Counting or Localizing? Evaluating Cell Counting and Detection in Microscopy Images

Luca Ciampi, Fabio Carrara, Giuseppe Amato and Claudio Gennaro

Institute of Information Science and Technologies, National Research Council, Pisa, Italy

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Abstract: Image-based automatic cell counting is an essential yet challenging task, crucial for the diagnosing of many diseases. Current solutions rely on Convolutional Neural Networks and provide astonishing results. However, their performance is often measured only considering counting errors, which can lead to masked mistaken estimations; a low counting error can be obtained with a high but equal number of false positives and false negatives. Consequently, it is hard to determine which solution truly performs best. In this work, we investigate three general counting approaches that have been successfully adopted in the literature for counting several different categories of objects. Through an experimental evaluation over three public collections of microscopy images containing marked cells, we assess not only their counting performance compared to several state-of-the-art methods but also their ability to correctly localize the counted cells. We show that commonly adopted counting metrics do not always agree with the localization performance of the tested models, and thus we suggest integrating the proposed evaluation protocol when developing novel cell counting solutions.

1 INTRODUCTION

Microscopy medical images analysis comprises several challenging Computer Vision problems involving a wide variety of tasks. Among them, cell localization (Lugagne et al., 2019) and counting (Falk et al., 2018) are essential steps for basic research, like disease diagnosis via the evaluation of cell growth kinetics, the estimation of cytotoxicity (i.e., the quality of being toxic to cells) (Kotoura et al., 1985), the quantification of perineuronal nets (Fawcett et al., 2019), the discovery of the role of particular genes in cell biology, microbiology, and immunology (Zhang et al., 2015), and many more. Manual cell counting is still conducted in many laboratories, often with the aid of a hemocytometer and its variants, which has been commonly used due to its low cost and versatility (Johnston, 2010). However, the procedure is time-consuming and error-prone, being subject to inter-user variation depending on the degree of expertise of the analyst (Altman et al., 1993). Therefore, there is a need to count cells automatically to facilitate this tedious and challenging task.

Recently, several vision models (mostly based on Convolutional Neural Networks) have been successfully adopted to count cells and other biological structures from microscopy images. However, the performance of these techniques is often measured only considering the counting errors occurring at inference time (i.e., the difference between the predicted and the actual cell numbers), which often leads to masked mistaken estimations. Indeed, counting errors do not take into account where the cells have been localized in the images and, consequently, counting models might achieve low values of errors while providing wrong predictions (e.g., a high number of false positives and false negatives). Therefore, it is hard to perform a fair comparison between the different state-of-the-art cell counting approaches to determine which performs best.

In this work, we investigate three baseline solutions belonging to the three main counting methodologies — a segmentation-based approach, a localization-based approach, and a count-density estimation approach — that have been successfully exploited for counting several different categories of objects, such as people and vehicles, and that repre-
sent the conceptual basis also for the cell counting techniques. We conduct experiments on three public datasets containing different cell types and characterized by distinct peculiarities. In addition to comparing the performance of investigated methods against state-of-the-art cell counters using established counting evaluation metrics, we also measure the ability of the models to localize the counted cells correctly. Specifically, we adopt two additional metrics: a) the Grid Average Mean absolute Error (GAME) metric, a hybrid metric that simultaneously considers errors in the object count and in their coarse location, and b) the mean Average Precision (mAP), that summarizes the cell precise localization performance. We show that commonly adopted counting metrics (like mean absolute error) do not always agree with the localization performance of the tested models, and thus we suggest measuring both whenever possible to facilitate the practitioner in picking the most suitable solution.

We organize the paper as follows. We review related work in Section 2. In Section 3, we describe the datasets used for our experiments. Section 4 describes the investigated methodologies, while Section 5 outlines the performed experiments and the obtained results. Finally, Section 6 concludes the paper. The code and the trained models are publicly available at https://github.com/ciampluca/counting_perineuronal_nets/tree/visapp-counting-cells.

2 RELATED WORKS

This section reviews some works concerning the counting task in its generality and specifically tailored to estimating the number of cells in microscopy images.

Visual Counting. The goal of the visual counting task is to estimate the number of object instances in still images or video frames (Lempitsky and Zisserman, 2010). Due to its interdisciplinary and widespread applicability to many real-world applications, like calculating the number of people present at an event (Boomninathan et al., 2016), evaluating the number of vehicles in urban scenarios (Ciampi et al., 2021a), or counting animals in ecological surveys (Arteta et al., 2016b), visual counting has recently drawn the attention of researchers. Current solutions address this task as a supervised deep learning-based process. They fall into two main categories: counting by detection (Amato et al., 2019; Amato et al., 2018; Laradji et al., 2018; Ciampi et al., 2018) that requires prior detection or segmentation of the single instances of objects, and counting by regression (Oñoro-Rubio and López-Sastre, 2016; Li et al., 2018; Ciampi et al., 2020; Ciampi et al., 2021b) that instead tries to establish a direct mapping between the image features and the number of objects in the scene, either directly or via the estimation of a density map (i.e., a continuous-valued function). Regression techniques have demonstrated superior performance in crowded scenarios where the objects’ instances are sometimes not well visible due to occlusions and clumps. However, they cannot precisely localize the objects present in the scene, eventually providing only a coarse position of the area in which they are distributed.

Microscope Cell Counting. Because of its paramount importance, several cell counting deep learning-based methods have been proposed in the last years. They belong to both the detection-based and the regression-based approaches, each having the advantages and the drawbacks already discussed above. A relevant example belonging to the former category is (Paulauskaite-Taraseviciene et al., 2019), where authors exploited the popular Mask R-CNN (He et al., 2017) instance segmentation framework to detect overlapping cells. On the other hand, a notable regression-based work is (Aich and Stavness, 2018), where the authors regulated activation maps from the final convolutional layer of the network by exploiting coarse ground-truth activation maps generated from simple dot annotations. Authors in (Xie et al., 2016), instead, introduced a CNN-based regression approach that maps the image features with an associated density map, providing also a coarse localization of the cells by finding its peak values. Another example is represented by (Segui et al., 2015), where the authors proposed a regression-based technique and explored the features that are learned to understand their underlying representation. In (Cohen et al., 2017), another regression-based deep neural network architecture (named Count-ception) is presented, inspired by the Inception family (Szegedy et al., 2015). More, in (Guo et al., 2021), another density-based deep network framework designed to solve the cell counting task is introduced. Specifically, the authors propose SAU-Net, extending the segmentation network U-Net (Ronneberger et al., 2015) with a Self-Attention module. Finally, in (He et al., 2021) the authors exploited auxiliary CNNs to assist the training of the intermediate layers of a density regressor. Hybrid strategies have also been devised to deal with densely concentrated cells but still generating individual cell detections, such as (Falk et al., 2018; Xie et al.,
These approaches first generate intermediate maps that indicate the likelihood of each pixel being the center of a cell in the image. Then, they convert these maps into detections by applying some form of Non-Maximum Suppression (NMS).

Most of these works measure the counting performance by computing the error between the predicted and the actual cell number, hiding potentially mistaken localization. In this work, we consider three general counting approaches on which cell-specific techniques rely, and we also evaluate the quality of the produced detections.

3 DATASETS

In this section, we describe the datasets employed in this work, summarized in Table 1; in particular, we consider three publicly available collections of microscopy images widely used in the context of the cell counting task, presenting different peculiarities and challenges.

3.1 VGG Cells Dataset

The VGG Cells dataset, introduced in (Lempitsky and Zisserman, 2010), comprises 200 RGB highly-realistic synthetic emulations of fluorescence microscopy images of bacterial cells. Images have a fixed size of $256 \times 256 \times 3$ pixels, and the cells are clustered in specific regions and occluded with each other. It is worth noting that the annotation procedure is performed automatically and so labels are free of errors. We show a sample of this dataset in Figure 1.

3.2 MBM Cells Dataset

The Modified Bone Marrow (MBM) Cells has been initially collected by the authors of (Kainz et al., 2015) from 11 RGB microscopy images (having a fixed size of $1200 \times 1200 \times 3$ pixels) of the human bone marrow tissues pertaining to 8 different patients. The marked cells belonging to this dataset have a significant shape variance; furthermore, non-homogeneous tissue background makes their localization more difficult. In a subsequent work (Cohen et al., 2017), the authors divided each image into four patches of $600 \times 600 \times 3$ pixels, for a total of 44 images. A sample of this dataset is reported in Figure 1.

3.3 Nuclei Cells Dataset

This dataset has been presented in (Sirinukunwattana et al., 2016) and comprises 100 RGB microscopy H&E stained histology images of colorectal adenocarcinomas having a common size of $500 \times 500 \times 3$. The images refer to 9 different patients. They have been cropped from non-overlapping areas representing a variety of tissue appearances from normal and malignant regions. Still, they also comprise areas with artifacts, over-staining, and failed autofocussing to simulate realistic outliers. Another peculiarity of this dataset is that the nuclei of the cells belong to four different categories, presenting different visual characteristics; some experts have manually annotated them by putting a dot over the centroids of each biological structure for a total of 29,756 nuclei marked. In the following, we refer to this dataset as Nuclei Cells dataset. We report a sample of this dataset in Figure 1.

4 METHOD

We assume to have a labeled collection of $N$ microscopy images $X = \{(I_1, L_1), \ldots, (I_N, L_N)\}$, where $L_i$ is the set of 2D-point annotations associated to the $i$-th image $I_i$. Each image has been manually annotated by a human expert, and the annotations are in the form of dots, i.e., coordinates localizing the centroids of the cells present in the region of interest, as is usually the case in the counting task.

We define a localization model $f_0$ as a Deep Learning-based algorithm that takes as input an image $I$ and produces as output an associated set of coordinates $L = \{p_1, \ldots, p_C | p_j \in \mathbb{R}^2 \}$ localizing the centroids of the cells to be counted. This model is trained using location data $X$ and can be implemented following several different strategies; here, we test three successful approaches from the literature, that are segmentation, detection, and density estimation, described below.

4.1 Foreground/Background Segmentation

Proposed by (Falk et al., 2018), in this approach we locate cells on the basis of a binary segmentation map $S \in \{0, 1\}^{H \times W}$ where ones represent pixels of objects of interest, while zeros are considered background. Each connected component in the segmentation map represents a single object; the positions of the objects $L$ are set to the coordinate of the centroids of the connected components. As the implementation of the model $f_0$, we adopt the original U-Net architecture (Ronneberger et al., 2015) commonly used in segmentation tasks. The model is trained to produce a real-valued segmentation map $S = f_0(I) \in [0, 1]^{H \times W}$.
Table 1: Summary of datasets. We show the different peculiarities that characterize the three datasets exploited in this work.

| Dataset                  | N.Img | Size       | N.Objs   | Objs/Img |
|--------------------------|-------|------------|----------|----------|
| VGG (Lempitsky and Zisserman, 2010) | 200   | 256×256    | 35,192   | 176 ± 61 |
| MBM (Kainz et al., 2015; Cohen et al., 2017) | 44    | 600×600    | 5,553    | 126 ± 33 |
| Nuclei (Sirinukunwattana et al., 2016)   | 100   | 500×500    | 29,756   | 297 ± 218 |

Figure 1: Samples and Targets. We show a dataset sample (1st column) and the corresponding targets used when training i) the detection-based method FRCNN (2nd column), ii) the density-based method D-CSRNet (3rd column), and iii) the segmentation-based method S-UNet (4th column).

that is then thresholded to obtain $S$. The target segmentation maps are generated drawing discs at the annotated positions and carefully separating overlapping discs with a background ridge (see the fourth column of Figure 1 for examples of targets). We minimize the weighted binary cross-entropy between pixels of the output and target maps as specified in (Falk et al., 2018); more important pixels (near ridges and foreground objects) are given an increased weight in the total loss computation. We will refer to this approach as S-UNet.

4.2 Bounding Box Regression

For this approach, we employ the standard Faster-RCNN detector (Ren et al., 2017). This deep neural network takes images as input and produces a list of bounding boxes localizing the objects as output. The detection pipeline follows the two-stage paradigm. In the first stage, the network generates a bunch of region proposals likely to contain objects, exploiting a set of anchors (i.e., pre-defined boxes) that are sliced over the image; in the second stage, these priors are refined and, for each of them, a score is assigned expressing the likelihood to really containing the object. We consider the centers of the final boxes as the lo-
calization of the entities we want to consider. We produce the targets by generating squared bounding boxes centered in the dot-annotated data and having fixed sides, again, depending on the typical object size in the dataset. A sample of a target is shown in the second column of Figure 1. We implement \( f_0 \) as a Faster-RCNN network with a Feature Pyramid Network module and a ResNet-50 backbone. From now on, we will refer to this method as FRCNN.

### 4.3 Density Estimation

We also account for density-estimation approaches that have shown superior counting performances in very “crowded” scenarios. In this case, the goal is to learn a regression between the features of an input image having height \( H \) and width \( W \) to a density map \( D = f_0(I) \in \mathbb{R}^{H \times W} \). The notion of density map is close to the physical/mathematical notion of density; specifically, each pixel of \( D \) corresponds to the quantity of the objects present at that precise location. The number of the objects \( n \) present in an image sub-region \( P \subseteq I \) is estimated by summing up pixel values in the region of interest, i.e., \( n = \sum_{p \in P} D_p \). Although these approaches are not suited for precisely localize objects, a coarse localization can be obtained by analyzing the estimated density map, in particular by finding the top-\( n \) maximum local peaks of it, as already done in (Xie et al., 2016). We train the model by minimizing the mean squared error loss between target and predicted density maps. Following previous works, we generate the target density maps by superimposing Gaussian kernels \( G_\sigma \) centered in the dot-annotated locations; the spread parameter \( \sigma \) is fixed, and it has been estimated depending on the typical object size in the considered dataset. We show an example of a target density map in the third column of Figure 1. We implement \( f_0 \) exploiting the Congested Scene Recognition Network (CSRNet), proposed in (Li et al., 2018), a CNN for accurate density estimation of congested scenes, comprising two major components. Specifically, it uses a modified version of the popular VGG-16 network (Simonyan and Zisserman, 2015) to extract the image features; stacked upon this, the authors built a back-end composed of dilated convolutional (Yu and Koltun, 2016) layers to extract deeper information of saliency and, at the same time, maintain the output resolution. We will refer to this method as D-CSRNet.

### 5 EXPERIMENTS AND RESULTS

In this section, we describe the experiments performed to validate our approach and discuss the obtained results. First, we evaluate the three adopted general counting solutions, i.e., the segmentation-based S-UNet, the detection-based FRCNN, and the density-based D-CSRNet approaches, over the three standard cell counting benchmarks described above to verify that the obtained counting errors are comparable with the ones provided by state-of-the-art cell-specific counting methods. Then, we perform additional experiments evaluating the quality of the localization of the cells, an aspect that is not taken into account by counting metrics.

#### 5.1 Comparison with the State-of-the-Art

We evaluate the three adopted counting methodologies over the VGG Cells, the MBM Cells, and the Nuclei Cells counting benchmarks described in Section 3, and we compare their performances with other state-of-the-art approaches. For the VGG Cells and the MBM Cells datasets, we follow the evaluation protocol introduced by (Lempitsky and Zisserman, 2010) and adopted by most subsequent works. Specifically, we consider a testing subset fixed for all the experiments (100 and 10 images for VGG Cells and MBM Cells, respectively) and training and validation subsets of varying size (\( N \) images for each subset) to simulate lower or higher numbers of labeled examples. This evaluation protocol simulates the real scenario in which scientists often have a significant variance regarding the number of available microscopy images. Following previous work, we set \( N \) to 16, 32, and 50 for VGG Cells and to 5, 10, 15 for MBM Cells. Concerning the Nuclei Cells dataset, we instead use two-fold cross-validation, with 50 images for testing, according to (Sirinukunwattana et al., 2016) and subsequent works. Following standard counting benchmarks, we use the Mean Absolute Error (MAE) to measure the counting performance. Specifically, it is defined as:

\[
\text{MAE} = \frac{1}{N} \sum_{n=1}^{N} |c_{gt}^n - c_{pred}^n|,
\]

where \( N \) is the number of test images, \( c_{gt}^n \) is the actual count (i.e., the ground truth), and \( c_{pred}^n \) is the predicted count of the \( n \)-th image. For the VGG Cells and the MBM Cells, we repeat the experiment 10 times, randomly sampling ten different splits for each configuration, and we report the mean and standard
deviation of the MAE computed between the different runs. On the other hand, concerning the Nuclei dataset, we report the mean and the standard deviation of the MAE calculated between the 100 images comprising the two test splits.

Table 2 reports the obtained results. The density-based solution performs best among the VGG Cells dataset and, more strongly, with the Nuclei Cells dataset, comparatively to the state of the art. The other two adopted methods, i.e., the segmentation-based S-UNet and the detection-based FRCNN, show larger errors, according to their intrinsic limitations when employed in highly “crowded” scenarios with occluded objects like the VGG Cells dataset and, especially, the Nuclei Cells dataset. On the other hand, considering the MBM Cells dataset, characterized by challenges more related to the object shape variations, all the approaches show competitive results, in some cases also outperforming state-of-the-art solutions.

5.2 Localization Analysis

Although the MAE is a fair metric for establishing a comparative in terms of counting, it can often lead to masking erroneous estimations. The reason is that the MAE does not take into account where the estimations have been done in the images. In other words, the MAE does not capture localization errors; models might achieve low values of MAE while providing wrong predictions (e.g., a high number of false positives and false negatives in detection-based techniques, or a bad allocation of density values in predicted maps of density-based methods). Hence, picking up the best counting model basing the decision only on the MAE metric can lead to blunders.

In this section, we conduct experiments to assess the ability of the three adopted solutions to localize the counted cells correctly. Specifically, we consider two additional metrics described in the following paragraphs.

**Grid Average Mean absolute Error (GAME).** (Guerrero-Gómez-Olmedo et al., 2015) is a hybrid metric that simultaneously considers the object count and the estimated locations of the cells. Specifically, it is computed by sub-dividing the image in $4^2$ non-overlapping regions and summing the MAE computed in each of these sub-regions. Formally:

$$GAME(L) = \frac{1}{N} \sum_{n=1}^{N} \left( \sum_{l=1}^{4^2} |c_{gt}^l - c_{pred}^l| \right),$$  

where $N$ is the total number of test images, $c_{pred}^l$ is the estimated count in a region $l$ of the n-th image, and $c_{gt}^l$ is the ground truth for the same region in the same image. The higher $L$, the more restrictive the GAME metric will be. Note that the MAE can be obtained as a particular case of the GAME when $L = 0$.

**Mean Average Precision (mAP).** is an established metric for the localization performances of object detectors. We compute the average precision for an image as follows. i) We assign a score to each detected cell in the image. Detection scores are obtained differently for the three tested methods. For the S-UNet model, the detection score of an object is set to the value of the predicted segmentation map $\hat{S}$ at the location of the centroid of the corresponding connected component. For D-CSRNNet, the location of an object is set to a local peak in the predicted density map, and its score is set to the value at its location. For FRCNN, the detection score is already part of the output of the Faster-RCNN model. ii) We filter out weak detections using a threshold on the detection scores. iii) We match the filtered detections with the ground-truth object positions using the Hungarian algorithm with a constraint on the maximum accepted displacement in pixels between predicted and real locations; once matches are found, we obtain the number of true positives (matched detection and ground-truth pairs), false positives (unmatched detections), and false negatives (unmatched ground-truth positions) locations. iv) We repeat these steps for several threshold values to obtain the precision-recall curve and the average precision (i.e., the area under the curve).

In Table 3, we report the MAE, the GAME, and the mAP metrics for all the tested solutions and the adopted datasets. Here, we consider only the splits having $N$ to 50 and 15 for the VGG and the MBM datasets, respectively, and the same two-fold cross-validation with 50 images for testing concerning the Nuclei dataset. Note that the density-based solution D-CSRNNet shines in the Nuclei benchmark where very dense regions of overlapped cells are common and strain non-density solutions, obtaining the best counting metrics (MAE, GAME) among the tested models. However, the denser the cells in the benchmark, the less the density-based solution can recover the exact locations of the counted cells, thus achieving lower mAP values. On the other hand, the detection-based solution FRCNN performs sufficiently well only when counting cells in the less crowded MBM and VGG benchmarks. Still, it is able to recover the exact position of more counted cells, as can be seen from the higher mAP values obtained. Last, the segmentation-based model sits in the middle of these two extremes, providing intermediate counting and localization performance.
Table 2: Comparison on Standard Benchmarks. For VGG and MBM datasets, we vary the training and validation subsets (\(N\) images for each subset), repeating the experiments 10 times. For Nuclei, we perform 2-fold cross-validation (\(N = 50\) images per fold). Mean±st.dev. of MAE is reported.

**VGG Cells** (Lempitsky and Zisserman, 2010). (200 images in total - 100 test images)

| Method                  | \(N = 16\) | \(N = 32\) | \(N = 50\) |
|-------------------------|------------|------------|------------|
| (Arteta et al., 2016a)  | N/A        | 5.06 ± 0.2 | N/A        |
| GMN (Lu et al., 2019)   | N/A        | 3.6 ± 0.3  | N/A        |
| (Lempitsky and Zisserman, 2010) | 3.8 ± 0.2 | 3.5 ± 0.2  | N/A        |
| VGG-GAP-HR (Aich and Stavness, 2018) * | N/A | 2.95** | 2.67 |
| SAU-Net (Guo et al., 2021) | N/A | N/A | 2.6 ± 0.4 † |
| FCRN-A (Xie et al., 2016) | 3.4 ± 0.2 | 2.9 ± 0.2  | 2.9 ± 0.2² |
| Count-Ception (Cohen et al., 2017) | 2.9 ± 0.5 | 2.4 ± 0.4  | 2.3 ± 0.4  |
| CCF (Jiang and Yu, 2020) | 2.8 ± 0.1 | 2.6 ± 0.1  | 2.6 ± 0.1  |
| C-FCRN+Aux (He et al., 2021) | N/A | N/A | 2.3 ± 2.2 * |
| S-UNet (Falk et al., 2018) | 8.3 ± 2.3 | 5.6 ± 1.1  | 4.5 ± 0.5  |
| D-CSRNet (Li et al., 2018) | 4.0 ± 0.2 | 3.2 ± 0.2  | 3.0 ± 0.1  |
| FRCNN (Ren et al., 2017) | 9.3 ± 0.7 | 8.2 ± 0.6  | 7.4 ± 1.0  |

* They did not report standard deviation. ** They used a validation subset of 100 – \(N\) images. † They did not use a test subset, but only a 100 – \(N\) images validation subset. ‡ They used a 5-fold cross validation-based evaluation protocol considering the whole dataset.

**MBM Cells** (Kainz et al., 2015; Cohen et al., 2017). (44 images in total - 10 test images)

| Method                  | \(N = 5\) | \(N = 10\) | \(N = 15\) |
|-------------------------|------------|------------|------------|
| (Xie et al., 2018)      | 28.9 ± 22.6| 22.2 ± 11.6| 21.3 ± 9.4 |
| FCRN-A (Xie et al., 2016) | 23.6 ± 4.6 | 21.5 ± 4.2 | 20.5 ± 3.5 |
| (Marsden et al., 2018)  | 12.6 ± 3.0 | 10.7 ± 2.5 | 8.8 ± 2.3  |
| CCF (Jiang and Yu, 2020) | 9.3 ± 1.4  | 8.9 ± 0.9  | 8.6 ± 0.3  |
| C-FCRN+Aux (He et al., 2021) | 6.5 ± 5.2 * | 6.5 ± 5.2 ** | N/A |
| SAU-Net (Guo et al., 2021) | N/A | N/A | 5.7 ± 1.2 † |
| S-UNet (Falk et al., 2018) | 9.0 ± 1.9  | 7.0 ± 1.6  | 6.7 ± 2.5  |
| D-CSRNet (Li et al., 2018) | 10.8 ± 2.5 | 8.0 ± 1.3  | 7.0 ± 1.3  |
| FRCNN (Ren et al., 2017) | 8.8 ± 1.4  | 9.9 ± 1.5  | 8.3 ± 1.9  |

* They used 14 test images. ** They used a 5-fold cross validation-based evaluation protocol considering the whole dataset. † They did not use a test subset, but only a 44 – \(N\) images validation subset. § They used a train/test split of 8/3 using full-size images.

**Nuclei Cells** (Sirinukunwattana et al., 2016). (100 images in total - 50 test images)

| Method                  | \(N = 50\) |
|-------------------------|------------|
| DeepFeat (Segui et al., 2015) | 71.8 ± 51.4|
| (Lempitsky and Zisserman, 2010) | 51.4 ± 39.8|
| StructRegNet (Xie et al., 2018) | 45.9 ± 47.9|
| FCRN-A (Xie et al., 2016) | 42.5 ± 33.5|
| Count-Ception (Cohen et al., 2017) | 34.1 ± 29.0|
| C-FCRN+Aux (He et al., 2021) | 29.3 ± 25.4|
| S-UNet (Falk et al., 2018) | 62.4 ± 55.4|
| D-CSRNet (Li et al., 2018) | 37.3 ± 41.0|
| FRCNN (Ren et al., 2017) | 96.5 ± 128.0|
Table 3: Counting and Localization Performance. The MAE measures global counting performance independently of localization. The mAP summarizes localization performances in terms of precision and recall of localized cells. The GAME(L) measure counting performance while being aware of the location of cells; the higher $L$, the more localization errors are penalized. Regarding the VGG and the MBM datasets, we consider the splits having $N$ to 50 and 15, respectively.

| Method    | MAE ↓ | GAME(L) ↓ | L = 1 | L = 2 | L = 3 | L = 4 | mAP (%) ↑ |
|-----------|-------|-----------|-------|-------|-------|-------|-----------|
| VGG Cells |       |           |       |       |       |       |           |
| S-UNet    | 4.5 ± 0.5 | 7.7 ± 1.3 | 12.8 ± 1.5 | 21.6 ± 2.4 | 38.0 ± 4.1 | 75.3 ± 15.8 |
| D-CSRNet  | 3.0 ± 0.1 | 6.5 ± 0.2 | 11.3 ± 0.4 | 18.9 ± 0.6 | 28.7 ± 1.0 | 43.2 ± 1.6 |
| FRCNN     | 7.4 ± 1.0 | 11.1 ± 0.9 | 18.3 ± 1.3 | 29.7 ± 2.0 | 43.3 ± 3.2 | 93.3 ± 0.6 |
| MBM Cells |       |           |       |       |       |       |           |
| S-UNet    | 6.7 ± 2.5 | 10.4 ± 2.5 | 17.3 ± 1.9 | 27.6 ± 2.0 | 40.9 ± 3.1 | 53.5 ± 5.3 |
| D-CSRNet  | 7.0 ± 1.3 | 10.8 ± 1.2 | 16.7 ± 1.3 | 27.0 ± 1.6 | 41.5 ± 2.2 | 67.9 ± 1.2 |
| FRCNN     | 8.3 ± 1.9 | 12.7 ± 2.4 | 20.4 ± 3.9 | 32.5 ± 4.7 | 47.2 ± 8.7 | 87.4 ± 1.9 |
| Nuclei Cells |       |           |       |       |       |       |           |
| S-UNet    | 62.4 ± 55.4 | 66.9 ± 51.7 | 75.1 ± 50.6 | 95.3 ± 54.1 | 138.4 ± 75.2 | 66.8 ± 11.7 |
| D-CSRNet  | 37.3 ± 41.0 | 45.7 ± 38.8 | 58.2 ± 38.5 | 77.6 ± 39.8 | 100.5 ± 45.0 | 27.7 ± 8.5 |
| FRCNN     | 96.5 ± 128.0 | 103.8 ± 125.3 | 112.6 ± 121.9 | 133.9 ± 118.7 | 168.2 ± 123.2 | 57.9 ± 10.8 |

Finally, in Figure 2, we show some examples of predictions with very low absolute counting errors (suggesting good performance) but in which the average precision metric indicates erroneous predictions instead. Note that in the detection-based solution FRCNN, the disagreement between the two metrics is less pronounced, as this methodology is usually adopted to optimize AP. Thus, we suggest integrating the mean average precision, or at least a GAME-L metric with a high-enough L, when optimizing and evaluating novel cell counting solutions. We deem the additional evaluation protocol would help practitioners to better characterize the performance of developed solutions.

6 CONCLUSIONS

In this work, we consider the cell counting task in microscopy images, investigating the ability of three general counting methodologies not only in estimating the number of the biological structures but also in localizing them. Indeed, most state-of-the-art solutions tailored to cell counting are evaluated merely considering the difference between the predicted and the actual number of the cells, skipping a further analysis focused on the quality of the provided estimations. We show that relying only on the counting metrics can lead to models producing incorrect cell localization. We performed experiments on three cell counting benchmarks, and we assessed that counting errors do not always agree with the localization performance. Thus, we suggest measuring and reporting also the mean average precision (or at least a grid average mean absolute error) whenever possible to help practitioners developing better models and to guide users to choose the model most tailored to their scenario.

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Figure 2: Absolute Error (AE) can be misleading. For each considered model (one per column), we show predictions obtaining a low AE, but also a low Average Precision (AP) due to high numbers of false positive and false negatives. The AP can discern cases where the MAE fails to capture poor model outputs. We indicate false positives in purple, false negatives in cyan, and true positives in green, with the corresponding ground-truth position drawn in red and connected via a thin yellow line. (Best viewed in electronic format.)
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