

# PseudoCell: Hard Negative Mining as Pseudo Labeling for Deep Learning-Based Centroblast Cell Detection

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**Abstract—Background:** Deep learning models for patch classification in whole-slide images (WSIs) have shown promise in assisting follicular lymphoma grading. However, these models often require pathologists to identify centroblasts and manually provide refined labels for model optimization. **Objective:** To address this limitation, we propose *PseudoCell*, an object detection framework for automated centroblast detection in WSI, eliminating the need for extensive pathologist’s refined labels. **Methods:** *PseudoCell* leverages a combination of pathologist-provided centroblast labels and pseudo-negative labels generated from undersampled false-positive predictions based on cell morphology features. This approach reduces the reliance on time-consuming manual annotations. **Results:** Our framework significantly reduces the workload for pathologists by accurately identifying and narrowing down areas of

interest containing centroblasts. Depending on the confidence threshold, *PseudoCell* can eliminate 58.18–99.35% of irrelevant tissue areas on WSI, streamlining the diagnostic process. **Conclusion:** This study presents *PseudoCell* as a practical and efficient prescreening method for centroblast detection, eliminating the need for refined labels from pathologists. The discussion section provides detailed guidance for implementing *PseudoCell* in clinical practice.

**Index Terms—**Centroblast cell detection, deep convolutional neural network, follicular lymphoma, hard negative mining, morphological features.

**Impact Statement—***PseudoCell* automates CB detection in H&E WSIs without requiring refined labels. It reduces pathologists’ workload by accurately eliminating 58.18–99.35% of non-centroblast regions and suggesting CB candidates at the cell level.

## I. INTRODUCTION

Follicular lymphoma (FL) is the second most prevalent lymphoid malignancy in Western and Asian countries. It is responsible for 5–35% of non-Hodgkin lymphoma (NHL) [1], [2], [3]. Most FL carries the translocation t(14;18), which causes the overexpression of the BCL-2 protein. FL patients usually present with lymphadenopathy, infrequent B-symptoms, systemic fever symptoms, night sweats, and weight loss. The progression of a disease can be predicted using a combination of clinical and laboratory findings and the histopathological grade of the disease [4].

The World Health Organization (WHO) classification system is currently the gold standard for grading follicular lymphoma (FL). This system relies on the number of centroblast cells (CBs), large neoplastic cells, identified within a tissue sample [5]. Traditionally, pathologists manually count CBs under a microscope using hematoxylin and eosin (H&E) stained tissue sections. However, this process is time-consuming and laborious due to the vast size difference between whole slide images (WSIs) and individual cells (as illustrated in Fig. 1). Additionally, it suffers from subjectivity and variability among experts, leading to inter- and intra-observer variability ranging from 61–73% [6]. This high variability introduces sampling bias,

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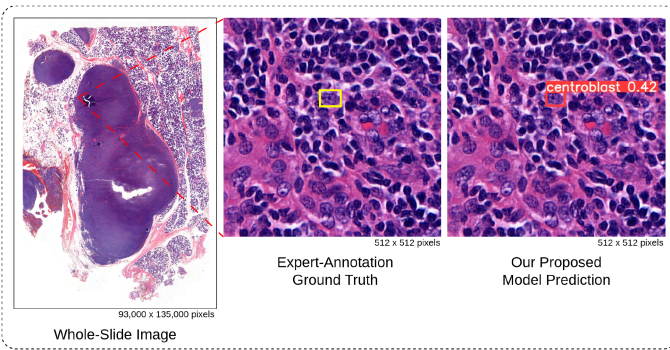
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**Fig. 1.** Automated detection of centroblast cells in whole-slide images. The tiny red square within WSI represents the patch image. Experts examined WSI under a patch-by-patch microscope to identify CB. In contrast, our proposed model can immediately identify CB with a confidence score.

hinders reproducibility, and ultimately impacts patient care due to a lack of consensus among pathologists [6]. Consequently, there is a crucial need to enhance the precision, reliability, and reproducibility of histological grading in FL.

Numerous studies have proposed automated methods to localize and classify FL by using whole-slide images (WSI), scanned images from the tissue samples, aiming to facilitate the work of pathologists [7], [8], [9], [10], [11], [12], [13], [14], [15], [16], [17]. The techniques can be categorized into two groups: 1) machine learning (ML)-based approaches with human-engineered features [7], [8], [9], [10], [11], [12], [13] and 2) deep learning (DL)-based approaches [14], [15], [16], [17]. ML-based approaches have been explored for classifying and detecting CBs. However, their performance was often limited by the reliance on hand-engineered features, particularly those based on color distributions and morphological characteristics. This approach can lead to overfitting, high false-positive (FP) prediction rates, and difficulties in generalizing to new datasets [8], [9], [10], [11], [12]. The heavy dependence on the specific combination of chosen features further restricts model performance. Consequently, recent research has shifted towards a DL-based approach, which eliminates the need for hand-engineered features and can automatically extract essential features from the training data.

DL-based models, especially Convolutional Neural Networks (CNN), have been recently applied to detect and classify lymph nodes on H&E-stained WSIs. To detect lymphocytes in breast cancer (BC), Liu et al. [14] addressed the tumor class imbalance problem by applying random sampling and data augmentation on patches (i.e., cropped images from WSI) before training the InceptionV3 [18]. Their method achieved the best sensitivity on the Camelyon16 dataset. Then, Lu et al. [15] proposed an automated pipeline to achieve a robust model for a new cohort. Their approach employed cascade training on a U-Net architecture [19], involving an iterative process of model fine-tuning. Specifically, the model was initially trained on a source dataset and then fine-tuned on the new cohort using its predicted lymphocyte masks. These masks were subsequently evaluated and refined by pathologists before being used for further model

training. This cascade training process was repeated for two iterations, ultimately yielding a model with an F1-score of 0.927. However, while demonstrating promising results, this method introduces additional workload for pathologists during the mask refinement stage, which contradicts the goal of reducing their workload.

In contrast to BC research, where DL has been extensively explored for WSI analysis, most DL studies in FL have primarily focused on patch-level classification, specifically identifying whether patches contain CBs. This patch-level approach offers limited interpretability for directly grading FL. Somaratne et al. [16] addressed this by proposing a one-class training approach to minimize the generalization gap between two FL datasets. Their method involved incorporating images from the target dataset into the training set and then applying transfer learning with AlexNet [20]. This resulted in a 12% improvement in patch classification accuracy compared to training from scratch. Syrykh et al. [17] employed a CNN-based model to differentiate between FL and follicular hyperplasia (FH) at four different resolutions. While their model achieved accurate patch-level classification at the highest resolution, the study also highlighted the sensitivity of DL approaches to pre-processing steps, particularly stain normalization (SN). This was evidenced by a significant drop in the area under the curve (AUC) from 0.92–0.99 on the internal dataset to 0.63–0.69 on an external dataset. This accentuates the importance of stain normalization in the pre-processing phase for robust and generalizable DL models in FL.

According to the limitations mentioned above: (1) Deep Learning (DL) is sensitive to the variation of stain color in WSIs; (2) the need for expert-refined labels during training, and (3) class imbalance between CB and non-CB cells. These limitations restricted DL's improvement on FL WSIs to cell-level prediction.

To overcome these limitations, we proposed a framework called *PseudoCell* to explore the feasibility of DL-based object detection models on CB detection tasks. We aim to use the state-of-the-art object detection model, YOLOv8 [21], as our backbone model. Firstly, we compare the consistency of two Stain Normalization (SN) methods on our dataset to prevent the effect of color variation from WSI. Secondly, the need for expertly refined labels during training will be imitated through the hard negative mining technique (HNM) [22], i.e., retrieving false-positive (FP) predictions from the trained model, afterward incorporating them into the training set as pseudo-negative labels (non-CB class), then use the new training set to train a new model. Since the number of pseudo-negative labels is higher than the number of CB labels from pathologists, the imbalance class issue must be addressed before incorporating pseudo-negative labels. Thirdly, three distinct undersampling approaches were explored to mitigate the class imbalance issue before incorporating pseudo-negative labels into the training set.

To our knowledge, HNM was initially introduced in the field of computer vision and has yet to be utilized in the context of digital histopathological image recognition. While previous work on cancer cells sought refined labels from experts to enhance the model, we instead attempted to imitate it through the HNM.

This framework allows us to improve the model autonomously without relying on additional work from pathologists. Therefore, the comparison between different HNM approaches was mainly investigated.

Lastly, we have provided a practical guideline based on high-power field selection and CB identification in WSI for applying our *PseudoCell* as a pre-screening tool for FL patients. Integrating this framework with histopathological workflow can reduce experts' workload by narrowing down the region experts focus on while examining the tissue. Potential real-world applications (such as quality control, training, and education tools) are also discussed to benefit human-machine collaboration.

## II. MATERIALS AND METHODS

### A. Data Collection

This study included 75,245 patches (512x512 pixels) of follicular lymphoma (FL) admitted for treatment at the Faculty of Medicine Siriraj Hospital between 2016 and 2020. No significant correlation between clinicopathological parameters was observed (data not shown). The Siriraj Institutional Review Board (SIRB) (MU-MOU CoA No. 973/2020) has approved the procedures for obtaining and using tissue. Formalin-fixed paraffin-embedded (FFPE) tissue samples with a thickness of 3-5 microns were prepared for automated hematoxylin and eosin (H&E) staining and scanned at a resolution of 0.12 microns per pixel using a 3Dhistech Panoramic 1000 microscope with a 40x objective lens. The resulting images were saved in NRXS format.

From a total of 75,245 patches, 1203 patches contain Centroblast (CB) cells, and 3045 patches without CB were selected and annotated by a consensus of two doctors (one of them is a pathologist). The annotation is manually drawn around CB as a bounding box (bbox), Fig. 2(a).

### B. The Proposed Framework

Based on the challenge of CB cell detection, we proposed a framework in Fig. 2(b)–2(d) that gives reproducible cell-level predictions. Our proposed framework comprises three parts: 1) Train original model, 2) Hard Negative Mining pipeline, and 3) Train model with pseudo-negative label.

**1) Train Original Model:** As shown in Fig. 2(b), three steps comprise this part to obtain a one-class dataset and a CB detection model: 1.1) Stain normalization selection; 1.2) Data preprocessing; 1.3) Model training.

**a) Stain normalization selection:** Even though our WSIs came from the same lab and scanner, the WSIs still have different stain colors. So, Stain normalization was applied to our preprocessing step.

Stain normalization (SN) is the color distribution transformation from a source image  $I$  into a target image  $I'$ . The transformation can be described through the operation  $I' = f(I, \theta)$  where  $\theta$  is a collection of parameters derived from the template image, and  $f$  is the function that maps the visual appearance of a given image  $I$  to the template image. Generally,  $\theta$  is designed to capture the color information of the primary stain components (e.g., hematoxylin and eosin). Consequently, stain-normalized

images will have a color distribution similar to the template image [23].

In this work, we consider two state-of-the-art SN methods:

- **Structure Preserving Color Normalization (SPCN):** Vahadane et al. proposed in [24], which tackled the stain separation problem with the assumption that stain density is non-negative, and the color basis is sparse. The sparseness constraint reduces the solution space of the color decomposition problem. Then, the color basis of a source image is replaced with those from a template image while maintaining its original stain concentrations.
- **Deep convolutional Gaussian mixture models (DCGMM):** Zanjani et al. proposed in [25]. This method first converts the source image into the HSD color system. Then, a GMM is fitted to the color distribution individually per tissue class. To train the DCGMM, *E-step* and *M-step* of the EM-algorithm are replaced by gradient descent and the back-propagation algorithm. The advantage of this approach is that it does not need any assumptions about the H&E image content.

We conduct an experiment, detailed in Section II-C1, to compare and select the most appropriate SN method for our dataset (i.e., one that produces processed images with low color variation and minimal background error).

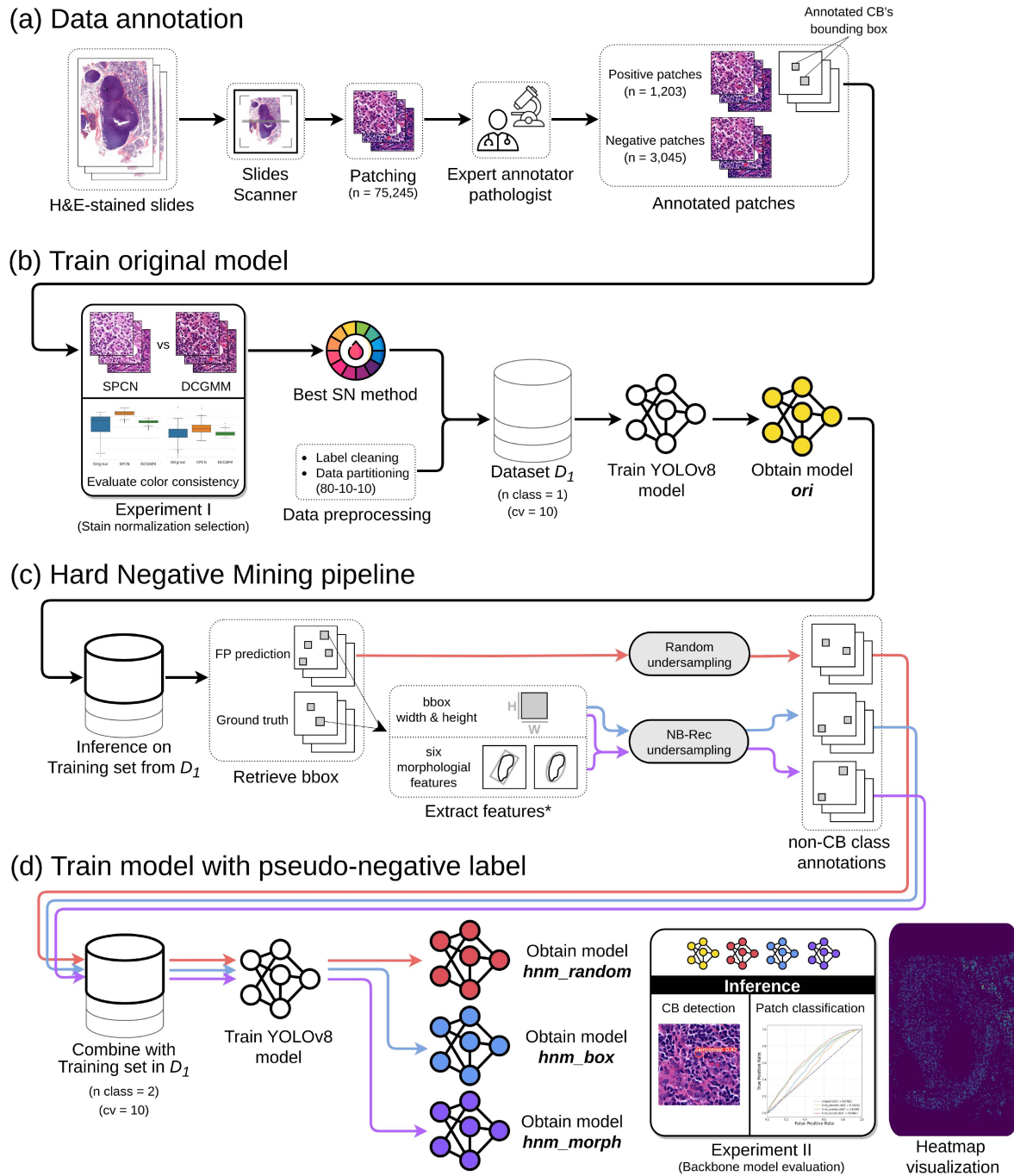
**b) Data preprocessing:** Due to the considerable human errors during annotation, label cleaning was necessary before feeding data into the model. Our dataset's two most prevalent errors were 1) bbox annotations with zero areas and 2) repeated bbox annotations on a single CB cell. Since the annotator may have accidentally generated a bbox with zero areas by clicking the mouse, we removed all box annotations with zero areas from our dataset. Regarding the second error, we first calculated the center of each bbox and then retrieved the groups of bounding boxes whose center-to-center distance is within a constant. If bbox annotations share the same CB cell, we select the bbox that best fits the cell based on manual inspection of each bbox group.

Then, we will apply the stain normalization method from the previous experiment to the annotated positive patches to standardize our dataset's color variation. Lastly, 80%, 10%, and 10% of the normalized positive patches were separated into train, validation, and test sets to create dataset  $D_1$ .

**c) Model training:** Before feeding the training set into the model, five augmentation methods (flip up-down, flip left-right, rotate 90 degrees, rotate 180 degrees, and rotate 270 degrees) were applied to the training set.

We trained and validated a YOLOv8 model with the X6 architecture using 10-fold cross-validation on the augmented dataset  $D_1$ . The model, consisting of 350 layers and approximately 97 million parameters, was trained under the default hyperparameter configuration. We employed stochastic gradient descent (SGD) to minimize cross-entropy loss during training, which ran for a maximum of 500 epochs with early stopping implemented to prevent overfitting. This resulted in the original model, referred to as "*ori*" throughout this work.

**2) Hard Negative Mining Pipeline:** In histopathological image recognition, pathologists typically annotate only target



**Fig. 2.** Overview of the *PseudoCell* framework. (a) Data annotation: H&E-stained slides were scanned and patched into 512x512 pixels. An expert pathologist annotated centroblast (CB) cells on selected patches by locating a rectangle bound to each CB while another expert pathologist reviewed the annotations. (b) Train original model: The CB-annotated patches were cleaned and then normalized the color using the best stain normalization method from *Experiment I*. Then, 80%, 10%, and 10% of the data were separated into train, validation, and test sets to create dataset  $D_1$ . Five methods augmented the training set before being fed into YOLOv8. YOLOv8 was trained and validated in a 10-fold cross-validation manner to generate *ori* model. (c) Hard Negative Mining pipeline: The *ori* model was then applied to infer the training set to retrieve false-positive (FP) samples. We then employed three undersampling strategies (red, blue, and purple paths) to avoid the imbalance class issue. \*: the six morphological features [28] were calculated using a binary image segmented by a trained HoverNet model [29]. (d) Train model with pseudo-negative label: The undersampled FP samples from each path were combined with the training set of dataset  $D_1$  as a new class. Consequently, each path had its own training set of two classes with identical validation and test sets from  $D_1$ . YOLOv8 was trained and validated with similar manner as *ori* model to obtain *hnm\_random*, *hnm\_box*, and *hnm\_morph* models. Finally, all models were compared in *Experiment II*, and the best approach was applied to visualize the heatmap of WSI.

cells (i.e., CB cells) and leave other cells unannotated to minimize the annotation cost. It causes DL-based models to typically perform poorly due to many false-positive (FP) predictions.

We hypothesize that distinguishing CB cells from other cells that look like CB cells (non-CB cells) is the key to improving the model. One approach is incrementing the non-CB labels as a new class in the dataset. In practice, we retrieve the FP bbox (i.e., non-CB annotation) from the *ori* model inference on the training set and add them to the training set as a new class. As shown in Fig. 2(c), the following three steps were employed to generate a dataset with pseudo-negative labels: 2.1) Retrieve FP predictions; 2.2) Undersample; and 2.3) Combine the non-CB class with the training set.

**a) Retrieve FP predictions:** To obtain FP annotations, we applied the *ori* model to the training set within each cross-validation fold using a low confidence threshold of 0.001. This ensured the model predicted all possible negative cases. Since CB bounding boxes (bbox) in the training set had a long side smaller than 100 pixels, we further filtered out any predicted FP bbox with a side length exceeding this threshold. This step helped to eliminate spurious detections of large objects that were unlikely to be true CBs.

**b) Undersample:** As the number of identified FP predictions remained significantly higher than the actual number of CB cells, directly incorporating these negatives into the training set would lead to a class imbalance problem. To address this issue and ensure balanced representation during model training, we considered two undersampling strategies: Random undersampling and Neighborhood-based Recursive search undersampling (NB-Rec) [26].

- *Random undersampling* is a popular non-heuristic technique due to its simplicity of application. Despite its simplicity, there is a significant disadvantage that must be considered. Given that balanced class distribution is a stopping criterion, random undersampling may eliminate potentially useful samples to achieve this balance [27].
- *NB-Rec* eliminates the majority class sample, which may overlap with the minority class. As described in Algorithm 1, the majority sample is considered overlapping when it is in the neighborhood of more than one minority sample. Since the NB-Rec uses K-Nearest Neighbor (KNN), we must search for  $k$  before execution to produce several negatives approximately equal to the actual CB.

As the NB-Rec undersampling method requires coordinate information, we extracted features from both the ground truth and FP bbox. The width and height of each bbox were directly obtained. Additionally, six morphological features, as described in [28], were calculated for each cell within the bbox. To achieve this, we first segmented the cells using a trained HoverNet model based on the *PanNuke* architecture [29], resulting in binary images of individual cells within each bbox. These binary images were then used to compute the morphological features.

As depicted in Fig. 2(c) by the red, blue, and purple paths, we obtain three sets of undersampled FP predictions: (1) the set from random undersampling, (2) the set from applied NB-Rec undersampling to bbox width and height, and (3) the set from NB-Rec undersampling applied to the first- and second-principal

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### Algorithm 1: NB-Rec.

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**Input:** training set,  $k$

**Output:** undersampled training set

**begin**

```

 $T \leftarrow$  training set
 $T_{pos} \leftarrow$  positive instances in  $T$ 
 $A \leftarrow$  frequency table
for  $x \in T_{pos}$  do
     $NN \leftarrow k$  nearest neighbours
     $NN_{neg} \leftarrow$  negative members of  $NN$ 
    for  $y \in NN_{neg}$  do
         $A_y.freq \leftarrow A_y.freq + 1$ 
for  $x \in A.instance$  do
    if  $A_x.freq > 1$  then
         $X \leftarrow X \cup x$ 

 $A' \leftarrow$  frequency table
for  $x_1 \in X$  do
     $NN_2 \leftarrow k$  nearest neighbours
     $NN_{2neg} \leftarrow$  negative members of  $NN_2$ 
    for  $y \in NN_{2neg}$  do
         $A'_y.freq \leftarrow A'_y.freq + 1$ 
for  $x_2 \in A'.instance$  do
    if  $A'_{x_2}.freq > 1$  then
         $X_2 \leftarrow X_2 \cup x_2$ 
 $T' \leftarrow T \setminus (X \cup X_2)$ 
return  $T'$ 

```

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components of bbox width, bbox height, and six morphological features using the Principal Component Analysis (PCA) method.

**c) Combine the non-CB class with the training set:** Following the undersampling procedures described in the previous step, we created three new datasets by incorporating the respective sets of undersampled FP predictions into the original training set of  $D_1$ . These new datasets have similar images to  $D_1$  but contain two classes (CB and non-CB) in the training set. However, to maintain consistency and evaluate the models' ability to generalize to unseen data, we retained only the CB class in the validation and testing sets for all three datasets.

**3) Train Model With Pseudo-Negative Label:** As shown in Fig. 2(d), similar to the part 1) *Training original model*, we use the same setup *model training step* on the dataset of red, blue, and purple paths to get model *hnm\_random*, *hnm\_box*, and *hnm\_morph*, respectively.

## C. Experiment Setup:

All experiments were performed with an NVIDIA Tesla V100-SXM2 graphic card.

**1) Experiment I: Stain Normalization Selection:** This experiment aimed to ensure color consistency within our dataset by comparing two stain normalization methods: SPCN and DCGMM. We selected a template image, preferred by an expert, from the available patches. Both stain normalization methods

were then applied to the remaining images in the dataset. The resulting normalized images were evaluated using metrics described in *Supplementary Materials, Section I-A1*. The stain normalization method that demonstrated superior performance will be employed in the pre-processing phase of this work.

**2) Experiment II: Backbone Model Evaluation:** This experiment aims to compare the performance of models from different training approaches (i.e., conventional and HNM approaches) on both object detection and image classification tasks. We use the training pipeline described in Section II-B to obtain four models:

- *Original (ori) model:* Conventional object detection approach with one class annotation.
- *Model trained with random HNM (hnm\_random):* Randomly add FP samples from *ori* prediction on the training set into the training set as a new class. Then, it trains the model with the same setup as *ori*.
- *Model trained with HNM of bbox features (hnm\_box):* Instead of randomly sampling, this approach undersamples the FP samples using NB-Rec on the width and height of the FP bounding box, then adds them into the training set.
- *Model trained with HNM of morphological features (hnm\_morph):* Similar to *hnm\_box* but using NB-Rec on first- and second-principal components from six morphological features and width and height of FP bounding box.

We use metrics from *Supplementary Materials, Section I-A2* to evaluate the performance models on object-level prediction.

To evaluate the model’s performance on the image classification task, we mapped the cell-level predictions to patch-level classifications using the following criteria: *a patch was classified as positive if it contained at least one predicted CB cell; otherwise, it was classified as negative*. Given that the test set of each cross-validation folds comprised 120-121 images containing CBs (positive images), we augmented each test set with additional negative images randomly selected from our database. This ensured a balanced class distribution for robust model classification performance evaluation.

### III. RESULTS

#### A. Experimental Results

**1) Experiment I. Stain Normalization Selection:** Deep convolutional Gaussian mixture models (DCGMM) yielded the lowest standard deviation (SD) and coefficient of variation (CV) for both Normalized Median Intensity (NMI) and Normalized Median Hue (NMH) metrics, as indicated in Table I. Since NMI qualifies the color consistency of the nuclei [30] and NMH quantifies the global color variation of an image population [31]. Thus, the results indicate that DCGMM provides qualitatively similar color distributions for nuclei with less color variation within the image population (see Fig. 3). Comparing the original and the Structure Preserving Color Normalization (SPCN), the box plots in Fig. 4 demonstrate that DCGMM has the smallest spread of NMI and NMH values around the median (interquartile range) with variance statistical significance ( $p < 0.01$ ).

TABLE I

STANDARD DEVIATION AND COEFFICIENT OF VARIATION FOR NMI AND NMH, ALONG WITH MEAN VALUES AND STANDARD DEVIATION FOR AMCE, FOR THREE DIFFERENT IMAGE POPULATIONS

Method	NMI		NMH		AMCE <sub>α</sub>	AMCE <sub>β</sub>
	SD	CV	SD	CV	Mean ± SD	
Original	0.082	0.137	0.039	0.042	5.30 ± 5.06	17.28 ± 9.34
SPCN	0.057	0.086	0.012	0.012	<b>4.67</b> ± 3.34	<b>1.64</b> ± 1.33
DCGMM	<b>0.026</b>	<b>0.044</b>	<b>0.008</b>	<b>0.008</b>	6.43 ± 2.69	18.47 ± 1.33

The bold values indicate the best score among the three stain normalization methods (Original, SPCN, DCGMM) in each metric.

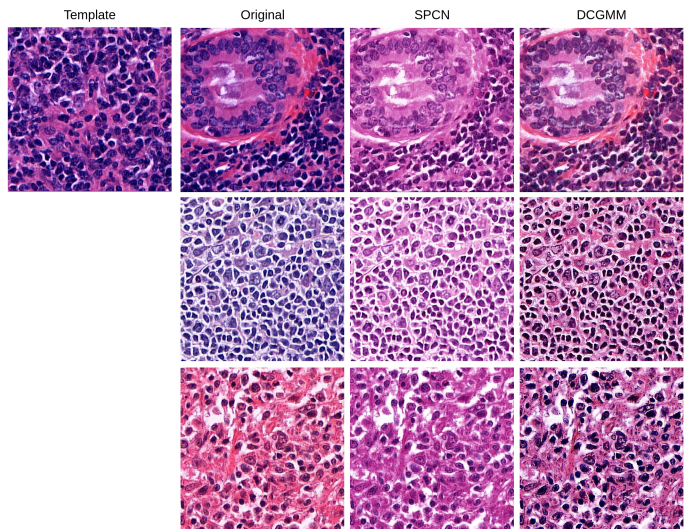


Fig. 3. Illustration of the performance of different stain normalization methods. The top-left image is the template image. The next column is the images sampled from the original images, followed by the results of normalization using SPCN and DCGMM, respectively.

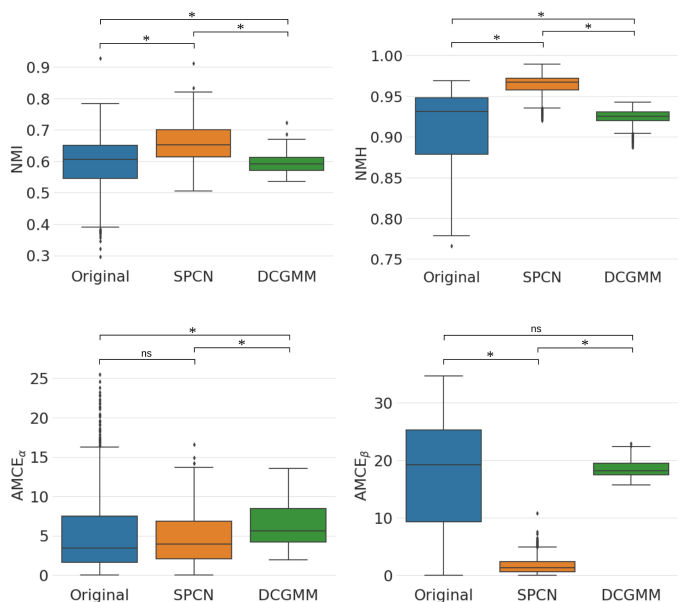
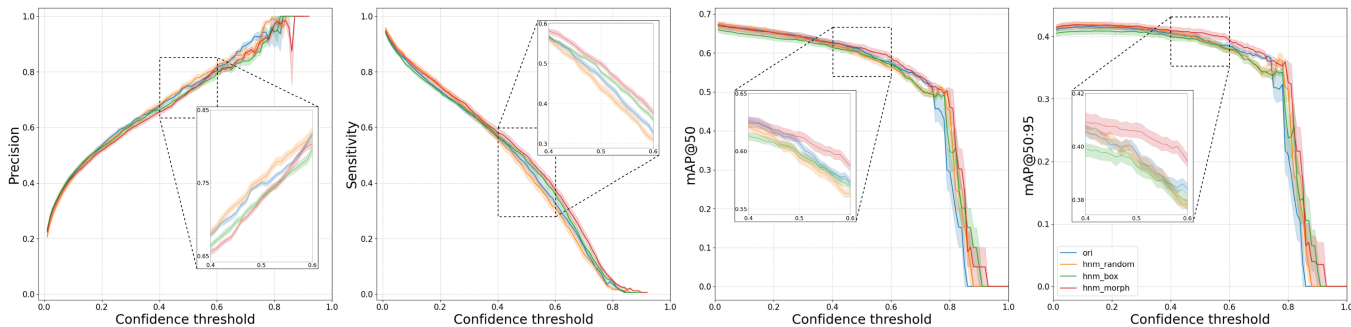
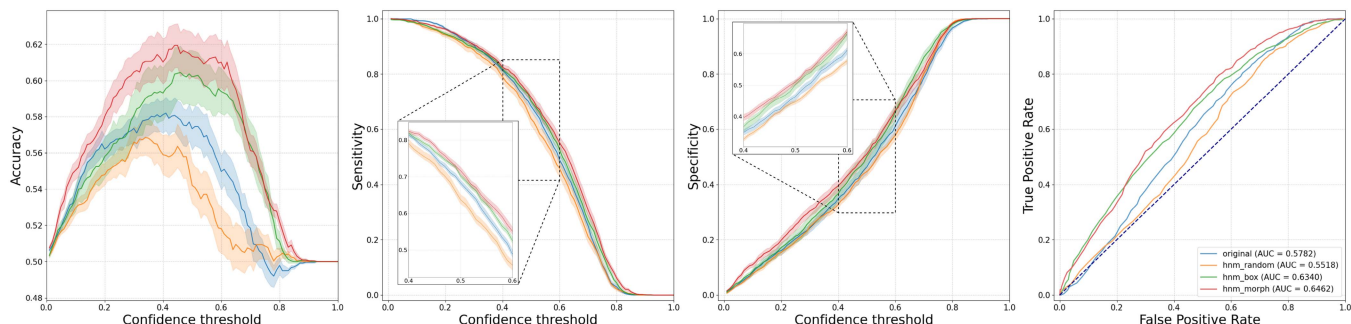


Fig. 4. Box plots of NMI, NMH, AMCE<sub>α</sub> and AMCE<sub>β</sub> values for all stain normalization methods in Experiment I. \* denotes the statistical significance of  $p < 0.01$ , and ns denotes not statistical significance.



**Fig. 5.** Centroblast detection results: Precision, Sensitivity, mAP at 0.5 IOU threshold (mAP@50), and mAP at 0.5 to 0.95 IOU threshold (mAP@50:95) of each model on test dataset with standard deviation error bar.



**Fig. 6.** Patch classification results: Accuracy, Sensitivity, Specificity, and Receiver Operating Characteristics (ROC) curves of each model on test dataset with standard deviation error bar.

Next, we evaluate the background error of each image population. DCGMM has a significantly higher mean Absolute Mean Color Error in  $\alpha$  space ( $AMCE_{\alpha}$ ) than the original image population. For  $\beta$  space ( $AMCE_{\beta}$ ), DCGMM does not have statistical significance compared to the original. It suggests that the DCGMM-processed images contain more or equal background errors than the original images, contradicting the goal of reducing color variations.

Even though SPCN does not have statistical significance with the original at  $AMCE_{\alpha}$ , at  $AMCE_{\beta}$ , SPCN provides significantly less error than the original. Moreover, both values of SPCN are statistically significantly less than DCGMM.

Therefore, we decided to implement SPCN in our framework pipeline, as it offers lower SD and CV in NMI and NMH values than the original and better AMCE values for both  $\alpha$  and  $\beta$  spaces than DCGMM.

**2) Experiment II. Backbone Model Evaluation:** This experiment aimed to benchmark the performance of models trained using both conventional and HNM approaches for object detection and image classification tasks. Prior to comparing their performance, we first analyzed the optimization process of each model during training to gain insights into their convergence behavior and stability. As indicated in *Supplementary Fig. S1*, it was observed that the validation loss of the original (*ori*) model reached the lowest value when converged. Other models (i.e., all HNM approaches) exhibited a similar converging pattern more rapidly than the *ori* model, albeit with a higher loss. Despite the model trained with HNM of morphological

features (*hnm\_morph*) going with the same trend as other HNM approaches, it achieved the highest performance in terms of mean average precision (mAP@50) and accuracy, as shown in Figs. 5 and 6, respectively.

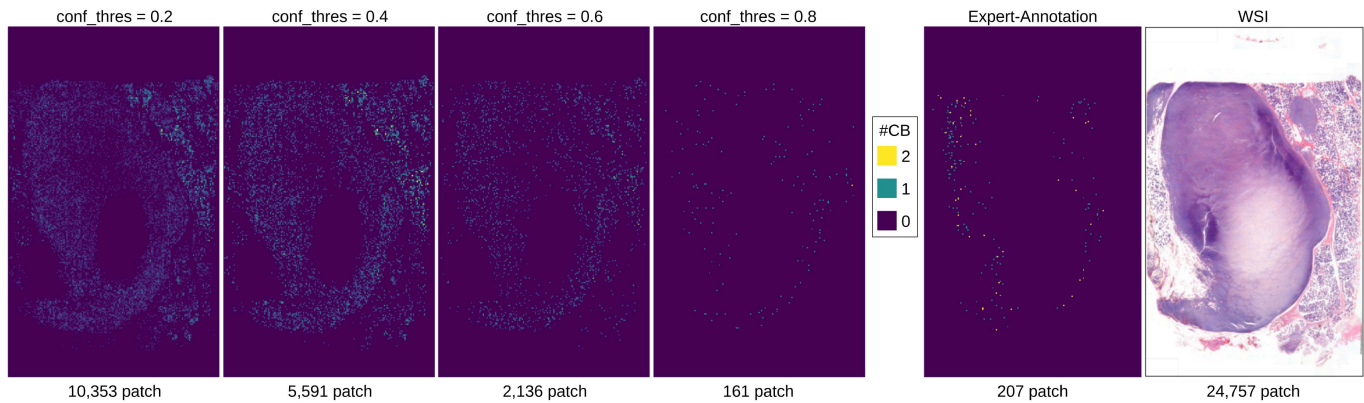
For the object detection task, Fig. 5 provided an overview of model performance on each metric with a confidence interval. All models follow the same trend. The *hnm\_morph* achieved slightly better sensitivity, but in contrast, the trade-off appears to be on precision. Nevertheless, its performance is superior to other models in mAP@50 and mAP@50:95.

For the image classification task, the *hnm\_morph* outperformed all other models, especially the model trained with random HNM (*hnm\_random*). Notice that the performance of *hnm\_random* dropped from *ori* on all metrics. In contrast, *hnm\_morph*, which was trained with the same approach but with a more reasonable undersampling method, improved its performance over the *ori*.

## IV. DISCUSSION

### A. Effect of Training With Pseudo-Negative Labels on the Model Performance

Concerning the impact of training with pseudo-negative labels from the hard negative mining (HNM) technique over the conventional training approach, we demonstrated improvements in centroblast (CB) detection and patch classification. However, the validation loss of these models is higher than that of the conventional training method (*Supplementary Materials, Fig.*



**Fig. 7.** Heatmap visualization of detected centroblast in each patch on an unseen WSI. As `conf_thres` increases, the bright patch becomes sparser and more prone to expert annotation.

S1). This is because the softmax function divides the total probability mass across multiple classes (i.e., CB and non-CB) rather than just one class of CB. When computing each class’s confidence score in the bounding box using the softmax function, the number of divisors is increased to two, resulting in the confidence scores for each class becoming smaller on average as the number of classes increases. Finally, the class loss of the YOLO model, which is the cross-entropy loss, in models trained with HNM is higher than in the *ori* model.

### B. Effect of Undersampling Approaches on the Model Performance

Since our framework was designed to imitate the training loop with the pathologist’s refined labels, identifying CB based on cell color and morphology, we retrieved false-positive samples and fed them to the model. We obtain the following models based on three undersampling approaches: *hnm\_random*, *hnm\_box*, and *hnm\_morph*. As the result of object detection and image classification tasks, the model with the morphological features (*hnm\_morph*) performs best. Suggests that the design of the undersampling approach is essential to take advantage of the HNM technique and that pathologists’ intuition still provides some information for deep learning to distinguish between non-CB cells and actual CB cells.

### C. Guidance for Clinical Implementation and Future Works

In conventional histopathological workflow, pathologists count the number of centroblasts (CB) in ten randomly selected high-power fields (HPF), leading to high inter- and intra-observer variability and being vulnerable to sampling bias. The inter- and intra-observer variability among pathologists is crucial since it directly impacts patient grading and management [6]. To reduce the variability between pathologists, a solid guideline for finding potential HPF in WSI is one solution.

With *PseudoCell* framework, pathologists will obtain two guidelines: (i) heatmap visualization for potential CB regions in WSI and (ii) CB annotations at the HPF level for identifying CB. Pathologists’ remaining job is to select the HPF and then

accept the annotation or self-identify CB cells. Pathologists must only set the confidence threshold (`conf_thres`), ranging from zero to one when using *PseudoCell*. The `conf_thres` parameter determines the initial confidence level of CB annotations reported to pathologists. A low `conf_thres` (`conf_thres = 0.2`) produces a dense heatmap, as in Fig. 7, whereas a high `conf_thres` (`conf_thres = 0.8`) produces a sparse heatmap that more closely resembles the expert-annotation.

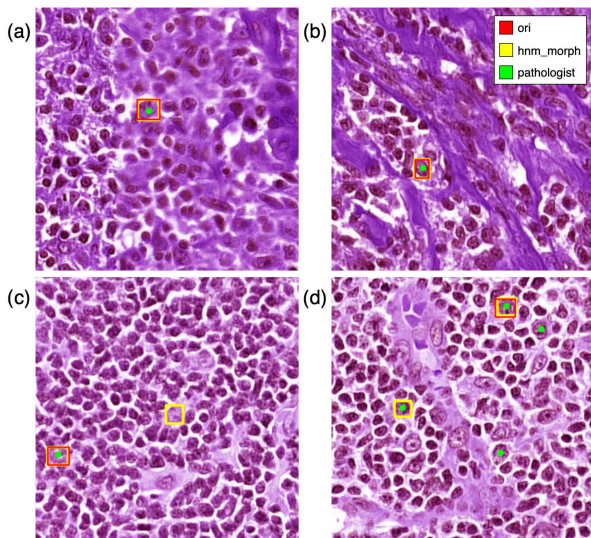
We will divide the histopathological process into the HPF selection and CB identification phases. In each phase, the real-world adjustment of `conf_thres` to facilitate pathologist preference could take the form of the following suggestion:

**1) HPF Selection Phase:** With a high `conf_thres`, *PseudoCell* offers a sparse HPF that is still sufficient to grade FL, which is suitable for pathologists who wish to complete the grading task rapidly. In contrast, when `conf_thres` is low, *PseudoCell* generates a dense heatmap that identifies the region containing intensive CB and regions with less CB. This approach is suitable for pathologists who wish to determine the HPF independently.

**2) CB Identification Phase:** A high `conf_thres` is advantageous for pathologists who prefer to self-identify on CB with some framework-suggested CB annotation. In contrast, a low `conf_thres` will enable the framework to recommend more CB annotation, which is ideal for pathologists who wish to check off the annotation.

The pathologists’ workload can be reduced by *PseudoCell* accurately narrowing down the areas requiring their attention during examining tissue as in Fig. 7. From all 24,757 patches with tissue in WSI, the framework highlights 10,353 and 161 patches that contain potential CB candidates based on `conf_thres` with an inference time of approximately 0.03 seconds per patch. In other words, the framework can eliminate 58.18 to 99.35% of all WSI patches that do not appear to be CB candidates at the `conf_thres`. Pathologists can, therefore, focus on identifying CB on the slide. We anticipated that the inter- and intra-variability of pathologists would decrease after implementing our framework in the real world. Meanwhile, the machine can benefit from pathologists’s actual CB as refined labels. These labels can be used to improve the model’s performance in the future. This





**Fig. 8.** Comparison between *ori*, *hnm\_morph*, and pathologist's annotation in identifying centroblast cells. The illustration shows that *ori* and *hnm\_morph* correctly identify CB in (a) and (b), which are partially blurred and mixed types of cells, respectively. For (c), *hnm\_morph* had a false positive. For (d), *hnm\_morph* performed better than *ori* but still had some false negative predictions.

cycle leads to human-machine collaboration in the real world, which is one of the objectives of this work.

*Pseudocell* can also provide a second opinion that offers additional patient safety and instills greater confidence in doctors, enhancing their efficiency and reducing the likelihood of errors. For instance, when there is a need to distinguish between an infection and follicular cell lymphoma, particularly in its early stages, Thai pathologists, already handling a heavy workload, could issue a false negative, especially when the pathological area is small.

Integrating *Pseudocell* into the histopathological workflow offers several benefits. Firstly, the model assists pathologists by highlighting regions or suggesting potential CB cell candidates within the tissue, narrowing the examination focus. It serves as an additional quality control mechanism, flagging areas that may contain CB cells and assisting pathologists in not overlooking significant findings, thereby reducing diagnostic errors. In addition, pathologists can use the model's predictions as a benchmark to compare and contrast their observations. This iterative process improves their recognition of centroblast cells, enhancing diagnostic precision. Incorporating *Pseudocell* contributes to improved efficiency, quality control, and training and education for identifying centroblast cells in histopathology.

While implementing the *Pseudocell* framework offers a cost-effective alternative to the ongoing expenses of maintaining a pathology team, its integration into clinical practice faces workflow optimization and trust-building challenges. The complexity of healthcare systems necessitates careful integration to ensure seamless operation, potentially requiring adjustments to existing infrastructure and the deep learning pipeline. Additionally, addressing the "black box" nature of deep learning models through enhanced interpretability and transparency is crucial for gaining the confidence of clinicians and patients in *Pseudocell*'s predictions and recommendations.

In future work, if additional object detection models or updated versions exist, the *Pseudocell* framework permits their implementation by modifying the backbone model. Furthermore, dealing with data limitations and model transparency is crucial for pathologists to understand and have confidence in the model's decision-making. Combining weakly supervised paradigms (e.g., MIL or Attention Map) with explainability techniques (e.g., LIME, SHAP, and CAM) is a promising next step to investigate.

## V. CONCLUSION

In conclusion, our study introduces the *Pseudocell* framework for centroblast (CB) cell detection, which enhances the performance of the backbone model by using false-positive samples from the Hard Negative Mining (HNM) method as pseudo-negative labels. *Pseudocell* effectively distinguishes between actual CB and non-CB cells in patches from whole-slide images (WSI). Our experiments and evaluations demonstrate that model training from HNM on Neighborhood-based Recursive search undersampling using morphological features achieves the best results in CB detection and patch classification tasks. *Pseudocell* can reduce pathologists' workload by accurately identifying tissue areas requiring attention during examination. Depending on the confidence threshold, *Pseudocell* can eliminate 58.18–99.35% of non-CB tissue areas on WSI. Furthermore, *Pseudocell* can serve as a second opinion to differentiate between infection and follicular cell lymphoma, particularly in the early stages, making it cost-efficient for quality control and educational purposes in CB recognition. This study presents a practical centroblast prescreening method that does not rely on pathologists' refined labels for improvement. It suggests the potential for human-machine collaboration in CB identification, alleviating the burden on clinicians by focusing their labeling efforts on regions suggested by *Pseudocell* rather than manual labeling as conventionally done.

## APPENDIX

Although our findings are promising, the limited sample size used for model training and testing may affect its accuracy and generalizability. Additionally, the *Pseudocell* model demonstrates sensitivity to WSI stain color variations, leading to false positives and false negatives in cell detection, as illustrated in Fig. 8. For example, Fig. 8(c) shows a false positive identification by the *hnm\_morph* model compared to the *ori* model. In contrast, Fig. 8(d) demonstrates some false negatives, potentially due to stain variations that cause missing cell structures. Further investigation is needed to fully understand the cause of these false negatives and explore potential mitigating strategies. Despite these limitations, the model accurately identified CBs in blurred images and those with mixed cell types, Fig. 8(a) and 8(b). This suggests that data augmentation techniques and a larger, more diverse dataset could improve the model's robustness. Our study demonstrates the potential of object detection models trained with pseudo-negative labels to enhance cell- and slide-level prediction performance, ultimately aiming to reduce pathologist workload.

## SUPPLEMENTARY MATERIALS

Supplementary materials explain all evaluation metrics, visualize validation losses for each model during training, and illustrate the models performance in challenging cases.

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