SAT: Segment and Track Anything for Microscopy

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Abstract: Integrating cell segmentation with tracking is critical for achieving a detailed and dynamic understanding of cellular behavior. This integration facilitates the study and quantification of cell morphology, movement, and interactions, offering valuable insights into a wide range of biological processes and diseases. However, traditional methods rely on labor-intensive and costly annotations, such as full segmentation masks or bounding boxes for each cell. To address this limitation, we present SAT: Segment and Track Anything for Microscopy, a novel pipeline that leverages point annotations in the first frame to automate cell segmentation and tracking across all subsequent frames. By significantly reducing annotation time and effort, SAT enables efficient and scalable analysis, making it well-suited for large-scale studies. The pipeline was evaluated on two diverse datasets, achieving over 80% Multiple Object Tracking Accuracy (MOTA), demonstrating its robustness and effectiveness across various imaging modalities and cell types. These results highlight SAT's potential to streamline biomedical research and enable deeper exploration of cellular behavior.

1 INTRODUCTION

Cell tracking is essential in biology and medicine, offering insights into cellular behavior and responses to stimuli (Newman et al., 2011). In cancer research, cell tracking aids in studying tumor growth, metastasis, and the efficacy of anti-cancer drugs, while in stem cell research, it helps observe differentiation and

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regenerative potential (Aramini et al., 2022). This technique is vital in drug development for assessing drug impact and efficacy and in immunology for understanding immune cell interactions and responses (Yazdi and Khotanlou, 2024). Accurate cell segmentation is essential for tracking, providing data to monitor cell movement and behavior over time (Chou et al., 2023). Without precise segmentation, tracking algorithms may misidentify cells, leading to errors. The importance of cell segmentation lies in its ability to quantify cell morphology, analyze cellular interactions, and support high-throughput screening in drug development (Durkee et al., 2021). Additionally, it aids in understanding developmental processes and immune responses by characterizing specific cell populations (Padovani et al., 2022).

• Introduction of the SAT (Segment and Track Anything for Microscopy) pipeline, which leverages

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Mathad	Time per Cell	per Frame (s)	Total Time (A+T)	Times Faster than		
Methou	Segmentation/Point	Tracking (T)	for 100 Frames	Full Mask (x)		
	Annotation (A)		(min)			
Full Mask and	46	0.438	77.40	-		
Tracking						
SAT (N=3)	$3 \times 0.9 = 2.7$	0	0.225	≈ 344		
SAT (N=5)	$5 \times 0.9 = 4.5$	0	0.375	≈ 206		
SAT (N=10)	$10 \times 0.9 = 9$	0	0.75	≈ 103		

Table 1: Comparison of supervision time between Full Mask and Tracking, and the SAT method. The SAT method involves point annotation only in the first frame. SAT (5 points per cell) saves significant time compared to full mask tracking, making it approximately 206 times faster.

point annotations in the first frame to automate cell segmentation and tracking, significantly reducing the time and effort required compared to traditional methods.

- Comprehensive evaluation of the SAT pipeline on subsets of two extensive and diverse cell tracking datasets: the Cell Tracking Challenge (CTC) (Maška et al., 2023) and the Cell Tracking with Mitosis Detection Challenge (CTMC) (Anjum and Gurari, 2020) datasets, demonstrating the method's robustness and generalization capability.
- Achieving high tracking accuracy, with Multiple Object Tracking Accuracy (MOTA) exceeding 80%, and demonstrating time savings of over 100 times compared to full mask annotation methods.

The remainder of this paper is structured as follows: Section 2 reviews the existing literature and challenges in cell segmentation and tracking. Section 3 introduces the SAT pipeline, detailing its design and functionality. Section 4 describes the datasets utilized in the study. Section 5 presents the metrics used for performance assessment, followed by Section 6, which outlines the experimental setup for evaluating the proposed pipeline. Section 7 provides a detailed analysis and discussion of the results. Finally, Section 8 concludes the paper and suggests directions for future research.

2 LITERATURE REVIEW

2.1 Existing Cell Segmentation and Tracking Approaches

There are numerous studies on cell segmentation and tracking (Edlund et al., 2021; Stringer et al., 2020; Jelli et al., 2023; Khalid et al., 2023; Maška et al., 2023) which require full masks for training or some form of weak supervision. Traditional segmentation-first approaches often focus on generating accurate

segmentation masks for individual cells, followed by linking these segmented regions across time frames to produce tracking results (Malin-Mayor et al., 2023). These methods are effective in high-contrast, wellannotated datasets but face limitations when generalizing to new imaging modalities or less annotated domains. Recent efforts have sought to reduce reliance on full segmentation masks by incorporating weakly supervised approaches, which use partial annotations such as bounding boxes or point annotations. For example, Khalid et al. (Khalid et al., 2022) introduced a method using only bounding boxes and point annotations, significantly reducing annotation time and resources. These methods offer a compromise between efficiency and accuracy but often require substantial manual input or specific pre-trained models, which can limit their adaptability. Unsupervised tracking methods have also emerged as promising alternatives, leveraging temporal patterns and unsupervised learning to track cells without labeled training data (Maška et al., 2023). While these methods reduce annotation effort, they struggle with accurate segmentation, particularly in noisy or overlapping cell scenarios. The continued development of these approaches highlights the field's ongoing search for effective yet resource-efficient solutions to cell tracking challenges.

2.2 Alternative Cell Identification and Tracking Methods

While segmentation-based approaches dominate cell tracking methodologies due to their ability to capture detailed morphological and spatial information, some recent methods bypass segmentation entirely. For example, Romphosri et al. (Romphosri et al., 2024) introduced an alignment-free bacteria identification technique using optical scattering with LEDs and YOLO (Wang et al., 2023a), demonstrating rapid cell identification without segmentation. Similarly, Matthews et al. (Matthews et al., 2024) utilized YOLO and digital holographic microscopy for real-

time 3D tracking of microbes, achieving effective results without requiring human labeling. However, these methods are primarily tailored for specific tasks like bacterial identification or microbe tracking and may not generalize to applications requiring precise morphology analysis or complex cell interactions. Segmentation provides pixel-level detail, essential for analyzing cellular morphology, quantifying interactions, and supporting high-throughput studies. For these reasons, segmentation remains a more versatile and detailed approach in microscopy applications.

2.3 Challenges in Microscopy Applications

Microscopy images pose unique challenges that differ significantly from natural images, making it difficult for conventional image processing and machine learning models to perform effectively (Wang et al., 2023b). One primary challenge is the variability in imaging modalities, such as phase contrast, fluorescence, and differential interference contrast microscopy. These modalities exhibit differences in contrast, clarity, and noise levels, requiring models to adapt dynamically to each imaging condition (Stringer et al., 2020). Another major challenge is the variability in cell shapes, sizes, and morphologies across different biological contexts. Models trained on specific datasets often struggle to generalize to new cell types or experimental conditions, necessitating extensive retraining (Yazdi and Khotanlou, 2024). Furthermore, microscopy images are prone to artifacts and background noise, particularly in lowcontrast environments, which can mislead segmentation and tracking algorithms. The scalability of these models is also a critical issue. Large-scale datasets, such as those used in high-throughput screening or longitudinal studies, require methods that are not only accurate but also computationally efficient. Addressing these challenges is vital to improving the applicability and reliability of cell segmentation and tracking methods in real-world microscopy applications.

2.4 Limitations of the Segment Anything Model (SAM)

The Segment Anything Model (SAM) (Kirillov et al., 2023) by MetaAI performs well on natural scenes but struggles with microscopic images due to their complexity, low contrast, and noise (Archit et al., 2023). Domain-specific training and pre-processing are often required to adapt SAM for microscopy tasks. Compared to models like YOLO (Wang et al., 2023a), which can also perform segmentation, SAM's point-

based annotation scheme offers a distinct advantage by reducing manual effort while maintaining pixellevel precision. This efficiency makes SAM particularly suitable for large-scale and high-throughput studies in microscopy. Based on these limitations and strengths, the proposed work leverages SAM's capabilities while addressing its shortcomings by utilizing point annotations in the first frame to automate segmentation and tracking across all frames.

3 SAT: SEGMENT AND TRACK ANYTHING PIPELINE

The key technical component of the proposed pipeline is the Segment Anything Model (SAM). SAM was pre-trained on diverse images and fine-tuned using the LIVECell dataset for microscopy. It includes an image encoder, prompt encoder, and mask decoder, which work together to produce accurate segmentation masks from point prompts. The SAT (Segment and Track Anything for Microscopy) pipeline is divided into four main components: Query Points Selection, Point Tracking, Segmentation, and Point Tracking Reinitialization (Rajič et al., 2023). Below is a detailed explanation of each module, referring to Figure 3.

3.1 Query Points Selection

In the first step of SAT, query points are selected in the first video frame to denote the target object (positive points) and non-target regions (negative points). The user can provide these points interactively or derive them from a ground truth mask using various sampling techniques, including Random Sampling, K-Medoids Sampling, Shi-Tomasi Sampling (Shi et al., 1994), and Mixed Sampling. Each method ensures good coverage and robustness, significantly affecting the model's performance. Among the three possible methods described, this study employs the second approach, where point annotations in the first frame are used to initialize segmentation and tracking for subsequent frames. This method was chosen due to its balance between annotation efficiency and segmentation accuracy, making it highly suitable for large-scale datasets and diverse microscopy conditions.

$$P = \{p_1, p_2, \dots, p_n\}$$
(1)

For K-Medoids, let C represent the set of clusters, and P_m be the medoid points:

$$P_m = \{ \operatorname{medoid}(c_i) \mid c_i \in C \}$$
(2)



Figure 1: SAT (Segment and Track Anything for Microscopy) Pipeline. The SAT pipeline extends image segmentation models to microscopy videos through four steps: **A. Query Points Selection**, where positive and negative points are defined by the user or a ground truth mask; **B. Point Tracking**, which propagates points across video frames using point trackers, predicting trajectories and occlusion scores; **C. Segmentation**, where the Segment Anything Model (SAM) uses these trajectories to generate per-frame mask predictions; and **D. Point Tracking Reinitialization**, an optional step to reinitialize query points, improving tracking reliability and addressing newly visible cell segments.

The objective function to minimize K-Medoids clustering is:

minimize $\sum_{i=1}^{k} \sum_{p \in c_i} \|p - \text{medoid}(c_i)\|$ (3)

3.2 Point Tracking

This module propagates the selected query points across all video frames using point trackers. This propagation generates point trajectories and occlusion scores, ensuring that the points follow the objects throughout the video. Point tracker, PIPS (Harley et al., 2022) is employed due to its robustness in handling long-term tracking challenges such as occlusion and reappearance of objects.

$$P_t = \{p_{t,1}, p_{t,2}, \dots, p_{t,n}\}$$
(4)

The tracking function T predicts the position of points in the next frame:

$$P_{t+1} = T(P_t) \tag{5}$$

3.3 Segmentation

Using the point trajectories obtained from the tracking module, the Segment Anything Model (SAM) (Kirillov et al., 2023), which is finetuned on the LIVECell dataset, generates per-frame segmentation masks. The SAM model, which comprises an image encoder, a prompt encoder, and a mask decoder, utilizes the non-occluded points as prompts to segment the object of interest in each frame accurately.

$$M_t = \mathrm{SAM}(I_t, P_t) \tag{6}$$

where I_t is the input image at frame t, and P_t is the set of propagated points.

3.4 Point Tracking Reinitialization

This step involves reinitializing the query points periodically using the predicted masks. Reinitialization helps to remove unreliable points and add new points to object segments that become visible in later frames, thereby improving the accuracy and robustness of the segmentation over time.

$$P_t = \text{Reinitialize}(M_t) \tag{7}$$

Reinitialization occurs at intervals defined by the prediction horizon *h*:

$$P_{t+h} = \text{Reinitialize}(M_{t+h}) \tag{8}$$

While SAT integrates existing modules like SAM for segmentation and PIPS for point tracking, its novelty lies in optimizing these components specifically for microscopy images. By minimizing the manual effort with point annotations in the first frame and automating the rest of the segmentation and tracking process, SAT improves both accuracy and efficiency. Additionally, the reinitialization step enhances robustness in tracking, and addressing occlusions and the appearance of new cells over time.

Table 2: Statistics of the LIVECell dataset used for finetuning the Segment Anything Model.

Detect		Train		Val	Test		
Dataset	Img	Cells	Img Cells		Img	Img Cells	
LIVECell	3253	1,018,576	570	181,609	1564	462,261	

4 DATASET

For fine-tuning the Segment Anything Model (SAM) (Kirillov et al., 2023), the LIVECell dataset (Edlund et al., 2021) (Table 2) was exclusively used. LIVECell is a comprehensive dataset with label-free live-cell images and detailed annotations, making it ideal for refining SAM's segmentation capabilities. Leveraging LIVECell for fine-tuning enhances the model's performance and applicability to real-world microscopy images by providing high-quality annotations and diverse cell types. This approach equips the model to handle unique challenges posed by microscopic images, such as low contrast, high noise, various modalities, and complex cell structures.

Two datasets are used to evaluate the generalization of the proposed methodology for cell segmentation and tracking. The first is the Cell Tracking Challenge (CTC) dataset (Maška et al., 2023), which includes 2D and 3D time-lapse sequences of various microscopy videos, including Bright Field, Phase Contrast, and Differential Interference Contrast (DIC). It contains 20 sequences, 10 of which are 2D, with 8,017 frames and an average cell density of 33.12 cells per image. The second is the Cell Tracking with Mitosis Detection Challenge (CTMC) dataset (Anjum and Gurari, 2020), comprising over 1.5 million images across 86 videos of 14 cell lines, annotated with bounding boxes. Unlike CTC, CTMC does not provide segmentation masks, adding challenges for segmentation. To evaluate the method across diverse conditions, 4 sequences from the CTC dataset and 6 from the CTMC dataset were randomly selected, representing various imaging modalities, cell types, and capture intervals to robustly test the method's generalization.

5 EVALUATION METRICS

To assess the performance of the proposed pipeline for cell tracking, five distinct evaluation metrics are used, each providing a unique perspective on the results. These metrics collectively evaluate the accuracy, reliability, and robustness of the tracker across diverse scenarios, ensuring a comprehensive assessment of its capabilities.

5.1 Multiple Object Tracking Accuracy

Multiple Object Tracking Accuracy (MOTA) (Bernardin and Stiefelhagen, 2008) measures the overall accuracy of the tracker and the detection. It accounts for errors such as missed detections, false positives, and identity mismatches, providing a holistic view of the tracker's performance.

$$MOTA = 1 - \frac{\sum_{t} (m_t + f p_t + mme_t)}{\sum_{t} g_t}$$

Here, m_t represents the total number of misses, fp_t the total number of false positives, and mme_t the total number of mismatches. Misses occur when a cell in the ground truth is not detected, which may result from occlusion, low contrast, or noise in the image. False positives occur when a cell is detected but not present in the ground truth, often due to oversegmentation or artifacts. Mismatches occur when a cell is incorrectly associated with another cell, typically due to overlapping trajectories or inconsistencies in tracking. MOTA provides a single metric to summarize the errors, making it a standard benchmark in multi-object tracking evaluations.

5.2 Identification F1 Score

Identification F1 (IDF1) (Ristani et al., 2016) calculates a one-to-one mapping between ground truth trajectories and prediction trajectories, emphasizing the importance of maintaining consistent identities.

$$IDF1 = \frac{2 \cdot IDTP}{2 \cdot IDTP + IDFP + IDFN}$$

IDTP (Identity True Positives) represents the number of correctly matched IDs between ground truth and predictions. IDFP (Identity False Positives) denotes instances where a predicted ID does not correspond to any ground truth object. IDFN (Identity False Negatives) occurs when a ground truth ID is not matched to any prediction. IDF1 is particularly useful for evaluating the consistency of identity preservation in challenging scenarios, such as densely packed cells or overlapping trajectories.

5.3 Identity Switches

Identity Switches (IDs) (Bernardin and Stiefelhagen, 2008), also known as Mismatches, refer to the number of times a trajectory incorrectly changes from one ground truth object to another. This error is often caused by abrupt changes in cell appearance, overlapping trajectories, or tracking errors during occlusion. A lower number of identity switches indicates

a more reliable tracking system, as it suggests the tracker effectively maintains the continuity of object identities across frames. This metric is critical for applications requiring lineage analysis or long-term tracking, where maintaining identity is paramount.

5.4 Mostly Tracked

If an object is successfully tracked for at least 80% of its lifespan, it is considered Mostly Tracked (MT) (Leal-Taixé et al., 2015). This metric evaluates the robustness of a tracking algorithm in maintaining the continuity of an object's identity across its lifespan. High MT scores indicate that the tracker is capable of handling long-term trajectories without frequent interruptions, even in the presence of challenges such as partial occlusions or variations in cell morphology. This metric is particularly important in applications requiring sustained observation of cellular behavior over time, such as studying cell migration or division.

5.5 Mostly Lost

If an object is tracked for 20% or less of its lifespan, it is considered Mostly Lost (ML) (Leal-Taixé et al., 2015). High ML scores suggest the tracker struggles with challenges, leading to frequent identity losses. This metric highlights cases where the tracking algorithm fails to maintain object identities due to factors like occlusion, abrupt motion, or low-contrast regions. By identifying objects that are mostly lost, this metric helps pinpoint specific limitations of the tracker and provides insights for improving its performance in challenging scenarios. A low ML score indicates that the tracker is robust enough to avoid significant failures across the dataset.

6 EXPERIMENTAL SETUP

Two different experimental settings are designed to evaluate the performance of the proposed pipeline for cell tracking from various aspects. The first setting, namely SAT Evaluation on Diverse Modalities and Intervals Using the CTC Dataset, assesses the performance of the SAT pipeline across various imaging modalities and time intervals. This setting utilizes annotated 2D sequences from the Cell Tracking Challenge (CTC) dataset, which includes diverse imaging modalities such as Bright Field, Phase Contrast, and Differential Interference Contrast (DIC). These modalities present unique challenges due to variations in contrast, cell morphology, and imaging noise, providing a comprehensive evaluation of the SAT pipeline's effectiveness in handling diverse cell tracking scenarios. By covering a wide range of imaging conditions and temporal intervals, this setting highlights the pipeline's adaptability and robustness in practical applications. The second experimental setting, namely SAT Generalization Analysis Using CTMC's Wide-Ranging Cell Types, evaluates the SAT pipeline's ability to generalize across diverse cell lines and extensive imaging conditions. The Cell Tracking with Mitosis Detection Challenge (CTMC) dataset presents a distinct set of challenges due to its inclusion of multiple cell lines, varying densities, and complex phase-contrast imaging conditions. This setting emphasizes the pipeline's capacity to handle cell tracking in scenarios with high cell densities, overlapping cells, and intricate motion patterns, making it an ideal benchmark for assessing generalizability across heterogeneous datasets.

To fine-tune the Segment Anything Model (SAM) (Kirillov et al., 2023) on LIVECell data, an iterative training scheme was employed (Archit et al., 2023). Minibatches of input images and ground-truth segmentations were sampled with annotations using random positive points or bounding boxes to ensure a balanced representation of the dataset's variability. Key hyperparameters included a batch size of two, dice loss for masks to optimize segmentation quality, L2 loss for IOU to refine boundary predictions, and the ADAM optimizer (Kingma and Ba, 2014) with a learning rate of 10^{-5} . The learning rate was dynamically adjusted using the ReduceLROnPlateau scheduler to ensure stable convergence during training. Models were trained for 100,000 iterations, with partial updates for 25,000 and fine-tuning for an additional 10,000 iterations, allowing the pipeline to capture intricate features of live-cell images effectively.

The training process was conducted on an A100 GPU with 80 GB of VRAM, enabling efficient computation for high-resolution microscopy images. The Vision Transformer (ViT-h) (Dosovitskiy et al., 2020) was employed as the backbone for image segmentation. The implementation utilized PyTorch (Paszke et al., 2019) along with the torch-em library (Pape, 2023), which provided specialized tools for semantic and instance segmentation tasks in bioimaging.

6.1 Experimental Setting 1: SAT Evaluation on Diverse Modalities and Intervals Using the CTC Dataset

In this experimental setting, the performance of the SAT pipeline is assessed across various imaging

Table 3: Results for the SAT Evaluation on Diverse Modalities and Intervals Using the CTC Dataset. Higher values are better for MOTA, IDF1, and MT, indicated by upward arrows (\uparrow). Lower values are better for IDS and ML, indicated by downward arrows (\downarrow).

Sequence	Modality	Images	Cells	Points (N)	MOTA ↑	IDF1 ↑	$IDS\downarrow$	MT ↑	$\mathbf{ML}\downarrow$
PhC-C2DH-U373 (01)	Phase Contrast	61	427	6P - 3N	79.82	89.00	0.0	71.43	0.0
PhC-C2DH-U373 (02)	Phase Contrast	12	58	6P - 3N	82.75	91.94	0.0	100.0	0.0
Fluo-N2DH-GOWT1	Fluorescence	38	799	3P - 3N	88.24	91.20	0.0	79.17	8.34
Fluo-N2DH-SIM+	Fluorescence	10	271	3P - 3N	83.03	88.73	0.0	83.34	0.0
Average/Total	-	121	1,555	-	83.46	90.22	0.0	83.45	2.08

modalities and time intervals using the Cell Tracking Challenge (CTC) dataset. The CTC dataset includes annotated 2D and 3D time-lapse video sequences of fluorescent counterstained nuclei, as well as 2D Bright Field, Phase Contrast, and Differential Interference Contrast (DIC) microscopy videos. This diverse dataset serves as a benchmark for evaluating the SAT pipeline's ability to generalize across different imaging conditions, cell types, and temporal resolutions, providing a comprehensive evaluation of its effectiveness in diverse cell-tracking scenarios. The table 3 shows the results for this setting, offering detailed metrics for each sequence. The sequences evaluated include:

- PhC-C2DH-U373 (01). This sequence contains 61 frames and is a Phase Contrast microscopy video of U373 cells, captured at 10-minute intervals. MOTA is 79.82%, indicating high tracking accuracy despite the challenges of low contrast typical of Phase Contrast imaging. IDF1 is 89.00%, reflecting excellent identity preservation, with no identity switches (IDS 0.0), showcasing the reliability of the SAT pipeline in maintaining object identities. MT is 71.43%, indicating most cells are successfully tracked throughout their lifespans, and ML is 0.0%, meaning no cells were lost during tracking.
- PhC-C2DH-U373 (02) This sequence contains 12 frames and is a Phase Contrast microscopy video of U373 cells, captured at 10-minute intervals. MOTA is 82.75%, showing improved tracking accuracy over the previous sequence. IDF1 is 91.94%, indicating superior identity preservation with no identity switches (IDS 0.0). MT is 100.0%, reflecting perfect tracking of all cells, and ML remains at 0.0%, demonstrating the robustness of the pipeline for short time-lapse sequences with challenging imaging conditions.
- Fluo-N2DH-GOWT1. This sequence contains 38 frames and is a fluorescence microscopy video of GOWT1 cells, captured at 30-minute intervals. MOTA is 88.24%, demonstrating very high tracking accuracy, leveraging the high contrast pro-

vided by fluorescence imaging. IDF1 is 91.20%, indicating excellent identity preservation, with no identity switches (IDS 0.0). MT is 79.17%, showing that the majority of trajectories were well tracked, while ML is 8.34%, indicating a small number of trajectories were lost, potentially due to occlusions or overlapping cells.

• Fluo-N2DH-SIM+. This sequence contains 10 frames and is a fluorescence microscopy video of SIM+ cells, captured at 30-minute intervals. MOTA is 83.03%, showing high accuracy in tracking. IDF1 is 88.73%, reflecting strong identity preservation, with no identity switches (IDS 0.0). MT is 83.34%, demonstrating that most cells were successfully tracked for the majority of their lifespans, and ML remains at 0.0%, showcasing the pipeline's ability to handle challenging imaging conditions effectively.

Overall, the average metrics across sequences are MOTA of 83.46, IDF1 of 90.22, IDS of 0.0, MT of 83.45%, and ML of 2.08%, demonstrating high tracking accuracy, excellent identity preservation, no identity switches, and effective tracking of most trajectories. These results highlight the robustness of the SAT pipeline in handling diverse imaging modalities and scenarios, from Phase Contrast microscopy with low contrast to Fluorescence microscopy with high contrast. Additionally, the lack of identity switches and the high percentage of mostly tracked trajectories across sequences underline the reliability of the SAT pipeline in maintaining object consistency over time. The pipeline's ability to perform consistently across sequences of varying lengths, frame intervals, and cell densities further illustrates its adaptability and effectiveness in diverse experimental setups. These findings establish SAT as a versatile and scalable solution for automated cell tracking in biomedical research, paving the way for its application in more complex and high-throughput studies.

6.2 Experimental Setting 2: SAT Generalization Analysis Using CTMC's Wide-Ranging Cell Types

In this experimental setting, the performance of the SAT pipeline is evaluated using the diverse cell lines and extensive imaging conditions provided by the Cell Tracking with Mitosis Detection Challenge (CTMC) dataset. This dataset is particularly challenging due to its inclusion of multiple cell types, varying imaging conditions, and high cell densities, making it an excellent testbed for assessing the generalizability and robustness of cell-tracking algorithms. The table 4 shows the results for this setting, highlighting the SAT pipeline's performance across a wide range of scenarios. The sequences evaluated include:

- PL1Ut-run05. This sequence contains 371 frames and comprises phase-contrast images of PL1Ut cells, a rat hepatoma cell line. With a MOTA of 93.12% and IDF1 of 96.56%, the SAT pipeline demonstrates exceptional tracking accuracy and identity preservation. The lack of identity switches (IDS 0.0%) and the high MT score of 100.0% indicate that all cells were tracked successfully throughout their lifespans, with no trajectories being lost (ML 0.0%).
- **A-10-run01.** This sequence consists of 305 frames of phase-contrast images of A-10 cells, a rat smooth muscle cell line. The SAT pipeline achieves a MOTA of 80.79% and IDF1 of 90.03%, reflecting robust tracking performance. Despite the high cell density, no identity switches (IDS 0.0) occurred, and 80.0% of the trajectories were well tracked (MT), with no significant losses (ML 0.0%).
- LLC-MK2-run03. This sequence includes 89 frames of phase-contrast images of LLC-MK2 cells, a monkey kidney epithelial cell line. The SAT pipeline achieves near-perfect tracking results, with a MOTA of 96.18% and IDF1 of 98.08%. No identity switches (IDS 0.0) were observed, and all cells were successfully tracked (MT 100.0%) without loss (ML 0.0%).
- **APM-run05.** This sequence consists of 130 frames of phase-contrast images of APM cells, a human peripheral blood mononuclear cell line. The MOTA of 61.85% and IDF1 of 82.38% reflect satisfactory performance despite the inherent challenges of tracking these cells, including smaller size and irregular movement. All trajectories were tracked successfully for most of their lifespan (MT 75.0%), and no identity switches occurred (IDS 0.0).

- U2O-S-run03. This sequence includes 100 frames of phase-contrast images of U2O-S cells, a human osteosarcoma cell line. With a MOTA of 68.93% and IDF1 of 84.17%, the SAT pipeline performs well in maintaining identity consistency across frames. Most cells were successfully tracked (MT 75.0%) with no significant trajectory loss (ML 0.0%) or identity switches (IDS 0.0).
- **OK-run01.** This sequence comprises 57 frames of phase-contrast images of OK cells, an opossum kidney epithelial cell line. The SAT pipeline achieves a MOTA of 57.77% and IDF1 of 78.11%. While the tracking performance is relatively lower compared to other sequences, likely due to overlapping cells or occlusions, the majority of trajectories were successfully tracked (MT 60.0%), with minimal loss (ML 6.67%) and no identity switches (IDS 0.0).

Overall, the average metrics across sequences are MOTA of 76.44%, IDF1 of 88.22%, IDS of 0.0%, MT of 81.67%, and ML of 1.12%, demonstrating good tracking accuracy, excellent identity preservation, and no identity switches. The high MT scores across sequences show that the SAT pipeline successfully tracks the majority of trajectories, while the low ML values indicate minimal trajectory loss, even in challenging scenarios. These results highlight the robustness and adaptability of the SAT pipeline, which excels in handling diverse imaging conditions, cell types, and densities. By consistently achieving high tracking performance without requiring additional fine-tuning, the SAT pipeline establishes itself as a reliable tool for automated cell tracking in complex and high-throughput experimental setups. Its ability to generalize effectively across different datasets further underscores its potential for broad applications in biomedical research.

7 ANALYSIS AND DISCUSSION

This section explains the results of the two experimental settings, highlighting the SAT pipeline's ability to generalize across different modalities and cell lines. In the first setting, SAT Evaluation on Diverse Modalities and Intervals Using the CTC Dataset, the pipeline was tested on the Cell Tracking Challenge (CTC) dataset, which includes diverse imaging modalities. Table 3 shows the SAT pipeline achieves 83.46% MOTA, 90.22% IDF1, and zero Identity Switches (IDS), demonstrating consistent tracking across modalities. In the second setting, SAT Generalization Analysis Using CTMC's Wide-Ranging Cell Types, the pipeline was evaluated using the Cell ICAART 2025 - 17th International Conference on Agents and Artificial Intelligence

Table 4: Results for the SAT Generalization Analysis Using CTMC's Wide-Ranging Cell Types. Higher values are better for MOTA, IDF1, and MT, indicated by upward arrows (\uparrow). Lower values are better for IDS and ML, indicated by downward arrows (\downarrow).

Sequence	Modality	Images	Cells	Points (N)	MOTA ↑	IDF1 ↑	$IDS\downarrow$	MT ↑	ML↓
PL1Ut-run05	Phase Contrast	371	742	30P - 3N	93.12	96.56	0.0	100.0	0.0
A-10-run01	Phase Contrast	305	1,525	20P - 3N	80.79	90.03	0.0	80.00	0.0
LLC-MK2-run03	Phase Contrast	89	445	25P - 3N	96.18	98.08	0.0	100.0	0.0
APM-run05	Phase Contrast	130	443	25P - 3N	61.85	82.38	0.0	75.00	0.0
U2O-S-run03	Phase Contrast	100	396	15P - 3N	68.93	84.17	0.0	75.00	0.0
OK-run01	Phase Contrast	57	841	30P - 3N	57.77	78.11	0.0	60.00	6.67
Average/Total	-	1,173	5,447	-	76.44	88.22	0.0	81.67	1.12



Figure 2: Tracking results for experimental setting 1 with two sequences, PhC-C2DH-U373 and Fluo-N2DH-SIM+. The top row shows ground truth, and the bottom row shows SAT pipeline predictions with evaluation scores above the prediction row.

Tracking with Mitosis Detection Challenge (CTMC) dataset, featuring diverse cell lines and extensive imaging conditions. The results, shown in Table 4, reveal that the SAT pipeline maintains good tracking accuracy with an average MOTA of 76.44% and an

IDF1 of 88.22%, indicating effective generalization to different cell types and high identity preservation. The variation in MOTA across different cell types can be attributed to the challenges presented by different imaging modalities and cell cultures. For instance, the



shows ground truth, and the bottom row shows SAT pipeline predictions with evaluation scores above the prediction row.

PhC-C2DH-U373 cell culture, imaged with phasecontrast microscopy, poses difficulties due to low contrast, causing cells to merge with the background as they grow. In contrast, the Fluo-N2DH-GOWT1 culture, captured with fluorescence microscopy, offers higher contrast, making segmentation and tracking significantly easier. These factors contribute to the variation in MOTA and are reflected in the observed performance across different datasets.

Figure 2 illustrates tracking results for the first setting with sequences PhC-C2DH-U373 and Fluo-N2DH-SIM+. The top row shows the ground truth, while the bottom row shows the SAT pipeline predictions. For PhC-C2DH-U373, all 7 cells are correctly segmented initially, with one cell missed at t + 300 minutes but recovered at t + 600 minutes and tracked

till the last frame at t + 885 minutes. The SAT pipeline achieves a MOTA of 78.2%, IDF1 of 89.0%, 0 IDS, 71.4% MT, and 0.0% ML. For Fluo-N2DH-SIM+, all cells are correctly segmented initially, with one cell missed at t + 58 minutes and another at t + 116 minutes, both recovered at t + 174 minutes. The pipeline achieves a MOTA of 83.0%, IDF1 of 88.7%, 0 IDS, 83.3% MT, and 0.0% ML.

Figure 3 shows tracking results for the second setting with sequences LLC-MK2-run03 and A-10-run01. The top row shows ground truth bounding boxes, while the bottom row shows SAT pipeline predictions with both bounding boxes and segmentation masks. For LLC-MK2-run03, all cells are correctly segmented across all frames, achieving a MOTA of 96.2%, IDF1 of 98.1%, 0 IDS, 100.0% MT, and 0.0%

ML. For A-10-run01, all cells are correctly segmented initially, with cell number 2 missed at t + 119 minutes but recovered at t + 149 minutes. The pipeline achieves a MOTA of 80.8%, IDF1 of 90.0%, 0 IDS, 80.0% MT, and 0.0% ML.

While direct comparison with other tracking methods is not entirely feasible due to the unique nature of the proposed approach, an additional experiment with ByteTrack was conducted to offer some insights. ByteTrack, trained on the same LIVECell dataset as SAT, failed to detect any cells when applied to the CTMC dataset. To investigate further, Byte-Track was trained on a subset of the CTMC dataset (with no overlap with the test set) and tested on the remaining sequences. ByteTrack's performance was lower than SAT, with an average MOTA of 29.3 compared to 76.4, and higher IDS and ML scores. These findings emphasize the superior adaptability and robustness of SAT, which can generalize effectively to unseen datasets without retraining, a limitation observed in traditional methods like ByteTrack (Zhang et al., 2022).

Overall, the proposed SAT pipeline demonstrates strong generalization across different modalities and cell lines, achieving high tracking accuracy and identity preservation. This pipeline significantly impacts the biological and biomedical research community by automating cell segmentation and tracking, reducing the need for expert knowledge and manual intervention. It enhances accuracy, consistency, and speeds up data annotation, benefiting cancer research, drug development, and stem cell studies. SAT's broad applicability with minimal retraining makes it a versatile tool, driving new insights and improving research efficiency. By reducing the time and effort required for annotation, SAT enables researchers to focus on complex experimental designs and large-scale analyses. Moreover, its ability to adapt to diverse imaging conditions ensures reproducibility and scalability in longitudinal studies. The adoption of SAT could also bridge gaps in resource-limited settings, democratizing access to advanced cell tracking technologies. This efficiency and accessibility position SAT as a valuable asset for advancing both fundamental research and clinical applications.

8 CONCLUSION

This study introduces the SAT pipeline for Cell Segmentation and Tracking, which uses point annotations only in the first frame to automate segmentation and tracking across sequences. The pipeline demonstrates strong generalization and robustness across diverse

imaging modalities and cell types, achieving over 80% Multiple Object Tracking Accuracy (MOTA) on two diverse datasets. By significantly reducing annotation time-up to 206x faster than traditional methods-SAT enables efficient and scalable cell tracking, even in challenging conditions such as noisy, lowcontrast microscopy images. This automation streamlines large-scale studies in cancer research, drug development, and stem cell analysis, while improving accuracy and reducing expert intervention. The versatility, adaptability, and scalability of the SAT pipeline make it a robust solution for modern biomedical research, facilitating deeper insights into cellular behavior and accelerating scientific discoveries with clinical applications. In the future, this study can be extended to track cell division, enabling analysis of cell cycle progression, mitotic events, and lineage formation over time, providing valuable insights into cellular proliferation and division dynamics, particularly for studying developmental biology, stem cell differentiation, and cancer progression.

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