</td>
</tr>
</tbody>
</table>

- Batch normalization, dropout, Gaussian noise and dense layer with 200 units
- Batch normalization, dropout, Gaussian noise and dense layer with 200 units
- Batch normalization, dropout, Gaussian noise and dense layer with 200 units
- Batch normalization, dropout and dense layer with 200 units
- Dense output layer with 2 units

The DNN was able to obtain a G-mean loss of 0.2843, accuracy of 0.7703, sensitivity of 0.8103 and specificity of 0.625 on the test set, as shown in Table 3. The results obtained are significantly better than that obtained from the improved CCNN, which had a G-mean loss of 0.5417. It was also able to achieve this with a noticeably faster training time. However, the CCNN model was able to obtain its results requiring less memory, as it constructed a network with a total of 47 hidden layers with 2 units each based on the improved variant. But with today's state of available hardware the memory usage from the DNN is not of a concern.

4.3.1 Convolutional Neural Network

Convolutional Neural Networks (CNNs) have shown great results for detecting breast cancer using various imaging data (Ciresan et al., 2013; Spanhol et al., 2016; Arajo et al., 2017). In an attempt to further improve the results of the DNN, a CNN model was also explored. Its design was based on the previously obtained results with higher focus around its ability to generalize.

Building a CNN implies that the input will be a 2D image with one or more channels (usually color). Thus, the input vector was transformed to a 2D image with two channels. The channels were used to represent the measured data at the skin and at a depth. The image itself will be of size 13 x 6 containing the normalized measurements from both glands and axillary points. The positioning on the image resembles closely to that of the Figure 1, while also obtaining the average of neighboring positions to better represent the overlap as depicted in Figure 3. The values, before being formed to an image representation, they were centered using a robust scaler based on the interquartile range to maintain outliers.

Figure 2: The average batch-wise G-mean loss and accuracy of the deep neural network as it is trained.

As seen in Figure 2, the regularization techniques prevented overfitting the training data against the validation. While the network has extracted all possible information from the training set need to classify those samples, it does not cover all possible cases in the validation set. The limitation of the model is derived again from the limited available data in expressing an accurate distribution of the problem within the three sets.

Image Transformation Calculations

<table>
<thead>
<tr>
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Figure 3: Methodology in transforming a vector of temperature measurements from the mammary glands, for both at the skin and at a depth, to a 2D array. The L represents the left gland and R the right gland. Any cells left blank have a value of 0.

The training set used for the CNN was oversampled, as described in section 3.3, and then applying image augmentations. The intention of this was to
obtain a more rotation invariance network when detecting features on the glands, which in turn should further improve generalization. The type of augmentations applied were image flipping on the vertical axis and rotations of the outer pointer of the mammary glands. The result was a total of 5472 samples split equally between low and high risk for the training set.

The hidden layers of the network used ReLu activation functions and Adam optimizer with a learning rate of 0.0000005, decay of first-order gradient at 0.9, decay of second-order gradient at 0.999 and epsilon at 1e-8. Additionally, the weights of all the layers were initialized using the Xavier method from a uniform distribution. The type of layers used were dense, convolutional (Lecun et al., 2015), separable convolution (Chollet, 2016), max pooling (Lecun et al., 2015), global average pooling (Lin et al., 2013), dropout, spatial dropout (Tompson et al., 2014), batch normalization and Gaussian noise. The convolution and pooling layers used a kernel of size 3x3, stride of 1 with the exception of spatial which used 2, padding set to same and no bias value. Lastly, all dropout layers had a dropout percentage of 20%. The full network architecture was defined as following: Input layer of size 13x6x2 Convolutional with 64 units
- Batch normalization, ReLu activation and convolutional with 64 units
- Batch normalization, ReLu activation, Gaussian noise with standard deviation of 0.01, max pooling and convolutional with 128 units
- Batch normalization, ReLu activation, convolutional with 128 units
- Batch normalization, ReLu activation, Gaussian noise with deviation of 0.001, max pooling and convolutional with 256 units
- Batch normalization, ReLu activation and convolutional with 256 units
- Batch normalization, ReLu activation, Gaussian noise with deviation of 0.001, spatial dropout, max pooling and separable convolution with 512 units
- Gaussian noise with deviation of 0.0001, batch normalization, ReLu activation and separable convolution with 512 units. Repeated four times Gaussian noise with deviation of 0.0001, batch normalization, ReLu activation, global average pool, dense with 512 units
- Batch normalization, ReLu activation, dropout, Gaussian noise with deviation of 0.1 and dense with 2 units

Figure 4: The average batch-wise G-mean loss and accuracy of the convolutional neural network as it is trained.

The CNN obtained a G-mean loss value of 0.3637, accuracy of 0.681, sensitivity of 0.5862 and specificity of 0.6875, which are included in Table 3. The network was not able to outperform that of the DNN based on the G-mean loss but did obtain the highest specificity rate from all other models. Additionally, the Figure 4 shows the training and validation G-mean loss and accuracy as the training of the network progresses. With the addition of augmentation it should of helped with generalization, but there was still some slight overfitting of the training set against the validation. Finally, there is a similar pattern as before where limited information from the training set can be generalized to the validation.

5 CONCLUSIONS

The best performing model introduced in this paper is the DNN, described in section 4.2.2, with a weighted G-mean loss of 0.2843 on the test set. Followed was the CNN with a loss value of 0.3637 and in third place XGBoost with a value of 0.3994. The three variants of CCNN were not able to outperform the non-neural network models, with the exception of SVM with RBF kernel.

The results of the DNN indicate the potential in automating readings from radiometry for the purposes of diagnosis or monitoring cancer patients, which is not only limited to breast cancer. The models here used only temperature readings so that the effectiveness of microwave radiometry in medical applications can be evaluated. Taking that into account, by including additional information from other systems and what is recorded from a clinical professional about the physiological condition of each person (Zenovich et al., 2016), it is expected to further
improve the capabilities of such a system. In addition, a more exhaustive search on deep neural networks should be conducted as they show promising results and can potentially bring further improvements.

As a potential diagnostic system to aid clinical professionals in making decisions it currently returns limited information, low or high risk of cancer with a prediction confidence. The problem is oversimplified due to limited amount of data. As more descriptive and broader data becomes available it can be expanded from a binary to a multi-class task. Some possible labels are benign and malignant tumors, noncancerous tumors, inflammation, infection and healthy patients. Additionally, as a future scope, the networks can be reevaluated and altered, as needed, when readings are obtained for cancerous tumors at various body locations.

The proposed variants of the CCNNs can be even further improved on the aspect of dynamic construction of the architecture. Additional suggestion is to allow the network to try against various activation and optimization functions and conduct hyperparameter optimization through an online grid search. However, this drastically increases the number of possible combinations for the network to explore, hence in-creased training time, for a possibly small improvement. Moreover, there is a risk with the preferential selection of units against the validation set. With such a fine level of selection and with the added high amount of noise it is possible to overfit the validation set. Hence, shifting the problem from generalizing from the training set to generalizing from the valida tion set.

REFERENCES


