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# A Guided-ensembling Approach for Cell Counting in Fluorescence Microscopy Images

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**ABSTRACT** Although deep learning and computer vision-based approaches have demonstrated success in the field of cell counting and detection in microscopic images, they continue to have certain limitations. More specifically, they experience an overall increase in false positives when dealing with cell populations that show high density and heterogeneity. Existing approaches require the reselection of parameters for each new data set to improve the accuracy of cell counting. Therefore, it is necessary to revise the fundamental models for each new microscopic image. This study introduces a novel neural network-based method that eliminates the need for retraining by combining the pretrained Cellpose and Stardist models. The accuracy of our proposed approach was evaluated on a variety of microscopic images. Despite variations in cell densities, our proposed approach demonstrated a notably improved cell counting performance in comparison to solely utilizing the Cellpose and Stardist models.

**INDEX TERMS** cell counting, cell detection, deep learning, ensemble learning

#### I. INTRODUCTION

THE traditional way of cell analysis in clinical practice involves microscopic observation of a tissue sample. The process allows us to precisely classify different cellular characteristics and evaluate cellular structure. The initial step of such analysis usually requires cell segmentation [1]-[9], cell detection [10], [11], or cell counting [12]. Although experts can extract limited information through counting cells, it is still an acceptable approach in clinical practice due to its practicality [13]. As one might expect, manual counting of cells is a very time-consuming and labor-intensive task with several potential drawbacks, including (i) risk of human error, (ii) lack of objectivity, (iii) poor reproducibility, and (iv) low throughput. In terms of human error, the issues include misidentification and missed cells due to fatigue or distraction. Moreover, it is subjective and dependent on the expertise of the observer, resulting in differences between observers [14]. Reproducibility is another problem that leads to variability between observers and differences across different laboratories or studies. It is one of the critical concerns in research, and several studies have highlighted the variability

in manual cell counting results [15]. Moreover, manual cell counting may not be practical for studies on a large scale or experiments requiring high throughput. Due to these significant limitations, semi-or fully-autonomous techniques based on image processing and neural networks for automated cell counting have been developed. The development of computerized cell counting software will significantly impact biological research and clinical practice, as it will facilitate faster, more objective, and more standardized cell counting than manual methods [16].

#### **II. RELATED WORK**

Conventional image processing-based techniques are straightforward methodologies that do not require labeled data and can resolve a specific problem with fewer lines of code compared to neural network-based methods [8], [9]. Nevertheless, they are susceptible to hyperparameter settings and necessitate manual adjustments when applied to diverse datasets. Furthermore, when confronted with cell populations exhibiting high density and heterogeneity, these methods generally tend to have a decline in precision or an increase in false positives [8], [17], [18]. Unlike traditional methods, state-ofthe-art cell detection techniques rely on training deep neural networks [4]–[6], [11]. Achieving a high level of accuracy in cell counting using neural network-based models demands extensive and data-intensive training. Despite the need for a substantial amount of data and comprehensive annotation, Cellpose [4], [5] and Stardist [6] have demonstrated particularly promising outcomes and have garnered considerable attention recently. Their popularity is primarily due to their remarkable accuracy on specific datasets. Nevertheless, they are resistant to interpretability, and the performance highly depends on the quality and appropriateness of the training data set.

Despite being popular, Cellpose and Stardist are still susceptible to variations in cell density, shape, and size. For instance, Kleinberg et al. [19] revealed that the Stardist trained model showed greater precision in estimating cell counts, particularly in regions where cells were in close proximity or even over-clumped, in comparison to Cellpose. However, in situations where the cell distribution was sparse, Stardist had a higher likelihood of producing inaccurately detected cells, whereas Cellpose appeared to exhibit greater resilience under such conditions. These findings suggest that utilizing an ensembling technique can harness the strengths of individual methods, leading to a substantial improvement in cell detection performance.

Ensemble approaches are a sophisticated method for constructing multiple models with the goal of achieving improved results [20]. There exists only a limited number of studies on the subject of cell counting through an ensembling approach [21]–[26]. For instance, [25] ensembles the output of three models, where each model utilized the same architecture but trained on different image batches. As another example, [26] uses an ensembling methodology to combine the outputs of five models to segment cells from 3D volume. Each model was trained with a different number of frames, i.e., a model seeing one frame, a model seeing three frames, etc. This way of ensembling helps eliminate low-confidence markers and merge overlapping segmentations to detect and count mimicked circulating tumor cells (mCTC). On the other hand, all of these studies necessitate training or fine-tuning of every model to enhance predictive accuracy, which is timeconsuming and laborious. Due to the challenges associated with training, there exists a significant demand for a novel approach, particularly in the context of ensemble approaches.

There are two main highlights of this study:

- We introduce a novel guided-ensembling technique that leverages two state-of-the-art cell detection methods, which have been trained with billions of cell images, with no additional retraining or fine-tuning. Experiments on three different cell counting datasets have shown that the guided-ensembling approach outperformed the separately used techniques.
- Contrary to most of the existing studies in this domain, we compare the results in images with high congestion

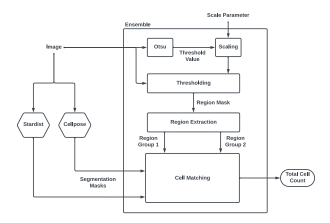


FIGURE 1. Top level view of the process.

diversity and tissue noise, which creates many artifacts in sparse regions and causes vague segregation of cells in congested regions. We show the effects of the heuristic approach on these harder-to-process images.

#### **III. METHODOLOGY**

This approach involves the integration of state-of-the-art cell segmentation models with a hard decision-making mechanism that relies on the size of the cell clusters. Fig. 1 shows the block diagram of the proposed approach. The objective of the proposed approach is to take advantage of the need for additional training by finding optimal regions for each model. Due to variations in training sets and network structures across models, each model is expected to show outstanding detection performance for specific areas in a given image. To maximize the performance of different models, the proposed study attempts to solve the cell counting problem by implementing a two-step approach. In the first phase, the algorithm produces segmentation masks of an image. The next step includes the utilization of Otsu's algorithm to extract areas containing cellular presence and then assigning each area to a model based on cellular density.

For convenience, in the rest of the paper  $M_{CP}$ ,  $M_{SD}$ , and  $M_E$  will denote three different detection methods, Cellpose, Stardist, and the proposed guided ensembling methods, respectively. The proposed approach will be explained comprehensively in the following sections.

# A. CELL SEGMENTATION VIA CELLPOSE AND STARDIST

For  $M_{CP}$ , we used the pretrained "cyto" model, which predicts the probability of a pixel being inside a cell, the flows of pixels toward the center of a cell in X and Y for each pixel. It utilizes a standard U-Net backbone with 32 layers of blocks. Each block consists of 3x3 convolution and max pooling (or upsampling).  $M_{CP}$  was trained with 540 images that have more than 70,000 cells.

As for  $M_{SD}$ , we used the pre-trained model "2D\_versatile\_fluo" as , which was trained with a subset of the DSB2018 nuclei

segmentation dataset [27]. It predicts for every pixel *a star*convex polygon for the cell instance at that position. On top of the popular U-Net architecture [28], an additional 33 convolutional layer with 128 channels (and relu activations) is added to avoid *fight over features*. Both methods accept a grayscale image (R-channel for  $D_{ORIG}$ ), *I*, as input.

### **B. GUIDED-ENSEMBLING APPROACH**

The ensemble stage uses the original image I with the segmentation masks  $S_{CP}$  and  $S_{SD}$ . Users can fine-tune the algorithms' decision-making mechanisms by adjusting two external parameters, namely  $P_{scale}$  and  $P_{threshold}$ . The variable  $P_{scale}$ is a numerical value within the interval [0,1], which represents the scaling factor applied to the threshold value obtained through Otsu's algorithm, as utilized in Section III-B1. The variable  $P_{threshold}$  is a parameter that can assume any positive integer value and serves to denote the threshold level for the pixel area covered by a connected component. Sections III-B2 and IV-C provide an additional explanation of the utilization of the parameters.

Initially, the algorithm produces a grayscale version of the original image and then employs thresholding methodologies to isolate areas that are densely occupied by cells. The congested cell groups can be identified by the threshold algorithm and indicated by each connected component produced by the threshold algorithm. The presence of a significant cluster is indicated by a large number of cells showing minimal contrast variation with the surrounding background and being near neighboring cells within the corresponding area. The algorithm's cell matching phase involves iterating through each region and determining the cells to be included from the input segmentation masks. The total number of cells in all areas is then computed by adding up the counted cells.

# 1) Binary Mask Generation and Region Extraction

At first, the ensembling procedure will determine a threshold value by means of Otsu's algorithm. Then, this value will be adjusted proportionally using an external parameter denoted as  $P_{scale}$ . The threshold value needs to be adjusted to include cells that have lower brightness values. This procedure results in the creation of certain insignificant artifacts that are related to tissue noise. The algorithm prevents artifacts by eliminating regions that do not meet the minimum size threshold to be categorized as cells. Following the generation of the mask, the region props tool in MATLAB is used to extract each region.

#### 2) Matching and Counting Cells

The algorithm evaluates each region and determines the appropriate segmentation mask to include the cells. The algorithm analyzes the area size of each cell and then examines the overlapping cells from  $S_{SD}$  if the area size exceeds the specified threshold parameter. If the value falls below the specified threshold parameter  $P_{threshold}$ , the algorithm proceeds to search for overlapping cells in  $S_{CP}$ . For a cell to be considered part of a given region, it is necessary that it has a minimum overlap of 50%. This method additionally

assigns new labels to all cells obtained from the segmentation masks and transfers them to an intermediary segmentation mask. The algorithm provides the total number of cells of the intermediate mask.

# **IV. EXPERIMENTS**

This section presents the experimental findings for each dataset.

# A. DATASET

Three data sets were used in the experimentation. The Broad Bioimage Benchmark Collection website offers two publicly accessible datasets, namely BBBC004 [29], [30] and BBBC039, as documented in [31]. Each of the mentioned datasets consists of images obtained through fluorescence microscopy. From now on, the data sets shall be denoted as  $D_{ORIG}^{1}$ ,  $D_{C4}$ , and  $D_{C39}$ , respectively. Table 1 presents an overview of the data sets mentioned.

| Dataset Name      | Number of Images | Image Resolution |
|-------------------|------------------|------------------|
| D <sub>ORIG</sub> | 290              | 1924x2572        |
| $D_{C39}$         | 200              | 520x696          |
| $D_{C4}$          | 300              | 950x950          |

TABLE 1. Dataset names, number of images, and image resolutions.

The sample denoted as  $D_{ORIG}$  includes fish brain cells that have been magnified by a factor of 20. Cells show significant variability in terms of counts, congestion, and tissue noise. The images provided have high resolution, measuring 2572 pixels in width and 1924 pixels in height. On average, the diameter of the cells measures 20 pixels. All cellular structures have uniform sizes and shapes. A total of 290 images have been reported, each with a reported count of the total number of cells in existence. A total of twelve images in the data set have point annotations.

The datasets denoted as  $D_{C4}$  and  $D_{C39}$  include fluorescent microscopy images that are supported by annotations of the cell count and foreground. These properties provide consistency to the outcomes in relation to  $D_{ORIG}$ , thus enhancing the reliability of the following comparisons. The images show different cell congestion patterns that match the characteristics of the  $D_{ORIG}$  dataset. The images obtained from  $D_{C4}$ and  $D_{C39}$  show comparatively lower levels of congestion and tissue noise, making them easier to process using state-of-theart cell counting models in contrast to  $D_{ORIG}$ .

The dataset denoted by  $D_{C4}$  comprises artificially produced cellular images showing different levels of congestion probabilities, namely 0%, 15%, 30%, 45%, and 60%. One hundred images have been distributed into five subsets, each showing distinct levels of congestion. Each image contains a total of 300 individual units. Foreground segmentation can

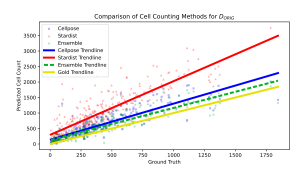
<sup>&</sup>lt;sup>1</sup>Bilkent University Local Animal Ethics Committee (HADYEK) approved the animal ethics protocols of this data with the following approval dates and numbers: July 10, 2015, with protocol number 2015/31 and February 21, 2018, with protocol number 2018/4.

also be obtained. The dimensions of each image are 950 pixels for both width and height.

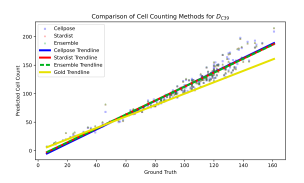
The dataset denoted by  $D_{C39}$  includes a total of 200 U2OS cellular images. The resolution of each image is 520x696 pixels. The ground truth data related to foreground segmentations, outlines, and cell counts are accessible.

#### **B.** RESULTS

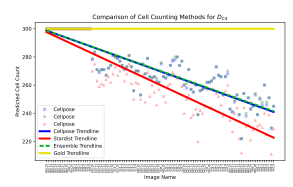
This subsection provides an analysis of the results obtained from the proposed approach. Trendlines are included in the cell count plots for each method to enable a comprehensive comparison. The results of the cell counts obtained by each method for each manual cell count can be seen in Fig. 2. Note that some images show identical manual cell counts, which allows for the possibility of a method that includes multiple markers that are vertically aligned. Each method has a trendline that fits its results. The trendline for the variable  $M_E$  shows a higher degree of closeness to the results of manual counting, in contrast to the trendlines of  $M_{CP}$  and  $M_{SD}$ . This implies that the accuracy of  $M_E$  is better in general. The values of  $P_{scale}$  and  $P_{threshold}$  for the given dataset have been set as 0.8 and 10000, respectively.



**FIGURE 2.** Cell count results for *D<sub>ORIG</sub>*. The X-axis denotes manual cell counts, and Y-axis denotes calculated cell counts for images. Red triangles indicate Stardist results. Blue squares indicate Cellpose results. Green circles indicate ensemble results. The gold trendline indicates the manual counts.



**FIGURE 3.** Cell count results for  $D_{C39}$ . The X-axis denotes manual cell counts, and Y-axis denotes calculated cell counts for images. Red triangles indicate Stardist results. Blue squares indicate Cellpose results. Green circles indicate ensemble results. The gold trendline indicates the manual counts.



**FIGURE 4.** Cell count results for  $D_{C4}$ . The X-axis denotes each image in the data set, and Y-axis denotes calculated cell counts for images. Red triangles indicate Stardist results. Blue squares indicate Cellpose results. Green circles indicate ensemble results. The gold trendline indicates the manual counts.

The results of the calculated cell count for each manual cell count are presented in Fig. 3. The graphical representation shows characteristics identical to those of Fig. 2. The closeness between the green trendline ( $M_E$ ) and the gold trendline (manual counts) is visible. The present data set has values of 0.8 and 800 for  $P_{scale}$  and  $P_{threshold}$ , respectively.

The performance of all methods for the  $D_{C4}$  data set is shown in Fig. 4. Given that the initial subset showed a congestion probability of exactly 0%, all methods effectively achieved cell segmentation. As the congestion rate increased, all techniques showed a deviation from the ground truth values. It is notable that the trendlines for  $M_E$  and  $M_{CP}$  show a significant degree of overlap. Although the proposed approach did not result in a statistically significant improvement in this particular dataset, the results indicate that overall performance is limited by the benchmark methods. The values of  $P_{scale}$  and  $P_{threshold}$  for the given dataset have been set as 0.7 and 500, respectively.

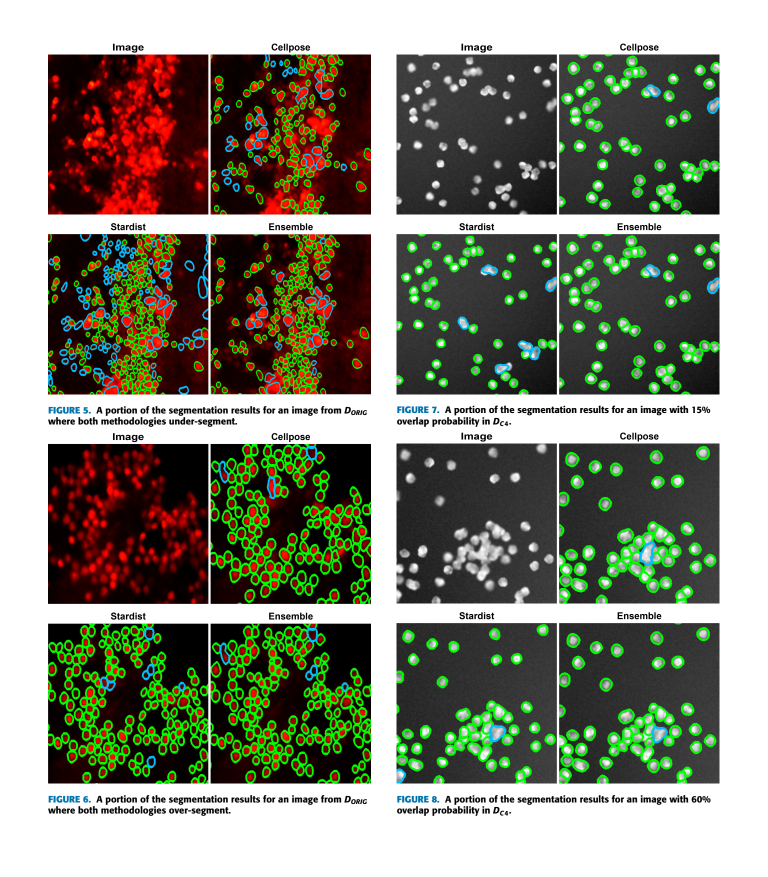
The segmentation masks corresponding to each method are represented by the green outlines in their respective segmentation figures in Figs. 5, 6, 7 and 8. The blue outlines denote segmentations that are problematic, covering cases of false positives, over-segmentations, and under-segmentations.

The graph shown in Fig. 5 illustrates the point at which  $M_{SD}$  begins to consider densely populated cells as a single unit, whereas  $M_{CP}$  ignores these regions entirely. The utilization of  $M_{SD}$  is intended to improve cell counts in densely populated regions as the primary goal of this study.

Both Fig. 5 and Fig. 6 show false-positive cell detection in low-intensity areas due to background interference. However, the impact of these extra cells is negligible since they can be readily eliminated through thresholding or similar methodologies.

The segmentation outcomes of all methodologies for the  $D_{C4}$  data set are shown in Fig. 7 and Fig. 8. Despite a few variations in the count results, the segmentation results show a high degree of similarities. It is challenging to make comparisons or observe enhancements as both  $M_{CP}$  and  $M_{SD}$  show near-perfect performance for images that are relatively

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simple to process.

#### C. PARAMETER ANALYSIS

Despite the observations of enhancements across all datasets, the proposed approach showed constraints for specific images. In the context of our experimentation, it was observed that there were cases where the accuracy of the results obtained from  $M_E$  was comparatively lower than those obtained from  $M_{CP}$  and  $M_{SD}$ . This section will discuss the external parameters that we used and their effects on the ensemble process.

As stated previously, the scaling factor  $P_{scale}$  is used to reduce the value of Otsu's threshold. This phenomenon improves the ability of  $M_E$  to detect cell clusters showing lower brightness levels, thereby increasing the recall rate of the algorithm. As a result, the proposed approach is expected to improve the detection of a more significant number of cells that also exist in the ground truth. Nonetheless, a lower scale may increase the algorithm's potential for tissue noise. Regions with high background brightness (and a lower contrast difference in the cells) are more responsible for generating false positives.

The usage rates of  $S_{CP}$  and  $S_{SD}$  are determined by the value of  $P_{threshold}$ . Selecting a threshold value that is either excessively high or excessively low may result in the nullification of the proposed approach's effects, as the algorithm will show a preference for one segmentation result over the other. If the value is set too high, the counting results will converge to  $M_{CP}$  results, whereas if it is set too low, the count results will converge to  $M_{SD}$  results.

# **V. CONCLUSION**

Cellpose and Stardist are two of the best state-of-the-art image segmentation models currently available. Although they are capable of meeting the majority of a researcher's requirements, they need help with processing images with densely populated cell clusters and tissue interference. It is often not feasible to provide further training to these models in order to address their errors, as this process can be prohibitively costly due to the need for extensive annotation procedures.

This study shows the effectiveness of ensembling state-ofthe-art models to reduce errors in cell counting assignments where additional training could not be possible. The proposed ensemble approach includes the identification of cellular regions and the subsequent matching of their characteristics with their corresponding models. This guarantees that the outcomes will solely comprise scenarios where each model performs optimally.

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