Deep Attention Model for the Hierarchical Diagnosis of Skin Lesions

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Abstract

Deep learning has played a major role in the recent advances in the dermoscopy image analysis field. However, such advances came at the cost of reducing the interpretability of the developed diagnostic systems, which do not comply with the requirements of the medical community nor with the most recent laws on machine learning explainability. Recent advances in the deep learning field, namely attention maps, improved the interpretability of these methods. Incorporating medical knowledge in the systems has also proved useful to increase their performance. In this work we propose to combine these two approaches in a formulation that: i) makes use of the hierarchical organization of skin lesions, as identified by dermatologists, to develop a classification model; and ii) uses an attention module to identify relevant regions in the skin lesions and guide the classification decisions. We demonstrate the potential of the proposed approach in two state-of-the-art dermoscopy sets (ISIC 2017 and ISIC 2018).

1. Introduction

The deep learning revolution has brought significant advances in various fields, particularly in medical image analysis [18]. In particular, the field of dermoscopy image analysis has seen a boom in the past few years, with several works relying on deep convolution neural networks (CNNs) to achieve increasingly better results in lesion diagnosis, which have started to rival the performance of trained dermatologists [12]. On par with CNNs, the release of public image data sets such as the ISIC archive [8] or PH2 [20], has also contributed to the growing interest on this topic.

There has been an effort to provide more insights on the factors that influence the performance of CNNs on lesion diagnosis, namely: i) performing data augmentation, both on the training [30] and test sets [23]; ii) using different architectures [14], as well as combining them in ensembles; iii) studying the best strategies for transfer learning; and iv) assessing the importance of image dimensions and other related properties [29].

Despite the aforementioned efforts, there are still aspects of the learning procedure that are not fully understood, while the decision process is not self-explainable. This hampers the transition of deep learning methods to the clinical practice, especially if one takes into consideration the recent European Unions General Data Protection Regulation, which mandates that any machine learning based method that is designed to interact with humans, such as a computer aided-diagnosis systems must be self-explainable. Therefore, it is important to design strategies that take advantage of the representation and classification power of CNNs, and at the same time are transparent and understandable [6].

Various strategies have been used to give interpretability to CNNs, such as: i) use of class activation (CAM) or attention maps that highlight the most relevant regions in the lesion for a decision and guide the network regarding the most discriminating feature maps [33, 25, 34]; ii) incorporation of the detection of relevant dermoscopic criteria in the CNNs (high level features) [16, 13]; and iii) incorporation of CNN features into a content-based image retrieval system (CBIR), which provides the user with examples similar to the one being diagnosed [9]. However, there are few evidences that these approaches lead to significantly better performance w.r.t to less transparent deep learning methods. Notorious examples are methods based on CBIR that demonstrated: i) the ability to be extended to the diagnosis of unknown diseases [27]; and ii) the capability to improve the diagnostic confidence of dermatologists [24].

In this work we propose a different approach, which takes into account the hierarchical organization of skin lesions (see Fig. 1). Lesions may be organized into hierarchical groups according to their origin (melanocytic or non-melanocytic), degree of malignancy (benign or malignant), and finally their diagnosis (e.g., melanoma, basal cell carcinoma, vascular lesion, or nevi) [2]. We explore this structure using as inspiration recent works on caption generation and attention models that use deep neural networks [32]. The developed deep learning system is able to sequen-
2. Related Work

Some works tried to introduce interpretability to CNNs in dermoscopy image analysis. Yang et al. [33] relied on the use of a multi-task CNN with CAMs, to simultaneously diagnose and locate the skin lesions in dermoscopy images. The obtained CAMs highlight the region of interest in the image, which usually matches the whole lesion. Zhang et al. [34] added attention modules to residual networks, to obtain improved feature maps between consecutive layers. These authors also resort to CAMs, and show that the proposed network focuses on the skin lesion and disregards most of the surrounding skin. Although some measure of interpretability may be obtained from CAMs, these maps usually highlight the entire lesion and are not able to specify smaller regions of interest, which may be associated with different dermoscopic structures. Van Simoens and Dhoedt [25] inspected the learned filter maps for a small CNN, and concluded that the model was able to learn filters that were sensitive to: border, lesion and skin color, hair, and artifacts. However, no diagnosis scores were provided.

Díaz [13] and Kawahara et al. [16] both used CNNs to detect relevant dermoscopic structures. The former proposed a dermoscopic segmentation structure network, whose output was a set of segmentation masks of different dermoscopic structures. These masks were then combined with the features maps obtained from a ResNet50 to guide the final diagnosis. The latter proposed a multi-task CNN that simultaneously evaluated the seven criteria of the 7-point checklist method [1] and provided a diagnosis. Both approaches showed promising results and allowed the integration of medical knowledge into the deep learning models. Nonetheless, the classification performance of such systems may be hampered by the incorrect identification of the dermoscopic structures. Moreover, these systems did not cover dermoscopic structures usually associated with non-melanocytic lesions, such as milia-like cysts [2].

Recently, Codella et al. [9] proposed a CBIR system to diagnose skin lesions based on deep learning features. This system was trained in a collaborative fashion, where human feedback regarding the similarity of the lesions was used as an input. Their results showed that the collaborative approach led to better results as well as the retrieval of lesions similar to the one being diagnosed, both in terms of visual properties and diagnosis.

None of the aforementioned works takes into consideration the inherent structure of skin lesions, i.e., the fact that they are organized into hierarchical groups. Usually, the diagnosis is treated as a one-vs-all problem, where the distinction between all the classes is performed in the same fully-connected layer. However, dermatologists must be able to distinguish between melanocytic and non-melanocytic lesions, prior to applying methods such as the ABCD rule or the 7-point checklist [2], which have been designed to diagnose only melanocytic lesions. Thus, adding this structure to the deep learning systems may lead to better results, as was recently demonstrated in [4]. Unfortunately, these systems lack interpretability.

In this paper we propose a deep learning system that is able to classify skin lesions in a hierarchical way and that at the same time is able to provide the user with information that may help interpret the classification results. The concept of hierarchy has been adopted before [12]. However, to the best of our knowledge, this is the first system that aims use hierarchy to leverage its interpretability.

3. Proposed Model

This paper proposes a new system for the classification of skin lesions, which is trained to mimic their sequential grouping from more general (melanocytic/non-melanocytic) to more specific (diagnostic) classes, as shown in Fig. 2.

We will treat the problem of sequential classification as one of caption generation [31], where the goal is to find
the correct sequence of words \( S = \{S_1, ..., S_T\} \) (\( T \) is the length of the sequence) that better describes image \( I \). In terms of inference, caption generation methods aim to find the sequence \( S \) that maximizes

\[
\log p(S|I) = \sum_{t=0}^{T} \log p(S_t|I, S_1, ..., S_{t-1}). \tag{1}
\]

This formulation allows us to condition a word \( S_t \) on all or a subset of previous words, up to \( t - 1 \). The probabilities \( p(S_t|I, S_1, ..., S_{t-1}) \) are commonly modeled with a recurrent neural network (RNN), where the hidden state \( h_t \) is responsible for propagating the condition on previous words. Interpretability may be incorporated into deep caption models, through the use of visual attention [32], making it possible to teach the network to focus on different salient regions to generate each word, and at the same time show what the model is “seeing”.

It is possible to establish a relation between caption models and the concept of hierarchical classification. In this type of classification scheme, the classes are organized in a tree structure, such that the ones at the lower levels are conditioned on those at the upper levels. Thus, we can formulate our classification problem as in (1): we want to find the best sequence of classes \( S \) for an image \( I \), imposing that each of the generated classes \( S_t \) is conditioned on the previous ones \( \{S_1, ..., S_{t-1}\} \). We also incorporate an attention module into our system to identify regions of interest in the images (see Fig. 2). In Section 4, we detail the proposed model.

### 4. Hierarchical Attention CNN-LSTM

The proposed model may be divided into three main blocks, as shown in Fig. 2: i) the encoder that receives a raw image and extracts features; ii) the decoder that sequentially generates the classes of the lesion; and iii) the attention module, which is responsible for guiding the decoder to attend at different locations and improve the transparency of the network. In the following subsections, we explain each of the blocks.

#### 4.1. Image Encoder

The encoding block receives as input a 299 \( \times \) 299 RGB image \( I \) and uses a bank of CNNs to extract a set of feature maps. In our experiments, we vectorize and concatenate the feature maps from the lower convolutional layers of two popular architectures: DenseNet161 [15] and ResNet-Inception [26]. This produces a set of \( L \) vector descriptors \( x = \{x_1, ..., x_L\}, x_l \in \mathbb{R}^D \), where \( \sqrt{D} \times \sqrt{D} \) is the shape of each of the feature maps. It is possible to associate these feature maps with portions of the image \( I \), which will allow us to identify regions of interest using the attention module, as described in sub-section 4.3.

#### 4.2. Decoder - Hierarchical Diagnosis

The decoder sequentially diagnoses the dermoscopy image, using a long short-term memory network (LSTM) of dimensionality \( P \), as proposed in recent works on image captioning [31, 32]. This network sequentially produces the hierarchical classes of a skin lesion, conditioned on a context vector \( z_t \), the previous hidden state \( h_{t-1} \), and the previously generated class \( S_{t-1} \), according to the following implementation

\[
i_t = \sigma(W_{iw}E_{S_{t-1}} + W_{ih}h_{t-1} + W_{iz}z_t + b_i),
\]

\[
f_t = \sigma(W_{fw}E_{S_{t-1}} + W_{fh}h_{t-1} + W_{fz}z_t + b_f),
\]

\[
c_t = f_t c_{t-1} + i_t \tanh( W_{cz}E_{S_{t-1}} + W_{ch}h_{t-1} + W_{cz}z_t + b_c),
\]

\[
o_t = \sigma(W_{ow}E_{S_{t-1}} + W_{oh}h_{t-1} + W_{oz}z_t + b_o),
\]

\[
h_t = o_t \tanh(c_t), \tag{2}
\]

where \( i_t, f_t, c_t, o_t \), and \( h_t \) are the input, forget, memory, output, and hidden state of the LSTM, respectively, at time-step \( t \). The weights \( W \) and biases \( b \) are learned during the training phase, as well as the embedding matrix \( E \in \mathbb{R}^{M \times K} \), where \( M \) is the word-embedding dimension and \( K \) is the number of words/classes. The context vector \( z_t \) is computed using the attention module, as described in the next section.

The memory state \( c_0 \) and the hidden state \( h_0 \) must be initialized. We adopted the strategy proposed in [32], and apply two separate perceptrons to predict the initial values given the average of the feature maps:

\[
p(S_t|x, S_{t-1}) = \text{softmax}(W_o(E_{S_{t-1}} + W_z z_t + W_h h_t)), \tag{3}
\]

At each time step, we can predict the next class \( S^*_t \) as the one that maximizes

\[
p(S_t|x, S_{t-1}) = \text{softmax}(W_o(E_{S_{t-1}} + W_z z_t + W_h h_t)), \tag{3}
\]
where $W_o \in \mathbb{R}^{K \times M}$, $W_z \in \mathbb{R}^{M \times D}$, and $W_h \in \mathbb{R}^{M \times P}$, where $P$ is the dimension of the hidden state, are trainable weight matrices.

### 4.3. Attention Module

This module estimates the context vector $z_t$, using the feature maps $x$ and a set of positive weights $\alpha_t = \{\alpha_{t1}, ..., \alpha_{tL}\}$. These weights are computed by an attention model, which we define as a multi-layer perceptron (MLP)

$$
\alpha_t = \text{softmax}(W_a(\tanh(W_{ax}x + W_{ah}h_{t-1}))),
$$

where $W_a \in \mathbb{R}^{L \times M}$, $W_{ax} \in \mathbb{R}^{M \times DL}$, and $W_{ah} \in \mathbb{R}^{M \times M}$ are all trainable weight matrices. Since $\alpha_t$ depends on the hidden state $h_{t-1}$ of the decoder, this allows us to simultaneously enforce the network to attend different locations in the image at each time instant, and at the same time condition it on the previously predicted classes.

Given the attention vector $\alpha_t$, it is possible to compute the context vector $z_t$. In this work we adopt the soft attention mechanism proposed by Chorowski et al. [7], which assumes that the weights express the relative importance of each feature map. Thus, the context vector may be defined as

$$
z_t = \sum_{i=1}^{L} \alpha_{ti}x_i.
$$

### 5. Experimental Setup

#### 5.1. Data sets and Experiments

The proposed model is evaluated on the publicly available ISIC 2017 data set [8]. This set contains 2,750 images each belonging to one of the following three classes: melanoma and nevus, both melanocytic, and seborrheic keratosis (non-melanocytic). All of the images were pre-processed in order to normalize their colors using the method proposed in [3].

The images are divided into three sets: training (2,000 images), validation (150 images), and test (600 images). The first subset was used to train different models, while the second subset was used to select the best combination of parameters. No external data sets were used to complement the training set.

Two main experiments were performed considering two configurations of the dermoscopy images: with and without cropping, where in the former case we use the segmentation masks of the lesions to compute a tight bounding box around them.

We also carried out a preliminary experiment on the ISIC 2018 [28] set, which comprises more than 12k images, divided into seven classes: melanoma and nevus (melanocytic), and basal cell carcinoma (BCC), actinic keratosis, benign keratosis, dermatofibroma, and vascular lesions, all of them non-melanocytic. Out of these images, a subset of 10,015 is available with the corresponding ground truth diagnosis. The remaining images are available for online scoring of developed methods. To train our model and select the best configuration, we randomly split the available data into training (80%) and validation sets (20%). Similarly to the previous experiments, we do not augment the training set with external data. We also submitted the model to the online platform, for scoring.

#### 5.2. Model Training

A set of dermoscopy images $I = \{I^1, ..., I^N\}$, for which we have the corresponding ground truth sequences of hierarchical labels $S = \{S^1, ..., S^N\}$, is used to train the model end-to-end to minimize the following loss

$$
L(I, S) = -\sum_{i=1}^{T} \log p(S_i),
$$

where $p(S_i)$ is given by (3) and $T$ is the length of the sequence (in our case, $T = 2$). This loss is optimized using the Adam variant of the stochastic gradient descent method [17], with mini-batches of size 10 and an adaptive learning rate with an initial value of $10^{-6}$. Additionally, we set the model hyperparameters to be the following: $D = 81$, $M = 50$, and $P$ tuned within $\{2^7, ..., 2^9\}$. Each model was trained for 500 epochs, with an early stopping criterion.

We adopted several strategies to improve generalization and reduce overfitting. First, several of the model weights $W_o$ were carefully initialized. In particular, we initialized the weights of the image encoders (DenseNet-161 and ResNet-Inception) from pre-trained versions on ImageNet. Although other works have stated that they do not need to retrain the encoders to obtain good captions [32], we found it beneficial to fine-tune these weights. The word encoding weights $E$ were also initialized from GloVe [22] and fine-tuned using Mittens [11].

Online data augmentation was also used to improve the performance of the model. In particular, at each epoch the training images were sequentially transformed using a random crop, random flip, and random color transformation. Finally, dropout with probability 50% was incorporated in several layers. The model was implemented in Tensorflow and executed on a NVIDIA Titan Xp.

#### 5.3. Model Evaluation

We evaluated the proposed model on a test set of dermoscopy images and quantified its performance using various metrics derived from the confusion matrix $C$. In particular, we evaluate the models regarding the class-specific sensitivity ($SE$) and specificity ($SP$), and the overall balanced accuracy ($BACC$). The area under the curve (AUC) for each class was also computed.

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1The source code to train and evaluate is available on https://github.com/catarina-barata/skin-hierarchy/
Table 1. Best performance scores for the ISIC 2017 test set, using full images.

<table>
<thead>
<tr>
<th>Lesion Class</th>
<th>SE</th>
<th>SP</th>
<th>BACC</th>
<th>AUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Melanocytic/Non-Melanocytic</td>
<td>92.5%</td>
<td>70.0%</td>
<td>81.3%</td>
<td>91.9%</td>
</tr>
<tr>
<td>Keratosis (1:90)</td>
<td>97.8%</td>
<td>92.0%</td>
<td>-</td>
<td>94.2%</td>
</tr>
<tr>
<td>Melanoma (117)</td>
<td>65.8%</td>
<td>88.6%</td>
<td>-</td>
<td>85.9%</td>
</tr>
<tr>
<td>Nevus (393)</td>
<td>82.2%</td>
<td>78.7%</td>
<td>-</td>
<td>88.9%</td>
</tr>
<tr>
<td>Average (600)</td>
<td>71.9%</td>
<td>86.5%</td>
<td>71.9%</td>
<td>87.9%</td>
</tr>
</tbody>
</table>

6. Results

6.1. ISIC 2017 Data set

6.1.1 Full Images

Table 1 and Figures 3-4 show the performance of the best proposed model ($P = 512$) on the ISIC 2017 test set, considering the non-cropped (full) images. The first decision (melanocytic/non-melanocytic) may be seen as a binary problem, while the specific diagnosis may be compared to a multi class one.

Overall, the model achieves a $BACC$ of 81.3% for the first decision and 71.9% for the second one, which is very promising and shows that both diagnoses are challenging. This difference in the scores is expected due to two main reasons: i) if the lesion was misdiagnosed in the first step, it will probably receive an incorrect label in the second step; and ii) even if the lesion is correctly diagnosed in the first step, it was observed that distinguishing between melanoma and nevus was also difficult. Interestingly, two non-melanocytic lesions were correctly diagnosed but then the model misdiagnosed them as nevi, which justifies the difference in the $SP$ of the first diagnosis (70%) and the $SE$ of keratosis (67.8%). We observed that this was due to an acquisition artifact (see Figure 4, last row) that will be discussed later.

Figure 3 shows that different regions are responsible for generating each of the hierarchical labels, as expected. In particular, for the melanocytic/non-melanocytic decision, the attention is mainly focused on the surrounding skin and the lesion, while for the final diagnosis, the attention is more specialized, giving variable importance to different regions of the lesions. These maps also provide medical interpretation, e.g., in the 2nd row, we can see that for the nevus diagnosis, the attention is mainly on regions that comprise many dots [2] and in the 5th row the model pays attention to a regression area with white and red, which is usually associated with melanoma [2].

Figure 4 shows some classification errors. The first row shows an example where the model correctly predicted the melanocytic label, but failed to perform the more specific diagnosis, since the lesion is a nevus. With the help of the attention map, it is possible to assess that the model was guided by the region with an atypical pigmentation. On the second row, the model incorrectly predicted the two labels. In the 3rd row, the model accurately predicted the classes of the lesion. However, the decision was guided by the existence of an acquisition artifact: a white reflection. Therefore, these maps may also provided relevant information about the existence of such problems with the data set.

6.1.2 Cropped Images

To evaluate the impact of the aforementioned artifact, which was present in several keratosis images, we have also
trained several models using cropped images. Here, the segmentation mask of the lesion was used to compute a bounding box around the lesion. The experimental results with the best model ($P = 512$) are shown in Table 2 and Fig. 5.

In this case, the performance of the model increased significantly for the melanoma class and marginally for the nevi class. On the other hand, the performance for the keratosis class dropped and now the $SE$ of this class equals the $SP$ of the melanocytic/non-melanocytic diagnosis, suggesting that the acquisition artifact shown in Figure 4 created a bias for this class. Nonetheless, this configuration is the one that achieves the best overall performance for the specific diagnosis (see the last row of Table 1 and 2), with a $BACC = 72.3\%$ and $AUC = 89.0\%$.

Figure 5 shows some examples of correctly diagnosed lesions. Note that, in this case, the cropped versions of the images are shown. As before, the attention varies according to the label that is being predicted and it is possible to see that different parts of the lesions play different roles in the diagnosis.

6.1.3 Comparison With Other Works

We compared our model with two kinds of works: i) the participants of the challenge that rank above and below us w.r.t the challenge evaluation metric - mean $AUC$ for the melanoma and keratosis classes; and ii) three recently published works that use the ISIC 2017 data set [33, 14, 34]. Since the challenge ranking was only performed using the performance for the melanoma and keratosis classes, we will adopt the same strategy to establish a comparison. Additionally, we have identified two criteria that influence the performance of a classification model: i) the use of external data sets to complement the training set; and ii) the use of ensembles of networks.

Table 3 shows the comparisons. Both configurations of the proposed model ranked in favorable positions, with the cropped formulation being slightly better. Additionally, our models achieve some of the best trade-off between $SE$ and $SP$. This not only shows the potential of using a hierarchical strategy to diagnose skin lesions, but also that it is still possible to further improve the model by: i) adding external data to the training set, as some of the best models do; and ii) increase the number of CNN in the ensemble of the image encoder, since we are using only two in this paper (DenseNet-161 and ResNet-Inception).
We believe that further experiments must be conducted on this data set, which may lead to an improvement in the scores. In particular, we plan to tune the model hyperparameters $M$ and $P$ and assess their influence in the performance. Additionally, we believe that incorporating more parameters of the model were not tuned and we selected the best hyperparameter configuration obtained for the ISIC 2018 data set. In this part, the hyperparameter tuning was done using the validation set, which may lead to an improvement in the obtained performance.

Table 5 shows the performance on the test set. In this section, we will discuss the results obtained on the ISIC 2018 data set. The obtained performances are significantly worse than those obtained on the validation set. This may be due to: i) the selected validation set not being representative enough and created a bias; ii) the used hyperparameters were not adequate and they require further tuning.

Table 3. Comparison with other works on the ISIC 2017 test set. * means that some information is missing from the paper.

<table>
<thead>
<tr>
<th>Lesion Class</th>
<th>SE</th>
<th>SP</th>
<th>BACC</th>
<th>AUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Melanoma</td>
<td>60.8%</td>
<td>90.9%</td>
<td>-</td>
<td>88.1%</td>
</tr>
<tr>
<td>Nevus</td>
<td>84.6%</td>
<td>90.5%</td>
<td>-</td>
<td>94.9%</td>
</tr>
<tr>
<td>Actinic</td>
<td>44.2%</td>
<td>99.0%</td>
<td>-</td>
<td>94.4%</td>
</tr>
<tr>
<td>BCC</td>
<td>60.2%</td>
<td>98.4%</td>
<td>-</td>
<td>96.5%</td>
</tr>
<tr>
<td>Keratosis</td>
<td>70.0%</td>
<td>91.7%</td>
<td>-</td>
<td>91.0%</td>
</tr>
<tr>
<td>Dermatofibroma</td>
<td>65.9%</td>
<td>99.4%</td>
<td>-</td>
<td>94.7%</td>
</tr>
<tr>
<td>Average (#1512)</td>
<td>63.7%</td>
<td>95.6%</td>
<td>64.1%</td>
<td>93.9%</td>
</tr>
</tbody>
</table>

Table 4. Best performance scores for our validation set randomly selected from ISIC 2018.

<table>
<thead>
<tr>
<th>Lesion Class</th>
<th>SE</th>
<th>SP</th>
<th>BACC</th>
<th>AUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Melanoma/Non-Melanocytic (±1±2)</td>
<td>95.7%</td>
<td>90.7%</td>
<td>92.2%</td>
<td>97.6%</td>
</tr>
<tr>
<td>Nevus (±312)</td>
<td>86.4%</td>
<td>94.1%</td>
<td>-</td>
<td>97.2%</td>
</tr>
<tr>
<td>Actinic (±456)</td>
<td>61.5%</td>
<td>99.4%</td>
<td>-</td>
<td>94.4%</td>
</tr>
<tr>
<td>BCC (±1201)</td>
<td>84.7%</td>
<td>98.9%</td>
<td>-</td>
<td>96.5%</td>
</tr>
<tr>
<td>Keratosis (±201)</td>
<td>93.4%</td>
<td>90.3%</td>
<td>-</td>
<td>97.2%</td>
</tr>
<tr>
<td>Dermatofibroma (±241)</td>
<td>66.1%</td>
<td>99.5%</td>
<td>-</td>
<td>93.0%</td>
</tr>
<tr>
<td>Average (±550)</td>
<td>89.2%</td>
<td>96.6%</td>
<td>-</td>
<td>93.0%</td>
</tr>
</tbody>
</table>

6.2. Preliminary Results on the ISIC 2018 Data Set

This section reports preliminary results for the proposed model on the ISIC 2018 data set. In this part, the hyperparameters of the model were not tuned and we selected the best hyperparameter configuration obtained for the ISIC 2017 data set ($M = 50$, $P = 512$, and $D = 81$). Table 4 reports the performance scores on our validation set.

ISIC 2018 is a much more complex data set than ISIC 2017, mainly because it comprises four more classes. Nonetheless, the proposed model is able to achieve a significantly good performance on the distinction between melanocytic and non-melanocytic lesions (BACC = 92.2%). The second level of diagnosis is more challenging, as expressed by the obtained scores. There is a significant unbalance between the different classes. Nonetheless, the model is still able to achieve SE above 60% for all of the classes and a BACC = 78.0%.

Table 5 shows the performance on the test set. In this case we do not have access to the ground truth diagnosis, thus it is not possible to evaluate the melanocytic/non-melanocytic diagnosis. The obtained performances are worse than those obtained on the validation set. This may be due to: i) the selected validation set not being representative enough and created a bias; ii) the used hyperparameters were not adequate and they require further tuning.

We believe that further experiments must be conducted on this data set, which may lead to an improvement in the scores. In particular, we plan to tune the model hyperparameters $M$ and $P$ and assess their influence in the performance. Additionally, we believe that incorporating more CNNs on the image encoding block may also improve the results, as many authors in the field of dermoscopy image analysis report the importance of using ensembles (e.g., [14]). Strategies to deal with the severe class imbalance, such as using weighted losses in the training, may also prove fruitful. Finally, as this data set comprises more lesion classes than ISIC 2017, it may be interesting to add an extra decision step, to distinguish between benign and malignant lesions, prior to the specific diagnosis (recall Figure 1), thus setting the number of the LSTM steps to $T = 3$.

7. Conclusions

This paper proposes a hierarchical classification model for the diagnosis of skin lesions that is able to provide interpretation through the use of attention maps. The proposed approach is inspired by recent works on deep models for caption generation, and combines CNNs with LSTM and attention modules. We evaluated our model on the ISIC 2017 and 2018 data sets and show that the hierarchical classification strategy has potential and that it is able to achieve competitive results with state-of-the-art methods, using a small number of ensembles and no external data. Moreover, the obtained attention maps show that the model is able to identify clinically relevant regions in the lesions, as well as to provide more insightful information on the importance of the different image regions in the diagnosis.
Acknowledgments

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