A Deep Learning Approach for Meibomian Gland Atrophy Evaluation in Meibography Images

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Purpose: To develop a deep learning approach to digitally segmenting meibomian gland atrophy area and computing percent atrophy in meibography images.

Methods: A total of 706 meibography images with corresponding meiboscores were collected and annotated for each one with eyelid and atrophy regions. The dataset was then divided into the development and evaluation sets. The development set was used to train and tune the deep learning model, while the evaluation set was used to evaluate the performance of the model.

Results: Four hundred ninety-seven meibography images were used for training and tuning the deep learning model while the remaining 209 images were used for evaluations. The algorithm achieves 95.6% meiboscore grading accuracy on average, largely outperforming the lead clinical investigator (LCI) by 16.0% and the clinical team by 40.6%. Our algorithm also achieves 97.6% and 95.4% accuracy for eyelid and atrophy segmentations, respectively, as well as 95.5% and 66.7% mean intersection over union accuracies (mean IU), respectively. The average root-mean-square deviation (RMSD) of the percent atrophy prediction is 6.7%.

Conclusions: The proposed deep learning approach can automatically segment the total eyelid and meibomian gland atrophy regions, as well as compute percent atrophy with high accuracy and consistency. This provides quantitative information of the gland atrophy severity based on meibography images.

Translational Relevance: Based on deep neural networks, the study presents an accurate and consistent gland atrophy evaluation method for meibography images, and may contribute to improved understanding of meibomian gland dysfunction.

Introduction

Dry eye disease is a multifactorial ocular surface disorder and is very common among adults. Meibomian glands (MGs) are believed to play a critical role in ocular surface health by secreting lipids into the tear film to slow down the rate of aqueous evaporation and minimize symptoms of dry eye. Dysfunction of MGs is the most frequent cause of dry eyes.1 The ability to visualize the glands and to monitor their changes with time or treatment is important for evaluating the risk of meibomian gland dysfunction (MGD) and dry eye diseases. Meibography, which is a photo documentation of MGs in the eyelids using either transillumination or infrared light, is commonly used in dry eye clinics for the diagnosis, treatment, and management of MGD.

The measurement of glandular loss is of significant clinical impact for the diagnosis of MGD.2,3 Percent MG atrophy, the ratio of gland loss area to the total eyelid area, may be an important clinical factor for assessing MGD severity. Using a standardized MG atrophy grading scale, the percent atrophy can be classified based on severity.4,5 In this paper, a previously published clinical-grading system6 (i.e., meiboscore) was applied (Table 1). Figure 1 depicts some sample meibography images with varying percent atrophy and corresponding meiboscores to help readers gain insights on the relationship between the percent atrophy and meiboscore. Currently,
Clinicians subjectively estimate the degree of MG atrophy severity. They assign a severity score after grossly estimating the relative ratio between MG atrophy area and total eyelid area. This subjective assessment has several limitations: (1) it may have high inter- and intraobserver variability and low repeatability; (2) it is based on qualitative judgments, therefore lacking quantitative evaluations to accurately track longitudinal changes; and (3) it may take a longer time and more costs to manually process a large number of images.

Recent advances in deep learning, a particular form of artificial intelligence (AI), show the ability of deep neural networks to learn predictive features directly from a large dataset of labeled images, without explicitly specifying rules or features. Additionally, deep learning has shown great success in medical imaging, such as diabetic retinopathy, breast cancer, melanoma, and others. It is of interest to use deep learning methods to benefit the process of evaluating atrophy in meibography images. Specifically, clinicians can use such methods to automatically segment the eyelid and atrophy areas in meibography images, and then compute the percent atrophy for the purpose of evaluating MG atrophy. Therefore, an automated method will potentially provide valuable and timely information on MG atrophy.

This study aimed primarily to develop and validate an automated deep learning system for evaluating MG atrophy severity from meibography images with clinician-verified annotations of eyelid and atrophy areas, for deep learning algorithm development and evaluation.

### Methods

#### Development and Evaluation Dataset

This study was based on the utilization of a meibography image dataset with clinician-verified annotations of eyelid and atrophy areas, for deep learning algorithm development and evaluation.

### Table 1. Percent MG Atrophy to Meiboscore Conversion Criteria

| MG Atrophy, % | Meiboscore |
|---------------|------------|
| 0             | 0          |
| 0–33          | 1          |
| 33–66         | 2          |
| >66           | 3          |

### Figure 1. Meibography images with ground-truth percent atrophies (%) and ground-truth meiboscores. Rows 1 to 4 refer to images with meiboscores of grade 0 to 3, respectively. Given a meibography image, the area of gland atrophy (marked in red) and eyelid (marked in blue) are compared by our deep learning algorithm to estimate the percent atrophy, and are further converted to meiboscore according to the criteria in Table 1.
by multiple clinicians. All clinicians were masked from the subject’s ocular surface health status. The most experienced clinician (TNY) also provided a separate set of the clinical meiboscore data for evaluation purposes (i.e., comparing scores among group clinicians, the lead clinical investigator [LCI] and our machine-learning algorithm against the ground-truth data generated by the machine algorithms). Subject demographics can be found in Table 2.

Data Annotations
A team of trained individuals labeled and measured the total eyelid and atrophy regions using the polygon tool in Fiji (ImageJ version 2.0.0-rc-59; Bethesda, MD). For labeling the total eyelid region, the upper border was defined at the MG orifices, the lower border was set at the edge of proximal tarsal plate, and the horizontal borders were where the top and bottom borders intersected. For labeling the total atrophy region, the upper border was drawn at the proximal ends of normal glands, the lower border was at the edge of proximal tarsal plate, and the horizontal borders were where the upper and lower borders intersected. Portions of glands that appeared atrophied (e.g., fainter, thinner) compared with other glands on the same eyelid were included in the atrophy region. The areas measured for the regions of interest were captured by selecting “Analyze>Measure” when the regions of interest were active. Final annotations were verified by an LCI (TNY) before they were made available for the machine-learning algorithm to minimize variability in the ground-truth data. From the fine-grained MG atrophy and total eyelid area annotation masks, these ground-truth annotated data were used to calculate the percent atrophy, which was then converted to meiboscore according to Table 1, for generating both “ground-truth percent atrophy” and the “ground-truth meiboscore.” Figure 1 depicts examples of atrophy- and eyelid-area annotations, along with corresponding ground-truth percent atrophy and ground-truth meiboscores. Algorithms were considered to achieve 100% accuracy if they predicted results exactly the same as the ground-truth annotations. Note that machine-learning systems can “predict” the MG atrophy region and percent atrophy from an meibography image not seen in the training phase. Machine predictions are different from medical predictions, which usually refer to predicting the future status of a disease or condition.

Data Allocations
All meibography images were randomly allocated into the following two sub-datasets: development and evaluation. The former was used for developing the deep learning algorithm, while the latter was for evaluating the performance of the algorithm. The percent atrophy distributions of the two datasets were very similar as shown in Figure 2. This minimized the scenario differences between training and evaluation.

For algorithm development, the development dataset was further divided randomly into 2 subsets, a train and validation set. The images in the train set were used to train the deep learning model, while the validation set was used for tuning the model hyper-parameters (e.g., network architectures, learning rate, etc.). The evaluation dataset, which did not have any overlapping image with the development dataset, was evaluated using the model that achieved the best performance from the validation set. The patient demographics and atrophy severity of the develop-

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### Table 2. Subject Demographics and Meiboscores of the Meibography Image Datasets

|                      | Development Train | Development Validation | Evaluation |
|----------------------|-------------------|------------------------|------------|
| Images, N            | 398               | 99                     | 209        |
| Patient demographics |                   |                        |            |
| Unique individuals, N| 308               | 77                     | 191        |
| Age, average ± SD    | 25.5 ± 10.9       | 27.0 ± 12.6            | 26.4 ± 11.6|
| Female/total patients, % | 63.5           | 66.6                   | 68.3       |

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ment and evaluation datasets can be found in Table 2 and Figure 3.

Algorithm Design and Training

In computer vision, image segmentation is the process of partitioning an image into multiple segments.\textsuperscript{23,24} Earlier methods first extracted hand-engineered features of the image, and then used the features to classify pixels independently.\textsuperscript{25–27} More recently, deep learning methods incorporated feature extraction and classification together into a unified framework, and achieved the state-of-the-art in image segmentation.\textsuperscript{11,28} A deep learning algorithm was built upon the pyramid scene parsing network\textsuperscript{29} to segment the atrophy and eyelid region of a given meibography image, and then the percent atrophy was calculated (Fig. 4).

The input to the neural network was a meibography image. The network could be considered as several stages of computations, parameterized by millions of parameters, and mapped the input image to the output segmentation masks of MG atrophy and eyelid area. Additionally, the network generated a vector indicating if MG atrophy existed in a given meibography image. The auxiliary output helped improve the segmentation performance by providing additional information. The final optimization goal was to have both correct segmentation and atrophy existence vector prediction.

Parameters of a neural network were determined by training the network on the development dataset. The network was repeatedly given images with known ground truth (the segmentation masks of atrophy and eyelid area, as well as atrophy existence vectors in our model). The model predicted the segmentation masks and atrophy existence vector of the given meibography image, and adjusted its parameters over the training process to make predictions increasingly more similar to the ground truth. The parameters of the model were optimized using stochastic gradient descent\textsuperscript{30} and the model performance was evaluated by the validation set every epoch. When atrophy area and eyelid area were predicted, the percent atrophy $A$ can be calculated as follows:

$$A = \frac{\text{area of atrophy}}{\text{area of eyelid}}$$ \hspace{1cm} (1)

A hyperparameter search was performed on the network architectures, data-augmentation techniques, learning rate, auxiliary loss ratio, and learning-rate decreasing policy. The hyperparameters, which attained the best performance over the validation set, was selected. The convolutional neural networks were developed using the PyTorch\textsuperscript{31} deep learning framework. The proposed networks were repeatedly trained and evaluated on two NVIDIA GeForce GTX 1080 GPUs with NVIDIA CUDA v8.0 and NVIDIA
cuDNN v5.1 acceleration (NVIDIA, Santa Clara, CA).

**Evaluation Protocol**

Evaluating the performance of the trained deep model was necessary. First, when performing the extensive tuning of hyperparameters on the training set, the model was evaluated on the validation set to select the hyperparameters that achieved the best performance. Additionally, once the best-performance model was obtained, the performance on the evaluation set was further evaluated to obtain the final performance. The algorithmic performance, including segmentation, percent atrophy, and meiboscore grading performance, was also examined.

**Atrophy Segmentation**

Two evaluation metrics were adopted to comprehensively assess the similarity between the predicted MG atrophy region and the ground-truth atrophy region.

**Accuracy (or pixel accuracy).** In atrophy segmentation example, the atrophy region is referred to as our region of interest (ROI). To evaluate the similarity between network predictions and the ground truth, the label of ROI is denoted as class $i$, while the rest as class $j$. In Table 3, $n_{ij}$ is denoted as the number of pixels of class $i$ predicted to belong to class $j$, $n_{ii}$ as the number of correctly classified ROI pixels (true positives, TP), $n_{ij}$ as the number of pixels wrongly classified as ROI (false positives, FP), $n_{ji}$ as true negative (TN), and $n_{jj}$ as false negative (FN). Thus, accuracy (abbreviated as ACC) is defined as follows:

\[
\text{ACC} = \frac{n_{ii} + n_{jj}}{n} = \frac{n_{ii} + n_{jj}}{n_{ii} + n_{ij} + n_{ji} + n_{jj}} = \frac{TP + TN}{FP + TP + TN + FN}\]

(2)

Similarly, in total eyelid area segmentation example, the eyelid region would be the ROI. ACC reports the percentage of pixels in the image that is correctly classified. However, this metric sometimes provides misleading results when ROI is small, as the measure would be biased by mainly reporting how well non-ROI cases are identified.

**Mean IU.** Mean intersection over union (mean IU, or Jaccard index) quantifies the percent overlap between the target mask and our prediction output. It measures the number of pixels common between the target and prediction masks divided by the total number of pixels present across both masks. Denoting ground truth ROI segmentation mask as ground-truth (GT), network predicted segmentation mask as prediction, mean IU $\text{IU}^{32}$ could be written as following:

\[
\text{mean IU} = \frac{\text{GT} \cap \text{prediction}}{\text{GT} \cup \text{prediction}} = \frac{n_{ii}}{n_{ii} + n_{ij} + n_{ji}} = \frac{TP}{FP + FN + TP}\]

(3)

Intuitively, Equation 3 is analogous to harmonic average of the precision and recall, F1-score, which was defined as $\frac{2TP}{2TP + FP + FN}$, and provided a more “comprehensive” evaluation—the method considered both precision and recall. For segmentation tasks, higher mean IU value indicated higher alignment of the algorithm prediction with the ground-truth.

In summary, ACC reflects the performance of pixel-wise classification accuracy, while mean IU for how the predicted ROI segmentation mask overlaps with the real counterpart. Mean IU is considered as a stricter evaluation metric than ACC in terms of segmentation. Both evaluation metrics for each image in the evaluation dataset and the calculated average score of all images were reported.

**Percent Atrophy**

Percent atrophy from the atrophy masks (Equation 1) can be calculated, and the percent atrophy performance of algorithm prediction can also be evaluated against the ground-truth. One standard way is to compute the atrophy ratio difference over the evaluation set and compute the root-mean-square deviation (RMSD) as follows:

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**Table 3. Confusion Matrix for Illustrating Segmentation Evaluation Metric Mean IU**

| Class | Ground-Truth | Predicted |
|-------|--------------|------------|
|       | $i$          | $j$        |
| $n_{ij}$: False positive # (FP) | $n_{ij}$: True positive # (TP) |
| $n_{ii}$: True negative # (TN) | $n_{jj}$: False negative # (FN) |

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\[
\text{RMSD}(\hat{A}) = \sqrt{\frac{1}{N} \sum_{i=1}^{N} (\hat{A}_i - A_i)^2}
\]  

(4)

where \(A_i\) is the real atrophy ratio of the \(i\)th image defined in Equation 1, \(\hat{A}_i\) represent the \(i\)th atrophy ratio predicted by the neural network. \(N\) is the total number of images in evaluation set. Intuitively, Equation 4 reflects the mean difference between predicted percent atrophy and ground truth over all the images in the evaluation set. RMSD could also be considered as the error made by the algorithm when predicting the percent atrophy of an image.

Converting to Meiboscores

Percent atrophy, as a numeric indicator, provides substantial information on the MG atrophy severity of a meibography image. In order to compare with human clinicians’ performance, the numeric ratios were converted back to the meiboscore.

From the meibography images with different percent atrophy in Figure 1, images near the grading transition limits (0%, 33%, 66%) were very similar and difficult to classify. A tolerance threshold near the grading transition limit was necessary. The converting criteria in Table 1 was applied with a relaxed standard. As illustrated in Figure 5, the tolerance threshold was set at 0.25%, so classifying images with percent atrophy 0% to 0.25%, 32.75% to 33.25%, and 65.75% to 66.25% either to its ground-truth or adjacent meiboscores were both considered as correct prediction. The same relaxed converting criteria for both human clinicians and algorithm in the experiments were followed for fair comparison.

**Results**

**Network Training Details**

Each meibography image and its corresponding segmentation mask(s) were resized to the size of 420 \(\times\) 420 pixels. During training, 400 \(\times\) 400 pixels were randomly cropped out of a given meibography image and corresponding annotations in every training epoch for data augmentation. A center crop of 400 \(\times\) 400 pixels was made to a given meibography image and corresponding annotations during the evaluation process for both validation and evaluation datasets.

Different network architectures (SqueezeNet, resnet18, resnet34, resnet50), auxiliary loss ratio (for 0–1.0 with grid of 0.1), learning rate, and learning-rate decreasing policy were carefully assessed to obtain the best performance of the network on the validation dataset (the best performance of the network over the validation set was resnet50, auxiliary loss ratio 0.1, learning rate 1e-3, 200 epochs in total, with learning rate decrease at 100, 150, and 180, respectively). The algorithm performance of the model on the evaluation dataset is reported.

**Algorithm Performance**

The baseline characteristics of the training and evaluation dataset were described in Table 2. Development and evaluation dataset had similar characteristics regarding patient demographics and MG atrophy severity.

Table 4 shows the ACC, mean IU, and RMSD of meibography images with different meiboscores. Note that images with meiboscore of grade 0 do not have atrophy so there are no corresponding ACC and mean IU. The performance of meiboscore of grade 0 can however be measured by RMSD and meiboscore grading accuracy. Regarding atrophy segmentation, ACC was higher than mean IU. The instance average ACC values were 97.6% and 95.5% for eyelid and atrophy, respectively, while mean IU values were 95.4% and 66.7% for eyelid and atrophy, respectively. Regarding mean IU for different gland atrophy severity, mean IU was the lowest for meiboscore of grade 3 samples, which only included five images in the training set and only two in evaluation set.

Regarding percent atrophy prediction performance, the instance average error was 6.7%. Although images with different meiboscores had relatively similar RMSD, the RMSD value was the highest at 9.0% for meiboscore of grade 1 and the lowest at 5.7% for
Although the intention was to capture each subject for one single visit, there were three subjects (providing a total of 10 images) from 467 subjects (0.6% subjects) returned to the research facility for two visits within at least a 2-year time lapse. Therefore, a 10-fold cross validation was performed to confirm that the images obtained from the repeated visits would not bias the study results. The results (e.g., mean and standard deviation of ACC, mean IU, and RMSD) were reported in Table 5. Specifically, the development and evaluation sets were randomly split 10 times according to the number of images of each meiboscore category presented in Table 2. The algorithm was trained on different development sets and evaluated on the corresponding evaluation sets for 10 times. Figure 6 plots the predicted percent atrophy versus ground-truth percent atrophy. While most of the points fall on the ideal line (percent atrophy prediction equals to ground-truth), the deep learning algorithm tends to give higher percent atrophy for some cases of meiboscore of grade 0. This is because the algorithm has been trained to be sensitive to even small atrophy, which might have been ignored by clinicians. The errors are greatly reduced when converting to meiboscores using the relaxed criteria as described in the “evaluation protocols” section.

Figure 7 visualizes the atrophy region and eyelid region segmentation results of clinician team and computer. From the visualization, the human segmentation and computer segmentation appear to be very similar, especially for eyelid region segmentations.

### Human Clinician’s Performance

The confusion matrix is a specific table layout that allows visualizations of how two identities perform the same classification task. Table 6 shows the confusion matrices of study clinician and single clinician. The highest agreement percentage was 69% for meiboscore of grade 1, while the lowest was 40% for meiboscore of grade 3. The kappa score is 0.324 for clinical team and the LCI, which led to a fair agreement according to Landis et al.33 In other words, clinicians’ ratings have high variability.

### Comparing Meiboscores

The meiboscore grading performance of the algorithm was compared against the ground-truth meiboscores. The ground-truth meiboscores were obtained from the percent atrophy (calculated from human-annotated segmentation masks) using Table 1. Tables 7 and Figure 8 show the meiboscore grading performance by the algorithm, clinical team (clinical meiboscore), and the LCI meiboscore. The algorithm

| Meiboscore | Percent Eyelid Area | Percent Atrophy Area | Percent Atrophy, RMSD |
|------------|---------------------|----------------------|-----------------------|
|            | ACC (Mean IU)       | ACC (Mean IU)        | ACC (Mean IU)         | ACC (Mean IU) |
| Meiboscore 0 | 98.0 ± 0.4 (96.0 ± 0.5) | / (96.1 ± 0.5) | / (9.0 ± 0.5) |
| Meiboscore 1 | 97.6 ± 0.3 (95.4 ± 0.6) | 96.1 ± 1.7 (65.9 ± 1.8) | 5.6 ± 0.6 |
| Meiboscore 2 | 97.9 ± 0.5 (96.0 ± 0.9) | 94.6 ± 1.9 (77.3 ± 1.8) | 7.4 ± 1.6 |
| Meiboscore 3 | 92.4 ± 2.0 (83.8 ± 4.0) | 85.1 ± 3.9 (61.6 ± 4.5) | 7.3 ± 2.0 |
| Class average accuracy | 96.4 ± 0.8 (92.8 ± 1.5) | 91.9 ± 2.5 (68.3 ± 2.7) | 7.5 ± 1.3 |
| Instance average accuracy | 97.6 ± 0.4 (95.5 ± 0.7) | 95.7 ± 1.8 (67.6 ± 1.8) | 6.6 ± 0.8 |
achieves 95.6\% overall grading accuracy, which outperforms the LCI meiboscore by 16.0\% and clinical team meiboscore by 40.6\%. For each meiboscore, the algorithm also largely outperforms human clinicians. Figure 9 depicts some failure cases in meiboscore grading of human clinicians and our algorithm. Failure cases appear for both human-assigned and algorithm meiboscores when percent atrophy is near the meiboscore grading transition limits.

The present work introduces a deep learning approach to automatically predict the MG atrophy region and compute percent atrophy in the meibography image. The proposed method has the following three advantages: (1) low variability and high repeatability (test–retest reliability); (2) output quantitative result rather than qualitative description (e.g., specific gland atrophy region and numeric percent atrophy prediction); and (3) efficient and low cost. The average processing time per meibography image was approximately 0.29 seconds (experiments were performed on one NVIDIA GeForce GTX 1080 GPU). This means that more than 1000 unprocessed or raw meibography images can be evaluated for atrophy severity in 5 minutes without additional human resource needed.

The algorithm also has very high performance. Accuracies of eyelid area and atrophy area achieve 97.6\% and 95.4\%, respectively, and the overall mean IUs are 95.5\% and 66.7\%, respectively. Our algorithm achieves a 95.6\% overall grading accuracy and outperforms the LCI meiboscore grading accuracy by 16\%. From the visualization of the predicted eyelid and atrophy segmentation, the algorithm predictions have high visual similarity with the ground-truth annotations.

In all, the proposed algorithm achieves very high performance with low variability and high repeatability in evaluating MG atrophy from meibography images. Additionally, the proposed algorithm could
potentially be applied to other similar image segmentation tasks in the clinical community.

Furthermore, regarding MG atrophy evaluation, numeric percent atrophy reporting is a better assessment than simple meiboscore grading. It is challenging to distinguish different meiboscores when the percent atrophy is near the grading transition limits (0%, 33%, 66%), because no distinct changes observed for meibography images in these regions. Forcing a strict meiboscore near the boundary can lead to grading inconsistency and variability. When applying numeric percent atrophy, however, clinicians would not need to worry about the images with percent atrophy near these grading transition limits. Additionally, there are several different grading scales for gland atrophy. Conversion between different grading systems is impossible as the percentage information already got lost. Numeric percent atrophy adopted in this study overcomes the above-mentioned problems.

In conclusion, a deep learning approach to automatically evaluate the MG atrophy in meibography images has been developed. The system has high accuracy, repeatability, and low variability, as well as outperforming human clinicians by a significant margin. The quantitative outputs (specific atrophy region and percent atrophy) provide valuable information of MG atrophy severity of the meibography image. In the present work, our deep learning system could only predict MG atrophy region, but not individual MG morphology. Future work can explore how deep learning can automatically analyze MG morphologic characteristics (e.g., gland number, width, intensity, and tortuosity), which can be potentially important for advancing the efficiency of MGD treatment and management. Capturing human expert knowledge with data-driven, deep learning systems is the future of image-based medical diagnosis. The possibility of using deep learning methods for clinical diagnosis in ocular surface diseases is shown. New surveys can be enabled by automatic and quantitative evaluations, opening up many exciting

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Table 6. Confusion Matrix of LCI and Clinical Meiboscore (%)

| Clinical Meiboscore | LCI Meiboscore |
|---------------------|----------------|
|                      | Meibo 0 | Meibo 1 | Meibo 2 | Meibo 3 |
| 96 Meibo 0          | 45%     | 51%     | 4%      | 0%      |
| 84 Meibo 1          | 11%     | 69%     | 18%     | 2%      |
| 22 Meibo 2          | 0%      | 32%     | 50%     | 18%     |
| 5 Meibo 3           | 0%      | 20%     | 40%     | 40%     |

Note: Warm colors with small values indicate less agreement.

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Figure 8. Meiboscore grading performance of the clinical team, the LCI, and the proposed algorithm. The algorithm outperforms the LCI meiboscore by 16.0% and clinical team meiboscore by 40.6%.

Figure 9. Examples of failure cases in meiboscore (MS) grading: human clinicians versus machine algorithm. GT refers to the ground-truth. Failure cases occur for both clinicians and the algorithm for the images with percent atrophy near the meiboscore grading boundaries.
opportunities for targeted medical treatment and drug discoveries.

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Table 7. Meiboscore Grading Performance of Clinicians and Algorithm (%)

| Meiboscore | Percent Clinical Team | Percent Lead Clinical Investigator | Percent Algorithm |
|------------|-----------------------|-----------------------------------|-------------------|
| 0          | 86.8                  | 100.0                             | 100.0             |
| 1          | 48.2                  | 78.0                              | 95.7              |
| 2          | 46.2                  | 57.7                              | 88.5              |
| 3          | 50.0                  | 100.0                             | 100.0             |
| Class average accuracy | 57.8                  | 83.9                              | 96.1              |
| Instance average accuracy | 55.0                  | 79.6                              | 95.6              |
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