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## ARTICLE

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# Label-Free Imaging Flow Cytometry with Rare Cell Classification using Motion-Sensitive-Triggered Interferometry

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We present a label-free imaging flow cytometry system that integrates a microfluidic chip imaged by a motion-sensitive (event-based) camera and an interferometric phase microscopy module using a simple frame-based camera. The event camera captures activity from the flowing cells corresponding to thousands of frames per second and triggers the significantly slower interferometric camera when a rare cell, requiring more sensitive analysis, is detected via a single raw-interferogram analysis, significantly reducing data volume. The raw interferometric data serves as an input to a deep neural network for rare-cell classification. We demonstrate the use of this system to detect and grade rare cancer cells in blood, where the event camera is used to rapidly classify between the common white blood cells and the rare cancer cells, and the interferometric camera is used to grade the cancer cell type (primary/metastatic), as a human model for detecting and grading circulating tumor cells in liquid biopsies. This hybrid approach enables efficient data acquisition, rapid processing, and high sensitivity, significantly reducing computational load and is expected to find various applications in detecting and processing rare cells in imaging flow cytometry.

## Introduction

In recent years, motion-sensitive (event-based) sensors have emerged as bio-inspired imaging devices, mimicking the asynchronous and change-sensitive nature of the human retina. Unlike frame-based cameras that capture entire scenes at a fixed rate, event-based cameras independently record pixel level changes in brightness with high temporal resolution on the order of sub-microseconds and low latency<sup>1,2</sup>. These sensors detect an event when a significant change in light intensity occurs. This event contains the pixel address, the timestamp of the change of brightness, and the polarity, which indicates an increase or decrease in brightness<sup>1,3</sup>. In addition to high temporal resolution, event-based cameras generally acquire a highly sparse data stream compared to the dense pixel data produced by frame-based cameras<sup>1-4</sup>, significantly lowering the data load at any given time interval<sup>1,3</sup>. This leads to more efficient processing, as algorithms can be designed to operate directly on the asynchronous stream of events<sup>4</sup>. Typically, such algorithms rely on event aggregation or filtering within local spatial and temporal neighbourhoods, which is considerably less computationally demanding than processing full image matrices<sup>3</sup>. This sparse imaging nature of event data translates directly into reduced computational resource requirements. For these reasons, event cameras are well suited for detecting fast events, including moving objects.

Flow cytometry is a high-throughput technique capable of analysing up to hundreds of thousands of biological cells per second as they pass through laser beams. It can provide

information on cell size and internal complexity using forward and side light scatter, respectively<sup>5</sup>. While conventional flow cytometry lacks imaging capabilities, imaging flow cytometry combines high-throughput analysis with cellular imaging during flow, thus allowing morphological and functional profiling of single cells<sup>6-8</sup>. Event cameras are well suited for imaging flow cytometry since they generate events only when a cell passes through the sensor's field of view<sup>3,4</sup>.

To achieve molecular specificity and reliable identification in flow cytometry, fluorescence based labelling with antibodies is typically required<sup>9</sup>. This approach is widely used across immunology, cancer biology, and infectious disease research<sup>10-15</sup>, but relies on the presence of distinct surface markers. Challenges arise when target cells lack unique antigens or share markers with other cell types. Moreover, attaching labels to the cell membrane may alter cell behaviour or compromise measurement integrity<sup>16</sup>.

Label-free imaging methods overcome these limitations by leveraging intrinsic optical properties of cells. One such parameter, the refractive index, reflects intracellular composition, including the cell dry mass and concentration. Interferometric phase microscopy (IPM) noninvasively measures the optical path delay (OPD) profile of cells, or its integral refractive-index profile, enabling analysis of cell morphology and subcellular contents, without using external labelling agents<sup>17-20</sup>. We have previously shown that IPM is sensitive enough to grade cancer cells (healthy/primary cancer/metastatic cancer)<sup>21</sup>, even during flow on the background of blood cells<sup>22</sup>. IPM typically requires the acquisition of full-frame off-axis imaging interferograms, generally dictating a low throughput of tens to hundreds of frames per second when using modest-budget cameras.

Several high-throughput quantitative phase imaging strategies have been developed for rapid imaging flow cytometry. For example, multi-ATOM enables blur-free

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quantitative phase imaging of over 10,000 cells per second using asymmetric detection and real-time FPGA processing<sup>23</sup>, while Coherent-STEAM applies dispersive Fourier transform for high-speed, label-free phase mapping of cells in flow<sup>24</sup>. However, these techniques require complicated optical setups, such as laser scanning modules, and meticulous alignments, which are not suitable for seamless integration into existing imaging flow cytometry setups.

In this paper, we propose a new concept for rapid and efficient quantitative label-free imaging and grading of rare cells within liquid biological samples, by utilizing a rapid event-based camera acquiring the cells during flow in a microfluidic chip and triggering a slower interferometric frame-based camera when rare cells that require a more sensitive analysis are detected. We demonstrate this approach's potential for detecting circulating tumor cells (CTCs) in liquid biopsies obtained in routine lab tests by using a human model of blood spiked with colorectal cancer cells.

Finding cancer in its early curable stages represents a critical unmet need in oncology. Certain cancers, such as pancreatic and colorectal cancers, can progress without noticeable symptoms, thereby bypassing early diagnosis<sup>25,26</sup>. Liquid biopsies, such as peripheral routine blood samples, can be analysed to detect CTCs, as a means of detection and monitoring of cancer. Currently, however, this procedure's sensitivity for analysing and grading the cells remains limited due to the extremely low concentration of CTCs; for example, there are typically 1–10 cells per 10 mL of peripheral blood<sup>27–29</sup>. In our label-free imaging flow cytometry setup, detecting a cell can be accomplished by monitoring multiple cells during flow within a confined region and time window<sup>3,30</sup>, followed by immediate triggering activation of the frame-based camera for more sensitive IPM analysis for cell grading, as soon as a candidate cancer cell is detected by the event-based camera. IPM cell grading utilises a deep learning network, which is trained to classify between the rare cell types (primary/metastatic cancer), such that during inference, when IPM is triggered by the event camera, it can predict the rare cell type directly based on the raw interferograms, further increasing the process throughput.

Several sensor approaches have been previously explored to combine high-speed detection with high-resolution imaging. High-speed CMOS cameras can record at kilohertz rates, yet they require full-frame readout, producing terabytes of data quickly and heavy post-processing, which limits real-time scalability. Hybrid sensors like DAVIS346<sup>31</sup>, offer simultaneous events and frames but are constrained by lower dynamic range and reduced spatial resolution. Shared readout pathways further degrade temporal precision. These limitations underscore the need for our dual-sensor design, which preserves both sparse, low-latency detection and high-quality interferometric imaging.

## Methods and Materials

### Integrated Imaging Flow Cytometry Setup

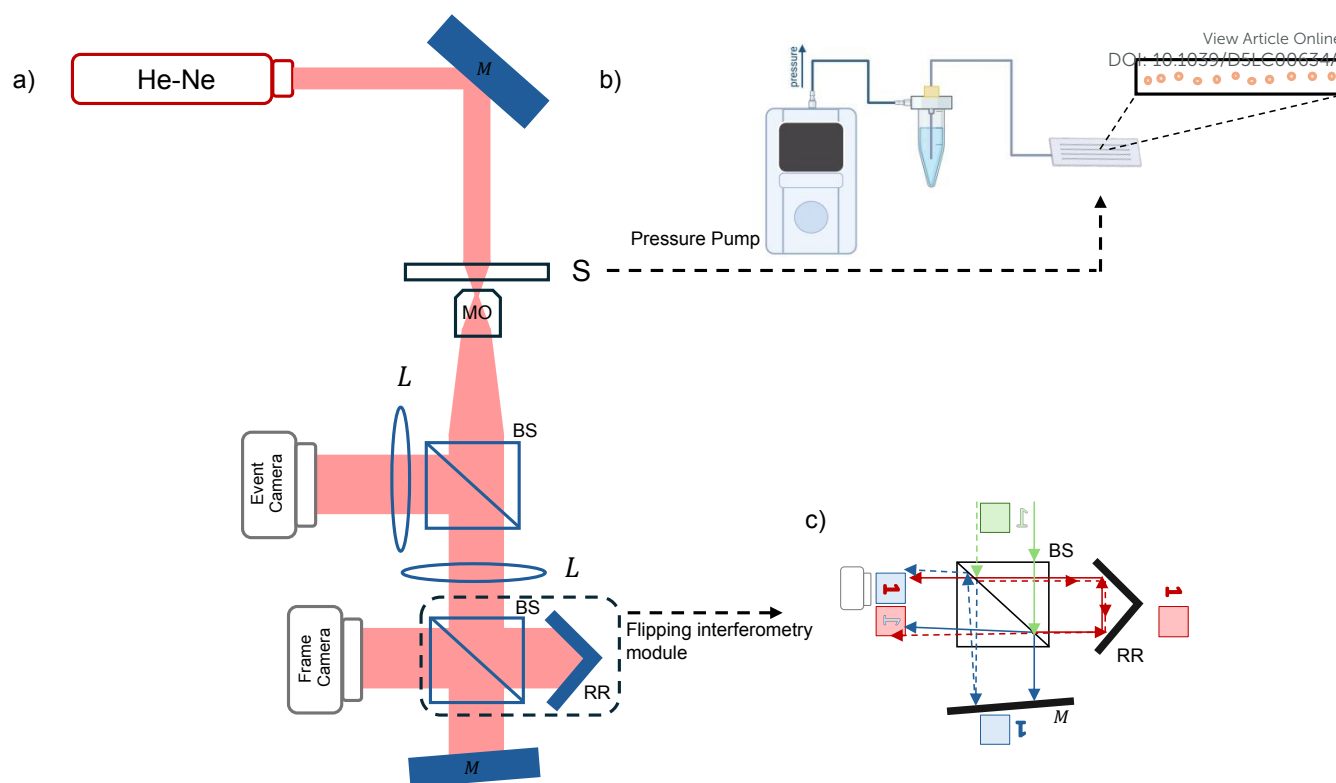
The proposed integrated imaging flow cytometry is shown in Fig. 1. For biological cell flow, we use a microfluidic chip (width: 100  $\mu\text{m}$ , height: 37  $\mu\text{m}$ , #10001444, ChipShop) containing a liquid sample with biological cells. The selected chip height minimizes cell focus variation across the field of view. A pressure pump (LU-FEZ-2000, LineUp Flow EZ, Fluigent) is used to control the flow rate, which is maintained between 5–15 millibar. The microfluidic chip is mounted on a three-axis stage, which enables fine alignment of the sample plane with the optical depth of focus. To further ensure focused imaging, the pressure is adjusted to maintain a steady flow velocity, so that cells move consistently within the focal plane on the floor of the channel. Under laminar flow conditions, cells are confined to a narrow z-range, further reducing the likelihood of defocusing.

The microfluidic chip is simultaneously imaged by two cameras, a rapid event camera and a simple CMOS camera. It combines an inverted microscope illuminated by a Helium-Neon (He-Ne) laser ( $\lambda = 632.8 \text{ nm}$ , 17 mW), which provides coherent light necessary for interferometric imaging. The laser beam is directed through the microfluidic chip positioned on a three-axes microscope stage. Cells flow in the microfluidic chip is induced via a pressure-controlled pump system. The event camera (IDS, UE-39BOXCP-E) is used for rapid preliminary cell detection and the frame-based CMOS camera (IDS, UI-3060CP-Rev.2) is used for further and more sensitive cell analysis via interferometric detection. In this setup, the event camera acts as a real-time trigger, detecting rare, rapidly flowing cells on localized bursts of events (sparse imaging). The event camera was configured with polarity thresholds of 80 for both ON and OFF events. When such a rare cell target is identified, the system is designed to initiate capture on the frame-based camera via the IPM module, recording off-axis imaging interferograms only upon these rare events, thus allowing much slower acquisition frame rate processing suitable for these rare events and just across the areas of the candidate cell. The optical path includes a 60 $\times$ , 0.85 numerical aperture (NA) infinity corrected microscope objective (Newport), which is shared by both the event-based and frame-based cameras, where a beam splitter positioned after the objective divides the magnified image between two imaging channels. Tube lenses positioned in the event-based and frame-based camera beam paths create total magnifications of 77 $\times$  and 70 $\times$ , respectively. This dual-path design allows both cameras to image the same scene with appropriate magnifications for their respective sensing modalities. The event camera captures the magnified image responses to motion only, via changes in brightness at the pixel level and outputs a continuous stream of events indicating the presence of moving cells. In our case, the event stream is grouped into temporal windows of  $\Delta t = 1000 \mu\text{s}$ , and we preliminarily detect rare cancer cells, which are larger than the background blood cells. This is done by converting the event stream into a binary image and applying morphological dilation to connect nearby events. The Density-Based Spatial Clustering of Applications with Noise (DBSCAN) algorithm is applied to

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**Fig. 1.** Integrated imaging flow cytometry setup combining event camera for high-throughput sparse imaging of flowing biological cells for preliminary detection of rare cells and triggering a slower interferometric camera for cell grading. a) The entire setup. b) The microfluidic chip and pump. c) The off-axis interferometric module (flipping interferometry) with beam tracing. BS, beam splitter; L, tube lens; M, mirror; MO, microscope objective; RR, retroreflector; S, sample.

group events into clusters representing individual cells. A diameter-based threshold of 180 pixels is then used to filter out small clusters and retain only those corresponding to larger cells. This threshold was selected based on the minimum cancer cell diameter measured in the training dataset and is consistent with the morphological filtering parameters (kernel size  $10 \times 10$  with 2 iterations). The value matches the chosen event accumulation window of  $\Delta t$  and could vary if a smaller temporal window is employed. Together with the selected temporal window, morphological dilation reduces missed detections by ensuring that weak or fragmented events are aggregated into candidate cells. Once a candidate cancer cell is identified, its centroid is computed and projected onto the frame-based camera coordinate space using a homography transformation derived from prior calibration. Thus, the frame-based camera is activated only when it receives a trigger that a rare cell is detected by the event camera. When multiple clusters exceed the threshold, each centroid is projected through the homography transformation to capture all relevant areas of interest simultaneously. Cell aggregates are filtered out using a maximum-size threshold and roundness criteria, as they are not considered reliable inputs for classification. Missed triggers of the interferometric camera are unlikely, as the inherent delay between event detection and trigger activation significantly exceeds the camera exposure time, ensuring readiness for the next trigger. This frame-based camera is positioned after a compact flipping interferometry (FI) module<sup>32</sup>. The module enables off-axis interferometric imaging by creating an interference pattern between the sample image and its flipped

version, thus allowing external interferometry that is mechanically stable. The FI module consists of a beam splitter that divides the magnified beam into two paths: one is directed toward a retro-reflector and the other toward a slightly tilted mirror. The interference between these two beams generates an off-axis image interferogram, containing amplitude and the quantitative phase map of the sample, capturing the sample morphology and content via its refractive index changes, thus allowing sensitive quantitative imaging without using chemical staining.

#### CTC Model, Cell Culture and Sample Preparation

The imaging flow cytometry concept explained above is relevant for detecting and grading rare cells on the background of common blood cells, where the event-based camera performs the initial classification to trigger the frame-based camera for more sensitive analysis and cell grading. We have chosen to demonstrate the system for detecting and grading of cancer cells on the background of white blood cells, as a human model of CTCs in liquid biopsy. For this, a pair of isogenic human colorectal adenocarcinoma cell lines was used: SW480 (ATCC CCL-228), derived from a primary tumor, and SW620 (ATCC CCL-227), derived from a metastatic lesion in a lymph node of the same patient. Cells were cultured in Dulbecco's Modified Eagle Medium (DMEM; ATCC, Cat. No. 30–2002) supplemented with 10% Fetal Bovine Serum (FBS; Biological Industries, Cat. No. 04-007-1A). Cultures were maintained at 37°C in a humidified incubator with 5% CO<sub>2</sub>. Cells were grown to approximately 80% confluence before harvesting. Prior to imaging, cells were





diluted in phosphate-buffered saline (PBS) at a concentration of  $2 \times 10^6$  cells/mL, which served as the buffer medium during the flow experiments. For preliminary data acquisition, cells were imaged in flow using both the frame-based CMOS camera and the event-based camera. The frame-based camera provided reference images for training and validation while the event camera images were used to explore data signatures and verify that the event-based data alone was not sufficient to distinguish between the two cell types, emphasizing the need for IPM.

For the actual CTC model, we used cancer-cell-spiked blood. Whole blood was obtained from the Israel National Blood Services, following the Tel Aviv University's IRB approval. First, red blood cells were depleted using a suitable depletion kit (EasySep, STEMCELL Technologies) following the manufacturer's protocol, leaving white blood cells (WBCs). To facilitate comparison between cancer cell types, two controlled samples were prepared: one spiked with SW480 cells and the other with SW620 cells, each at a 1:100 cancer cell to WBC ratio. These samples were then inserted into the microfluidic chip and imaged using the setup described above.

### Cancer Cell Classification using Interferometric Imaging

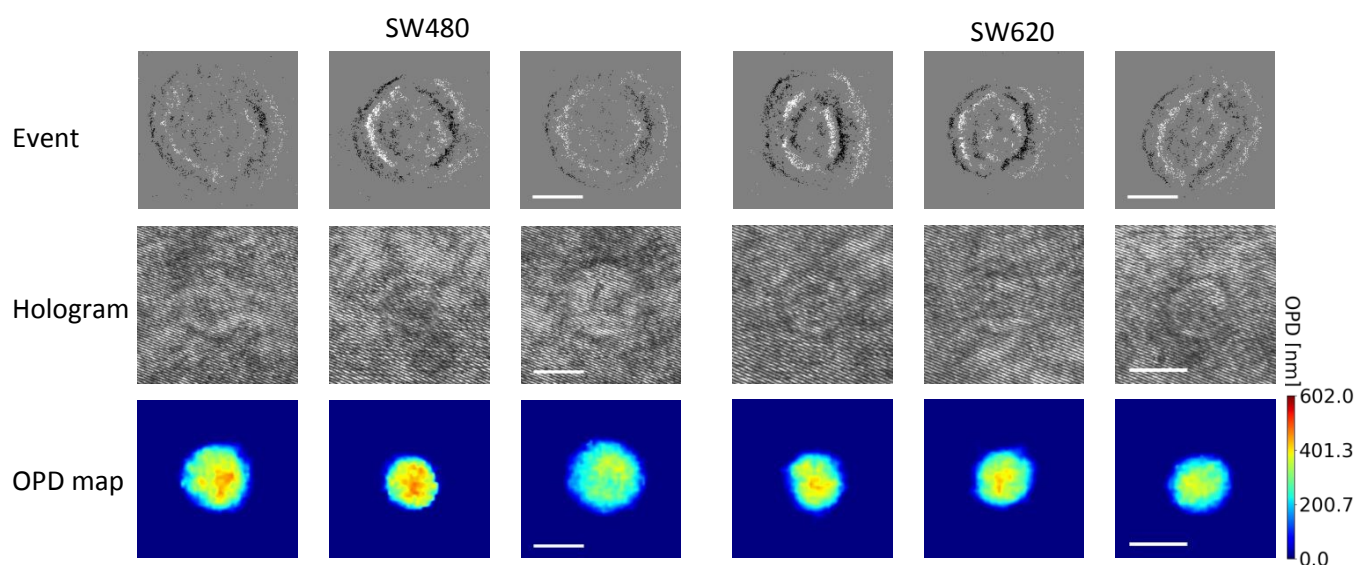
We collected raw off-axis interferograms of the two types of colorectal cancer cells for training and analysis. Following data acquisition, a convolutional neural network (CNN) based on the MobileNetV2 architecture. This model has previously been shown to outperform other commonly used CNNs for classifying similar colorectal cancer cell types<sup>33</sup>. We trained it for classification based on the acquired data. To further evaluate the system performance, a controlled sample was created by spiking WBCs with cancer cells. Cancer-cell detection on the background of WBCs was then performed using an event-based camera. The detected cells were subsequently analysed using the trained model to assess classification accuracy. The network

was trained directly on the raw interferograms without OPD extraction. A background interferogram frame was first prepared through averaging a sequence of frames without cells present and subtracted from the actual interferograms with the cells, enhancing cell visibility for segmentation. During training, cell label (primary/metastatic) was known since a one-type cell sample was imaged by the interferometric system. In this stage, 1400 images were collected for each cell type, and the database was divided into train and test sets in an 80:20 ratio, respectively. Since classification is done on the raw interferograms rather than on the OPD maps of the cells, the model was first trained on synthetic digital off-axis holograms having varying off-axis fringe spatial frequency, designed to improve robustness against interference patterns, thereby enhancing training efficiency and classification accuracy<sup>33</sup>. The model was trained using the Adam optimizer with a batch size of 32, 20 epochs, and a decreasing learning rate schedule to optimize convergence. All computations were performed on an Intel i7-10700 CPU. After training, the CNN can predict the cancer cell type based on the raw interferogram.

Figure 2 shows examples of cancer cells imaged by the event camera (first row), their off-axis interferogram as imaged by the CMOS camera through the interferometric module (second row), and their OPD extracted map (third row), as obtained by applying an off-axis hologram reconstruction process (although not used for cell classification)<sup>34</sup>.

### Results

The overall pipeline of our detection and classification is illustrated in Figure 3. The system is designed to operate in real time, beginning with rapidly capturing both WBCs and cancer cells in flow using an event-based camera.



**Fig. 2.** Primary cancer cells (SW480) and metastatic cancer cells (SW620) as imaged during flow by the event-based camera (first row), and off-axis image interferograms as imaged by frame-based camera through the interferometric module (second row). The corresponding OPD maps extracted from the interferograms are shown in the third row, although not used for cell classification. The white bar indicates 5  $\mu\text{m}$ . The color bar on the bottom right refers to the OPD maps.



The event data is processed to detect potential cancer cells based on spatial and temporal event patterns. Upon detection, a trigger signal activates the frame-based CMOS camera, which captures an off-axis hologram of the detected cell at a specific area of interest corresponding to the coordinates obtained from the event camera. This approach reduces the computational load by limiting processing to a 256×256-pixel region per cell, instead of the full 2-megapixel frame.

Supplementary Video S1 shows an illustration of the system at work, with both cameras imaging the cells at different frame rates. Table 1 summarizes the classification performance on the test set using different imaging modalities: event-based data, amplitude images, OPD maps, and raw off-axis interferograms. As expected, interferogram-based classification achieved the highest accuracy due to its ability to capture the cell quantitative phase information and amplitude information. Classification using the raw off-axis interferograms yielded perfect scores which can be attributed to the rich information content of the holograms, the use of a MobileNetV2 model pre-trained on a large synthetic dataset of the same cell types acquired with varying fringe frequencies (as detailed in the Methods), and we implemented 10-fold cross-validation. This approach ensured evaluation across diverse subsets, reducing overfitting risk and providing a robust estimate of model performance. In contrast, event-based data alone yielded limited performance, highlighting the importance of interferometric imaging for cancer cell classification, rather than just using the data acquired by the event camera for cancer cell classification. The event-based imaging did not miss detecting any cancer cell but could not classify the cancer cell type in high accuracy, necessitating utilizing interferometric-based classification. The average processing time required to detect a candidate cancer cell from the event stream was  $2.05 \pm 7.43$  milliseconds per frame, which enables near instant triggering of the frame-based interferometric camera.

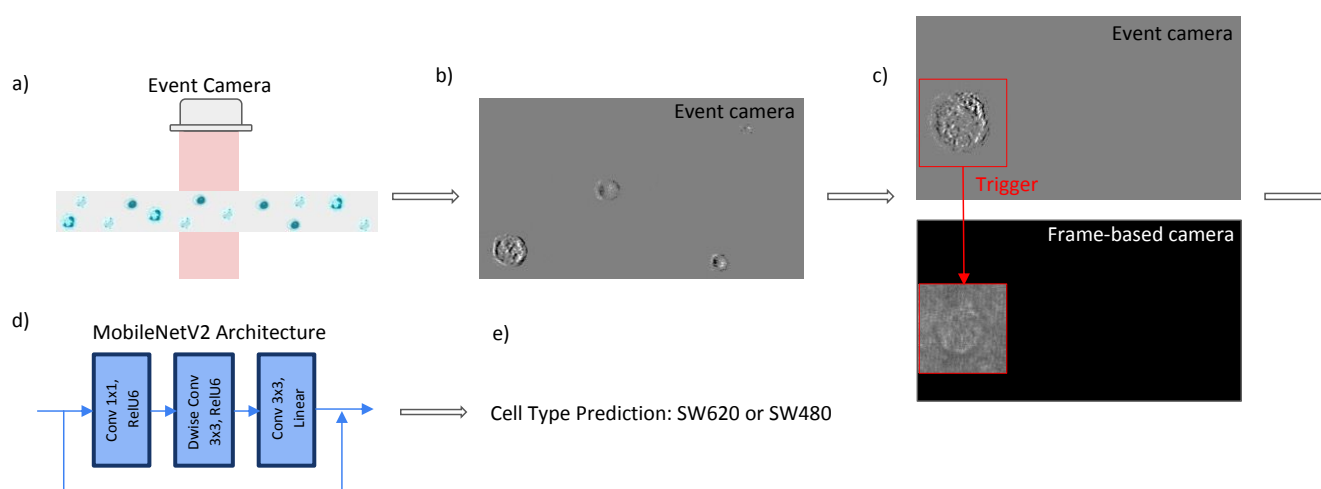
As part of our experimental validation, we tested the system on blood samples spiked with the known cancer cell type. Each sample contained a defined ratio of cancer cells to white blood cells (1:100). The validation consisted of two independent experiments: blood spiked with SW620 cells, and blood spiked with SW480 cells, enabling controlled evaluation.

Table 2 presents the number of WBCs and cancer cells detected in each controlled sample, along with the corresponding classification accuracy of the CNN model when given the respective interferograms as input after projecting the coordinates of the detected cancer cells to the interferogram coordinate space.

To further assess the classification performance based on the event data alone, the same cells identified in the event stream were also fed into a MobileNetV2 architecture model that was trained on event-derived input. Notably, these cells were misclassified in approximately 38% of cases, suggesting that the event-based data lacks the robustness required for accurate classification of complex cell types, and that the more sensitive imaging by IPM is needed to grade the rare cells.

## Conclusions

We introduced a hybrid system of imaging flow cytometry for rare cell detection and classification that combines the high temporal resolution of an event-based camera via sparse imaging for preliminary screening and a more sensitive secondary analysis for rare cell grading via off-axis IPM using a low temporal resolution frame-based camera. This integration enables real-time detection of rare, fast-moving cells in flow, triggering IPM only when needed. This selective acquisition approach reduces redundant data and minimizes computational load, offering a practical high-throughput solution for rare cell quantitative imaging flow cytometry.



**Fig. 3.** Imaging flow cytometry pipeline of the detection and classification of cancer cell types on the background of WBCs. a) WBCs and cancer cells are captured during flow using the event camera. b) The event camera data is processed to generate images only when detection of cancer cells occurs. c) Cancer cells are detected by the event camera based on size differences compared WBCs. The frame-based camera is triggered to capture an off-axis interferogram of the detected cancer cells by imaging the projected area of interest. e) The off-axis interferogram of a triggered-area containing a cancer cell is used as input for cell classification via a MobileNetV2 deep learning model. f) The output is a prediction of the cancer cell type (primary (SW480)/metastatic (SW620)).



**Table 1.** Performance metrics for cancer cell line classification based on the test set.[View Article Online](#)

Data type	Accuracy	Label	Precision	Recall	F1-score
Off-axis interferogram	1	SW480	1	1	1
		SW620	1	1	1
OPD	0.94	SW480	0.98	0.90	0.94
		SW620	0.91	0.98	0.95
Amplitude	0.90	SW480	0.90	0.88	0.89
		SW620	0.89	0.91	0.90
Event-based image	0.90	SW480	0.83	1	0.91
		SW620	1	0.80	0.89

**Table 2.** Detection counts and classification accuracy for WBCs and cancer cells in various spiking ratios

Sample Label	CTCs Detected	WBCs Detected	Detected Ratio (WBCs:CTCs)	Classification Accuracy
SW480	80	11,252	0.71:100	100%
SW620	160	10,833	1.48:100	100%

The processing time to detect a candidate cancer cell is influenced primarily by the number of activated pixels in each event frame, which scales with the spatial extent of the detected cells. In particular, larger cells generate denser event clusters, resulting in increased computational load. Notably, all processing was performed on a standard CPU, suggesting that further reductions in latency could be achieved through GPU acceleration or algorithmic optimization. These improvements would enhance real-time compatibility in high-throughput applications. Importantly, the trained model provided accurate predictions for every controlled sample, correctly distinguishing the cancer cell types. This successful classification between primary and metastatic colorectal cancer cell lines provides meaningful morphological insights that could be used for diagnosis of cancer from routine blood test, be used for cancer monitoring to inform personalized treatment decisions. In this proof-of-concept implementation, the acquisition of the cells was applied in real time, but the triggering mechanism was applied offline to retrospectively identify and extract relevant frames with rare cells using recorded event data, effectively demonstrating the potential to reduce redundant data acquisition. Validation was performed through two independent spiking experiments at a 1:100 cancer to blood cell ratio: one with SW620 cancer cells and another with SW480 cancer cells. These controlled experiments provided initial evidence of reproducibility and feasibility, while future clinical evaluation should assess performance at the lower CTC levels typically observed (<10 cells/10 mL), in addition to conducting further biological replicates. Future real-time triggering is possible by opening a specific area of interest via the application programming interface (API) of the frame-based camera. The event camera processing window was configured to match a 30 frames-per-second acquisition rate of the frame-based camera.

However, the system is designed to support faster cameras by adjusting  $\Delta t$  of the event-based camera accordingly in order to enable high-throughput operation driven by the event-based trigger. Beyond cancer detection in liquid biopsies, this framework opens new possibilities for rare event identification in biomedical and industrial applications, including stem cell studies for personalized medicine. The suggested approach's ability to detect rare phenomena with minimal processing supports the development of compact, low-power diagnostic tools suited for point-of-care or resource-limited settings, especially when label-free imaging is used. Apart from biomedical uses, the suggested dual-sensor approach could support various high-speed monitoring tasks such as industrial quality control through vibration analysis or analysis for fluid dynamics by triggering high-resolution imaging only during relevant events, thereby reducing data load while capturing key information.

In summary, the proposed system offers a fast, label-free, and computationally efficient quantitative imaging flow cytometry method for rare-cell detection and classification, with strong potential to impact early cancer diagnostics, as well as broader rare-event sensing domains.

## Author contributions

**Eden Dotan:** Conceptualization; methodology; software; formal analysis; writing – original draft; data curation; investigation. **Dana Yagoda-Aharoni:** Formal analysis; methodology; Writing – review & editing. **Eli Shapira:** formal analysis; methodology; software; **Natan T. Shaked:** Conceptualization; methodology; investigation; writing – original draft; supervision; resources.



## Conflicts of interest

There are no conflicts to declare.

## Data Availability

Data for this article, consisting of the training dataset for SW480 and SW620 cell classification using digital holography and event-based imaging, are available at Zenodo under the following DOI: <https://doi.org/10.5281/zenodo.15723128>

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## References

- G. Gallego, T. Delbruck, G. Orchard, C. Bartolozzi, B. Taba, A. Censi, S. Leutenegger, A. J. Davison, J. Conradt, K. Daniilidis and D. Scaramuzza, *IEEE Trans. Pattern Anal. Mach. Intell.*, 2022, **44**, 154–180.
- P. Lichtsteiner, C. Posch and T. Delbruck, *IEEE J. Solid-State Circuits*, 2008, **43**, 566–576.
- J. Howell, T. C. Hammarton, Y. Altmann and M. Jimenez, *Lab Chip*, 2020, **20**, 3024–3035.
- Z. Zhang, H. Yang, J. Li, S. W. Chong, J. K. Eshraghian, K.-T. Yong, D. Vigolo, H. M. McGuire and O. Kavehei, *Cell Biology*, 2024, preprint, DOI: 10.1101/2024.11.16.623904.
- A. L. Givan, in *Methods in Cell Biology*, Elsevier, 2001, vol. 63, pp. 19–50.
- S. D. Tanner, V. I. Baranov, O. I. Ornatsky, D. R. Bandura and T. C. George, *Cancer Immunol Immunother*, 2013, **62**, 955–965.
- M. Doan, I. Vorobjev, P. Rees, A. Filby, O. Wolkenhauer, A. E. Goldfeld, J. Lieberman, N. Barteneva, A. E. Carpenter and H. Hennig, *Trends in Biotechnology*, 2018, **36**, 649–652.
- N. T. Shaked, S. A. Boppart, L. V. Wang and J. Popp, *Nat. Photon.*, 2023, **17**, 1031–1041.
- M. Watanabe, Y. Uehara, N. Yamashita, Y. Fujimura, K. Nishio, T. Sawada, K. Takeda, F. Koizumi and Y. Koh, *Cytometry Pt A*, 2014, **85**, 206–213.
- R. E. Duque, M. Andreeff, R. C. Braylan, L. W. Diamond and S. C. Peiper, *Cytometry*, 1993, **14**, 492–496.
- M. Brown and C. Wittwer, *Clin Chem*, 2000, **46**, 1221–1229.
- L. G. Dressler and S. A. Bartow, *Semin Diagn Pathol*, 1989, **6**, 55–82.
- I. M. Bromilow and J. K. M. Duguid, *Clinical & Laboratory Haematology*, 1997, **19**, 137–142.
- B. H. Davis, S. Olsen, N. C. Bigelow and Jc. Chen, *Transfusion*, 1998, **38**, 749–756.
- J. M. Navenot, T. Merghoub, R. Ducrocq, J. Y. Muller, R. Krishnamoorthy and D. Blanchard, *Cytometry*, 1998, **32**, 186–190.
- E. Heitzer, M. Auer, C. Gasch, M. Pichler, P. Ulz, E. M. Hoffmann, S. Lax, J. Waldispuehl-Geigl, O. Mauermann, C. Lackner, G. Höfler, F. Eisner, H. Sill, H. Samonigg, K. Pantel, S. Riethdorf, T. Bauernhofer, J. B. Geigl and M. R. Speicher, *Cancer Research*, 2013, **73**, 2965–2975.
- M. Cristofanilli, G. T. Budd, M. J. Ellis, A. Stopeck, J. Matera, M. C. Miller, J. M. Reuben, G. V. Doyle, W. J. Allard, L. W. M. M. Terstappen and D. F. Hayes, *N Engl J Med*, 2004, **351**, 781–791.
- T. Yamauchi, H. Iwai and Y. Yamashita, *Opt. Express*, 2011, **19**, 5536. DOI: 10.1039/D5LC00634A
- M. Haifler, P. Girshovitz, G. Band, G. Dardikman, I. Madjar and N. T. Shaked, *Fertility and Sterility*, 2015, **104**, 43–47.e2.
- N. T. Shaked, L. L. Satterwhite, N. Bursac and A. Wax, *Biomed. Opt. Express*, 2010, **1**, 706.
- D. Roitshtain, L. Wolbromsky, E. Bal, H. Greenspan, L. L. Satterwhite and N. T. Shaked, *Cytometry Pt A*, 2017, **91**, 482–493.
- N. Nissim, M. Dudaie, I. Barnea and N. T. Shaked, *Cytometry Pt A*, 2021, **99**, 511–523.
- K. C. M. Lee, M. Wang, K. S. E. Cheah, G. C. F. Chan, H. K. H. So, K. K. Y. Wong and K. K. Tsia, *Cytometry Pt A*, 2019, **95**, 510–520.
- A. Mahjoubfar, C. Chen, K. R. Niazi, S. Rabizadeh and B. Jalali, *Biomed. Opt. Express*, 2013, **4**, 1618.
- R. C. Bast, B. Hennessy and G. B. Mills, *Nat Rev Cancer*, 2009, **9**, 415–428.
- A. Makohon-Moore and C. A. Iacobuzio-Donahue, *Nat Rev Cancer*, 2016, **16**, 553–565.
- A. Pulumati, A. Pulumati, B. S. Dwarakanath, A. Verma and R. V. L. Papineni, *Cancer Reports*, 2023, **6**, e1764.
- C. Alix-Panabières and K. Pantel, *Nat Rev Cancer*, 2014, **14**, 623–631.
- W. J. Allard, J. Matera, M. C. Miller, M. Repollet, M. C. Connelly, C. Rao, A. G. J. Tibbe, J. W. Uhr and L. W. M. M. Terstappen, *Clinical Cancer Research*, 2004, **10**, 6897–6904.
- S. Takatsuka, N. Miyamoto, H. Sato, Y. Morino, Y. Kurita, A. Yabuki, C. Chen and S. Kawagucci, *Ecology and Evolution*, 2024, **14**, e70150.
- C. Brandli, R. Berner, Minhao Yang, Shih-Chii Liu and T. Delbruck, *IEEE J. Solid-State Circuits*, 2014, **49**, 2333–2341.
- D. Roitshtain, N. A. Turko, B. Javidi and N. T. Shaked, *Opt. Lett.*, 2016, **41**, 2354.
- M. Dudaie, I. Barnea, N. Nissim and N. T. Shaked, *Sci Rep*, 2023, **13**, 12370.
- E. Cuche, P. Marquet and C. Depeursinge, *Appl. Opt.*, 1999, **38**, 6994.





## Data availability

Data for this article, consisting of the training dataset for SW480 and SW620 cell classification using digital holography and event-based imaging, are available at Zenodo under the following DOI: <https://doi.org/10.5281/zenodo.15723128>

