Single-cell cytometry via multiplexed fluorescence prediction by label-free reflectance microscopy

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Abstract

Traditional imaging cytometry uses fluorescence markers to identify specific structures, but is limited in throughput by the labeling process. Here we develop a label-free technique that alleviates the physical staining and provides highly multiplexed readout via a deep learning-augmented digital labeling method. We leverage the rich structural information and superior sensitivity in reflectance microscopy and show that digital labeling predicts highly accurate subcellular features after training on immunofluorescence images. We demonstrate up to 3× improvement in the prediction accuracy over the state-of-the-art. Beyond fluorescence prediction, we demonstrate that single-cell level structural phenotypes of cell cycles are correctly reproduced by the digital multiplexed images, including Golgi twins, Golgi haze during mitosis and DNA synthesis. We further show that the multiplexed readout enables accurate multi-parametric single-cell profiling across a large cell population. Our method can dramatically improve the throughput for imaging cytometry towards applications for phenotyping, pathology, and high-content screening.
Introduction

Cell morphology features are powerful phenotypical readout, which have been the basis for pathology for decades. They are also the underlying mechanisms for varieties of imaging cytometry and high-content screening platforms to characterize pathological changes and responses to drug treatments (1). The most widely used approach for imaging readout is fluorescence labeling that highlights subcellular components or cell functions through immunofluorescence (IF), fluorescent reporter cells or dyes. However, the throughput of these approaches is fundamentally limited by the physical process of labeling. The IF staining is labor-intensive and generally requires cell fixation that does not allow kinetic observations of live cells over time. Fluorescent reporter cells are permissive for longitudinal live cell imaging. However, the process of gene editing and validation takes a significant amount of time and can be difficult to introduce multiple markers within the same cells for multiplexed analysis. Regardless of the fluorescence labeling approaches, the overlapping of the fluorescence emission spectra further limits the multiplexing capability. To alleviate these limitations, here we develop a label-free single-cell cytometry that is highly multiplexed and can forgo the physical staining via a deep learning (DL)-augmented digital labeling method.

Our work relies on the premise that label-free scattering-based microscopy captures rich structural information and can be effective to characterize cell morphological features (2). Brightfield, phase contrast and differential interference contrast (DIC) microscopy have been routinely used for observing and quantifying cell morphology (3). Scattering-based microscopy and tomography techniques have been increasingly utilized to reconstruct cellular structures (4). Of particular interest is the reflectance-mode microscopy that provides exquisite sensitivity in detecting nanoscale structural changes beyond the diffraction limit (5–7). By capturing backscattering signals, reflectance imaging provides access to the highest spatial-frequency components in the reciprocal Fourier space and thus can provide higher structural contrast than the transmission techniques (2). Indeed, our recent work shows that backscattering signals allow resolving finer details than the transmission counterparts (8, 9). In this work, we further leverage the higher sensitivity provided by the reflectance-mode microscopy and demonstrate how enriched label-free information allows predicting highly accurate subcellular structural features.

The framework of this study is illustrated in Fig. 1. The angle-dependent backscattering features are captured with darkfield oblique illumination and paired with IF images (Fig. 1A). Using the IF images as the ground truths, multiple DL models are independently trained for individual IF labels (Fig. 1B). Once all the models are trained, we perform digital multiplexing by feeding the same reflectance input to each network and make different fluorescence predictions in parallel (Fig. 1C). By doing so, multiple subcellular structures and cell states can be revealed simultaneously without physical labeling. While previous work has shown that DL models can disentangle the complex structures captured in the label-free data and make *in-silico* fluorescence labeling with high accuracy (10–12) or holistically capture “hidden” structural features that are not easily perceived or described (13–22), these results are fundamentally limited by the weak structural contrast from the transmission modes that contain only forward scattering information. By exploiting the enhanced resolution and sensitivity in the backscattering data, we demonstrate a dramatic increase in the fluorescence prediction accuracy with up to 3× improvement as compared to the current state-of-the-art.
One distinct contribution of our work is to advance beyond the prediction of fluorescence images, and to demonstrate accurate structural phenotyping and quantitative single-cell cytometry using digitally multiplexed fluorescence images. Importantly, we show that our DL model can correctly capture and predict characteristic subcellular features during the cell cycle, including morphological changes of nuclei and Golgi apparatus. We also show that our DL model can capture the structural features of cell proliferation and recapitulate the DNA duplication through the cell cycle. Remarkably, the label-free structural elements identified in proliferating cells are not obvious by visual inspection of the raw images, demonstrating that our holistic DL model can potentially capture novel cellular attributes with high accuracy. Another distinct advantage of multiplexing several fluorescence markers is that it enables the development of multiple quantitative metrics for imaging cytometry and phenotyping across the large cell population and at the single-cell level (e.g. cell size, nuclear-cytoplasmic ratio, nuclear roughness, Golgi eccentricity, etc.). As a demonstration, we evaluate several cellular features, including morphology and fluorescence expressing intensity on the DL predicted digitally multiplexed readouts.

A common critic of DL-based method is the “black-box” nature of these models (23). To overcome this issue, we adapt the attention mechanism (24) to elucidate on the working mechanism of our DL model. We construct the saliency map that highlights the most important subcellular features contributing to each fluorescence prediction by the network. Our results show that the structural components in label-free reflectance input that correspond to the fluorescence labels can be correctly identified by the saliency map, and the “attention” is consistent across different cell batches when predicting all IF labels. This indicates that our network learns to extract the salient and specific structural information from the reflectance images matching the underlying subcellular component. In addition, the improved prediction accuracy is attributed to the enhanced resolution and sensitivity to subcellular structures from the backscattering information.

Results

**Oblique illumination-based reflectance microscopy captures rich morphological information**

The imaging platform is based on our recently developed LED-array reflectance microscope for capturing co-registered label-free reflectance and fluorescence images (8). By flexibly controlling the LED patterns, this new platform enables capturing multiple angle-dependent backscattering contrasts in the darkfield without any mechanical switching. Based on our prior work (8), we heuristically optimize the illumination strategy and implement half-annulus LED patterns along four different orientations (including top, bottom, left, and right) (see Fig. 1A). In addition, we compute the darkfield reflectance differential phase contrast (drDPC) based on the raw measurements (see Materials and Methods). The raw oblique-illumination darkfield and drDPC images contain complementary structural contrasts. In particular, sub-cellular structures are shown with high contrast in the raw darkfield measurements, including the nuclei, nucleoli, and hyper-reflective structures at the nuclear periphery. Cell membranes with sharp boundaries are highlighted in the drDPC images, with cytoplasm spreading on the substrate with thicker nuclei at the cells’ centers. The drDPC better exhibits the cell topography and height. These label-free images are used as the multi-channel input to our DL model. On the same platform, two-channel epi-fluorescence images are acquired on the same sample to serve as the ground-truth for training our DL models (Fig. 1A) (see Materials and Methods). The significance of this new
microscopy platform is that we capture enriched label-free information by multiple contrasts in the reflectance-mode. This empowers our label-free high-content cytometry technique to uncover highly sensitive and specific structural phenotypes at the single-cell level across large cell populations.

**Individual fluorescence prediction achieves state-of-the-art performance**

To evaluate the performance of our DL models, we take measurements on fixed HeLa cells containing in-total six IF labels, including DNA (Hoechst), Golgi apparatus (GM130), endosome (EEA1), actin (Phalloidin), proliferation (EdU), and apoptosis (TUNEL). Specifically, five separate batches of IF staining are performed with GM130, EEA1, Phalloidin, EdU, TUNEL, each of which is co-stained with Hoechst (see Materials and Methods). We then train six networks for performing individual IF label predictions using paired reflectance-fluorescence image dataset (Fig. 1B). Additional details about the network implementation and the data preprocessing procedure are provided in Materials and Methods and Supplementary Materials Figs. S1 and S2, respectively.

After training, we first evaluate each network’s prediction accuracy on unseen reflectance input from the same cell batch. Figure 2 shows the label-free input, the individual IF ground-truth and prediction for the all six labels. The predicted subcellular structures and cell states have excellent visual agreement with the ground-truths. Characteristic morphological features are clearly recovered, including rounded nuclei, cytoplasmic endosome, spreading cell membrane (actin), and Golgi apparatus at the nuclei periphery. Selective cellular events or functions such as proliferation and apoptosis are also captured by the DL predictions. Additional visualizations of the prediction results are shown in Supplementary Materials Fig. S3.

We quantify the prediction accuracy by computing the evaluation metrics on the network predictions based on the underlying cytometry tasks. Specifically, we formulate the predictions of the DNA, endosome, actin and Golgi apparatus as regression problems because they are pervasive in every cell. Accordingly, the regression performance is quantified by the Pearson correlation coefficient (PCC) (see Materials and Methods). We formulate the prediction of the proliferation and apoptosis as detection problems because they are highly selective to the underlying cell states. Accordingly, the detection performance is quantified by the Area Under the Receiver Operating Characteristic (ROC) Curve (AUC) (see Materials and Methods). The distribution of the sample-wise accuracy is shown in the violin plots in Fig. 3. Notably, all four of our regression models achieve higher accuracy as compared to the current state-of-the-art (10), with the mean PCCs on the DNA, endosome, actin, and Golgi apparatus label predictions reporting 95.32%, 90.43%, 89.11%, and 57.27%, respectively. This result agrees well with the visualizations in Fig. 2. The cellular features in the reflectance image associated with the corresponding fluorescence label are clearly visible. Although these scattering signals are entangled with other signals in the raw label-free images, our result shows that the DL model is able to recognize and distill these salient features with high accuracy. The overall average detection accuracy is shown to be 83.9% and 60.2% for the proliferation and apoptosis, respectively. Notably, the scattering features in the proliferating cells cannot be easily described from the raw reflectance images, yet our DL model can capture the salient structural features with high accuracy. Overall, these results validate our hypothesis that the improved sensitivity and resolution in reflectance images contain rich morphological features that can be utilized more effectively for structural phenotyping by DL.
Multiplexed prediction recovers biological accurate cellular structures

Next, we demonstrate the digital multiplexing capability by feeding the same reflectance input to each network and make six different IF predictions in parallel. By doing so, multiple subcellular structures and cell states are revealed simultaneously. In Fig. 4A, the image multiplexes the nucleus, Golgi apparatus, actin, and endosome virtual IF labels in a single large field-of-view (FOV) for a large cell population. In Fig. 4B, the virtual labels for proliferating and apoptotic cells are multiplexed with the darkfield reflectance input in the same FOV as Fig. 4A. These multiplexed predictions are performed on the cell batch under the Golgi apparatus staining condition. To further demonstrate the robustness of this digital multiplexing procedure, we show additional examples of multiplexed predictions performed on different cell batches/staining conditions in Supplementary Materials Fig. S4.

Importantly, our results show that the DL model can correctly capture and predict characteristic subcellular features during the cell cycle. During interphase, the nuclei have a regular rounded shape with nucleoli present, and Golgi apparatus is anchored primarily to one side of the nuclei (Figs. 4C-4E). At this stage, cells that have initiated or ongoing DNA/chromatin replications have a positive signal for proliferation (Fig. 4F). When cells enter mitosis, the chromosomes start to condense towards the centers of the cells, and nucleoli disappears. Golgi apparatus undergoes vesiculation and fragmentation, and its components are found scattered throughout the cytoplasm in the form of tiny (~50-nm) vesicles, often referred to as the “Golgi haze” (25). During metaphase, chromosomes align at the metaphase plate and the cell shape also changes dramatically, bulging into a sphere (Figs. 4G-4I). Golgi haze appears rounded, with a shaded center where chromosomes are located (Fig. 4H). During anaphase, the duplicated chromosomes separate from one another and move to opposite poles of the spindle (Fig. 4K-4M). During telophase, chromosomes start to de-condense, and begin to take on a more interphase-like shape (Figs. 4L-4Q). In this stage, Golgi apparatus has also completed replication, and reassembled into two closely-spaced cell bodies, referred to as “Golgi twins” (25) (Fig. 4P). There is no DNA replication during mitosis, so the marker for proliferation (incorporation of the fluorescent nucleoside, EdU) are absent for the metaphase, anaphase, and telophase (Figs. 4J, 4N, 4R). As shown in Fig. 4C-4R, these structural, subcellular, cell cycle-dependent features are accurately captured and predicted by our DL model, which validates our hypothesis that label-free reflectance imaging and DL enables structural phenotyping.

Cell profile analysis on multiplexed images allows phenotyping and quantitative cytometry

A distinct advantage of multiplexing several markers is that it enables the development of multi-variant quantitative metrics for imaging cytometry and phenotyping across the large cell population and at the single-cell level. As a demonstration, we evaluate several cellular features, including cell morphology and fluorescence expressing intensity, on the digitally multiplexed readouts. Figures 5B-5C show scatter plots, similar to those used in the flow cytometry, of EdU vs. Hoechst fluorescence intensity for each cell analyzed from both the ground-truth images and the digitally multiplexed images. The DL-multiplexed prediction matches well with the ground truth, both of which show the increase in EdU and doubling of Hoechst intensity in the S and G2/M phases of the cell cycle, respectively.

In Figs. 5D-5K, we extract several biologically relevant single-cell profile metrics using the predicted IF labels and compare them with the ground truths (see Materials and Methods), as visualized in the violin plots. In particular, we show the statistics of eight different morphological and subcellular structural parameters. Additional metrics are provided in the
Supplementary Materials Fig. S6. In Fig. 5D-5E, we gather statistical data about the DNA label to measure the nuclear size and intensity variance. In Fig. 5F-5G, we evaluate the actin size (i.e. cell size) and its compactness. In Fig. 5H, we compute the nuclear- cytoplasmic ratio (NCR), an important marker for cancers, as the area ratio between the nucleus and actin. In Fig. 5I, we measure the endosome size from its IF label. In Fig. 5J-5K, we collect morphological parameters about the Golgi apparatus, including the eccentricity and concentration. For all these single-cell profile metrics, the prediction and the ground truth show excellent agreement. These results clearly demonstrate that our DL-augmented label-free cytometry can provide comprehensive morphological quantifications with high accuracy at the single-cell level, which is the key element for phenotyping and high-content screening (26).

**Saliency map reveals inner mechanism of the deep neural network**

Deep neural networks have shown high expressivity for complex models, but suffer from poor explainability. Many theoretical explanations for the deep learning model have resorted to statistical perspective while treating the overall model as a “black box”. Instead, we utilize the “attention”-based technique (24) to elucidate on the specific label-free subcellular features that contribute to the fluorescence prediction. To do so, we treat the DL model as a mapping function between an input and the corresponding output. We then visualize the network’s gradient with respect to the input and extract the salient features (i.e. those having the largest gradients) the network pays attention to (see details in Materials and Methods). The resulting “saliency map” highlights the most important features contributing to the prediction. By doing so, the saliency map directly evaluates the specificity of the structural features extracted from the reflectance images and how they are transformed to the target fluorescence labels by our network.

Figure 6 shows the computed saliency maps for each network across different sample batches and labeling conditions. Importantly, distinct subcellular features not only are highlighted by the network’s saliency map, but also have good correspondence to the targeted fluorescence label. By inspecting different columns, we show morphologically distinct features from different networks, indicating that different networks can indeed learn to recognize and focus on specific features present in the label-free images. For example, the DNA saliency maps show emphases on nuclear boundaries and some subcellular structures. The actin saliency maps show concentration over the whole cell and spreading out to the membrane boundaries. The saliency maps for Golgi apparatus generally form shapes in “partial-moon” or circular lines. By contrast, the saliency maps for proliferation and apoptosis show that the network selectively pays attention to certain features around the nuclei. In addition, the saliency maps show that our network learns to extract invariant structural features specific to the underlying fluorescence label regardless of the cell preparation and labeling processes. Across different rows, we observe consistent saliency maps under different sample batches / staining conditions for the same labeling network.

**Discussion**

In this study, we have presented a DL-augmented label-free cytometry technique that accurately predicted six fluorescence targets in-parallel at the single-cell level. The accuracy has been improved up to 3x in predicting subcellular structures as compared to the current state-of-the-art. Remarkably, the DL model is able to accurately recognize subcellular organelles, such as Golgi apparatus reconfigure during the cycle of proliferation, as well as to distinguish subtle morphological differences between the proliferating and non-
proliferating cells. These results demonstrate the data-driven model’s unique capability of holistically extracting “non-intuitive” structural features from the label-free imaging data on a large cell population. The specificity for cellular features by our DL model is illuminated by the saliency maps. This analysis demonstrated the ability of the DL models in processing highly complex and entangled structural information from scattering images.

Beyond predicting IF labels, we have further demonstrated quantitative cytometry analysis based on the multiplexed digital output from our DL models. Importantly, our analysis has shown that a multitude of single-cell profile metrics can be accurately extracted from the DL predictions. The digital multiplexing enabled us to simultaneously quantify several morphological features on multiple subcellular components across a large cell population. This capability drastically improves the technique’s throughput for structural phenotyping in the application of imaging cytometry, such as high-content analysis/screening. Cell morphological features are effective phenotypes for different disease states and environmental influences. This phenomenon is well described and practiced in pathology and cell biology. Nuclear condensation, enlargement, and increased environmental hallmarks of cancers (5, 6). Cell morphology is distinct for different cell types, which is often denoted in their terminology (i.e. astrocytes, macrophage, squamous and columnar cells, etc.), and stem cells change structures along separate differentiation paths (13). It has been shown that cell morphological changes can be directly associated with changes of morphogenic gene expressions (27), and comprehensive morphological profiling can be used to detect genetic functions (28). Meanwhile, it has been shown that DL techniques can holistically capture complex structural features for classification. This has found broad applications in detecting cell types (12–14), cell states (15–18, 22), drug response (19), and stem cell lineage (20). By fully leveraging the label-free and high multiplexing nature of our technique, it can potentially generate significant impacts in imaging cytometry by offering unprecedented information content and discovering new compound morphological features necessitating multiplexed fluorescence readout.

Label-free, DL-augmented method of cell-morphology profiling is data-driven and ultimately relies on the rich information content in the images. Our LED-array reflectance microscopy enables multi-contrast imaging (i.e. angle-dependent dark field and drDPC) by detecting the angled-dependent backscattering signals by a programmable LED array without any mechanical moving part. The superior sensitivity in detecting subtle structures using backscattering than transmission-microscopy is well documented (2). The superb sensitivity of backscattering-based method has been demonstrated in a variety of techniques, such as partial wave spectroscopy (29), confocal light absorption and scattering spectroscopic microscopy (30), confocal reflectance quantitative phase microscopy (31), and spatial-domain low-coherence quantitative phase microscopy (32). In addition, the angle-dependent measurement has been used to measure characteristic structural length scale (6) and to enable 3D reconstruction of the refractive index distribution (33). Leveraging angle-dependent reflectance signal, we outperformed the state-of-the-art for predicting multiple subcellular components. Recently, the LED array microscopy has also been extensively explored in transmission that allows sampling the low spatial frequency components in the Fourier space (33). In addition, backscattering spectroscopic techniques further enable characterization of ultrastructural phenotypes with sensitivity down to mm-length scale (34). A potential future improvement of our imaging system is to incorporate additional transmission and multispectral LED-array illumination to fully exploit the angle- and wavelength-dependent scattering contrast with a single objective lens by versatile illumination engineering.
One limitation of our current work is that it is based on fixed cells that does not allow longitudinal imaging. This can be overcome by using fluorescent reporter cell lines or live cell dyes to provide the fluorescence ground-truth (10) and enable dynamic observation. The additional temporal dimension may further improve the model’s sensitivity in cell phenotyping and discover new label-free features by incorporating the information about the cell dynamics (20, 22, 35).

In summary, we have reported a label-free imaging cytometry technique that multiplexes six IF labels in-parallel with high accuracy via DL models. We have validated the fluorescence predictions by comparing them to the ground-truth IF images. In addition, we have conducted imaging cytometry studies on several quantitative morphological metrics on sub-cellular structures and phenotyping of cell proliferation. Finally, the specificity of the DL model is assessed by visualizing the saliency map at the single cell level across different staining and fixation conditions. With this unique combination of new capabilities, this new framework may find wide applications in image-based cytometry, in particular for high-content screening and analysis.
Materials and Methods

Cell preparation and immunofluorescence staining
HeLa cells were cultured in a Dulbecco's Modified Eagle Medium (DMEM) with 10% fetal bovine serum (Gibco, 10564011) and 5% penicillin streptomycin (Gibco, 15140122). The cells were trypsinized and passaged twice a week. Two days before the staining and imaging, cells were cultured on glass-bottom petri dishes (FluoroDish FD35-100) which were first treated with 10% poly-L-lysine (SigmaAldrich, RNBG0769) with PBS for ten mins in an incubator. The staining and imaging were performed on the glass-bottom dishes.

We follow the standard IF staining protocols. In total, six IF stains are used to label DNA (Hoechst), actin (Phalloidin, Alexa Fluor 488 Phalloidin, Invitrogen, A12379), endosome (EEA1, Santa Cruz, sc-137130 AFF488), Golgi apparatus (GM130, Cell Signaling, 12480), proliferation (EdU, Click-iT Plus EdU Alexa Fluor 488 kit, Invitrogen), and apoptosis (TUNEL, Click-iT TUNEL Alexa Fluor 488 kit, Invitrogen). The HeLa cells were first fixed with ice-cold methanol, washed three times (10 min each) in 0.05% PBST (0.05% Triton X-100 PBS solution), and incubated for 20 mins at room temperature in a blocking solution containing 0.25% Triton X-100 and 10% bovine serum albumin in PBS. Alexa-488 conjugated antibodies were diluted in the blocking solution with the recommended concentration by the manufacturers and incubated with cells to label the specific subcellular components (EEA1 for endosome, Phalloidin for actin). For actin staining, cells were fixed with ice-cold acetone to preserve the structures. For Golgi staining, a secondary antibody Anti-rabbit IgG (Santa Cruz, 4412S) was diluted in blocking solution and used to culture the cells for 1.5 hours at room temperature in dark. To stain cell proliferation and apoptosis, we used EdU and TUNEL assays, respectively, according to the recommended protocol by Invitrogen. The apoptosis was induced by culturing the cells with 1µM Staurosporine for 24 hours. In all the above stains, cell nuclei were counterstained with 1× Hoechst 33342.

Image data acquisition
We collect the data using our custom-built multimodal reflectance microscope (8), as shown in Fig. 1A. A custom-built LED array consisting of two LED rings is used for providing controllable darkfield illumination in reflection. We use commercially available LEDs (APTF1616SEEZQGBDC, Kingbright) that can provide three independent RGB color channels (central wavelength is 460, 515, and 630 nm, respectively). All the LEDs are individually addressable using two cascaded LED drivers (TLC5955, Texas Instruments). A microcontroller (Teensy 3.2, PJRC) provides the camera trigger signal through digital Input/Output pins and simultaneously controls the LED illumination pattern. The LED array is mounted around the objective lens (10× 0.3 NA, UPlanFL N, Olympus, Japan) using a 3D-printed adapter. The tube lens is a commercial SLR lens (Nikon AF DC-NIKKOR 135 mm f/2D) to maximize the FOV. The microscope provides an overall 7.5× magnification. An sCMOS camera (CS2100M-USB, Thorlabs, 1920×1080 pixels, 5.04-µm pixel size, 16-bit depth) is used to acquire the images. We capture four darkfield reflectance images by using half-anulus green LED patterns along different orientations, including top ($I_{\text{Top}}$), bottom ($I_{\text{Bottom}}$), left ($I_{\text{Left}}$), and right ($I_{\text{Right}}$). The exposure time is 700 ms. Two drDPC images along two orthogonal orientations are generated by

$$I_{\text{DPC1}} = \frac{I_{\text{Top}} - I_{\text{Bottom}}}{I_{\text{Top}} + I_{\text{Bottom}}}$$

$$I_{\text{DPC2}} = \frac{I_{\text{Left}} - I_{\text{Right}}}{I_{\text{Left}} + I_{\text{Right}}}$$  \hspace{1cm} (1)
The fluorescence excitations are provided by two LED sources (M365LP1 and M470L4, Thorlabs, central wavelength 365 nm, 470 nm, respectively) combined with a dichroic mirror (DMLP425R, Thorlabs). The epi-fluorescence illumination is introduced by a 50/50 beam splitter (CCM1-BS013, Thorlabs). The emission filters (MF460-60, MF525-39, Thorlabs) are placed on a filter wheel (CFW6, Thorlabs) for blue and green fluorescence emissions. Two-channel fluorescence images are acquired sequentially after acquiring the reflectance images. The exposure time is 400 ms for IF imaging. Specifically, the first green channel is for one of the five IF antibodies conjugated with the green fluorophores (Alexa 488) for endosome, actin, Golgi apparatus, proliferation, and apoptosis; the second blue channel is for the co-stained DNA. We capture 30 image stacks for each sample batch / IF stain.

**Data preprocessing procedure**

The raw reflectance and fluorescence images are preprocessed before feeding into our networks for training the networks. The preprocessing procedure consists of four steps, including flat-field cropping, image denoising, background correction, and intensity normalization. Since the fluorescence excitation illumination is not evenly distributed across the entire rectangular FOV, we first perform flat-field cropping by using only the central 1000 × 1080-pixel region for training, where the excitation is approximately uniform. Second, we perform image denoising on the measurements. We apply two denoising approaches. In the first approach, we apply an unsupervised DL-based denoising algorithm, noise2void (36), to suppress the sensor noise present in the images. To do so, each 1000 × 1080-pixel image is cropped into 256 × 256-pixel patches. Each image patch is fed to a blind-spot network to perform denoising. After denoising, the patches are then stitched back together by alpha blending. This unsupervised denoising algorithm is found to be effective in removing unstructured, signal-independent noise, including the sensor noise and isolated hot pixels, in particular for measurements with low signal-to-noise ratios (SNRs). The total training and inference (denoising) procedure takes more than 10 hours for processing the entire dataset containing 30 images. We find this denoising procedure is only necessary for processing the Golgi and proliferation fluorescence images, as well as for the reflectance images for the actin prediction where the sensor noise severely corrupts the images. In the second approach, when the SNRs are sufficiently high for the measurements on other cell batches, we use a computationally more efficient morphology opening operation to remove the hot pixels in the fluorescence images under the assumption that hot pixels are isolated pixels with extreme intensity values. The opening operation takes a square kernel of size 2 × 2 pixels. This hot-pixel removal procedure takes ~15 min to process the entire dataset containing 30 images. Third, we perform background correction on the fluorescence images by eliminating the potential background bias across the batches. To do so, we calculate the histogram of each fluorescence image and denote the mode value (i.e. the most frequent value) as the constant background. This background of each fluorescence image is subtracted; the negative values from the subtraction are clipped to zero. Fourth, we perform intensity normalization by normalizing the pixel values of both the input and output images to be between 0 and 1. Additional details about the data preprocessing steps are provided in Supplementary Materials Fig. S2.

Our network takes 256 × 256-pixel input images. Accordingly, we split the 30 pairs of images into 256 × 256-pixel patches to generate the training and testing data. For each IF prediction, 512 training samples and 128 testing samples are randomly generated from the full-FOV image pairs. Each input stack consists of four different channels combining darkfield reflectance images from different oblique illumination patterns (Fig. 1A). In
addition, we construct the two-direction drDPC images from the normalized four-channel darkfield images (Fig. 1A) for predicting the actin labels since they highlight the cell boundaries. Example comparisons of the prediction results with and without the drDPC inputs are shown in Supplementary Materials Fig. S5. When making the full-FOV predictions (e.g. Fig. 4), we use the entire 1080 × 1920-pixel reflectance images since they do not suffer from the non-uniform illumination issue.

**Neural network implementation**

We develop a convolutional neural network (CNN) to learn the highly complex nonlinear mapping between the morphology information contained in the multi-channel reflectance images and the fluorescence labels. The network structure follows the encoder-decoder “U-net” architecture and further incorporates the dense-block and skip-connection structures to enable high-resolution information prediction (37). The input of the preprocessed 256 × 256-pixel reflectance image stack passes through the “encoder” path consisting of four dense blocks followed by the max-pooling layers, and the bottleneck feature maps are then fed into the “decoder” path with four dense blocks followed by upsampling layers. The skip connections bridge the lower-level activation maps with higher-level activation maps and preserves the high-frequency information. More details about the network are provided in Supplementary Materials Fig. S1. We use the negative Pearson correlation coefficient (NPCC) as the training loss (38). We train our network using ADAM optimizer with 500 epochs and 0.1 training/validation splitting. No overfitting is observed during the training.

**Quantitative evaluation of network prediction**

The evaluation metrics are computed on the testing sample image, each contains a 256 × 256-pixel FOV. In total, the statistics from 128 testing samples at each condition are aggregated and shown in the violin plots in Fig. 3. We use two evaluation metrics, including the PCC for the regression problems and the AUC for the detection problems.

The PCC is used to quantify the prediction quality for pervasive subcellular features, including DNA, endosome, actin and Golgi apparatus labels. It computes the statistical correlation between the predicted and ground-truth IF images and is able to quantify the similarity on the fine subcellular features. The PCC between the prediction \( X \) and the ground truth \( Y \) (each image is reshaped to a \( N \) dimensional vector) is computed as

\[
PCC = \frac{\sum_{i=1}^{N}(X_i - \bar{X})(Y_i - \bar{Y})}{\sqrt{\sum_{i=1}^{N}(X_i - \bar{X})^2} \sqrt{\sum_{i=1}^{N}(Y_i - \bar{Y})^2} + \epsilon}
\]

where \( \epsilon = 10^{-10} \) is a small regularizer to prevent zero denominator, \( \bar{X} \) denotes the mean, and \( i \) is the index of each vector. The value of PCC ranges from \(-1\) to \(+1\), where \( \pm 1 \) indicates total positive or negative correlation and 0 indicates no correlation. The PCC computation is implemented by a custom code in Python.

The AUC is used to quantify selective cell events in proliferation and apoptosis. To plot the ROC of the detection label performance, the ground-truth labels are first binarized with certain thresholds. After visual inspection to find the optimal threshold that preserves the most important morphological features, such as cell nuclei, the thresholds are set to be 0.4 and 0.05 for the proliferation and apoptosis label, respectively. The pixels in the predicted images take continuous values that are regarded as the predicted probability of label expression. By using the binarized ground-truth images as the target, our predictor (neural network) provides different pixel-wise True Positive Rate (TPR) and False Positive Rate
(FPR) under different detection thresholds on the predicted images. By varying the detection thresholds, TPR and FPR as functions of the thresholds can be plotted on the ROC curve. The AUC measures the area under the ROC curve and provides an aggregated quantification of performance across all possible detection thresholds. The AUC is computed by the built-in functions ‘roc_curve’ and ‘auc’ in scikit-learn module in Python.

**Digital cytometry analysis**

We develop a digital cytometry analysis framework for exploring the interdependencies of different fluorescence markers on the multiplexed predictions. A commonly used flow cytometry analysis is performed by displaying the scatter plot of single-cell level proliferating DNA concentration in the log-scale against the DNA concentration in the linear-scale. Different from flow cytometry that directly collects the integrated fluorescence intensity from each cell, our method performs imaging with subcellular resolution across a large cell population. As a result, we first perform segmentation and aggregation of the fluorescence signals for each cell region to carry out the single-cell digital cytometry analysis. We perform the digital cytometry analysis to relate the cell-level Hoechst and EdU fluorescence concentration using CellProfiler (39). Since our data contain co-registered two-channel fluorescence images with co-stained Hoechst and EdU, we can directly compare the ground-truth cytometry scatter plot with that from our multiplexed prediction. To generate the scatter plots, we feed the co-registered Hoechst and EdU images to CellProfiler. First, we segment the nuclei in the Hoechst image by the ‘IdentifyPrimaryObject’ module and extract the corresponding nuclei masks. We then apply the masks to both fluorescence images and calculate the integrated intensity inside the mask by the ‘MeasureObjectIntensity’ module. One confounding factor of this pipeline we find is that using the pure image data can mis-detect certain M-phase cells during the segmentation. Specifically, in anaphase and telophase (sub-phases in M phase) when a cell is about to split into two, the typical morphology is that a cell is attaching or intimately close to a nearby one. While common flow cytometry readout treats the two cells as one (since they are likely to be flown through the fluidic channel simultaneously) and gives the right readings of DNA and proliferating DNA expression, our digital cytometry does not with a plain segmentation algorithm. To overcome this issue, we refine the segmentation procedure by the following additional steps. First, we set the segmenting size to 24-35 pixels to find normal-sized nuclei that are prevalent in S and G1/G2 phases. Next, we set the segmenting size to 16-23 pixels to find small-sized nuclei that exist in the M phase. Third, we use the ‘merge-object’ module to combine the small-sized nuclei masks that are within a distance of 2 pixels into a single mask. Finally, we combine the merged masks and the normal-sized nuclei masks for computing the cell-level fluorescence intensities. We find this procedure successfully reduces the misdetection of M-phase nuclei and provides a more accurate scatter plot. For the multiplexed predictions, we apply the same pipeline on the predictions from the DNA and proliferation networks. The paired cell-level fluorescence intensity data are then plotted in Matlab. A small value (10^-10) is added to the fluorescence intensity of the proliferating DNA before taking the log operation to avoid singularity at 0. Three distinct clusters representing the S, G1, and G2/M phase are clearly shown in the digital cytometry scatter plots.

**Cell profile analysis**

We use CellProfiler (39) to generate the single-cell profiles across each entire fluorescence image. We feed the ground truth and the predicted IF images of DNA, actin, endosome and Golgi apparatus to CellProfiler. After initial cell segmentation, single-cell level parameters of morphology and intensity distribution are computed automatically by different
measurement modules in CellProfiler. Specifically, we use the ‘MeasureObjectSizeShape’ module to measure the fluorescence marker size (area), compactness, and eccentricity, the ‘MeasureObjectIntensity’ module to measure the fluorescence concentration, the ‘MeasureTexture’ module to measure the single-cell-level fluorescence variance and contrast. The fluorescence marker size is measured by the number of pixels in the segmented cell region. The compactness is computed by the mean squared distance of the cell mask pixels from the centroid divided by the area. The eccentricity is defined by the ratio of the distance between the foci of the effective ellipse that has the same second-moments as the segmented region and its major axis length. The concentration is computed as the sum of the intensities within the segmented masks. The contrast is measured by the local variation in a cell region. The variance is measured by the variation of the intensity values. The cell-level parameters are extracted from CellProfiler and then imported to Matlab to generate the violin plots. NCR is a compound metric that involves multiplexed DNA and actin fluorescence labels. To compute this metric, co-registered ground-truth / prediction images containing the DNA and actin labels are processed individually in CellProfiler. NCR is computed as the ratio between the area of segmented nuclei and actin masks.

**Saliency map visualization**

The saliency map is computed as the gradient map of the output with respect to the given input. Specifically, we use the visualization package in Keras to compute the gradients of the output layer with respect to Layer 137 using the seed from the first layer of the network (40). We compute the modified gradient to visualize the absolute values of the gradient, which shows the regions in the input that contribute most to the change in the output regardless of the sign of the change (i.e. negative or positive). We also use the guided backpropagation to propagate only the positive gradients for positive activations to achieve a smoother gradient visualization. The saliency map is calculated for each network and on different sample input (for varying sample batches / fixation conditions). The inputs are 256 × 256-pixel image stacks randomly selected from the testing groups under the six sample conditions. The computed saliency maps are normalized to have a uniform range between 0 and 1 for visualization.
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**Author contributions:** L.T. and J.Y conceived the idea. S.F. and Y.K. prepared the cell sample culturing, fixation, staining and acquired all the imaging data. S.C., Y. L. and Y. X. conducted the image processing, network training, cell phenotype and profile analysis, quantitative evaluation and saliency map analysis. W.S. developed the LED-array reflectance microscope platform. L.T., J.Y. and S.C. further discussed the results and refined the deep learning model, cell profiling and cytometry pipeline. All authors contributed to the writing of the manuscript.

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Fig. 1. Overview of the deep learning augmented label-free cytometry technique. (A) A multimodal LED-array reflectance microscope is developed to acquire co-registered label-free reflectance and fluorescence images. Reflectance images from oblique darkfield illumination and computed differential phase contrast (DPC) contain rich morphological information and are the multi-channel input to the DL model. Two-channel epi-fluorescence images are acquired on the same sample to serve as the ground truth for training our DL model. (B) Individual DL models are trained independently with paired label-free and IF images. The saliency map is used to reveal specific label-free features captured by the model to perform the transformation. (C) To perform digital multiplexed predictions, the same reflectance input is fed to each network and makes six different IF predictions in parallel.
**Fig. 2. Visualization of the results from the six IF prediction networks.** The rows show a sample dark-field reflectance image from each input stack, the network’s prediction, and the ground-truth IF image, respectively. The columns represent six IF labels covering four different subcellular features, including nuclei (DNA), endosome, actin, and Golgi apparatus, as well as two different cell states, including proliferation and apoptosis. The predictions have excellent visual agreement with the ground-truths in all six cases.
Fig. 3. Quantitative evaluation of the DL prediction. The above violin plots show the quantitative metrics of 128 testing samples from each cell batch, the upper and lower dashed lines represent the 75% quantile and 25% quantile values respectively, while the middle dashed lines indicate the median values. (A) The Pearson correlation coefficient (PCC) is used to evaluate the similarity between the regression-type predictions and ground-truth subcellular features, including nuclei (DNA), endosome, actin, and Golgi apparatus. (B) The Area under curve (AUC) is used to assess the detection accuracy of the predictions for cell proliferation and apoptosis.
Fig. 4. Multiplexed prediction on six IF labels from the same label-free input. (A) Visualization of the Full-FOV multiplexed prediction including DNA (blue), endosome (red), actin (green), and Golgi apparatus (yellow), and (B) proliferation (cyan) and apoptosis (magenta) from the same the reflectance input (grayscale). (C-R) Zoomed-in of DNA, Golgi apparatus, multiplexed, and proliferation predictions. White circles indicate representative cell morphology during different phases of cell cycle, including (C-F) interphase, (G-J) metaphase, (K-N) anaphase, and (O-R) telophase.
Fig. 5. Cell profile analysis on digital multiplexed IF staining. (A) An illustration of the cell cycle. (B-C) The scatter plots between the cell-wise EdU (proliferating DNA) and Hoechst (DNA) concentrations from (B) the co-stained ground-truth and (C) the DL-prediction across the entire cell population in the cell batch prepared under the proliferation staining condition. (D-K) The comparisons of the statistics of eight different single-cell profile metrics extracted from the entire cell population in the ground truth (GT) and DL-predictions (pred), including (D) the nuclear size, (E) the DNA (nuclear) fluorescence intensity variance, (F) the cell (actin) size, (G) the compactness of actin, (H) the nuclear-cytoplasmic ratio (NCR) measured by the area ratio between the nuclei and actin, (I) the endosome size, (J) the eccentricity of the Golgi apparatus distribution, and (K) the concentration of the Golgi apparatus. All the single-cell profile metrics show good agreements between the predictions and the ground truths.
**Fig. 6. Saliency maps from each network across six cell batches with different staining conditions.** The columns show the label-free input (the first channel dark-field reflectance image) and the saliency maps computed for six different IF labels, including DNA (blue), endosome (red), actin (green), Golgi apparatus (yellow), proliferation (cyan), and apoptosis (magenta). The rows show the label-free input and the saliency maps from six cell batches with different staining conditions. The saliency maps show good consistency across different batches and highlight distinct morphological features.