eDetect: A Fast Error Detection and Correction Tool for Live Cell Imaging Data Analysis

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HIGHLIGHTS
- eDetect is a user-friendly software for live cell imaging data analysis
- It provides an easy solution to detect and correct errors in imaging data analysis
- It enables precise cell lineage reconstruction from long-term imaging datasets
- It is compatible with CellProfiler and performs well in different imaging datasets

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eDetect: A Fast Error Detection and Correction Tool for Live Cell Imaging Data Analysis

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SUMMARY
Live cell imaging has been widely used to generate data for quantitative understanding of cellular dynamics. Various applications have been developed to perform automated imaging data analysis, which often requires tedious manual correction. It remains a challenge to develop an efficient curation method that can analyze massive imaging datasets with high accuracy. Here, we present eDetect, a fast error detection and correction tool that provides a powerful and convenient solution for the curation of live cell imaging analysis results. In eDetect, we propose a gating strategy to distinguish correct and incorrect image analysis results by visualizing image features based on principal component analysis. We demonstrate that this approach can substantially accelerate the data correction process and improve the accuracy of imaging data analysis. eDetect is well documented and designed to be user friendly for non-expert users. It is freely available at https://sites.google.com/view/edetect/ and https://github.com/Zi-Lab/eDetect.

INTRODUCTION
Live cell imaging has been widely used in monitoring cellular processes and dynamics at the single-cell level (Ni et al., 2018; Specht et al., 2017; White et al., 2018). During the course of these studies, large volumes of time-lapse images are often generated by automatic microscopes, which poses a challenge in quantifying cellular dynamics from these datasets with high precision (Skylaki et al., 2016). In the past, computational tools have been developed to enable automated analysis of large image datasets (Blanchoud et al., 2015; Carpenter et al., 2006; Cooper et al., 2017; de Chaumont et al., 2012; Held et al., 2010; Hilsenbeck et al., 2017; Liepe et al., 2016; Winter et al., 2016). However, the errors in the automated analyses impaired the accuracy in quantifying long-term single-cell dynamics. In most cases, fully automatic algorithms cannot guarantee 100% accuracy (Ulman et al., 2017). Even a small portion of errors can result in a much larger amount of false cell lineages (Skylaki et al., 2016). Therefore manual inspection and correction are needed (Hilsenbeck et al., 2016).

Several tools have provided functions to manually correct image data analysis results. For example, CellProfiler and qTfy allow the users to manually edit the contours of segmented objects one by one (Carpenter et al., 2006; Hilsenbeck et al., 2016). In addition, CellTracker, NuclITrack, LEVER, and tTt support manual track editing (Cooper et al., 2017; Hilsenbeck et al., 2016; Piccinini et al., 2016; Winter et al., 2016). These tools are useful, but an approach is still needed to improve the efficiency of data curation.

Here, we report an error detection and correction tool for live cell imaging data analysis (eDetect). In eDetect, we proposed a new gating approach to classify flawed segmentation and tracking results by visualizing image features in scatterplots based on principal-component analysis (PCA). As cell objects with similar appearance often aggregate in the scatterplot, we can easily detect a group of errors from voluminous time-lapse images. Similar errors can be efficiently corrected with batch operations in eDetect. In addition, eDetect uses multiple quality control steps to achieve high accuracy in the final live cell imaging analysis results. In summary, eDetect is a comprehensive software tool that integrates cell segmentation, feature extraction, cell tracking, and efficient data curation.

RESULTS
eDetect Provides a Fast Way to Detect Segmentation Errors
We first used eDetect to perform cell segmentation on the live cell imaging data of mouse stem cells, which is a benchmark dataset (Fluo-N2DH-GOWT1) from the Cell Tracking Challenge (Maška et al., 2014; Ulman et al., 2017). The results of automatic segmentations are shown in the image display window (Figure 1A).
found that the application of automatic segmentation algorithms can result in different types of errors such as under-segmentations (multiple adjacent nuclei are labeled as one object), over-segmentations (one nucleus is divided into multiple objects), and irregular shapes. To efficiently detect these errors, eDetect provides a function of segmentation gating, which uses a scatterplot to visualize image features based on PCA. Each point in this plot represents a segmented object. As shown in Figure 1B, the correct and wrong segmentation objects are well separated and displayed as different groups. By clicking on the points in the scatterplot, the corresponding segmentation objects are highlighted in the image display window. In this way, the users can easily inspect and identify different groups of segmentation errors.

**eDetect Can Correct a Group of Errors with Batch Operations**

To further improve the efficiency of error correction, eDetect provides both manual and batch operations for splitting and deleting cell segmentations. When the same type of segmentation errors are grouped...
together in the PCA plot, eDetect can correct these types of errors with a batch operation after selecting them with a polygon. For example, the cluster of under-segmentations can be corrected by one click of splitting objects in eDetect (Figure 1C). Similarly, the irregular non-cell segmentations can be easily discarded with one “Delete objects” operation.

Multiple Quality Control Steps Improve the Accuracy of Live Cell Imaging Data Analysis

Although segmentation gating can assist the users to correct segmentation errors efficiently, some mistakes may still exist. Therefore eDetect provides additional steps to improve the accuracy of data analysis. To illustrate this feature, we applied eDetect to analyze a live cell imaging dataset (HaCaT-FUCCI) from human HaCaT cells that stably express the CFP-H2B nuclear marker and mCherry-Geminin FUCCI cell cycle indicator proteins (Sakaue-Sawano et al., 2008).

To improve the precision of cell tracking, eDetect uses cell pair gating to visualize identified cell divisions in another PCA-based scatterplot. eDetect automatically locates and marks the corresponding pair of cells in the original image when the user points at a cell division pair (Figure 1D). By inspecting the highlighted pairs of cells, the users can find and correct misassigned cell divisions.

As the last quality control module in eDetect, cell lineages display window assists the users to identify missed errors from previous steps. This is implemented by visualizing different measurements (e.g., nuclear area) of the objects in cell lineages with a heatmap. In general, nuclear area changes gradually under normal cell growth and only shows abrupt changes during cell divisions. The users can easily find abnormalities such as abnormal cell divisions, abrupt changes of nuclear sizes, and incomplete cell lineages (Figure 2). eDetect can assist us to check and correct these abnormalities in convenient ways. For example, eDetect has an automatic outlier detection function to mark abrupt changes within cell lineage branches. In addition, it can generate a synchrogram (a sequence of images of the selected cell lineage), which facilitates the check on the correctness of cell lineages (Sigal et al., 2006). After multiple error correction steps,
we can accurately quantify the dynamics of the cell cycle sensor protein mCherry-Geminin in individual HaCaT cells (Figure S1).

Compatibility of eDetect with CellProfiler

eDetect can be used together with CellProfiler to maximize its interoperability. The users can first perform image preprocessing and segmentation with CellProfiler and then import the results into eDetect for error detection and correction. As CellProfiler provides advanced algorithms and pipelines for image analysis, the compatibility of eDetect with CellProfiler will improve the broad application of eDetect for live cell imaging analysis.

Evaluation of eDetect Performance to Detect and Correct Errors in Live Cell Imaging Data Analysis

To evaluate the effectiveness and time cost of eDetect in error detection and correction, we first performed automatic data analysis in CellProfiler (CellProfiler*) and eDetect (eDetect*) without any corrections. In parallel, we imported the automatic analysis results from CellProfiler and then performed additional error correction with eDetect (CellProfiler+). In addition, we implemented automatic image analysis and error correction with eDetect (eDetect). Performance was quantified based on segmentation accuracy (SEG), tracking accuracy (TRA), complete tracks (CT) and recall of complete lineages (RCL). The RCL score measures the fraction of cells (including their descendants) that are correctly tracked from the first frame to the last frame of the movie. A detailed description of the performance metrics is provided in the Transparent Methods. The performance scores were calculated for four datasets, consisting of an imaging dataset HaCaT-FUCCI and three training datasets from the Cell Tracking Challenge.

As shown in Figure 3, eDetect can improve the performance of data analyses on different datasets by implementing error correction (comparing eDetect* and eDetect†). The average values of SEG and TRA for the tested four datasets are increased by 0.04 and 0.10, respectively. More importantly, the average values of CT and RCL are increased to a larger extent, by 0.55 and 0.38, respectively. Similar performance augmentation is observed by comparing CellProfiler* and CellProfiler+. The improvement of CT and RCL values is more apparent when a long-term live cell imaging dataset is analyzed. For the 2-day imaging dataset HaCaT-FUCCI, although automatic algorithms in CellProfiler* and CellProfiler+ reached very good SEG and TRA scores, the corresponding CT and RCL values are very low. This result indicates that a small portion of segmentation and tracking errors occurring in the automatic analysis can lead to a lot of errors in CT, which need to be corrected to reach valid conclusions. By performing error detection and correction in eDetect, the RCL value was increased from 0.111 to 1 with about 9 min of error correction on HaCaT-FUCCI dataset. The RCL value is especially important to draw biological conclusions when a reconstruction of complete cell lineages is needed. Therefore, eDetect is very useful especially for long-term imaging data analysis.

In general, eDetect is efficient and has a low time cost (Figure 3). The execution time (TIM) for the automatic analysis step is less than 1 min for six tested imaging videos running on windows with eight CPU cores (Intel Core i7-6,700 Processor, 3.4 GHz). The time used in the error detection and correction steps (TIM_EDC) depends on the performance of automatic segmentation algorithms and the size of the image dataset. The error correction steps took about 10-90 min for six of the analyzed videos with total number of frames ranging between 65 and 206. For the video #2 of Fluo-N2DH-SIM+ dataset, the performance measures are relative poor when automatic analyses were performed in both CellProfiler* and eDetect*. This might be caused by the artificial noise introduced in this simulated dataset. In this case, several hours of manual correction of segmentations are needed to substantially improve the accuracy of data analysis.

We next compared eDetect with the top-three performing methods from Cell Tracking Challenge using three benchmarking datasets (Figure 4). In terms of the SEG and TRA scores, eDetect has comparable performance to those of top-performing methods. It is important to note that, with the function of error detection and correction, eDetect could achieve the best performance in the measurements of complete tracks and recall of complete lineages. These results highlight the importance of error correction in long-term live cell imaging data analysis.

DISCUSSION

The quantification of long-term single-cell dynamics is crucial for understanding cell signaling dynamics and cell fate in development. Although improving the accuracy of automatic algorithms is very important...
for live cell imaging data analysis (Maśka et al., 2014; Ulman et al., 2017), an efficient curation tool is also urgently needed because a small fraction of errors, especially occurring earlier in the videos, could lead to a large fraction of errors in the complete cell lineages. In eDetect, we proposed a PCA-based approach to detect and correct errors, which substantially reduces the time for manually pinpointing errors from massive image datasets. In addition, eDetect can help us to correct or remove errors with one-click batch operations. Last, but not least, multiple quality control steps ensure a high precision in the final imaging analysis results.

Despite its effectiveness, segmentation gating in eDetect is inherently limited: it requires the users to customize PCA input variables. In practice, it is difficult to completely separate correct and incorrect
The users can achieve accurate segmentations by applying multiple gating steps with different PCA inputs even if each round of gating is not ideal. Moreover, overlooked errors can still be found in subsequent quality control steps. On the other hand, supervised learning methods have been previously used for phenotypic analysis of high-content imaging data (Piccinini et al., 2017; Ramo et al., 2009; Smith et al., 2018). In the future, these methods could be potentially used in eDetect to detect segmentation errors.

Here, we demonstrated eDetect’s capability to detect and correct errors in live cell imaging data analysis. Making sense of large imaging datasets is a demanding challenge that requires powerful and efficient computational tools. The development of eDetect will be valuable for improving the precision of live cell imaging data analysis.

**Limitations of This Study**

Current release of eDetect does not provide multiple algorithms for segmentation and tracking. The performance of automatic analysis in eDetect may not be suitable for certain datasets. The compatibility of eDetect with CellProfiler offers the choice of various segmentation and tracking methods from CellProfiler, but it requires additional pipeline setup work. In the future, eDetect could be improved by integrating...
more established methods for segmentation and tracking. In addition, there is also a strong demand for the analysis of high-throughput time-lapse datasets. Although eDetect can in principle handle these datasets, it would require a large amount of manual labor. To address this challenge, it would be helpful to apply machine learning methods for error detection and correction.

**METHODS**

All methods can be found in the accompanying Transparent Methods supplemental file.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Transparent Methods, one figure, and one table and can be found with this article online at https://doi.org/10.1016/j.isci.2019.02.004.

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**AUTHOR CONTRIBUTIONS**

H.H. developed the software and implemented performance evaluation on benchmark datasets. G.W. generated some live cell imaging data for testing this software. Y.L. tested the tool. Z.Z. conceived and supervised the project, provided input on the design and features of the application, and tested the tool. Z.Z. wrote the paper with input from H.H.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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Supplemental Information

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Supplemental Figures and Legends

Figure S1: The dynamics of mCherry-Geminin protein in HaCaT single cells, which were quantified with eDetect. Related to Figure 2.
## Supplementary Tables

### Table S1. Names and IDs of shape and additional features, Related to Figure 1.

| ID in Segmentation Gating and Cell Pair Gating | Feature Category | Feature Name          |
|----------------------------------------------|------------------|-----------------------|
| v{1}                                         | shape¹           | Area                  |
| v{2}                                         | shape            | Eccentricity          |
| v{3}                                         | shape            | Orientation           |
| v{4}                                         | shape            | Solidity              |
| v{5}                                         | shape            | Perimeter             |
| v{6}                                         | shape            | MeanIntensity         |
| v{7}                                         | additional²      | radial distribution   |

¹ The meanings of shape features can be found at:
https://www.mathworks.com/help/images/ref/regionprops.html

² The additional feature “radial distribution” describes how intensity varies as a function of distance from the nearest point in the border of the object.
Transparent Methods

Cell segmentation

eDetect uses an adaptive thresholding approach to segment cell nuclei from nuclear marker images (Bradley and Roth, 2007). To de-clump the clustered nuclei, we apply watershed transformation (Meyer, 1994) on a filtered distance map of nuclei contours (Schneider et al., 2012). Because single nuclei sometimes are wrongly split into smaller objects, we adopt a hierarchical procedure to merge directly adjacent objects based on the areas of minimum enclosing ellipses (Moshtagh).

Feature extraction

eDetect extracts cellular features from nuclei contours and nuclear marker images. It provides the options of computing 5 categories of features, i.e. shape, intensity, Haralick features (Haralick and Shanmugam, 1973), Zernike features (Boland et al., 1998) and additional features. By default, eDetect calculates shape and additional features, which are listed in Table S1. The calculations of intensity, Haralick and Zernike features are implemented using the source code of CellProfiler (Carpenter et al., 2006).

Cell tracking

eDetect adopts a constrained nearest-neighbor strategy for cell tracking. For each object (except the objects in the first frame), eDetect calculates its distances to all the objects in the previous frame. The closest object in the previous frame is the predecessor, unless this shortest distance is still larger than a predefined threshold.

Sometimes, consecutive frames are not perfectly aligned to each other in the x-y plane because of paraxial shifts of the field. We tackle this problem using an optimization-based approach. For each frame, before calculating distances, eDetect subtracts a “shift vector” from the coordinates of all objects in the current frame. Therefore, the assignment of predecessors depends on the value of this “shift vector”, and the sum of the aforementioned shortest distances is a function of the “shift vector”. eDetect searches for the “shift vector” (within a square whose size is user-defined) that minimizes this function. The assignment that is associated to this minimal is the tracking result.

Measurements

In eDetect, the following measurements are calculated for different fluorescent channels (the channels for signal extraction): nuclei median and mean intensities, cytoplasm median and mean intensities, and nuclear-cytoplasmic ratios of median and mean intensities.

To approximate nuclei regions, we shrink the nuclear segmentations by a user-defined distance to exclude nuclear membrane pixels. The cytoplasm regions are approximated using peri-nuclear ring regions, whose inner and outer contours are expanded from nuclear segmentations. In order to compensate medium illumination, background intensity is subtracted from both nuclei and cytoplasm intensities.

To generate the foreground region, we expand the nuclear segmentations by a large distance that is the maximum of the following two values: (1) the largest object diameter and (2) twice of the distance by which nuclear segmentations are expanded to generate the outer contour of the peri-nuclear ring regions. Afterwards, we derive the background region by removing the foreground region from the whole image.
In eDetect, to shrink an object by a distance of \( n \), we remove the contour pixels of the object \( n \) times iteratively. Similarly, to expand an object by a distance of \( n \), we add a layer of pixels to the exterior of the object \( n \) times iteratively. We implemented both shrinkage and expansion operations with an algorithm of morphological operations on binary images in Matlab.

**Segmentation gating and cell pair gating**

eDetect segmentation gating and cell pair gating employ principal component analysis (PCA) to project high dimensional cellular features onto a 2D plane, where the data is visualized with scatter plots. PCA converts a data table, in which the observations are described by a set of possibly correlated variables, into a new one, where the new variables (i.e.: principal components) are uncorrelated (Abdi and Williams, 2010). From the first principal component (PC) to the last one, each PC maximizes the data variance under the constraint that it is orthogonal to all the preceding ones. In eDetect, PC1 and PC2 are used as the x- and y-axis variables in the scatter plots. The meanings of imaging features for the PCA input are listed in Table S1.

In cell pair gating, apart from the variables specified in the formula, the input to PCA additionally includes 3 default variables: the 3 sharp angles between 3 lines. The 3 lines are: the major axes of the 2 objects and the line that passes through the centroids of the 2 objects.

**Outlier detection in cell lineages display**

In eDetect, abrupt changes of cellular measurements are detected based on the median absolute deviation (Leys et al., 2013). eDetect calculates relative difference of measurements between each cell and its predecessor. Among all the relative differences, those whose distances to the median value are larger than a certain predefined number of standard deviations are defined as outliers.

**The calculation of performance scores**

Segmentation accuracy (SEG), tracking accuracy (TRA) and complete tracks scores (CT) are calculated using the evaluation software packages provided in the Cell Tracking Challenge website. Recall of complete lineages (RCL) is calculated with a customized MATLAB script.

Complete lineages are lineages that start in the first frame and end in the last frame of the entire video. The formula of recall (or sensitivity) is given below (\( TP \): true positive, \( FP \): false positive, \( FN \): false negative.):

\[
Recall = \frac{TP}{TP + FN}
\]

The ground truth of HaCaT-FUCCI dataset was produced by manually correcting the result of eDetect automatic analysis until the tracks were correct and segmentations were accurate under human inspection.
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