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Microscopy cell counting and detection with fully convolutional regression networks

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ABSTRACT

This paper concerns automated cell counting and detection in microscopy images. The approach we take is to use convolutional neural networks (CNNs) to regress a *cell spatial density map* across the image. This is applicable to situations where traditional single-cell segmentation-based methods do not work well due to cell clumping or overlaps. We make the following contributions: (i) we develop and compare architectures for two fully convolutional regression networks (FCRNs) for this task; (ii) since the networks are fully convolutional, they can predict a density map for an input image of arbitrary size, and we exploit this to improve efficiency by end-to-end training on image patches; (iii) we show that FCRNs trained entirely on synthetic data are able to give excellent predictions on *microscopy images from real biological experiments* without fine-tuning, and that the performance can be further improved by fine-tuning on these real images. Finally, (iv) by inverting feature representations, we show to what extent the information from an input image has been encoded by feature responses in different layers. We set a new state-of-the-art performance for cell counting on standard synthetic image benchmarks and show that the FCRNs trained entirely with synthetic data can generalise well to real microscopy images both for cell counting and detections for the case of overlapping cells.

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Microscopy image analysis; cell counting; cell detection; fully convolutional regression networks; inverting feature representations

1. Introduction

Counting and detecting objects in crowded images or videos is an extremely tedious and time-consuming task encountered in many real-world applications, including biology (Arteta et al. 2012, 2014, 2015; Fiaschi et al. 2012), surveillance (Chan et al. 2008; Lempitsky & Zisserman 2010) and other applications (Barinova et al. 2012). In this paper, we focus on cell counting and detection in microscopy, but the developed methodology could equally be used in other counting or detection applications. Numerous procedures in biology and medicine require cell counting and detection, for instance: a patient's health can be inferred from the number of red blood cells and white blood cells; in clinical pathology, cell counts from images can be used for investigating hypotheses about developmental or pathological processes; and cell concentration is important in molecular biology, where it can be used to adjust the amount of chemicals to be applied in an experiment. While detection on its own, is able to determine the presence (and quantity) of an object of interest, such as cancer cells in a pathology image, furthermore, detection can be used as seeds for further segmentation or tracking.

Automatic cell counting can be approached from two directions, one is detection-based counting (Arteta et al. 2012, 2015; Girshick et al. 2014), which requires prior detection or segmentation; the other is based on density estimation without the need for prior object detection or segmentation (Lempitsky & Zisserman 2010; Fiaschi et al. 2012; Arteta et al. 2014). In our work, we take the latter approach, and show that cell detection can be a side benefit of the cell counting task. Following (Lempitsky & Zisserman 2010), we first cast the cell counting problem as a supervised learning problem that tries to learn a mapping between an image I(x) and a density map D(x), denoted as $F : I(x) \rightarrow D(x)$ ($I \in R^{m \times n}, D \in R^{m \times n}$) for a $m \times n$ pixel image, see Figure 1. During the inference, given the input test image, the density map and cell detections can be obtained, as shown in Figure 2.

We solve this mapping problem by adapting the convolutional neural networks (CNNs) (LeCun et al. 1998; Krizhevsky et al. 2012), which has re-emerged as a mainstream tool in the computer vision community. CNNs are also starting to become popular in biomedical image analysis and have achieved stateof-the-art performance in several areas, such as mitosis detection (Cireşan et al. 2013), neuronal membranes segmentation (Cireşan et al. 2012), analysis of developing C. elegans embryos (Ning et al. 2005), and cell segmentation (Ronneberger et al. 2015). However, they have not yet been applied to solve the target problem here of regression in microscopy cell for counting and detection simultaneously.

One of the issues we investigate is whether networks trained only on synthetic data can generalise to real microscopy images. (Jaderberg et al. 2014) showed that for text recognition a CNN trained only on synthetic data gave excellent results on real images. The great advantage of this is that it avoids the problem of obtaining large data-sets with manual annotation. These are available for natural images, e.g. ImageNet (Russakovsky et al. 2014) in the computer vision field, but *the* biomedical image data is limited, expensive, and time-consuming to annotate.



Figure 1. The training process aims to find a mapping between I(x) and the density map D(x). (a) Red crosses on I(x) are dot annotations near the cell centres. (b) The density map D(x) is a superposition of Gaussians at the position of each dot. Integration of the density map D(x) over specific region gives the count of cells.

In this paper, we develop a fully convolutional regression networks (FCRNs) approach for regression of a density map. In Section 2, we describe several related works. In Section 3, we design and compare two alternative architectures for the FCRNs, and discuss how the networks can be trained efficiently with images of arbitrary sizes in an end-to-end way. In Section 4, we present results on a standard synthetic data-set for counting, and show that the networks trained only on synthetic data can generalise for different kinds of microscopy images from real biological experiments, and the performance can be further improved by fine-tuning parameters with annotated real data. Overall, experimental results show that FCRNs can provide stateof-the-art cell counting for a standard synthetic data-set, as well as the capability for cell detection. And as an extension to our previous paper, which was published in the MICCAI 1st Deep Learning Workshop (Weidi et al. 2015), we also propose to visualise to what extent the information from input image has been encoded by feature responses of different layers in the trained networks.

2. Related work

We first review previous approaches to cell counting by density estimation, and then turn to CNN-based methods for cell detection. We also build on fully convolutional networks for semantic segmentation (Long et al. 2015), where the fully connected layers of a classification net are treated as convolutions, and upsampling filters combined with several skip layers are used to enable the network to take an input of arbitrary size and produce an output of the same size during training and inference.

2.1. Counting by density estimation

Cell counting in crowded microscopy images with density estimation avoids the difficult detection and segmentation of individual cells. It is a good alternative for tasks where only the number of cells *in an arbitrary region is required*. Over the recent years, several works have investigated this approach. In Lempitsky and Zisserman (2010), the problem was cast as density estimation with a supervised learning algorithm, $D(x) = c^T \phi(x)$, where D(x) represents the ground-truth density map, and $\phi(x)$ represents the local features. The parameters *c* are learned by minimising the error between the true and predicted density map with quadratic programming over all possible subwindows. In Fiaschi et al. (2012), a regression forest is used to exploit the patch-based idea to learn structured labels, then for a new input image, the density map is estimated by averaging over structured, patch-based predictions. In Arteta et al. (2014), an algorithm was proposed that allows fast interactive counting by simply solving ridge regression with various local features.

2.2. Detection by regression

Although there has been much recent work on detection in *natural* images, there has been little application so far to *microscopy* images. Approaches on natural images include detections based on region proposal and classification networks (Girshick et al. 2014; He et al. 2014; Ren et al. 2015), sliding window and classification networks (Sermanet et al. 2014), and using modes from heat map regression (Tompson et al. 2014; Pfister et al. 2015).

One work that has been developed independently and shares similar ideas to our own on detection is Yuanpu et al. (2015). In their work, they cast the detection task as a structured regression problem with the dot annotation near the cell centre. They train CNN model that takes an image patch of fixed size as input and predicts a "proximity patch" of half the resolution of the original input patch. During training, the defined proximity mask \mathcal{M} corresponding to image *I* is calculated as,

$$\mathcal{M}_{ij} = \begin{cases} \frac{1}{1+\alpha D(i,j)} & \text{if } D(i,j) \le \gamma, \\ 0 & \text{otherwise,} \end{cases}$$
(1)



(a) Test Image: I'(x)

(b) Estimated Density Map: D'(x)



Figure 2. During the inference process, given the test image l'(x) in (a). (b) The trained model aims to predict the density map D'(x). The integration of the density map D'(x) over a specific region gives the cell counts. (c) Cell detections in T'(x) can be obtained by taking local maxima on the density map D'(x). (*Yellow crosses*).

where D(i, j) represents the Euclidean distance from pixel (i, j) to the nearest manually annotated cell centre ($\alpha = 0.8$ and $\gamma = 5$ in their paper). Therefore, \mathcal{M}_{ij} gives value 1 for the cell centre, and decreases with distance from the centre. During inference, in order to calculate the proximity map for an entire testing image, they propose to fuse all the generated proximity patches together in a sliding window way. After this, the cell detection is obtained by finding the local maximum positions in this average proximity map.

In contrast to this approach, our paper focuses on models that enable end-to-end training and prediction of density maps for images of arbitrary size using FCRNs. Cell counting and detection in the specific region of microscopy images can then be obtained simultaneously from the predicted density map.

3. Fully convolutional regression networks

3.1. Architecture design

The problem scenario of cell counting and detection is illustrated in Figures 1 and 2. For training, the ground truth is provided by dot annotations, where each is represented by a Gaussian, and a density map D(x) is formed by the superposition of these Gaussians. The central task is to regress this density map from the corresponding cell image I(x), then the cell count in a specific region can be obtained by integrating over D(x) and cell detection by local maxima detection on D(x).

In this paper, we propose to solve this problem by training FCRNs. We present two network architectures, namely FCRN-A and FCRN-B, as shown in Figure 3. In designing the network architectures, we consider two points: (i) for cell counting and detection problems, cells are usually small compared to the whole image. Therefore, deep networks that can represent highly semantic information are not necessary; and (ii), based on this, we consider only simple architectures (no skip layers). However, since cell clumps can have very complicated shapes, we are interested in finding out if networks of this simplicity are able to deal with these complications and variety.

The popular CNN architecture for classification contains convolution–ReLU–pooling (Krizhevsky et al. 2012). ReLU refers

to rectified linear units. Pooling usually refers to max pooling and results in a shrinkage of the feature maps. However, in order to produce density maps that have same resolution as the input, we reinterpret the fully connected layers as convolutional layers and undo the spatial reduction by performing upsampling-convolution-ReLU, mapping the feature maps of dense representation back to the original resolution (Figure 3). During upsampling, we use bilinear interpolation, followed by trainable convolution kernels that can be learnt during end-toend training.

Inspired by the very deep VGG-net (Simonyan & Zisserman 2015), in both regression networks, we only use small kernels of size 3×3 or 5×5 pixels. The number of feature maps in the higher layers is increased to compensate for the loss of spatial information caused by max pooling. In FCRN-A, all of the kernels are of size 3×3 pixels, and three max-poolings are used to aggregate spatial information leading to an effective receptive field of size 38×38 pixels (i.e. the input footprint corresponding to each pixel in the output). FCRN-A provides an efficient way to increase the receptive field, while contains about 1.3 million trainable parameters. In contrast, max pooling is used after every two convolutional layers to avoid too much spatial information loss in FCRN-B. In this case, the number of feature maps is increased up to 256, with this number of feature maps then retained for the remaining layers. Comparing with FCRN-A, in FCRN-B we train 5×5 upsampling kernels leading to the effective receptive field of size 32×32 pixels. In total, FCRN-B contains about 3.6 million trainable parameters, which is about three times as many as those in FCRN-A.

3.2. Implementation details

The implementation is based on MatConvNet (Vedaldi & Lenc 2015). During training, we cut large images into patches, for instance, we randomly sample patches of size 100×100 pixels from 256×256 images. Simple data augmentation techniques are also used, e.g. small rotations, horizontal flipping. Before training, each patch is normalised by subtracting its own mean value and then dividing by the standard deviation.



(a) Fully Convolutional Regression Network - A (FCRN-A)



(b) Fully Convolutional Regression Network - B (FCRN-B)

Figure 3. FCRNs in this paper (FCRN-A & FCRN-B). (a) FCRN-A is designed to use small 3×3 kernels for every layer. Each convolutional layer is followed by pooling. (b) FCRN-B is designed to use fewer pooling layers than FCRN-A, 5×5 kernels are used. In each FCRN architecture: (1) The *size* of the input image or feature maps is shown on top of each block, indicating whether pooling has been used. (2) The *number of feature maps* in each layer is shown at the bottom of each block. (3) The *size* of kernels is shown beside the small black or red blocks. Conv – convolution; Pooling – 2×2 max pooling; ReLU – rectified linear units; Upsampling – bilinear upsampling.

The cost function is defined as:

$$I(W; X_0) = \frac{1}{M} \sum_{i=1}^{M} (Y_n - X_n^{(i)})^T (Y_n - X_n^{(i)}) \quad \text{(Mean Square Error)} \quad (2)$$

where *W* are all the trainable parameters, X_0 is the input patch, *Y* is the ground-truth annotation with Gaussians of $\sigma = 2$ and X_n is the predicted density map for the input patch.

The parameters of the convolution kernels are initialised with an orthogonal basis (Saxe et al. 2014). Stochastic gradient descent with momentum are used for optimisation. Then, the parameters *w* are updated by:

$$\Delta w_{t+1} = \beta \Delta w_t + (1 - \beta) \left(\alpha \frac{\partial I}{\partial w} \right) \quad \text{(Include Momentum)} \quad \text{(3)}$$

where β is the momentum parameter. The learning rate α is initialised as 0.01 and gradually decreased by a factor of 10.

The momentum is set to 0.9, weight decay is 0.0005 and no dropout is used in either network. Since the non-zero region in the ground-truth density map is really small, most of the pixels in ground-truth density map remains to be zero. Moreover, even for non-zero regions, the peak value of a Gaussian with $\sigma = 2$ is only about 0.07, the networks tend to be very difficult to train. To alleviate this problem, we simply scale the Gaussian-annotated ground truth (Figure 1(b)) by a factor of 100, forcing the network to fit the Gaussian shapes rather than background zeros.

After pretraining with patches, we fine-tune the parameters with whole images to smooth the estimated density map, since the 100 \times 100 image patches sometimes may only contain part of a cell on the boundary.

4. Experimental validation

In this section, we first determine how FCRN-A and FCRN-B are compared with previous work on cell counting using synthetic

data. Then, we apply the network trained only on synthetic data to a variety of real microscopy images without fine-tuning. Finally, we compare the performance before and after fine-tuning on real microscopy images.

In terms of cell detection, we consider it as a side benefit of our main counting task. The detection results are obtained simply by taking local maxima based on our predicted density map. We show detection results both on synthetic data and *microscopy images from real biological experiments*.

4.1. Data-set and evaluation protocol

4.1.1. Synthetic data

The synthetic data-set (Lempitsky & Zisserman 2010) consists of 200 images of cell nuclei on fluorescence microscopy generated with (Lehmussola et al. 2007). Each synthetic image has an average of 174 ± 64 cells. Severe overlap between instances are often observed in this data-set, which makes it challenging for counting or detection. As shown in Figure 4, under this situation, it even becomes impossible for a human expert to tell the difference between overlapping cells and a single cell. The synthetic data-set is divided into 100 images for training and 100 for testing, and several random splits of the training set are used. Such splits consist of five sets of N training images and N validation images, for N = 8, 16, 32, 64. We report the mean absolute errors and standard deviations for FCRN-A and FCRN-B.

4.1.2. Real data

We evaluated FCRN-A and FCRN-B on four different kinds of data; (1) retinal pigment epithelial (RPE) cell images. The quantitative anatomy of RPE can be important for physiology and pathophysiology of the visual process, especially in evaluating the effects of aging (Panda-Jonas et al. 1996); (2) embryonic stem cells. Cell counting is essential to monitor the differentiation process (Faustino et al. 2009); (3) plasma cell. The relative number of plasma cells in a bone marrow specimen is a clinical parameter important for the diagnosis and management of plasma cell dyscrasia (Went et al. 2006); (4) images of precursor T-Cell lymphoblastic lymphoma. Lymphoma is the most common blood cancer, usually occurs when cells of the immune system grow and multiply uncontrollably.

4.2. Evaluation on synthetic data

4.2.1. Network comparison

During testing, each image is mapped to a density map first, then integrating over the map for a specific region gives the count, or taking local maxima gives the cell detection of that region (Figure 5). The performances of the two networks for cell counting are compared in Table 1 as a function of the number of training images.

As shown in Table 1, FCRN-A performs slightly better than FCRN-B. The *size* of the receptive field turns out to be more important than being able to provide more detailed information *over* the receptive field, we hypothesis that this is because the real difficulty in cell counting lies in regression for large cell clumps, and a larger receptive field is required to span these. For both networks, the performance is observed to improve using

more training images from N = 8 to N = 32, and only a small additional increase for N to 64.

There are three key sources of error: first, from the dataset itself. As shown in Figure 4, the annotation for the data-set itself is noisy. In this case, the L2 regression loss tends to overpenalise. In future research, we will investigate other regression loss functions to address this; second, from the boundary effect due to bilinear up-sampling. Cells on the boundary of images tend to produce wrong predictions in this case; and the third source of error is from very large cell clumps, where four or more cells overlap. In this case, larger clumps can be more variable in shape than individual cells and so are harder to regress; further, regression for large cell clumps requires the network to have an even larger receptive field that can cover important parts of the entire clumps, like concavity information, or curved edges in specific directions. Since our networks are relatively shallow and only have a receptive field of size 38×38 pixels and 32×32 pixels, for elongated cell clumps, their curved edges can usually be covered, and correct predictions can be expected. However, for a roughly round cell clump with four or more cells, it can be bigger than our largest receptive field, and this usually leads to an incorrect prediction.

4.2.2. Comparison with state-of-the-art

Table 1 shows a comparison with previous methods on the synthetic cell data-set. FCRN-A shows about 9.4% improvement over the previous best method of Fiaschi et al. (2012) when N = 32.

4.3. Evaluation on real data

We test both regression networks on real data-sets for counting and detection. Here, we only show figures for results from FCRN-A in Figures 6, 7, 8 (without fine-tuning) and Figure 9 (before and after fine-tuning). During fine-tuning, two images of size 2500×2500 pixels, distinct from the test image, are used for finetuning pre-trained FCRNs in a patch-based manner, the same annotations following Figure 1(b) were performed manually by one individual, each image contains over 7000 cells. It can be seen that the performance of FCRN-A on real images can be improved by fine-tuning, reducing the error of 33 out of 1502 (before fine-tuning) to 17 out of 1502 (after fine-tuning).

When testing FCRN-B on two data-sets of real microscopy data, for RPE cells: Ground-truth/Estimated count = 705/699, and for Precursor T-Cell LBL cells: Ground-truth/Estimated count = 1502/1473 (without fine-tuning). Surprisingly, FCRN-B achieves slightly better performance on real data than FCRN-A. Our conjecture is that the real data contains smaller cell clumps than synthetic data, therefore, the shape of cell clumps will not vary a lot. The network is then able to give a good prediction even with a small receptive field.

5. Inverting feature representations

5.1. Problem description

In order to understand the features that have been captured by the deep networks, we considered the following question: "given an encoding of an image, to what extent is it possible to reconstruct that image?" In other words, we sought to visualise



(a) Synthetic Image: I(x)

(b) Groundtruth Density Map: D(x)

Figure 4. Annotation noise in the standard data-set. (a) The image from a standard synthetic data-set. For reader convenience, the rough boundaries of the cells have been manually drawn with a red ellipse. (b) Put a Gaussian at the centre of each generated cell. The highlighted *upper right* region contains one single cell, while the *lower left* region actually contains two overlapping cells.



(a) Synthetic Image: I(x)





Figure 5. Counting inference process for pre-trained FCRNs. (a) Input image from test set. (b) Ground-truth density map. Count: 18 (Upper left), 16 (Lower right). (c) Estimated density map from FCRN-A. Count: 17 (Upper left) 16 (Lower right). (d) Estimated density map from FCRN-B. Count: 19 (Upper left) 16 (Lower right). *Red crosses* on (c) and (d) indicate cell detection results.

Table 1. Mean absolute error and standard deviations for cell countin	ig on the standard s	vnthetic cell data-set	(Lehmussola et al. 2007:	Lempitsky	v & Zisserman 2010).
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Method	174 ± 64 cells				
	N = 8	<i>N</i> = 16	N = 32	N = 64	
Lempitsky and Zisserman (2010)	8.8 ± 1.5	6.4 ± 0.7	5.9 ± 0.5	N/A	
Lempitsky and Zisserman (2010)	4.9 ± 0.7	3.8 ± 0.2	3.5 ± 0.2	N/A	
Fiaschi et al. (2012)	3.4 ± 0.1	N/A	3.2 ± 0.1	N/A	
Arteta et al. (2014)	4.5 ± 0.6	3.8 ± 0.3	3.5 ± 0.1	N/A	
Proposed FCRN-A	3.9 ± 0.5	3.4 ± 0.2	2.9 ± 0.2	2.9 ± 0.2	
Proposed FCRN-B	4.1 ± 0.5	$.7\pm0.3$	$\textbf{3.3}\pm\textbf{0.2}$	3.2 ± 0.2	

The columns correspond to the number of training images. Standard deviation corresponds to five different draws of training and validation sets



Figure 6. FCRN-A applied on RPE cells made from stem cells. Only nucleus channel is used. *Cell Count:* Ground-truth vs. Estimated: 705/697. The data are from: http://sitn. hms.harvard.edu/waves/2014/a-stem-cell-milestone-2/. (a) RPE Cells. (b) Annotated density map. (c) Estimated density map. (d) Cell detection.



Figure 7. FCRN-A applied on Embryonic Stem Cells. Only nucleus channel is used. Cell Count: Ground-truth vs. Estimated: 535/530.

how much information of the input image has been captured by the feature representations of different layers in the deep networks (Mahendran & Vedaldi 2015). whe

The problem can be formalised as a reconstruction problem (Figure 10). Given a representation function $F : \mathbb{R}^{H \times W \times C} \to \mathbb{R}^d$ and a representation $\phi_0 = \phi(x_0)$ to be inverted, the reconstruction process aims to find another image $x \in \mathbb{R}^{H \times W \times C}$ that minimises the objective:

$$x^* = \underset{x \in \mathbb{R}^{H \times W \times C}}{\operatorname{argmin}} I(\phi(x), \phi_0) + \lambda L_2(x)$$
(4)

$$I(\phi(x), \phi_0) = \|\phi(x) - \phi_0\|^2 \quad \text{(Euclidean Distance)} \tag{5}$$

where the loss *I* compares the image representation $\phi(x)$ to the target ϕ_0 , and in our case, we choose the L_2 penalty to avoid the large pixel values.

5.2. Optimisation

Similar to training deep networks, the optimisation of Equation (4) is also a non-convex problem. However, simple gradient descent (GD) algorithms have been shown to be very effective.



(b) Estimated density map

(c) Cell detection

Figure 8. FCRN-A applied on Plasma Cells: Only greyscale image is used. Cell Count: Ground-truth vs. Estimated: 297/294.



- (a) Precursor T-Cell LBL
- (b) Estimated density map

(Without fine-tuning)

- (c) Estimated density map (After fine-tuning)
- (d) Cell detection

Figure 9. FCRN-A applied on Precursor T-Cell. Only greyscale image is used. textitCell Count: Ground-truth vs. No fine-tuning 1502 vs. 1469. Cell Count: Ground-truth vs. Fine-tuning 1502 vs. 1485.



Figure 10. Example of inverting feature representation for the cell clump based on layer ϕ_2 . Step 1: Feed an input image I(x) to the trained FCRN-A, and make a record of the feature representations $\phi_2(l(x))$. Step 2: Feed a random input image l'(x) to FCRN-A, similarly, calculate feature representations $\phi_2(l'(x))$. Step 3: Optimise the random input image l'(x) with gradient descent (GD), such that $\phi_2(l(x)) = \phi_2(l'(x))$. (Shown as the red arrows).



Figure 11. Reconstruction results from feature representations in different layers of FCRN-A.

In our implementation, *momentum* is also used to speed up the convergence:

$$\Delta x_{t+1} := \beta \Delta x_t - \eta_t \nabla E(x) \tag{6}$$

$$x_{t+1} := x_t + \Delta x_t \tag{7}$$

where $E(x) = l(\phi(x), \phi_0) + \lambda L_2(x)$ is the objective function, weight decaying factor $\beta = 0.9$, learning rate η_t is gradually reduced until convergence.

5.3. Reconstruction results

For simplicity, we only show the visualisation results from FCRN-A in this paper, but the same procedure can be performed for FCRN-B as well. In essence, CNNs were initially designed as an hierarchical model, which aimed to extract more semantic information as the networks get deeper. For our density map prediction tasks, the biggest challenge is caused by the highly overlapping cell clumps with various shapes. In Figure 11, we show to what extent the information from original cell clump can be encoded by the feature responses of different layers, and try to present an intuition about how the FCRNs make predictions.

As shown in Figure 11, when the networks get deeper, feature representations for this cell clump become increasingly abstract. Therefore, when we try to reconstruct the input image with feature representations, reconstruction quality decreases with the depth of networks, and only important information has been kept by deep layers, for instance, in *Conv3*, edges around the cell clump are captured, and for *Conv4*, which contains most abstract information in this network, only concavity information has been kept for prediction.

6. Conclusions

In this paper, we have proposed FCRNs for regressing density maps, which will later be used for both cell counting and detection tasks. The approach allows end-to-end training with images of arbitrary sizes, and is able to perform fast inference for microscopy images. Moreover, we provide intuitive understanding of feature representations from FCRNs by visualising to what extent the information has been encoded different layers.

Disclosure statement

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