Sperm Tracking and Trajectory Analysis in Fluorescence Microscopy Image Sequences

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Abstract: In this work, we analyze the performance of several tracking methods in the scenarios of low temporal sampling acquisition setup in fluorescence microscopy. Machine Learning methods were applied to classify and analyze the extracted trajectories of sperm samples and their motion parameters. The results were compared with the most widely used sperm motility classification methods. Analyzed image sequences include real sequences acquired by confocal fluorescence microscopy and synthetic sequences generated by in-house software. The complete framework runs as a standalone application and can be used with minimal training by users with no programming skills.

1 INTRODUCTION

Infertility is a worldwide increasing condition, affecting up to 12\% of couples of childbearing ages (Kumar and Singh, 2015). Although the male factor accounts for at least 30–40\% of the cases, there is a traditional lack of emphasis on the contribution of men to the field (Walters et al., 2020). As a consequence, the vast majority (>70\%) of male infertility cases are deemed idiopathic, a situation that severely limits treatment strategies to rescue fertility (Walters et al., 2020). Currently, there are no diagnostic methods that determine the condition of infertility and this categorization is based fundamentally on the patient’s clinical history and sperm analysis (spermiogram) (World Health Organization, 2010). Semen analysis is the gold standard test for analyzing male fertility status and it includes the analysis of sperm count, sperm motility, and sperm morphology. Sperm motility is one of the parameters closely related to the success of \textit{in vivo} fertilization. Consequently, there is an especial interest in tracking sperm movements to understand sperm biology as a marker of sperm’s ability to fertilize the egg (Mortimer et al., 2015).

Computer-Aided Sperm Analysis (CASA) technology was developed in the late 1980s for analyzing sperm movement based on the extraction and analysis of the spermatozoa’s trajectories (Mortimer et al., 2015; Walters et al., 2020). This technology is mostly used in the research area since in general, CASA’s confidence level is not enough to be applied in the clinical field (Mortimer et al., 2015). Moreover, different CASA instruments compute sperm motility parameters using different algorithms to classify sperm, thus, values may not be comparable among systems (World Health Organization, 2010).

CASA are developed to work with bright-field microscopy images, using phase-contrast optics to highlight the head of each spermatozoon that allows researchers to analyze the sperm motility (Alquézar-Baeta et al., 2019; Goodson et al., 2017; Mortimer et al., 2015), however, it is not usual to find a proper solution to be applied to fluorescence images.

On the other hand, there are many algorithms and implementations for tracking objects and particles in sequences. Some are free accessible and open source, such as TrackMate (Tinevez et al., 2017) for ImageJ (Schneider et al., 2012) or TrackPy (Allan et al., 2019). However, none of them were developed specifically for fluorescence microscopy images, where fluorescence intensity or distribution is quantified at every frame.
Beyond advances in optics and detectors, imaging has strongly benefited from the development of fluorescent probes. Fluorescent probes or genetically encoded fluorescent proteins can provide real-time monitoring of function on living cells. In the case of sperm cells, the use of fluorescent probes combined with computer analysis may provide a new tool to analyze and classify samples e.g. to differentiate fertile from infertile men. MitoTracker™ Red CMXRos is a red-fluorescent dye that stains mitochondria in live cells and its accumulation is dependent upon membrane potential (Zhang et al., 2019). In the sperm cells Mitotracker™ stains part of the flagella; the midpiece.

Although the particle-like object detection in fluorescence images is benefited from the specificity of the fluorophore marker, the achieved frame rate is an important variable for a tracking algorithm. It is already known that a low frame rate defies the process of association of the detected objects in frame \( I_{k+1} \) with the set \( Z_k \) of trajectories \( \zeta \), computed up to frame \( I_k \), \( Z_k = \{ \zeta \}_k \). In fact, in Laser Scanning Confocal Microscopy (LSCM) such as the one used in this work, the image formation process depends on several factors (e.g. size of pinhole, photobleaching, instrument characteristics, etc.) and it could determine errors in the classification of sperm. This bottleneck in the acquisition of higher frame rates sequences is produced because the image formation process is performed in a raster scanning pattern, where a photomultiplier quantifies the intensity of light only in one selected region of the image at a time, instead of in a parallel process as in the CCD or CMOS detectors (Dobrucki, 2013).

In this work, we analyzed the performance of five different particle tracking algorithms under different frame rates ranging from a challenging 6 fps to 60 fps. Also, we evaluated different Machine Learning methods for trajectory analysis and compared them with the WHO manual procedure. To obtain a gold standard for quantification, we developed software for generating synthetic sperm image sequences. The software allowed us to analyze the performance varying parameters such as sequence frame rate, spermatozoon size, shape, speed, head beating angle, among others. Also, real image sequences were acquired and processed.

The document is organized as follows. In Section 2 we review the main characteristics of the sample preparation and image formation process in LSCM. In Section 4.1 we present the studied algorithms among with the summary of results of performance analysis. In Section 4 we describe the Machine Learning methods studied for trajectory classification and the results obtained in synthetic and real sequences. Section 5 presents the conclusions of this work and future lines of research.

## 2 MICROSCOPY IMAGE SEQUENCE ACQUISITION

### 2.1 Sample Preparation

Semen samples of three donors were studied. Semen samples were spun at 300g for 10 minutes and supernatants were carefully removed. Pellets were resuspended in BWW Albumin 0.3% 25 mM NaHCO₃ HEPES medium. Approximately 5 million spermatozoa were incubated with 50 nM MitoTracker™ Red CMXRos (ThermoFisher) minutes at 37°C during 30 minutes. Volumes of 100-110 µl of samples were loaded in the ThermoFischer™ Nunc™ Lab-Tek™ Chambers and observed under the confocal microscope (Leica, TCS SP5 II). Acquisition parameters of the confocal microscope were: zoom 1.7×, magnification 40×, two lines average, excitation lasers HeNe (543 nm, 633 nm), 512 × 512 pixels 120 frames sequences. The confocal microscope’s pinhole aperture was maximized (3 Airy) to reduce the optical sectioning.

### 2.2 Image and Sequence Acquisition

Laser Scanning Confocal Microscopy (LSCM) image formation process is schematically presented in Figure 1. A laser emits a beam of light to the sample with a certain wavelength to excite the fluorophores. The beam of light passes through a lens, a beam splitter, and another lens to illuminate a specific part of the focal field in the sample. This beam excites the fluorophore molecules attached to the midpiece of the spermatozoon, and they go to an unstable state of energy. To go back to a stable state, the fluorophore

![Figure 1: Laser Scanning Confocal Microscope image formation schematic diagram.](image-url)
emits surplus energy by emitting photons in a certain wavelength. These photons are reflected by the beam splitter, pass through a pinhole to filter the out-of-focus planes’ light, and then are captured and converted using a photomultiplier (PMT) detector. The PMT records the intensity of light only in a selected region of the sample at a time, which then is converted to an intensity level in some pixels of the image (Dobrucki, 2013). These steps are repeated for each pixel composing the image. This process is slower than acquiring the image with a CCD or CMOS array detector where the photon-voltage conversion is performed in parallel for all the pixels.

Moreover, if more than one fluorophore is used in the sample, the previous process is repeated for each one of the lasers needed to excite each fluorophore. In addition to these steps, each region of the image could be visited many times, averaging the measures to increase the Signal to Noise Ratio (SNR). Besides this line averaging, several frames could be averaged to form only one image, too. This process reduces the frame rate that could be achieved by an LSCM, impacting the analysis of time-lapse sequences.

On the other hand, the size of the image could be reduced to increase the frame rate, which could produce a resolution compromise for a given region of the sample.

Post-acquisition image processing, including denoising and models of particle movements, could help to overcome these acquisition problems. In practice, the size of the objects and the bandwidth of their trajectories affect the spatial and temporal sampling rates, respectively. In the dataset analyzed in our case, as a general conclusion, the post-acquisition image processing of the noisy image sequences allowed us to increase the temporal sequence resolution with good detection of the spermatozoon location in each frame.

3 Sperm Tracking

A general framework for object tracking usually has two main blocks, preceded by the preprocessing operations. First, all the objects are detected in each frame; in this step, prior information about the objects’ appearance and image acquisition is used. Second, based on the trajectories computed up to frame $k$ a prediction of the position of each particle at frame $k + 1$ is computed, then these predictions and the detection are linked updating the trajectories (Pulford, 2005). This second process is schematically shown in Figure 2a.

The literature on object tracking in Computer Vision and Image Processing is extensive (see references in (Bar-Shalom and Daum, 2010)) given the existence of several issues related to this problem. Sperm tracking can be thought of as a specific case of particle tracking, in which each spermatozoon is a single particle. In this case, a modified Kalman Filter is used to model the sperm movement as in (Urbano, 2014). This model estimates the average path of the sperm trajectory. The deviation from the average path, which is mainly caused by the movement of the head, is modeled as random noise which represents the acceleration of the spermatozoon. This model can be good for low sample rates but not for high sample rates, where the deviation from the head predominates over the average path. Given the acquisition process, the sample rate of the image sequences taken by the confocal microscopy is rather low so the model fits the problem.

Since multiple spermatozoa are present in a sperm sample in each frame one Kalman Filter for each particle is needed. Therefore, once all the spermatozoa are detected in a frame, it is necessary to determine which measurement was originated by which sperma-
This is a key step in the tracking process since it is important to update the Kalman Filter of each spermatozoon with the right measurement; meaning, the one that was originated by that same particle.

The tracking analysis in this work is strongly based on the works of Urbano at al. (Urbano, 2014; Urbano et al., 2017) and Mortimer et al. (Mortimer et al., 1988; Mortimer and Swan, 1999). Based on these works we analyzed the performances of the following methods to make the association between the spermatozoa detections (trajectory linking).

- Nearest Neighbor (NN) (Vo et al., 2015) is a greedy method where particles are sequentially processed assigning the target whose position is closest to the predicted position.
- Global Nearest Neighbor (GNN) (Munkres, 1957) differs from the previous one on the targets are assigned considering all the detected objects using the Hungarian method of combinatorial optimization.
- Probabilistic Association Filter (PDAF) (Bar-Shalom and Daum, 2010) computes an association probability to the target for each detected position, then given an association, the state of a target is estimated by a filtering algorithm, and this conditional state is weighted by the association probability.
- Joint Probabilistic Association Filter (JPDAF) (Bar-Shalom and Daum, 2010) is the PDAF extension for multiple targets, and
- Exact Nearest Neighbor - Joint Probabilistic Association Filter (ENN-JPDAF) (Fitzgerald, 1986) computes an approximation of the probability of association used in JPDAF which can be solved in close form which is applied before a nearest-neighbor scheme.

Once the trajectories are determined for each spermatozoon, the next step in sperm analysis is to group them based on their motility characteristics. The traditional approach is to take some predefined parameters, representing the speed, traveled distance, linearity of the movement, etc. Guidelines for computing these descriptors and their thresholds for each group are published by the World Health Organization (WHO) (World Health Organization, 2010; Elia et al., 2010). Some of these trajectory descriptors, shown in Figure 2b, are:

- VCL (curvilinear velocity in $\mu$m/s): averaged speed of a spermatozoon along its 2D curvilinear path.
- VSL (straight-line velocity in $\mu$m/s): average speed of a spermatozoon along the line between its first and last detected positions.
- VAP (average path velocity in $\mu$m/s): average speed of a spermatozoon along its average path.
- ALH (amplitude of lateral head displacement in $\mu$m): lateral displacement of a spermatozoon about its average path.
- BCF (beat-cross frequency in Hz): average rate of the crosses between the curvilinear and the average path.
- MAD (mean angular displacement in degrees): average of absolute values of the turning angle of the spermatozoon.

### 3.1 Tracking Experiments

To analyze and quantify the performance of these methods under different sequences’ frame rates, simulation software was implemented. This software assumes an ellipsoidal shape for the spermatozoa midpieces and models its movement following the characteristics of real sequences as described in (Urbano, 2014).

The detection of the spermatozoo’s midpieces in fluorescence is obtained by automatic thresholding (Otsu’s method) followed by mathematical morphology operations for filling holes and close regions. In phase-contrast images, this step is substituted by a...
Two sequence scenarios were tested taking into account the density of spermatozoa in the images, considered low or high by the experts. Sequences were generated with a maximum frame rate of 60 fps and undersampled to obtain sequences at 40, 30, 20, 15, and 6 fps. For every frame rate, trajectories were extracted with the five methods described above and compared with the simulated trajectories as ground truth. Tracking methods’ performance is evaluated based on two criteria comparing the extracted and ground-truth trajectories sets. First, the Jaccard Index for measuring the similarities between sets, and second, the mean relative error in the computation of the WHO’s trajectory descriptors described above.

Results are consistent between the experiments. The performance for all the methods is significantly reduced at very low frame rates (6 fps). Results for GNN, JPDAF, and ENN-JPDAF are indistinguishable, and they improve with higher frame rates, except in very high frame rates (60 fps) where the performance decreases. Analyzing the 60 fps sequences and trajectories, this drop of performance is based on the high frequency and amplitude of lateral movement of the spermatozoa which are not well captured by the methods. On the other hand, NN and PDAF performance is poor and decreases for high frame rates. These performances are worse in high-density sequences where the presence of more particles produces more assignment errors.

Figure 4 shows the mean relative error in six selected WHO’s trajectory descriptors. Results are shown for the high-density sequences; the discussion and conclusions are similar for the low-density sequences. In general, as in the previous criteria, the 6 fps sequences’ mean relative errors are systematically higher than in higher frame rates. Also, NN and PDAF for frame rates higher than 15 fps have worse results than the rest of the methods. For GNN, JPDAF, and ENN-JPDAF results are almost uniform for frame rates between 15 and 40 fps, this is particularly true in the VSL where only the first and last points are used in the computation. The mean relative error for 60 fps increases, following the behavior of the previous criteria, in particular for the ALH and BCF which are strongly dependent on the frequency and amplitude of the lateral head displacement.

4 TRAJECTORIES CLASSIFICATION

A four-category system for grading motility is used in andrology laboratories, both from manual assessment of sperm motility as well as computer-aided sperm analysis. These categories are defined as (Elia et al., 2010): Straight-Line Progressive (SLP), Straight Slow Progressive (SSP), Non-Straight Progressive (NSP), and Non-Progressive (NP).

As mentioned before the WHO published a set of guidelines for sperm motility classification based on trajectory descriptors (VSL, VCL, ALH, etc.). It is known that different CASA instruments compute these parameters using different algorithms, therefore, the values may not be comparable among systems (World Health Organization, 2010). Another weakness of these parameters is that they tend to depend on the frame rate of the acquired sequence, this issue is well studied by Mortimer et al. (Mortimer et al., 1988; Mortimer and Swan, 1999) and our analysis confirms these results. In this section, a data driven approach is presented comparing a method based on Machine Learning with the classification obtained with the mentioned trajectory descriptors (Elia et al., 2010).

One of the challenges for a Machine Learning framework is to cope with input vectors with differ-
ent dimensions. This is the case of trajectories for objects appearing in different numbers of frames. In our framework, the trajectory for the $i$-th spermatozoon is represented as a $3N_i$-dimensional array

$$\mathbf{z}_i = \{[x_i^0, y_i^0, t_i^0], \ldots, [x_i^{N_i}, y_i^{N_i}, t_i^{N_i}]\},$$

where $N_i$ is the number of points detected in the trajectory, and $j = 0, \ldots, N_i - 1$.

To obtain a fixed dimensional input vector describing each trajectory, we follow the proposal of Yao et al. (Yao et al., 2017), where they studied a problem of trajectory classification applied to Global Positioning System (GPS) data. They propose a Behavior Feature Extraction Algorithm based on sliding windows through the trajectory points. These features are the input of a neural network based on encoder-decoder (autoencoder) to learn characteristics from the trajectories.

The classification is performed using a Support Vector Machine (SVM) trained with labeled trajectories. The ground-truth labels were computed based on the classification guidelines defined by Elia et al. (Elia et al., 2010) using the WHO trajectory descriptors (World Health Organization, 2010).

### 4.1 Classification Experiments

Following the procedure described in Section 3.1 we generated a set of synthetic sequences where 1658 simulated trajectories were extracted. The trajectories are balanced among the four groups: Straight-Line Progressive (SLP), Straight Slow Progressive (SSP), Non-Straight Progressive (NSP), and Non-Progressive (NP). Each trajectory $\mathbf{z}_i$ is preprocessed and the trained autoencoder computes the $\mathbf{h}_i$ feature vector. The ground truth labels are computed based on WHO descriptors for the extracted trajectories as described in Section and compared with the label assigned by the trained classifier.

A $k$-fold training of the SVM is performed splitting the set of feature vectors $\{\mathbf{h}_i\}$ in training and testing sets with 80% and 20% of the samples respectively. The obtained confusion matrix is shown in Table 1. The Accuracy obtained in this case is 96.4%, Average Specificity (Recall) of 96.2%, and Average Sensitivity (Precision) of 96.6%. The Specificity and Sensitivity for each class are shown in Table 3. Figure 5 shows a 2D and 3D embedding via Principal Component Analysis (PCA) decomposition.

The results are very good, confirming that the latent space learned by the autoencoder is a good feature representation space for the trajectories.

Taking into account the embedding obtained in Figure 5 we observed similar behavior for both classifiers. They show well-defined regions for the Non-Progressive and Non-Straight Progressive. The regions for the Straight-Line Progressive and Straight-Slow Progressive are very close between each other, and far from the NSP and NP. In the case of the WHO classification, there are more NP trajectories sparsely distributed in all the embedding.

This lower distance observed between the SSP and SLP classes is remarkable since it is well known that in practice for fertility evaluation the spermatozoa considered are only these two classes without discrimination among them. Categorization of progressively motile spermatozoa as rapid or slow, (with a speed cut off of 25 μm/sec at 37°C) is constantly under debate in the field. This is based mainly on the difficulty to define the forward progression so accurately without bias (Cooper and Yeung, 2006). As a consequence, the last WHO recommendations suggest that clinicians may use total motility (PR + NP) or progressive motility (PR) to establish reference values in infertility (World Health Organization, 2010).

### Table 1: Confusion matrix for synthetic sequences (in percentage, normalized by row).

<table>
<thead>
<tr>
<th>True label</th>
<th>SLP</th>
<th>SSP</th>
<th>NSP</th>
<th>NP</th>
</tr>
</thead>
<tbody>
<tr>
<td>SLP</td>
<td>88.5</td>
<td>10.2</td>
<td>0.0</td>
<td>1.3</td>
</tr>
<tr>
<td>SSP</td>
<td>0.0</td>
<td>99.0</td>
<td>0.0</td>
<td>1.3</td>
</tr>
<tr>
<td>NSP</td>
<td>0.0</td>
<td>0.0</td>
<td>99.0</td>
<td>1.1</td>
</tr>
<tr>
<td>NP</td>
<td>0.0</td>
<td>1.2</td>
<td>0.0</td>
<td>98.8</td>
</tr>
</tbody>
</table>

### Table 2: Confusion matrix for real sequences (in percentage, normalized by row).

<table>
<thead>
<tr>
<th>True label</th>
<th>SLP</th>
<th>SSP</th>
<th>NSP</th>
<th>NP</th>
</tr>
</thead>
<tbody>
<tr>
<td>SLP</td>
<td>54.5</td>
<td>9.1</td>
<td>0.0</td>
<td>36.4</td>
</tr>
<tr>
<td>SSP</td>
<td>30.0</td>
<td>60.0</td>
<td>0.0</td>
<td>10.0</td>
</tr>
<tr>
<td>NSP</td>
<td>22.2</td>
<td>22.2</td>
<td>0.0</td>
<td>55.6</td>
</tr>
<tr>
<td>NP</td>
<td>4.7</td>
<td>28.6</td>
<td>0.0</td>
<td>66.7</td>
</tr>
</tbody>
</table>

### Table 3: Specificity (Recall) and Sensitivity (Precision) for synthetic sequences.

<table>
<thead>
<tr>
<th></th>
<th>SLP</th>
<th>SSP</th>
<th>NSP</th>
<th>NP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specificity</td>
<td>88.46</td>
<td>98.73</td>
<td>98.89</td>
<td>98.82</td>
</tr>
<tr>
<td>Sensitivity</td>
<td>100</td>
<td>89.65</td>
<td>100</td>
<td>96.55</td>
</tr>
</tbody>
</table>
Figure 5: 2D and 3D embedding via Principal Component Analysis (PCA) decomposition for the synthetic trajectories. Colors are assigned by the SVM (left) or ground truth (right) classification.

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Table 4: Specificity (Recall) and Sensitivity (Precision) for real sequences.

<table>
<thead>
<tr>
<th></th>
<th>SLP</th>
<th>SSP</th>
<th>NSP</th>
<th>NP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specificity</td>
<td>54.55</td>
<td>60.00</td>
<td>0</td>
<td>66.67</td>
</tr>
<tr>
<td>Sensitivity</td>
<td>40.00</td>
<td>57.14</td>
<td>-</td>
<td>56.00</td>
</tr>
</tbody>
</table>

5 CONCLUSIONS AND FUTURE WORK

In this work, we presented an analysis of several methods for sperm samples tracking in sequences with different frame rates in fluorescence microscopy. Three of these methods achieved similar performances, these were Global Nearest Neighbor (GNN), Joint Probabilistic Association Filter (JPDAF), and Exact Nearest Neighbor Joint Probabilistic Association Filter (ENN-JPDAF). Performance was measured using the Jaccard Index, and the mean relative error in several trajectory descriptors. The performance was similar for frame rates spanning from 15 to 40 frames per second, but it drops significantly for 6 frames per second.

We also analyzed the performance of a Support Vector Machine classification of the extracted trajectories based on an intermediate representation given by an autoencoder latent space. The performance achieved in synthetic sequences was very high. For real sequences, we observed several drawbacks leading to new lines of work.

The lack of labeled data in real sequences in fluorescence microscopy is one of the main issues and it will become part of a future line of work. Increasing the amount of data will help in the training process of machine learning methods, including modern methods based on convolutional networks or deep learning.

The developed framework allows us to extract information about the sperm trajectories while quantifying the fluorescence in the spermatozoa midpieces. In the future, this will provide an extremely useful tool for studying the health of each spermatozoon, measured by its fluorescence, and its motility parameters simultaneously. Here we presented the track-
ing analysis of sperm labeled with MitoTracker™ that specifically binds to mitochondria in the sperm flagella. The possibility to track sperm movements using other fluorescent probes may have multiple applications. Several fluorescence compounds have been developed that are able to detect changes in calcium variations, production of reactive oxygen species, mitochondrial activity, etc. meaning that it will be possible to match either of these functions with sperm motility patterns. The tool may be useful to test drugs that modify the motility patterns of subpopulations of sperm. Moreover, multiple probes may be used at the same time to explain different biological effects.

The developed framework is publicly available in our GitLab repository 1.

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REFERENCES


1https://gitlab.fing.edu.uy/pfc-tde/testing-docker