Automated Quantification of Morphodynamics for High-Throughput Live Cell Imaging Datasets

Anonymous submission

No Institute Given

Abstract. We present a fully automatic method to track and quantify the morphodynamics of differentiating neurons in fluorescence time-lapse microscopy datasets. While previous efforts have successfully quantified the dynamics of organelles such as the cell body, nucleus, or chromosomes of cultured cells, neurons have proved to be uniquely challenging due to their highly deformable neurites which expand, branch, and collapse. Our approach is capable of robustly detecting, tracking, and segmenting all the components of each neuron present in the sequence including the nucleus, soma, neurites, and filopodia. To meet the demands required for high-throughput processing, our framework is designed to be extremely efficient, capable of processing a single image in approximately two seconds on a conventional notebook computer. For validation of our approach, we analyzed neuronal differentiation datasets in which a set of genes was perturbed using RNA interference. Our analysis confirms previous quantitative findings measured from static images, as well as previous qualitative observations of morphodynamic phenotypes that could not be measured on a large scale. Finally, we present new observations about the behavior of neurons made possible by our quantitative analysis, which are not immediately obvious to a human observer.

1 Introduction and Related Work

The process of forming functional connections between neurons is complex and dynamic. Time-lapse microscopy has revealed that differentiating neurons undergo a large range of dynamic processes including cell body motility, filopodial dynamics, and repeated cycles of neurite growth and retraction. Of critical importance is the process by which axons and dendrites are formed in which a neurite ceases retracting, extends a long distance, and eventually forms a connection. These dynamic events are governed by a complex protein network that coordinates cellular dynamic functions within the cytoskeleton, membrane, etc.

Recently, powerful tools have become available to help investigate the molecular interactions that govern such complex morphogenetic processes at the systems biology level. RNA interference (RNAi) technology, fluorescent protein labeling, image processing, and automated high-throughput microscopy have opened the door for large scale perturbation studies. RNAi screens have already led to novel insights into a number of cellular processes such as cell migration [1] and endocytosis [2]. However, limitations in image processing technology have restricted most studies to steady-state analysis using single images.

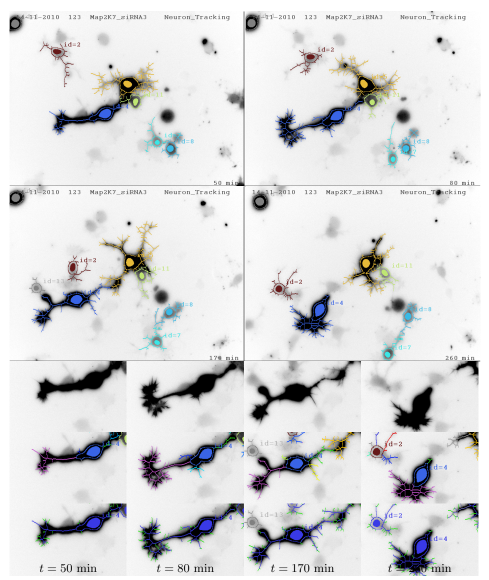


Fig. 1: Neuron Tracking Results. The top two rows contain frames from an experiment where MAP2K7 gene functions are inhibited. For visibility we enhanced the image contrast. Tracked neurons are marked by a unique color and id tag. Nuclei are denoted by filled ellipsoids, somata as contours, and neurites as trees. Bottom rows show details from above: 1) enhanced original image 2) tracked neurites marked with a different colors 3) detected filopodia marked in green. Our approach performs well even in challenging situations where neurons appear in close proximity. Videos are provided in supplementary materials. Note: faintly stained cells are ignored for robustness.

Knowledge of dynamics is essential if we are to understand complex processes such as neuron morphogenesis. However, designing algorithms to quantify dynamic behaviors is challenging, and automatic methods have appeared only very recently. State-of-the-art high-throughput techniques have successfully quantified morphodynamics of HeLA cancer cells in an effort to understand the process of mitosis [3–5]. However, the morphology and dynamics of these cell types are relatively simple in comparison to the wide range of morphodynamics exhibited by differentiating neurons.

We propose a fully automatic method for detecting, tracking, and segmenting the entire neuron including the nucleus, soma, neurites, and filopodia. We begin by detecting nuclei in each time step. Next, a greedy tracking algorithm associates detected nuclei in different time steps, forming lists of detections corresponding to individual neurons. Using the detected nuclei as seed points, a region-growing algorithm segments the soma of each neuron. The somata are used to initialize a joint segmentation of the entire structure of all neurons appearing in a image using a probabilistic method based on shortest path computations. We extract a graph describing the morphology of the neurites from the segmentation. The greedy tracking algorithm also tracks individual neurites over time, and filopodia are detected by analyzing the topology of each neurite.

As demonstrated in Fig. 1 and our supplementary materials, our approach produces reliable segmentations that capture complex neuron dynamics. For validation, we applied it to a small-scale siRNA screen of 5 genes (3 siRNAs/gene). Steady-state phenotypes associated with these genes were previously analyzed in MetaMorphTM [6] with simple image measurements. In addition, a few dynamic phenotypes were qualitatively observed but not quantified. Quantitative analysis of features extracted from our segmentations confirms the effects described in [6], as well as the observed dynamic behaviors. Our analysis also uncovered new dynamic behaviors which were previously unquantifiable.

While our greedy tracking and probabilistic segmentation algorithms are novel, they are designed to be efficient and thus are relatively simple. The main contribution of this paper is the system as a whole, which is capable of high-throughput processing of videos, tracking individual parts of neurons, and quantifying their dynamic behaviors in ways that were previously not possible.

2 High-throughput Tracking and Segmentation

The input to our approach is a series of T images $\mathcal{I} = \{I_1, \dots, I_t, \dots, I_T\}$ from which we extract K nucleus detections d_t^k . The tracking step described in Sec. 2.2 associates valid detections across time steps while rejecting spurious detections. Since each neuron contains only one nucleus, there is a one-to-one mapping between each valid nucleus detection c_t^i and a neuron X_t^i . Thus, the tracking task is to provide a set of neuron detections $\mathcal{X}^i = \{X_a^1, \dots, X_t^i, \dots, X_b^i\}$ defining an individual neuron i from time t=a to t=b. As depicted in Fig. 2, each neuron detection X_t^i is composed of a nucleus c_t^i , a soma s_t^i , a set of J neurites $\{n_t^{i,1}, \dots, n_t^{i,j}, \dots, n_t^{i,j}\}$, and a set of L filopodia associated with each neurite $F_t^{i,j} = \{f_t^{i,j,1}, \dots, f_t^{i,j,l}, \dots, f_t^{i,j,L}\}$ so that $N_t^i = \{(n_t^{i,1}, F_t^{i,1}), \dots, (n_t^{i,j}, F_t^{i,j})\}$. Thus, a complete neuron i is described by $X_t^i = \{c_t^i, s_t^i, N_t^i\}$ at time step t.

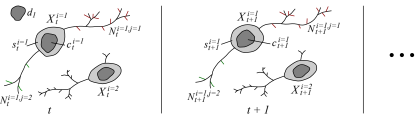


Fig. 2: Neuron Tracking Notation. A neuron i is defined by a time-series of neuron detections $\mathcal{X}^i = \{X_a^1, \dots, X_t^i, \dots, X_b^i\}$. The tracking returns a set \mathcal{X}^i for each neuron i. At time t a neuron detection $X_t^i = \{c_t^i, s_t^i, N_t^i\}$ contains a nucleus c_t^i , a soma s_t^i , and a set of neurite-filopodia tuples $N_t^i = \{(n_t^{i,1}, F_t^{i,1}), \dots, (n_t^{i,j}, F_t^{i,j}), \dots, (n_t^{i,J}, F_t^{i,J})\}$ which contains J neurites and their associated filopodia shown in red for j = 1 and green for j = 2. A spurious nucleus detection d_1 is also shown.

2.1 Nuclei and Somata Detection and Segmentation

The first step in our approach is to extract a set of nucleus detections $\{d^1, \ldots, d^K\}$ over the image series. We worked with two-channel images, one in which the cytoskeleton is marked with Lifeact-GFP. In the other, the nuclei are marked with NLS-mCherry. Thus, we can reliably detect and segment nuclei by simply thresholding the NLS-mCherry channel and performing a morphological filling operation. In the absence of such a channel, one could apply a fast machine-learning nucleus detector such as the one described in [7].

Using the nuclei as seed points, the somata are segmented by an iterative region growing algorithm. A list of pixels neighboring the current soma segmentation is maintained. At each iteration, the neighbor with the smallest weighted distance to the centroid of the seed nucleus detection $D = \lambda ||u - d^k|| + |I(u) - \hat{I}(d^k)|$ is added to the soma so long as D < T, where u is a location in the image, I(u) is the pixel intensity at that location, $\hat{I}(d^k)$ is the mean intensity of detection d^k , and T is a threshold.

2.2 Fast Greedy Tracking of Nucleus Detections

The tracking algorithm searches through the full set of nuclei detections and iteratively associates the most similar pairs of detections, returning lists of valid detections corresponding to each neuron \mathcal{X}^i . This is accomplished by constructing a graph $\mathcal{G}=(\mathcal{D},\mathcal{E})$ where each node $d_t^k\in\mathcal{D}$ corresponds to a detection. For each detection d_t^k in time step t, edges $e\in\mathcal{E}$ are formed between d_t^k and all past and future detections within a time window W. A weight w_e is assigned to each edge according to spatial and temporal distances, and a shape measure $w_e=\alpha||d_{t1}^k-d_{t2}^l||+\beta|t1-t2|+\gamma f(\nu_{t1}^k,\nu_{t2}^l)$ where $e^{k,l}$ connects d_t^k and d_t^l , and ν^k is a shape feature vector containing d_t^k 's area, perimeter, mean intensity, and major and minor axis lengths of a fitted ellipse. f evaluates differences between a feature a extracted from d_t^k and d_t^l as $f(a^k,a^l)=\frac{|a^k-a^l|}{|a^k+a^l|}$. The tracking solution corresponds to a set of edges $\mathcal{E}'\subset\mathcal{E}$ that minimizes the cost $\sum_{e\in\mathcal{E}'}w_e$.

To minimize this cost function, we adopt a greedy selection algorithm outlined in Table 1 and summarized in Fig. 3 that iteratively selects an edge with minimum cost \hat{w}_e and adds it to the set \mathcal{E}' , removing future and past connections

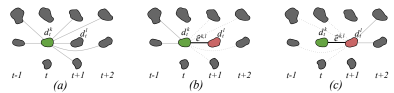


Fig. 3: Greedy Tracking. (a) The algorithm begins with each detection fully connected to all future and past detections within a time window W. Above, only d_t^k 's edges are shown. (b) Each iteration, the edge $\hat{e}^{k,l}$ with minimum cost \hat{w}_e is added to \mathcal{E}' . Edges connecting d_t^k to future detections are removed from \mathcal{E} . (c) Edges connecting d_t^l to the past are removed from \mathcal{E} . The process is repeated until $\hat{w}_e > T$.

from the detections $e^{k,l}$ connects. The algorithm iterates until the minimum cost \hat{w}_e is greater than a threshold T. The track for neuron i is extracted from \mathcal{E}' by traversing the graph $(\mathcal{G}, \mathcal{E}')$ and appending linked nucleus detections to \mathcal{X}^i .

Algorithm 1 Greedy tracking association algorithm

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Start with an empty set \mathcal{E}'.

repeat

Find edge \hat{e}^{k,l} with minimum cost \hat{w}_e.

Add \hat{e}^{k,l} to \mathcal{E}', linking detections d^k_{t1} and d^l_{t2}.

Remove \hat{e}^{k,l} from \mathcal{E}.

if t1 < t2 then

Remove edges between d^k_{t1} and future detections (where t > t1) from \mathcal{E}

Remove edges between d^l_{t2} and past detections (where t < t2) from \mathcal{E}

else

Remove edges between d^k_{t1} and past detections (where t < t1) from \mathcal{E}

Remove edges between d^l_{t2} and future detections (where t < t1) from \mathcal{E}

end if

until \hat{w}_e > T
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2.3 Probabilistic Neuron Segmentation and Neurite Tree Extraction

Given an image I_t and the set of somata present in it $S_t = \{s_t^1 \dots s_t^m\}$, our goal is to associate to each pixel u a label $J_t(u)$ that indicates to which soma it belongs. The probability of $J_t(u)$ can be deduced using Bayes' rule,

$$P(J_t(u) = i|S_t, I_t) = \frac{P(S_t, I_t|J_t(u) = i)}{\sum_{n=1}^m P(S_t, I_t|J_t(u) = \eta)},$$
(1)

where we have assumed a uniform distribution on $P(J_t(u))$. The numerator is modeled as the probability of the path L that connects maximally the voxel u to the soma s_t^i , $P(S_t, I_t | J_t(u) = i) = \max_{L: u \to s_t^i} \prod_{\{l_r\} \in L} P(I_t(r) | l_r)$, where l_r are indicator variables for the locations forming the path L. We chose this model since an optimal maxima can be found by minimizing its negative likelihood using Dijkstra's shortest path and because it produces connected components.

To optimize this function, we map the image I_t to a graph $\mathcal{G}_t^i = (V, E)$ whose vertices u are the pixels in I_t and whose directed edges $e_{r,v}$ connect each pixel to its four neighbors. We assign to each edge a weight $w_{r,v} = -logP(I_t(v)|v)$.

 $P(I_t(v)|v)$ represents the probability that a neurite traverses a node v and is obtained by applying a sigmoid function to the output of the tubularity filter of [8]. The parameters of the sigmoid function are estimated using maximum likelihood. Finally, we define the set of neurite pixels U_n^t as those that connect to any soma with a higher probability than ϵ . We predict their labels as the ones that maximize Eq. 1. The set of pixels associated to neuron X_i^t is the union of the neurites and the soma associated with i, $U_i^t = \{u \in U_n^t | J_t(u) = i\} \cup s_i^t$. To reduce the neurite segmentation to a tree, we skeletonize the neuron and define as root node the pixel of the skeleton closest to the centroid of the nucleus. We instantiate a Minimum Spanning Tree from the root and create a neurite tree every time the the spanning tree exits the soma.

2.4 Neurite Tracking and Filopodia Detection

Neurites are tracked by applying the algorithm described in Sec 2.2 using the centroids of the neurite trees instead of nucleus centroids, with the additional constraint that edges may only exist between neurites that emanate from the same soma. Filopodia are detected by starting at each leaf node in a neurite and traversing the tree until a branch point is reached. If the distance traversed is less than a threshold T_f , the traversed locations are considered to be filopodia.

3 Extracting Morphodynamic Features

Our tracking and segmentation method produces sets of graphs linking detections, contours, and trees to define each neuron over time. This data structure is not immediately useful for quantifying dynamic behaviors. To facilitate the analysis, we extract a set of 156 meaningful features from our data structure to quantify morphodynamics, which are too numerous to list here. A few examples for the nucleus and soma include: area, perimeter, Lifeact-GFP intensity, NLS-mCherry intensity, speed, acceleration, total distance traveled, time spent expanding/contracting, frequency of expansion. For neurites: number of branches, distance from tip to soma, filopodia length, number of filopodia, major axis, minor axis and eccentricity of an ellipse fitted to the neurite, total length, time spent expanding/contracting, frequency of expansion (and Δ 's for all above).

4 Results and Conclusion

Experimental Setup and Methodology – We used our method to perform a small-scale siRNA screen in which the functions of 5 genes were inhibited with three goals in mind: to reproduce the findings of [6], to confirm previous qualitatively observed morphodynamics, and to uncover new dynamic behaviors. Three siRNAs were applied for each gene (SrGAP2, MAP2K7, RhoA, Trio, and Net), for a total of 17 experiments including 2 controls. 30 videos per experiment were obtained over the course of 3 days, with images taken in 10 minute intervals for a total of 510 videos, each containing approximately 100 2-channel images of 696×520 resolution. A total of 7,298 neurons and 33,213 neurites were tracked and segmented. Our method processes a single video in 210 seconds on a Lenovo W510 notebook computer (2.1 seconds per image), and the entire screen was processed in parallel in just a few hours using conventional PCs.

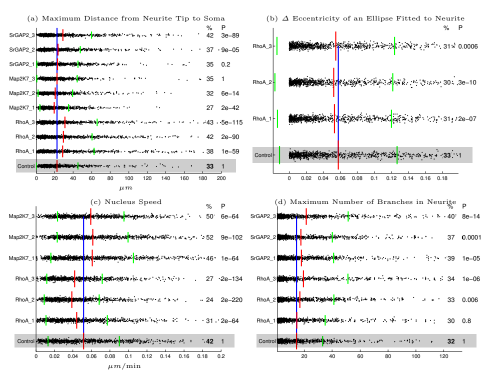


Fig. 4: Quantitative Morphodynamic Analysis. Four informative morphodynamic features are plotted, where the control experiment is marked in gray below the siRNA targets. Black dots represent collected data points. Red bars indicate the mean, green bars indicate standard deviation. The blue line shows the control's mean for comparison. Values under the % column are the percentage of data points above the control's mean. Values under P indicate the student-t test p-value. See text for details.

Our quantitative analysis investigated the effects of inhibiting various gene functions on each of the 156 morphodynamic features we extracted from our segmentations as described in Sec. 3. We summarize our findings below and in Fig 4. We also provide a detailed table and video results in the supplementary material. Throughout the pipeline, many potential sources of noise exist, including variability in neuron behavior, transfection effects, mistakes in our segmentation, and imaging conditions. The sheer number of cells and neurites analyzed helps to average out the noise. Our findings reported below are statistically significant, all having p-values << 0.05.

Analysis – Our analysis confirms the effects found through static image analysis in [6]. In particular, RhoA loss of function resulted in fewer but longer neurites than the control. SrGap loss was found to have longer neurites, and Map2K7 loss was found to have more neurites but with short length. These findings were confirmed by dynamic measures in our experiments: the mean longest neurite length – control $22.6\mu m$, RhoA-3 $32\mu m$, SrGap2-1 $28.9\mu m$, and Map2K7-1 $19.5\mu m$ (see

Fig. 4a)¹; and by a dynamic measure – the mean number of neurites belonging a neuron over its lifetime: control 3.4, RhoA 3.1, and Map2K7-1 3.9.

It was qualitatively observed, but never quantified, that loss of SrGap2 function produces a high number of filopodia, and that RhoA loss results in neurites that easily extend but have difficulty retracting. These morphodynamics were confirmed by our analysis. Mean number of filopodia detected per neurite over its lifetime was 6.69 in the control and 8.81 for SrGap-3¹. The mean change in elongation as measured by an ellipse fitted to the neurite was 5.7% for the control and 5.3% for RhoA-1 (see Fig. 4b)¹. While this difference may seem small, it is significant due to the large amount of data collected (p-value is 2×10^{-7}).

Our quantitative analysis revealed new morphodynamics which were not obvious to human observers. We found that RhoA function loss slowed neuron motility and Map2K7 increased it. Control cells moved at .30 $\mu m/min$, RhoA moved at .23 $\mu m/min$, and Map2K7-2 moved at .37 $\mu m/min$ (see Fig. 4c). We also found that RhoA and SrGap increased the branching of the neurites (see Fig. 4d). Over the course of a neurites lifetime, the maximum number of branches in a control neuron was 14.5, 19.44 for RhoA-3, and 21.39 for SrGap2-3¹.

Conclusion — We have described a set of algorithms which, as a system, are capable of robustly tracking and segmenting entire neurons including the nucleus, soma, neurites and filopodia. Our approach is efficient, and can analyze high-throughput datasets using meaningful dynamic features extracted from our segmentations. We validated our approach by reproducing previous findings, confirming anecdotal findings, and uncovering new dynamic phenotypes. It is beyond the scope of this work to comment on the biological significance of these findings, if there is any. We leave that for future work.

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¹ Only neurons containing the 10th percentile of longest neurites were considered.