An Interpretable Algorithm for Uveal Melanoma Subtyping from Whole Slide Cytology Images

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Abstract

Algorithmic decision support is rapidly becoming a staple of personalized medicine, especially for high-stakes recommendations in which access to certain information can drastically alter the course of treatment, and thus, patient outcome; a prominent example is radiomics for cancer subtyping. Because in these scenarios the stakes are high, it is desirable for decision systems to not only provide recommendations but supply transparent reasoning in support thereof. For learning-based systems, this can be achieved through an interpretable design of the inference pipeline. Herein we describe an automated yet interpretable system for uveal melanoma subtyping with digital cytology images from fine needle aspiration biopsies. Our method embeds every automatically segmented cell of a candidate cytology image as a point in a 2D manifold defined by many representative slides, which enables reasoning about the cell-level composition of the tissue sample, paving the way for interpretable subtyping of the biopsy. Finally, a rule-based slide-level classification algorithm is trained on the partitions of the circularly distorted 2D manifold. This process results in a simple rule set that is evaluated automatically but highly transparent for human verification. On our in house cytology dataset of 88 uveal melanoma patients, the proposed method achieves an accuracy of \[87.5\%\] that compares favorably to all competing approaches, including deep “black box” models. The method comes with a user interface to facilitate interaction with cell-level content, which may offer additional insights for pathological assessment.

1. Introduction

Uveal Melanoma (UM) is the most common primary intraocular malignancy in adults (Singh et al., 2011). As standard care for UM, Fine Needle Aspiration Biopsy (FNAB) is often performed to confirm the diagnosis and enable UM prognostication. To this end, a molecular test, Gene Expression Profile (GEP), is performed and microscopic Cytology of Fine Needle Aspirates images are created from the biopsy. According to a recent study, there exist two subtypes in UM that can be identified based on its GEP: The first subtype exhibits low metastatic risk, while the second subtype has been linked to high metastatic risk. There is a stark contrast in long-term survival between the two classes: the 92-month survival probability in class 1 patients is 95%, versus 31% in class 2 patients (Onken et al., 2004). It is evident that access to UM subtype information is critical for proper management of patients by providing appropriate recommendation for metastasis surveillance. However, even after 10 years of development, GEP is still only available in the United States. The technique is also expensive and has a long turn around time. A more accessible test for UM subtyping is, therefore, highly desirable.

There is increasing evidence that the underlying genetic profile affects cancer growth on multiple scales. Radiomics, for example, exploits this observation to develop imaging-derived biomarkers that are informative for prognosis (Grossmann et al., 2017). In the particular case of UM prognostication, there is huge potential in using imaging-derived biomarkers to determine GEP subtype and metastatic risk directly from cytology slides. While it is impossible even for highly trained pathologists to derive this information from cytology images, learning-based algorithms that discover associations between intensity patterns in cytology images and GEP subtype are promising (Liu et al., 2020; Chen et al., 2020). However, as “black box” models that perform a super-human task, these algorithms do not offer insights beyond the final recommendation to the human decision makers, which has been linked to automation bias and over-trust or dis-trust in such systems (Nourani et al., 2020; Gaube et al., 2021). A more transparent algorithm design may enable humans to better calibrate their trust in the recommendation, which would be an important...
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Figure 1. An overview of the automatic interpretable algorithm for uveal melanoma subtyping from cytology images. The algorithm consists of cell instance segmentation to extract cell appearance information that is used to cluster cells of similar appearance in a circular space. Based on a coarse partitioning of the embedding space, which we refer to as pie chart, shown in (a), we find simple rule sets (b) that enable uveal melanoma subtyping, which otherwise, requires gene analysis. A pie chart of a representative patient is shown in (c) - the patient is at high metastatic risk (GEP class 2).

2. Related work

The high resolution and complexity of whole slide images (WSIs) make cell-level annotations difficult or impossible to obtain, which is why many of the annotated datasets are limited to slide-level labels that correspond to the overall diagnosis. However, standard automatic cancer subtyping and analysis in WSIs is based on multiple small regions extracted from slides, that then need to be aggregated to a single prediction on the slide level. These methods include majority voting, coarse-to-fine techniques (Liu et al., 2017; Hou et al., 2016; Xu et al., 2017; Zhang et al., 2020), and multiple instance learning approaches (Chikontwe et al., 2020; Campanella et al., 2019). While most of the deep learning approaches in WSIs analysis consider black box models, recent works attempt to introduce features that enhance model understanding by mimicking the decision process of pathologists. For instance, content-based histopathological image retrieval (Peng et al., 2019; Hegde et al., 2019) contrasts a query image with a large database to determine the search results with more similar histological features. Making intelligent systems interpretable is another frontier in developing trustworthy medical decision support applications (Rudin, 2019). In contrast to explainable models that rely on post hoc analysis, interpretable models aim to explain the reasoning behind a prediction. In a histopathologic context, patch-based regions visualizations introduced in (Pirovano et al., 2020) display features related to tumor tissue, in addition to providing slide-level heatmaps that improve WSI classification. Different from previous methods that attempt to provide human-meaningful visualizations, either from learned representations or image retrieval, our method is interpretable by nature.

3. Method

Given high-quality Region of Interests (ROIs) extracted from cytology images, we create an interpretable system to analyze UM biopsy cytology and reveal GEP subtype based
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Figure 2. System overview of the proposed method. Cell-level features are obtained by aggregation over instance cell segmentation masks and then embedded into a 2D space. Several slides are embedded in this way to create a representative cell appearance space, and the 2D embedding space is subsequently distorted to a circle. For every other whole slide image (WSI), cells representations are extracted and projected with the same embedding process into the circular space, such that one density chart is generated for every slide. Finally, we find an interpretable rule set to classify UM biopsies based on the density charts.

on overall cell composition of the sample. Our learning-based system comprises of three parts: 1) instance-level cell segmentation, 2) cell feature embedding, and 3) interpretable classification. We note that the specifics of each component may easily be replaced by other techniques since aspects pertaining to the exact method choice are not the main focus of our work. In the remainder of this chapter, we describe a cost-efficient way of weakly labeling our dataset to enable supervised learning of the cell segmentation network (Section 3.2). Cell-level features are then generated and embedded into a 2D space for further classification as described in Section 3.3. Finally, we define an interpretable classification model within the 2D space to distinguish UM GEP classes by cell composition (Section 3.4). The system overview is shown in Figure 2.

3.1. Considerations around interpretability

Before we introduce the technical details of the proposed method, we first frame our model in the current interpretability definition. As proposed in (Murdoch et al., 2019), interpretable machine learning is defined to be the use of machine-learning models for the extraction of relevant knowledge about domain relationships contained in data. Knowledge is considered to be relevant if it provides insight for a particular audience into a chosen domain problem. Our proposed method aims to provide an interpretable model for UM GEP classification to pathologists, who already possess substantial expertise in reading cytology slides. Indeed, clinicopathologic features of UM tumors, e. g. epithelioid cell type and aggressiveness of cancer cells, can be readily estimated from cytology images and have been associated with worse patient prognosis and a higher incidence of metastatic disease (Worley et al., 2007). These risk factors are widely used by pathologists, however, their accuracy to predict metastatic potential has been shown to be limited (Schopper & Correa, 2016). Our interpretability mechanism is motivated by the fact that clinicopathologic features of UM tumors, e. g., cell appearance, are relevant knowledge for pathologists to predict UM metastatic risk. Instead of building interpretable models with the clinicopathologic information manually extracted by pathologists themselves, our system automatically extracts clinicopathologic features (cell appearance features) and summarizes the cell appearance distribution in a 2-dimensional space, which is further classified by a simple and interpretable rule set. While the algorithm was developed in close collaboration with ophthalmic oncologists and pathologists, this manuscript is limited to describing the proposed system and characterizing its performance and does not empirically demonstrate its interpretability on a larger user group. Doing so will remain subject of future work.

3.2. Instance cell segmentation

There exist no cell annotations for the high-quality ROIs that are automatically extracted from cytology images using the method described in (Chen et al., 2020). However, instance cell segmentation is essential for further cell-level analysis. Thus, we prepare annotations on a small sub-set
YOLACT because it can be easily modified to enable training with minimal manual labor to enable supervised training of an instance segmentation network. Figure 3 presents the annotation procedure. In detail, we randomly select 500 ROIs from the 131k pool and partially annotate super-pixels generated by SLIC (Achanta et al., 2010) to reduce the annotation workload. We group all super-pixels within any annotated cells to generate instance-level annotations. The ROI annotation procedure. (a) the extracted high-quality ROI; (b) the generated super-pixels.; (c) the annotations on super-pixels. Yellow and blue regions are annotated super-pixels for cancer cells and background, respectively.

### 3.3. Cell-level feature embedding

Previously, pathologists have attempted to quantify different cell components, such as nuclear size and nucleolar size, to predict the behavior of tumors. Our approach is similar to this process as it extracts network feature representations of cells, which we assume contain information about cell appearance. Cell-level features $F_c$ for cell $c$ are then extracted from the entire feature map $F$ using the segmentation mask $M_c$ with masked average pooling:

$$F_c = \text{Avg}(F[M_c]),$$

where $F$ is the output of the backbone network architecture. To prepare subtype classification based on cell composition, and improve classification performance, we embed all cell-level features in 20 GEP class 1 slides to create a 2D embedding space with UMAP (McInnes et al., 2020). All other slides are then embedded into that space, to represent the respective cell composition. We expect slides of distinct GEP classes have different cell composition, and thus distribution in the 2D embedding space. The embedding space is created using slides of one GEP class only to potentially maximize the difference of GEP class 1 and class 2 representation in the embedding space.

### 3.4. Interpretable Classification

Based on our hypothesis that slide-level cell composition, and thus distributions in the 2D cell appearance embedding space, should be different between GEP classes, we devise an interpretable algorithm that reasons based on these representations. Direct comparisons between distributions, e.g. chi-square test (Pearson, 1900) and Kolmogorov-Smirnov tests (Kolmogorov, 1933), are complicated and not usually interpretable. Instead, we partition the embedding space and analyze the region densities. Because cells with similar appearance, thus similar features, are close to each other in the embedding space, the density of each region represents the portion of cells with a specific kind of cell appearance in the slide. To make it easier to define the spatial partitioning of the embedding space, we first distort the space into a unit circle. We treat the center of gravity of all embedded cells as the origin. Then, we normalize to unity the scale of all embedded cells in every degree of angle in polar coordinate, so that the whole embedding space is distorted to a unit circle. Parameters in circular distortion are determined simultaneously with the embedding generation and are fixed when embedding new slides. Finally, we divide the unit circle equally into 12 regions, as shown in Figure 4. Since we posit that each GEP class will have different densities in distinct regions, in addition to the individual densities of these regions ($D_i$), we define the relative densities ($D_i/D_j$) as input variables for classification. Finally, an interpretable bayesian rule set algorithm (Wang et al., 2017) takes all these 78 input variables (12 values ($D_i$), and 66 relations ($D_i/D_j$)) for GEP classification.

Different from logistic regression (which is only interpretable in low dimensional problem settings because humans can handle at most $7 \pm 2$ cognitive entities at once (Miller, 2020; Cowan, 2010)), the rule set algorithm is not limited by the number of input variables. The number of arguments in each rule can be controlled, as simple as determining the largest allowed depth of a tree. In addition, it is different from a random forest (which uses majority vote, and is thus not interpretable) since here, the predicted output is determined once the sample obeys at least one rule in the rule set.

### 4. Experiments

We demonstrate that our proposed interpretable learning pipeline does not compromise on performance of UM subtyping when compared to deep black box models. We also demonstrate an additional, interactive tool for expert review of cell level composition through interaction with the 2D...
embedding space. By simply clicking on areas of interest in the density charts, users may retrieve and visually inspect cells that are representative of the appearance in that specific embedding location. Details are shown in Figure 5.

4.1. Experimental setting

Dataset: The dataset we use includes 100 cytology samples from 88 uveal melanoma patients. To the best of our knowledge, this is the largest dataset on UM cytology. The dataset contains 50 slides from 43 patients with GEP class 1 and 50 slides from 45 patients with GEP class 2. The cellular aspirates obtained from cytology of each tumor were submitted for cytology and GEP testing. The cytology specimen was flushed on a standard pathology glass slide, smeared, and stained with hematoxylin and eosin. The specimen submitted for GEP was flushed into a tube containing extraction buffer and submitted for DecisionDx-UM testing. Whole slide scanning was performed for each cytology slide at a magnification of 40x. Automatic ROI extraction is performed using SLIC (Achanta et al., 2010), where the number of components is 400, and the Euclidean distance ratio is 1. On average, each of the 500 randomly sampled ROI for manual annotation has 9 cells and 38 background super-pixels annotated. The number of prototypes in YOLACT is doubled to 64 to potentially segment more cells within every ROI. The segmentation model is optimized using Adam (Kingma & Ba, 2014) with a learning rate of $10^{-5}$ and 4000 iterations with a batch size of 1. We train the model on 450 annotated ROIs and validate on the other 50 ROIs. We empirically split the circular embedding space into 12 partitions, as shown in Figure 4, which in internal development was found to yield the best performance compared to other split approaches. All cells that map outside the circular embedding space are projected to the nearest region. For the interpretable classification, we use 80% of the projected slides in both class 1 and class 2 for training (64) and the other 20% for testing (16). The rule set algorithm is trained with simulated annealing procedure as described in (Wang et al., 2017). The maximal length of each rule in the rule set is set to 2 to preserve its intelligibility.

4.2. Cell Segmentation Performance

We use Mean Average Precision (mAP) as the main evaluation metric for cell segmentation performance. The mAP is about 70% when Intersection-Over-Union (IoU) is larger than 50%, which indicates that the segmentation process catches a fairly good number of cancer cells. However, mAP is low with high IoU threshold, because of the low quality of super-pixel-based annotations on the cells’ boundary. Table 1 and Figure 6 present both quantitative and qualitative results, respectively. The algorithm can easily tell apart cancer cells from blood cells, while some cancer cells with ambiguous boundaries are missed. We attribute this to the low quality of super-pixels for these cells during annotation. As a result, cells with ambiguous boundaries are usually skipped in annotation if more clear cells exist in the same ROI. Because there exist numerous cells in each slide, missing some cells at random will not significantly impact the overall cell composition, and further, the classification performance.
Figure 5. The additional tool for embedding space interaction. For every slide embedding chart, users can click any area of interest, i.e. the green/blue cross location. Several closest cells in the embedding space are visualized in native and scaled resolution.

Figure 6. Examples of segmentation results. The segmentation network is able to (a) separate cancer cells (purple, large) from blood cells (red, small); (b) segment cells with all sizes, but (c) misses some ambiguous cells. The numbers within the boxes correspond to confidence scores.

4.3. UM subtype classification

We compare our proposed method with a previously proposed deep black box model (Coudray et al., 2018; Liu et al., 2020) evaluated on the same dataset, which classifies UM subtype directly from ROIs. In (Coudray et al., 2018), slide-level subtype prediction is obtained by simply averaging class predictions for all corresponding ROIs. Both, the black box and our proposed method have the same backbone network architecture (ResNet-50 (He et al., 2015)) and the same training and testing split for a fair comparison. We find that the accuracy performance of our method (87.5\%) compares favorably to the black box approach following (Coudray et al., 2018) (83.3\%), and more importantly, is interpretable based on the following rule set over the density chart ($D_i$), and thus, over the cell appearance composition of the whole slide:

$$D_6/D_{11} > 1.5 \text{ AND } D_1 < 0.07$$

OR

$$D_7/D_{12} \leq 0.4$$

(2)

There only exist 3 arguments in the rule set, which makes algorithmic recommendations transparent and verifiable, while enabling users to understand overall cell composition. This rule set was visually represented already in Figure 1.

4.4. Ablation Study

We conduct an ablation study of the rule-based interpretable classification to benchmark its performance against other classification methods, i.e. logistic regression, Support Vector Machine (SVM) and Artificial Neural Network (ANN). We also compare different embeddings, by creating the initial UMAP embedding space with either, GEP class 1 or GEP class 2 slides. After the embedding space creation, only 80 slides remain to train and evaluate the classification models. Therefore, we also introduce an ensemble method to enrich the input data by creating synthetic cell compositions. To create a synthetic slide, we randomly selected 30\% cells from one slide and 1\% cells from all the other slides in the same class as all the cells in the synthetic slide. Then, the synthetic slide will represent the main pattern of one observed slide but also introduce other variations. We created 100 synthetic slides for each class using this approach, which is indicated as “Ensemble” in Table 2. The simple ANN we used is fc($n$) + ReLU + fc(1), where $n$ in fc($n$) means the number of output channels. To evaluate the methods, we then perform 100 random training/testing splits of our dataset on the patient-level and train all models on every split. The mean results and the corresponding confidence intervals are summarized in Table 2.

Logistic regression has the lowest testing accuracy (75.14\%) and the rule set achieves the highest performance (87.50\%), which is comparable to SVM (82.07\%) and ANN (83.71\%). Creating the embedding from distinct GEP classes results in similar accuracy of the rule set algorithm (87.50\% v.s. 84.33\%). As in the previous comparisons to black box models, the rule set approach has the added benefit of being

| IoU | box | mask |
|-----|-----|------|
| 0.5 | 70.67\% | 69.30\% |
| 0.6 | 64.41\% | 64.72\% |
| 0.7 | 49.20\% | 53.07\% |
| 0.8 | 27.52\% | 33.91\% |
| 0.9 | 3.24\% | 2.49\% |

Table 1. mAP for segmentation boxes and masks with different IoU threshold.
Table 2. Ablation study of interpretable classification with different methods and an ensemble technique. LR refers to logistic regression. Rule Set (class $k$, $k = 1, 2$) refers to results using the embedding created from class $k$ slides.

|               | w/o Ensemble | w/ Ensemble |
|---------------|--------------|-------------|
|               | Accuracy     | # of rules  | Accuracy     | # of rules  |
| LR            | 67.50 ± 5.56%| N/A         | 75.14 ± 9.00%| N/A         |
| SVM           | 83.00 ± 6.37%| N/A         | 82.07 ± 8.23%| N/A         |
| ANN           | 82.86 ± 8.33%| N/A         | 83.71 ± 10.15%| N/A        |
| Rule Set (class 1) | 86.36 ± 10.25% | 2.28 ± 0.57 | 87.50 ± 9.56% | 2.11 ± 0.37 |
| Rule Set (class 2) | 81.93 ± 8.02% | 2.06 ± 0.49 | 84.33 ± 10.68% | 1.96 ± 0.31 |

interpretable. Logistic regression and SVM models suffer in this regard due to the high dimensionality of the input representation (78). Dimensionality reduction techniques, e.g. principle component analysis (PCA), exist but are not applicable here because the number of input variables (78) is larger than the number of training samples (64). Finally, all models reach higher accuracy with the ensemble except SVM.

Due to the fact that our segmentation model is not perfect, we also evaluate the rule set model for different segmentation results. During early training, the segmentation model will first identify the most clear cancer cells, but along with plenty of false positives. As the optimization progresses, fewer cancer cells are segmented but much fewer false positives occur. The accuracy of the rule set algorithm for segmentation results after 2000, 3000 and 4000 training iterations is 77.23 ± 10.98%, 84.64 ± 10.46% and 87.50 ± 9.56%, which suggests that the algorithm favors the output of a highly specific cell segmentation algorithm.

### 5. Discussion

Our overall system utilizes the segmentation features to generate the embedding and classify UM subtypes based on slide-level cell composition. We assume that cell composition will be different across the two GEP classes, which will result in different cell density chart representations of slides from the two subtypes that can then be distinguished using an interpretable rule-based algorithm. This hypothesis is supported by our experiments. One aspect of the current approach is that we do not currently interpret the embedded features themselves, e.g., by classifying cell types, so that regions in the density chart do not immediately carry semantic information. This circumstance may limit the interpretability of our tool for non-subject matter experts, however, we emphasize that the tool is designed with pathologists as primary user group in mind who possess substantial domain expertise. While the algorithm does not currently identify specific cell types during embedding, pathologists are domain experts and will be able to explore and contextualize cell appearance in different embedding regions using the graphical user interface (Figure 5). Doing so as part of a training period may allow pathologists to understand and identify the major cell types in specific embedding regions, linking pie chart sectors to semantic cell types. Experience in observing how GEP class 1 and class 2 slides behave in the pie chart embedding space combined with the above training may further add to the interpretability of the model. Future work will investigate how this paradigm compares to other approaches and black box models in building trust and confidence in the user group.

In the current form, the circular space is evenly partitioned into 12 parts. However, this partitioning process could be further guided by other semantic information, e.g. clustering of specific cell types. If cell type annotations are available, the segmentation network could also output cell types for every extracted cell. We would expect to see clusters of cells types in the embedding space and the partitioning process could be further guided by the cell type clusters. However, such information is not available in our dataset and it is also unclear whether such approach would prove beneficial.

The boundary defined by the interpretable rule set could also be used as a criterion for user trust calibration. If a slide maps close to the boundary, a little variation could change the prediction result. Thus, the prediction of that slide may be perceived as less reliable. We will investigate in future work whether proximity to the decision boundary indeed correlates with prediction performance, and more importantly, other clinical outcome measures such as survival.

### 6. Conclusion

We have presented an automated yet interpretable system for UM subtyping from fine needle aspiration cytology images that does not compromise performance compared to conventional deep black box models. In future work, we will study how our interpretable model affects treatment decisions and user trust, as a next step to realize the huge potential of image-based tests for UM subtyping.
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