Prediction of Thyroid Malignancy Using Contextual Semantic Interpretability from Sonograms

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Abstract: The gold standard in thyroid nodule malignancy diagnosis consists of ultrasound (US or sonogram) guided fine needle aspiration biopsy. This procedure is ordered based on an assessment of malignancy risk by a trained radiologist, who uses US images and relies on experience and heuristics that are difficult to effectively systematize into a working algorithm. Artificial Intelligence (AI) methods for malignancy detection in sonograms are designed to either perform segmentation (highlight entire thyroid gland and/or nodule) or output a probability of malignancy. There is a gap between AI methods trained to perform a specific task using a black-box method, and the sonogram features (e.g., shape, size, echogenicity, echotexture) that a radiologist looks at. We aim to bridge this gap, using AI to reveal saliency in sonograms for features that are easily understood by clinicians. We propose a deep-learning model that performs two tasks important to radiologists: sonogram feature saliency detection, as well as probability of malignancy. We perform both a quantitative and qualitative evaluation of our method using an open dataset, the Thyroid Digital Image Database (TDID). Our framework achieves 72\% accuracy in the task of classifying thyroid nodules as benign or malignant.

1 INTRODUCTION

Thyroid cancer cases have been on a rising trend worldwide at a rate of around 3\% yearly increase in incidence, doubling every 30 years (Morris et al., 2013). While this increase may be attributed to better access to care, it is significant, and early detection leads to better outcomes and increased 5-year survival rates (Siegel et al., 2019). The most commonly used imaging modality for the thyroid is ultrasound, which is non-invasive, fast and relatively inexpensive. Usually, an initial malignancy risk evaluation is done for thyroid nodules, based on the sonogram features.

The American College of Radiology (ACR) assembled committees to develop a standardized risk-stratification system. The committees’ goals were to develop management guidelines for incidentally discovered nodules, produce a lexicon for sonogram nodule description, and to develop a scoring system, named TI-RADS (Tessler et al., 2017).

The TI-RADS system observes five sonogram feature categories. Each category contains attributes that receive a number of points to be added into a final score. The higher the final score (TR1 through TR5), the higher the risk of malignancy. This scoring system is useful to guide further clinical actions, such as watchful-waiting or fine-needle aspiration (FNA). The point system is defined as follows:

- **Composition** of the nodule. Cystic and spongiform compositions receive no points. Mixed cystic and solid receive one point, while solid or almost completely solid receive two points.
- **Echogenicity** refers to a tissue’s ability to bounce an echo from the probe. Anechoic tissues receive zero points. Hyperechoic and isoechoic tissues receive one point each. Hypoechoic or very hypoechoic echoes receive two and three points, respectively.
- **Shape** refers to the orientation of a mostly elliptical nodule in a sagittal and transverse sonogram. A nodule that is wider-than-tall receives zero points, while a nodule that is taller than wide receives three points.
- **Margins** refers to the edges of the nodule. Ill-
defined or smooth margins receive zero points. Lobulated margins receive two points. When extensions outside of the thyroid are observed, it is highly indicative of malignancy and this feature receives three points.

- **Echogenic foci** refer to bright spots seen on a sonogram. These can be not present, or very large V-shaped and receive zero points. Macrocalcifications receive 1 point. If foci are observed peripherally, this receives 2 points. Microcalcifications receive the highest score of three.

Existing work in the medical image analysis literature focuses heavily on providing additional automated diagnostic tools whose goal is to assist clinicians in their decision making processes. These systems are functionally black-box approaches, such as deep convolutional neural networks (CNNs, along with their many variants) that take images and metadata as the inputs and produce a probability of malignancy, a score or a combination thereof. These existing systems can easily generate an answer to *what?* but little attention has been given to the *why?*

Our work, inspired in part by (Marcos et al., 2020), focuses on a method to help humans understand the *why?* in decisions made by highly complex multi-billion parameter models. Our primary contribution is a method that automatically produces a human understandable explanation for a decision made by a complex system. We achieve this by explicitly modeling contextual information as defined by TI-RADS, and automatically discovering features that are meaningful to predict malignancy when linearly combined into a final score.

In this paper, we use the Thyroid Digital Image Database (TDID) (Pedraza et al., 2015) for our experiments. We propose a technique for the prediction of features or attributes of thyroid nodules from sonograms using deep learning. These features are then used to make interpretable predictions about the possibility of malignancy for a given thyroid nodule. Thus, the predicted features and the proposed framework provide insights into why a thyroid nodule is predicted as benign or malignant.

### 2 RELATED WORK

The recent successes of artificial intelligence (AI) systems has made it possible to design and implement various systems. However, their results are often difficult to interpret leading to mistrust. “Explainable Artificial Intelligence” or xAI is a field that aims to bridge this gap by developing methods that try to make the output of AI algorithms understandable.

**eXplainable AI.** A common distinction in xAI methods is made on the basis of the timing of the explanation with respect to training at which explainability is implemented (Speith, 2022). Ante-hoc methods are applied before training while post-hoc methods, like ours, rely on either pre-trained or end-to-end trained DNN models and generate explanations for a non-interpretable, black box model.

Among post-hoc methods for explanations, saliency-based methods (Simonyan et al., 2013) assign an importance to each pixel with respect to
the image’s predicted class. Saliency-based methods are often unreliable due to highlighting edges in images (Rudin, 2019).

**Class Activation Maps for Explainability.** In addition to the above techniques, Class Activation Mapping (CAM) is widely used to improve the explainability of CNN models. The method proposed by (Zhou et al., 2015) uses the features output from the final convolution layer to determine the important parts of an image that were identified by a CNN for classification. Gradient-weighted Class Activation Mapping (Grad-CAM) (Selvaraju et al., 2019) and GradCAM++ (Chattopadhay et al., 2018), use the gradients of a target to produce a localization map. Ablation-CAM (Desai and Ramaswamy, 2020) uses ablation studies to identify weights of individual feature maps for a specific class. Score-CAM (Wang et al., 2019) uses the global contribution of the corresponding input features. While SS-CAM (Wang et al., 2020) introduces smoothing to Score-CAM, producing sharper visualization and localization, IS-CAM (Naidu et al., 2020) introduces the use of integration in Score-CAM to generate sharper activation maps.

**Thyroid Ultrasound.** Wu et al. (Wu et al., 2016) showed echogenicity to be an important predictor of malignancy while Alexander et al. (Alexander et al., 2004) showed that spherical shape is associate with an increased rate of malignancy in solid nodules.

Thus, the diagnosis of whether a thyroid nodule is benign or malignant depends on a combination of features (attributes). There have been several attempts to directly classify sonograms of thyroid nodules as benign or malignant (Kwon et al., 2020) (Wang et al., 2022) (Koh et al., 2020a) (Liang et al., 2020), and some techniques use features extracted from CNNs to perform the classification using a separate model (Chi et al., 2017) (Lee et al., 2019). However, the use of post-hoc xAI and concept bottlenecks (Koh et al., 2020b) to identify and verify the relation of features or attributes such as echogenicity, margins, and composition in the automatic classification of thyroid nodules as benign or malignant has not been explored.

## 3 PROPOSED METHOD

### 3.1 Data

The Thyroid Digital Image Database (Pedraza et al., 2015) consists of 65 thyroid sonograms confirmed malignant and 33 sonograms of benign cases. Each image is annotated by a radiologist and contains classifications with respect to composition, echogenicity, margins, and calcifications. With respect to composition, the nodules are classified as cystic, spongiform in appearance, predominantly solid, or solid. With respect to echogenicity nodules are classified as hypoechoic, hyperechoic, marked hypoechogenic or isoechogenic. In terms of margins, the categories are ill-defined, micro-lobulated, speculated or well-defined smooth. Finally, the types of calcifications are micro-calcifications, macro-calcifications or no calcifications. Besides, each image is classified as benign or malignant. We randomly split the dataset of 98 images into a training and a test set. Our training set consists of 73 images and the test set consists of 25 images.

### 3.2 Proposed Deep Learning Architecture

We design an end-to-end neural network that is trained to classify images of thyroid nodules as benign or malignant. Our method takes the thyroid nodule sonograms as inputs and outputs the predicted attributes with respect to composition, echogenicity, margins and calcifications as well as a probabilistic prediction of malignancy.

Our framework consists of two parts that are trained in an end-to-end manner. The first part of the model predicts the classification of nodules in terms of the attributes (composition, echogenicity, margins and calcifications) and the second part of the framework uses the predicted attributes to predict nodule malignancy.

#### 3.2.1 Input and Data Augmentation

We use a two channel input, where the first channel consists of the original sonogram and the second channel consists of the masked sonogram. The mask blocks all parts of the image except the nodule(s). Our decision to use this 2 channel image is based on the work of Liang et al. (Liang et al., 2020) where the use of the mask improved the accuracy of detection of the type and the nature of the disease in comparison to using the original image.

In this work, the input training images undergo augmentations with a probability of 0.5. The augmentations used are left-right flip and up-down flip.

#### 3.2.2 Prediction of Attributes

We use a VGG-16 (Liu and Deng, 2015) backbone for our end-to-end model, pre-trained on ImageNet.
We fine-tune it for our application. Since the inputs are two channel images, we introduce an additional, convolutional layer to the original VGG-16 architecture that takes a two channel image as an input and outputs three feature maps.

Each attribute (e.g., echogenicity) has several potential values (e.g., iso, hyper and hypoechoic). We encode these attributes as groups of one-hot vectors. The output from this part of our framework consists of 15 (sum of all possible values of all attributes) probabilities, where each one of these indicates the probability that the nodule possesses a specific attribute value. If the predicted probability for a specific binary attribute is 0.5 or more, the nodule is said to possess the corresponding binary attribute. For this purpose, we use the sigmoid activation function instead of ReLU to re-scale the outputs of the final, linear layer, to values between 0 and 1, which indicate probabilities.

3.2.3 Generation of Activation Maps

We further modify the modified VGG-16 network from Section 3.2.2 to facilitate the generation of activation maps for each image. This indicates the part of the image that played an important role in making a prediction for each attribute. For this purpose, we implement the Class Activation Map (CAM) concept proposed in (Zhou et al., 2015). We remove two of the three linear layers in the VGG architecture, while retaining the final linear layer to compute the outputs. We introduce a Global Average Pooling (GAP) (Lin et al., 2013) layer before the final, linear layer in the VGG-16 model.

To generate the activation maps for a specific class, we multiply each feature map that is output by the layer preceding the GAP layer, by the corresponding weight in the final, linear layer. This gives us 512 weighted feature maps for each class. We then sum up the weighted feature maps for each class to get the class activation map for each binary attribute. Figure 2 contains a pictorial representation of how the CAM technique is used to generate the activation maps for the binary attributes.

3.2.4 Prediction of Nodule Type Using Binary Attributes

We use the predicted probabilities of each binary attribute to predict the tumor type using an interpretable function. Our final model uses a linear layer to compute the probability of malignancy for a thyroid nodule by using the predicted probabilities for each binary attribute as inputs. The linear layer is restricted to have zero bias and learns a matrix $W$, representing the weight of each of the binary attributes in deciding
whether the nodule is benign or malignant. After each training iteration, the weights are re-scaled to ensure that they are all positive and that they sum up to 1. We use Sankey plots to pictorially represent the weights in the matrix \( W \) and to facilitate the process of understanding the model. We use a sigmoid function to convert the output of the linear layer of this module into a value between 0 and 1 to represent the predicted probability of malignancy. Figure 3 provides a pictorial representation of our end-to-end architecture.

We attempt to use a second model that uses two linear layers to predict malignancy using the binary attribute predictions as the inputs, inspired by (Marcos et al., 2020). The first layer groups \( A \) binary attributes into \( Z \) groups. This operation is performed by learning the weights in a \( A \times Z \) matrix, \( G \), which outputs group probabilities to group binary attributes together into groups. The group presence probabilities, \( z \) are computed using \( z = G a \). We use another linear layer to compute the probability of a thyroid nodule being malignant by using the group presence probabilities, \( z \) as the inputs and identifying the relation between group probabilities and predictions about malignancy. For this, the values (weights) in a \( Z \times 1 \) matrix, \( Q \) are learnt. The probability of a tumor being malignant, \( y \) is computed using the equation \( y = Q z \).

4 EXPERIMENTS AND RESULTS

4.1 Experimental Setup

Our code is implemented in python, and we use PyTorch to implement our framework. Our code is available at: https://github.com/rpmihail/thyroid.git. We run our training and inference experiments on an NVIDIA GeForce RTX 3090 GPU with 24 GB memory. The processor is a 64-bit, 3.7GHz Intel i9 computer with 32 GB RAM. The OS used is Ubuntu 22.04 LTS.

4.2 Training

The framework described in Section 3.2, is trained in an end-to-end manner. It is trained for 1000 epochs using a learning rate of 0.000001 and the loss used is binary cross-entropy loss both for the binary attributes and for the prediction of nodule as benign or malignant. The optimizer used is Adam optimizer.

<table>
<thead>
<tr>
<th>Backbone</th>
<th>Average accuracy (binary attributes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>VGG-16</td>
<td>76%</td>
</tr>
<tr>
<td>ResNet-50</td>
<td>75%</td>
</tr>
</tbody>
</table>

4.3 Ablation Studies

We perform various ablation studies to identify the best set of augmentations, backbone and the number of layers for predicting type.

4.3.1 Identifying the Better Backbone

We tried ResNet50 (He et al., 2015) as an alternate backbone. However, as shown in Table 3, the VGG-16 backbone achieves marginally higher average accuracy in terms of the prediction of binary attributes.

4.3.2 Selecting Augmentations

We consider four types of augmentations for our framework: flip in left/right direction; flip in up/down direction; rotate by a random multiple of 90°; and random shift in one of 4 directions by 100 to 126 pixels. As shown in the results in Table 2, the use of right/left and up/down flip result in improvements in the binary attribute prediction accuracy.

<table>
<thead>
<tr>
<th>Augmentations</th>
<th>Average accuracy (binary attributes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>76%</td>
</tr>
<tr>
<td>Flip-ud</td>
<td>77%</td>
</tr>
<tr>
<td>Flip-ud + Flip-lr</td>
<td>79%</td>
</tr>
<tr>
<td>Flip-ud + Flip-lr + Rotate</td>
<td>76%</td>
</tr>
<tr>
<td>Flip-ud + Flip-lr + Random Shift</td>
<td>76%</td>
</tr>
</tbody>
</table>

4.3.3 Selecting the Number of Linear Layers for Type Prediction

While our work is inspired by (Marcos et al., 2020), the problem tackled in their work involves a concept that depends not only on the attributes, but on different combinations of the attributes. Hence, a 2-layer network is used for performing the final prediction from the attributes. However, our problem can be solved by a linear combination of the binary attributes. We experiment with one layer and two layer networks described in Section 3.2.4 to identify the better option. For the two layer networks, we vary the...
Figure 4: CAM for 4 different binary attributes using 4 sonograms. Each image has the mask for the nodule shown to the right. It is evident that the relevant parts of the image are indeed more significant for the prediction of binary attributes. Images are all for correct classification of images possessing the following attributes. a) Isoechogenicity b) Smooth margin c) No calcification d) Spongiform composition.

<table>
<thead>
<tr>
<th>Number of Layers</th>
<th>Type prediction accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>72%</td>
</tr>
<tr>
<td>2</td>
<td>72%</td>
</tr>
</tbody>
</table>

Table 3: Comparison of the type prediction accuracy when the prediction of type (benign or malignant) from binary attributes is performed using 1 and 2 layer linear networks.

Table 4: Binary attribute prediction accuracy for each binary attribute as well as the average binary attribute prediction accuracy and the type prediction accuracy.

<table>
<thead>
<tr>
<th>Binary Attribute</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cystic composition</td>
<td>96%</td>
</tr>
<tr>
<td>Predominantly solid composition</td>
<td>96%</td>
</tr>
<tr>
<td>Solid composition</td>
<td>64%</td>
</tr>
<tr>
<td>Spongiform composition</td>
<td>68%</td>
</tr>
<tr>
<td>Hyper echogenicity</td>
<td>96%</td>
</tr>
<tr>
<td>Hypo echogenicity</td>
<td>68%</td>
</tr>
<tr>
<td>Iso echogenicity</td>
<td>56%</td>
</tr>
<tr>
<td>Marked hypo echogenicity</td>
<td>96%</td>
</tr>
<tr>
<td>Ill-defined margin</td>
<td>68%</td>
</tr>
<tr>
<td>Microlobulated margin</td>
<td>92%</td>
</tr>
<tr>
<td>Spiculated margin</td>
<td>96%</td>
</tr>
<tr>
<td>Smooth margin</td>
<td>60%</td>
</tr>
<tr>
<td>Macro calcification</td>
<td>96%</td>
</tr>
<tr>
<td>Micro calcification</td>
<td>64%</td>
</tr>
<tr>
<td>No calcification</td>
<td>72%</td>
</tr>
<tr>
<td>Average binary attributes</td>
<td>79%</td>
</tr>
<tr>
<td>Benign / Malignant (type)</td>
<td>72%</td>
</tr>
</tbody>
</table>

4.4 Results

In Table 4, we tabulate the accuracy of our framework with respect to each of the binary attributes. We compute the average accuracy of prediction of the binary attributes by averaging the accuracy values we get for each of the binary attributes and also tabulate the type accuracy. As is evident from the table, there are a few binary attributes, such as isoechochogenicity and smooth margins that have lower prediction accuracy. Since deep learning models perform better when a large amount of data is used for training, the fact that our training set has only 73 samples is one of the major causes of this problem.

Our training set consists of only 29 different combinations of binary attributes. However, the total possible combinations of attributes are $4 \times 4 \times 4 \times 4 \times 3 = 192$. Besides, we identified that our randomly selected test set has 5 attribute combinations that are not present in the training set. This is a factor that is negatively affecting our type prediction accuracy. However, despite these challenges, our framework achieves a type prediction accuracy of 72%.

In Figure 4, we show some samples where we use CAM to identify the important parts of an image in making a decision about a specific binary attribute. For each image, we also show the masked version of
5 DISCUSSION AND LIMITATIONS

In Figure 5, the thickness of the edge connecting a binary attribute to the node for benign/malignant indicates its relative weight with respect to the other binary attributes. On matching our Sankey plot with the TI-RADS values in Figure 1, we see that the top 5 weights in the plot correspond to binary attributes that have a value of 2 or 3 according to TI-RADS. 6 of the bottom 7, including the missing binary attribute hyper echogenicity (which has a weight of 0), have a TI-RADS value of 0 or 1. However, mostly solid composition is incorrectly assigned a very low weight. Overall, while the model identifies important features for predicting the malignancy risk of nodules, there are some inconsistencies from TI-RADS. Our model is capable of separating the binary attributes with TI-RADS values of 0 and 1 from binary attributes with TI-RADS values of 2 and 3. However, it does not discover a decreasing order of importance according to TI-RADS.

The biggest and most important limitation of this study is the size of the dataset. The low number of sonogram samples is a significant limitation and contributes negatively to the prediction performance of our approach. However, the goal of this work was to add explainability to models in a domain where such work has not yet been attempted, to the best of our knowledge.

6 CONCLUSION

In this paper, we propose a framework for interpretable malignancy classification of thyroid nodules. Our framework uses deep learning to predict binary attributes from sonograms, then uses the predicted attributes in a linear layer to make predictions about the type of the tumor. We use CAM to visualize the parts of the images that are important for binary attribute prediction and Sankey plots to visualize the importance of each binary attribute in predictions about nodule type. The aim of this work is to ensure that deep learning is used to provide interpretable results in medical imaging.

In the future, we will explore different CAM techniques to generate more accurate maps. We will also work on improvements to the attribute prediction pipeline and attempt to use a larger dataset.

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