Reconstructing the flight kinematics of swarming and mating in wild mosquitoes

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We describe a novel tracking system for reconstructing three-dimensional tracks of individual mosquitoes in wild swarms and present the results of validating the system by filming swarms and mating events of the malaria mosquito Anopheles gambiae in Mali. The tracking system is designed to address noisy, low frame-rate (25 frames per second) video streams from a stereo camera system. Because flying A. gambiae move at 1–4 m s⁻¹, they appear as faded streaks in the images or sometimes do not appear at all. We provide an adaptive algorithm to search for missing streaks and a likelihood function that uses streak endpoints to extract velocity information. A modified multi-hypothesis tracker probabilistically addresses occlusions and a particle filter estimates the trajectories. The output of the tracking algorithm is a set of track segments with an average length of 0.6–1 s. The segments are verified and combined under human supervision to create individual tracks up to the duration of the video (90 s). We evaluate tracking performance using an established metric for multi-target tracking and validate the accuracy using independent stereo measurements of a single swarm. Three-dimensional reconstructions of A. gambiae swarming and mating events are presented.

Keywords: target tracking; Anopheles gambiae; mosquito swarms; mating mosquitoes

1. INTRODUCTION

Quantitative observations of the flight patterns of wild mosquitoes are critical to expanding our understanding of swarming and mating behaviour [1–6]. Female Anopheles gambiae find male swarms in order to mate [5,7]. A single mating event results in all of the fertilized eggs that a female mosquito lays in her lifetime [8,9]. Although the basis of mate selection has generated much interest [6–8,10,11], generation of three-dimensional trajectory data of mosquitoes in wild swarms has not been previously accomplished. As in earlier work on midges [12], such trajectory data can provide valuable insight into the dynamical aspects of collective behaviour [13,14]. Past studies on swarming insects [2,3,5,15] focused on two-dimensional trajectories or three-dimensional positions. Recent advancements in high-resolution filming, computer vision and estimation techniques have increased the degree of automation in data collection, and have made available large datasets for subsequent analysis, such as those developed for starlings [16] and fruitflies [17]. Similar analyses of malarial mosquitoes may inform the first steps towards strategies of vector control [7,8].

Multi-target tracking systems have been developed for other animals. In two dimensions, ants have been tracked using a joint-state particle filter and interaction models [18]. In three dimensions, up to a hundred bats have been tracked with three cameras using a Kalman filter in conjunction with a multi-dimensional assignment strategy [19,20]. Fruitflies have been tracked in an acrylic box by setting up the problem of data association across views and time in the form of a global optimization problem that is solved at every step [21]. Real-time tracking systems for flies were developed using an extended Kalman filter in Straw et al. [22] and Grover et al. [23]. Each of these tracking systems...
implements a nonlinear filtering or optimization method with specialized likelihood functions, data association strategies and/or experimental design. However, the targets are large with a dark centre or appear in an arena constructed to minimize noise, unlike wild mosquitoes.

Filming wild mosquitoes poses special challenges such as low natural lighting and a cluttered dynamic background. At least two cameras are needed to reconstruct the three-dimensional position of individual mosquitoes. A multi-camera set-up with a large baseline reconstructs positions accurately, but may be difficult to implement in the field. In a multi-target scenario, one must also address the data-association problem, which entails assigning image blobs to targets across multiple views and time steps. With the typical number of mosquitoes in the swarms we studied, the data-association problem is non-trivial. The challenge lies in tracking small, highly manoeuvrable targets that appear as dots or faded streaks in noisy images with frequent occlusions.

This paper describes an automated multi-target tracking system that reconstructs the three-dimensional flight kinematics of individual mosquitoes in wild swarms. We collect data using two cameras operating synchronously at 25 frames per second. (The frame rate is limited by the ambient light.) The cameras and a laptop are powered by an uninterrupted power supply (UPS) for up to 30 min. The mosquitoes appear as dark streaks or dots on a light background. At high speeds, the mosquito streaks fade, making them hard to detect and even harder to track. Because the swarms are dense, occlusions are frequent and often appear in both camera frames. We tested the system by filming swarms and mating events of *A. gambiae* in a rural village in Mali in August 2010. Figure 1 shows a pair of magnified and enhanced sample frames from this field experiment.

In order to track each insect in a wild swarm, we implemented a probabilistic multi-target tracking framework. Specifically, the contributions of this paper are as follows: (i) we provide a measurement likelihood function that uses the properties of image streaks such as midpoint and endpoint locations to extract insect position and velocity; (ii) we provide methods to improve data association in noisy images by adaptively seeking missing measurements and splitting occluded blobs into individual measurements; and (iii) we present validated tracking results in the form of three-dimensional trajectories of wild mosquito swarming and mating events filmed in Mali in August 2010. Although we describe the experimental method and tracking algorithm for mosquito swarms, the techniques presented in this paper may be beneficial for generating trajectory data for other insect swarms in the field or laboratory.

The tracking system is implemented in MATLAB and consists of two parts: an automated component that outputs track segments called tracklets and a human-supervised component that is used to verify and combine the tracklets into full-length tracks. Tracklets produced by the automated component typically range between 15 and 25 frames (0.6–1 s) long and can be used to extract position and velocity data for 80 per cent of the swarm. The human-supervised component uses a particle filter to combine tracklets into individual mosquito tracks. It takes up to 20 min to generate a 10 s track (250 frames). When validated using data filmed in Mali in August 2010, the tracking system produced 30–40 s trajectories of individual mosquitoes in swarms of 6–25 mosquitoes. We have reconstructed six swarms and six mating events from these data. We evaluate the performance of the automated component of the tracking system using an established metric based on position error and the number of targets tracked (cardinality); tracking accuracy was also evaluated using two independent rigs to simultaneously track the same swarm.

The paper is organized as follows: §2 provides a background on multi-target tracking, data association and tracking performance evaluation. Section 3 presents the novel components of the tracking system, including the likelihood function and occlusion reasoning. Section 4 describes the data collection, validation methods and performance evaluation; it also includes representative kinematic data for wild mosquito swarming and mating events. Section 5 summarizes the paper and our ongoing analyses of the kinematic data.

2. PROBABILISTIC TRACKING AND DATA ASSOCIATION

Our aim in designing the mosquito tracking system was to combine nearly indistinguishable measurements available from stereo images recorded at discrete times into trajectories that represent real mosquitoes (targets). We represent target *i* at time step *k* by the state vector $X_i[k] \in \mathbb{R}^6$, which contains the target’s instantaneous three-dimensional position and velocity. In a Bayesian framework, a tracking algorithm recursively iterates through two steps: the update step and the predict step. The update step uses a measurement model to revise the estimate based on new observations. The predict step integrates a motion model to obtain the target state at the time of the next measurement. The measurement $Z[k] \in \mathbb{R}^6$ in our case consists of the two-dimensional positions of the midpoint and two endpoints of an elongated blob in an image that corresponds to the motion-blurred silhouette of a flying mosquito in each of the two images. Assuming

![Figure 1](http://rsif.royalsocietypublishing.org/) The Pair of images above are magnified and enhanced versions of raw footage obtained from the authors’ fieldwork in Mali.
motion model $F$ and measurement model $H$, the state of target $i$ satisfies
\[
X_i[k] = F(X_i[k-1], u)
\]
and
\[
Z_i[k] = H(X_i[k], n),
\]
where $u$ and $n$ denote disturbance and noise values, respectively.

Because of noise, disturbances and approximations in $F$ and $H$, the state estimate is a random quantity represented in the form of a probability density function (pdf). We recursively construct the filtering pdf $p(X[k]|Z^k)$ of the joint state $X[k]$ at time step $k$ given the set $Z^k$ of all measurements up to $k$ using the conditional probabilities $p(Z_i[k]|X_i[k])$ and $p(X_i[k]|Z_i[k])$. The probability $p(Z_i[k]|X_i[k])$, known as a likelihood function, is the conditional probability of measurement $Z_i[k]$ given the state $X_i[k]$. The filtering pdf can be obtained with minimum mean-square error provided the models $F$ and $H$ are linear and the noise values $u$ and $n$ are Gaussian. Otherwise, it is possible to use suboptimal methods such as an extended Kalman filter [24], which predicts and updates the estimate using a first-order linearization of (2.1), or a particle filter [25,26], which represents the target state as a point-mass distribution. A particle filter is attractive in that it relaxes most restrictions on the target and measurement models and the disturbance and measurement noise, but a particle filter is computationally burdensome for a large, joint, state space. We address the computation-size problem by making the assumption that the targets do not interact at short timescales (less than 40 ms), which allows us to use a separate filter for each target. Particle-filtering methods also allow us to encode extra information such as the velocity of the mosquito using the streak endpoints. (See the electronic supplementary material for a description of particle filtering.)

A multi-target tracking system must associate measurements and targets. A target-based method associates each target to a measurement [24], whereas a measurement-based method associates each measurement to a target [27]. A measurement-based method can inherently handle a variable number of targets, which may appear and disappear from the field of view. The reliability of the association depends on the proximity of the actual measurement to the predicted measurement, which is produced from the target estimate using the motion model and the measurement model; measurement proximity is determined using the position likelihood function.

Our choice of a data-association strategy is based on speed, variability and density of targets in the image. A nearest-neighbour association is target-based and associates the predicted measurement to the nearest measurement. It works well in low-target densities with high frame rates [22], but results in duplicate tracks and incorrect associations at high target densities. A global nearest-neighbour (GNN) association avoids duplicate assignments by minimizing a global assignment [28]. GNN has been successful in tracking dense aggregations [17,29] in which the number of targets are fixed and move in two dimensions (so that target overlap is rare), however, the possibility of a variable number of targets and frequent occlusions make it difficult to use GNN without additional heuristics. Short-duration occlusions can be addressed using motion coherence [19,21,30], whereas long-duration occlusions can be overcome by methods that minimize a global cost function over all measurements in a sliding window [20]. However, it is not clear how an offline global optimization method might address a low probability of detection, which is common in our datasets. Instead, we selected a measurement-based method called the multiple-hypothesis tracker (MHT), which looks into future assignment probabilities before making a decision on the current assignment [27]. Within the MHT, we use a motion model at each step to search for missing measurements.

A hypothesis in MHT is a combination of measurement-target assignments that satisfy the following two rules [27]: (i) a target is not associated to more than one measurement and (ii) a target is only associated to a measurement that lies within its gating volume. The gating volume or validation region is generated from the measurement that lies within its gating volume. The gating volume or validation region is generated from the measurement that lies within its gating volume. The gating volume or validation region is generated from the measurement that lies within its gating volume. The gating volume or validation region is generated from the measurement that lies within its gating volume. The gating volume or validation region is generated from the measurement that lies within its gating volume. Let $S$ be the covariance of $v$, which is also called the innovation. If a measurement that is normally distributed about the true value lies within the validation region, the weighted norm $|w|^2 = v^T S^{-1} v$ satisfies $|v| < t_{gate}$. (The quantity $|v|$ is also called the Mahalanobis distance [31].) For example, a threshold value $t_{gate} = 16$ defines a region around a predicted measurement with 99.97 per cent probability of containing the actual measurement [24]. A measurement may be assigned to an existing target, a new target or a false alarm. As time progresses, each hypothesis gives rise to successive hypotheses resulting in an exponential growth in time. Hypothesis reduction strategies include applying a threshold on track probability, choosing a few best hypothesis [31], and clustering the targets [27]. (See the electronic supplementary material for a description of MHT.)

Measures of effectiveness to evaluate a multi-target tracking algorithm include the following [32]: track initiation delay (timeliness), position and velocity errors (accuracy), fragmentation and identity swaps (continuity) and number of targets tracked (cardinality). Performance evaluation also includes visual verification, running the algorithm on simulated data [19], comparison with manually generated ground truth and reconstructing the same event from independent camera systems [30]. We evaluate the performance of our tracking algorithm using manually generated ground-truth as well as filming using two independent camera systems. For comparing tracking results with ground truth, we use the optimal subpattern assignment (OSPA) metric [33], which is a well-established metric for evaluating multi-target tracking algorithms [32] that allows comparison of sets with differing cardinality. (See the electronic supplementary material for details on computing OSPA.)

3. THE MOSQUITO TRACKING SYSTEM

The mosquito tracking system takes a sequence of stereo image pairs as input and produces three-dimensional tracks as output. Figure 2 depicts a block diagram of

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the tracking system, which was created in MATLAB and includes an automated tracking algorithm and a human-supervised component. Image streaks are modelled as straight lines; we extract the midpoint and endpoints as measurements. We find missing measurements using a gating volume generated around predicted measurements. Measurement pairs, i.e. one from each camera, that satisfy the epipolar constraint [34] are selected for data association. We again use a gating volume to assign measurements and targets to independent sets called clusters. Instead of generating definite tracks, hypotheses connecting measurements to targets are propagated to the next step using a particle filter. Based on the probability of each hypothesis at the current time step, the number of hypotheses at a previous time step are reduced to a single assignment. A particle filter verifies and combines tracklets under human supervision and the combined tracks are passed through a Kalman smoother. The tracking algorithm is summarized in Table 1.

The remainder of this section describes the novel aspects of the tracking system that we designed to improve its accuracy and level of automation. First, we describe an image-processing technique to find missing measurements during image segmentation. We then describe the measurement model that is used to extract velocity information from streaks. Finally, we present the data-association method, including the strategy to detect and address mosquito occlusions.

3.1. Extracting measurements

During observation of mosquito swarms, which typically appear silhouetted in front of swaying trees under a cloudy sky, it may not be possible to use a static (mosquito-free) background to segment the mosquitoes out of the image stream. Instead, we create a dynamic background by choosing the highest intensity point within a sliding window [35]. Let \( B_{u,v} \) be the background image value at the pixel position \((u,v)\) and \( t_{\text{win}} = 2d + 1 \) be the width of the sliding window centred at time step \( k \). The background value at time \( k \) is

\[
B_{u,v}[k] = \max_{i \in [k-2d+1:k+2d]} B_{u,v}[i]. \quad (3.1)
\]

The foreground \( F \) is obtained by subtracting the background \( B \) from the current image \( I \) and applying an intensity threshold \( t_{\text{th}} \), i.e. \( F_{u,v}[k] = \max(I_{u,v}[k] - B_{u,v}[k], t_{\text{th}}) - t_{\text{th}} \). We automatically select the value of \( t_{\text{th}} \) by running the background subtraction algorithm recursively on different segments of the image sequence until the number of blobs detected are within an acceptable range of the expected number of mosquitoes. We extract blobs using the \texttt{regionprops} routine in MATLAB, which performs connected-component labelling to extract features such as centroid, area and bounding ellipse. We remove large insects and birds from the foreground by applying a threshold on the blob area. (See Table 2 for the values of the threshold parameters.)

Owing to the duration of the camera exposure (\( \delta t = 25 \text{ ms} \)),\(^2\) fast mosquitoes (1–4 m s\(^{-1}\)) appear as elongated image blobs or streaks. Depending on the mosquito speed, the streaks may fail to appear in the foreground for a given value of \( t_{\text{th}} \). Existing strategies for low signal-to-noise environments include the track-before-detect approach [36], which permits raw sensor data as the input. The success for track-before-detect relies on the low target density and relatively straight movement of targets in the measurement space [37]. However, using raw sensor data is not a viable option for mosquito tracking, because it generates more false targets than observed in a single noisy image. Instead,\(^2\)

\(^2\)The duration of exposure (25 ms) is less than the time between frames (40 ms). The remaining time (15 ms) is for image processing.
we search for the missing streak in a new foreground generated using a threshold equal to \( t_w = 0.75 t_{\text{int}} \).
The search is performed within the gating volume of the predicted measurement. If a missing measurement is found, it is added to the list of existing measurements.

A measurement \( Z^c = [e^c, \mathbf{w}^c, e^c]^T \) from camera \( c \) contains the image locations of a streak’s start \( e^c \), midpoint \( \mathbf{w}^c \) and end \( e^c_\text{end} \). These values are extracted from a blob by modelling it as a straight line along the major axis of the bounding ellipse. The streak, therefore, represents a perspective projection of the mosquito trajectory for the duration of exposure \( \delta t_c \). Let \( \bar{\mathbf{w}}^c = [[\mathbf{w}^1]^T, 1]^T \) be the homogeneous representation of \( \mathbf{w}^c \). Assuming without loss of generality that camera 1 is the origin, a pair of measurements with midpoints \( \mathbf{w}_1^c \) and \( \mathbf{w}_2^c \), one from each camera, must satisfy the epipolar constraint [34]:

\[
|([\mathbf{w}_2^c]^T F \bar{\mathbf{w}}_1^c)| < t_e, \quad (3.2)
\]

where \( F \in \mathbb{R}^{3 \times 3} \) is the fundamental matrix for the stereo camera calibration and \( t_e < 1 \) is a value that depends on calibration accuracy. Measurement pairs from a true target satisfy the above constraint; clutter or mismatched measurement pairs should not. We use the midpoint and endpoint locations to define likelihood functions for position and velocity.

### 3.2. Position and velocity likelihood functions

A constant-velocity model suffices to describe the mosquito motion during the exposure, \( \delta t_c = 25 \text{ ms} \). (The streaks are well approximated by straight lines on the image plane.) Let \( \mathbf{r} \in \mathbb{R}^3 \) be the three-dimensional location of the midpoint of a streak. The start and end of the streak are located at \( \mathbf{r}_s = \mathbf{r} - \delta \frac{\mathbf{w}_1^c}{t_e} \) and \( \mathbf{r}_e = \mathbf{r} + \delta \frac{\mathbf{w}_2^c}{t_e} \), respectively. The corresponding point on the image plane is given by the perspective projection model [34],

\[
f^c(\mathbf{r}) = \begin{pmatrix} w_1 \ w_2 \ w_3 \end{pmatrix},
\]

where \( \mathbf{w} = P \mathbf{r} \in \mathbb{R}^3 \) and \( P \) is the camera projection matrix. Let \( \mathcal{N}(\mathbf{w}; f(\mathbf{r}), \Sigma) \) denote a normal density function evaluated at \( \mathbf{w} \) with mean \( f(\mathbf{r}) \) and covariance matrix \( \Sigma \in \mathbb{R}^{3 \times 2} \). Assuming that the measurement is normally distributed about the true value, the likelihood of midpoint \( \mathbf{w}^c \) given \( \mathbf{r} \) is

\[
P_{\text{mp}}(\mathbf{w}^c|\mathbf{r}) = \mathcal{N}(\mathbf{w}^c; f^c(\mathbf{r}), \Sigma_{\text{mp}}), \quad (3.4)
\]

We set the diagonal entries of \( \Sigma_{\text{mp}} \) equal to the length of the major and minor axes of the streak’s bounding ellipse in the streak frame; the off-diagonal entries are zero.

As with the midpoint likelihood function, we assume the endpoint likelihood function is based on a normal density function. However, owing to uncertainty in the labelling of the start and end of the streak, the endpoint likelihood function is bimodal. The directional ambiguity is described by a sum of conditional
probabilities on the order of endpoints. Let $\Sigma_{ep}$ be the covariance of the endpoint position in pixels (computed empirically). The endpoint likelihood function is

$$P_{ep}(e^c \cdot e^f \mid r, \dot{r}) = 1 - (1 - N(e^c \cdot f^c(r) \cdot \Sigma_{ep}) N(e^f(r), \Sigma_{ep})) (1 - N(e^f \cdot f^c(r), \Sigma_{ep}) N(e^c(r), f^c(r), \Sigma_{ep})),$$

(3.5)

where $r_\perp = r \pm t(\delta t/2)$. The combined effect of using a pair of points in the endpoint likelihood function (3.5) is to reduce the set of velocity values along the camera axis, which is otherwise unobservable.

The combined position and velocity likelihood function is

$$P(Z \mid X) = \prod_{c=1,2} P(e^c \cdot u^c, e^f \cdot r, \dot{r}) = \prod_{c=1,2} P_{mp}(u^c \mid r) P_{ep}(e^c \cdot e^f \cdot r, \dot{r}).$$

(3.6)

Figure 3 shows the combined position and velocity likelihood function. The likelihood function (3.6) is used to weight the particles in the resample step of the particle filter. We update the position estimates using triangulation [38], thereby effectively marginalizing out the position from the combined position and velocity filtering pdf.

A velocity likelihood function improves the reliability of data association by placing predicted measurements closer to the actual measurements. We compared the absolute velocity estimation error between a stand-alone position likelihood function and the combined position and velocity likelihood function (3.6). To create ground-truth data, we isolated a single mosquito track in both camera frame for 8 s. We then interpolate the position values to every 1/800th of a second. These values were then used to create an artificial mosquito streak during the time of exposure $\delta t_e$ from a 1 cm sphere. We then project the streak on a pair of white synthetic left and right camera images with resolution $1392 \times 1024$ pixels. To achieve a faded-streak effect, we reduce the intensity value of a pixel by 30 every time it is visited on the screen during the time of exposure. Mosquito motion normal to the image plane results in darker, shorter streaks, whereas motion parallel to the image plane results in lighter, longer streaks (figure 4a). We tracked this dataset using multiple Monte-Carlo runs of a particle filter. The combined position and velocity likelihood function performed better than the stand-alone position likelihood function, with an average improvement in mean absolute velocity error of 27 per cent (figure 4b).

### 3.3. Data association and occlusion resolution

Prior to weighting a target distribution with a likelihood function, we must first address the data-association problem. The mosquito data-association problem is challenging owing to the variable number of targets. To mitigate the uncertainty in association (for example, did the paths of two mosquitoes cross each other, or was it a close encounter?), we use a deferred-logic method called the MHT [27]. Each assignment of measurements to targets is set aside as a hypothesis and acted upon in a future time step when we are more certain. The certainty is computed using the probability of a hypothesis that depends on the innovation $v^c = u^c - f^c(r)$ of each measurement-target assignment in the hypothesis, the probability of detection of actual targets, and the covariance of the predicted measurement $\Sigma_{ep}$.

We reduce the number of hypotheses by clustering and prune them by selecting a few best hypotheses based on their probability at each step. Clustering is performed by dividing the measurement and hypothesized targets at each step into independent sets. At each time step, measurements are associated with each cluster based on the combined gating volume of all targets within the cluster. Measurements that do not belong to any cluster form their own clusters. Two clusters that consist of the same measurement are combined to form a single cluster. Similarly, we split clusters that consist of targets only assigned to a single measurement. Hypotheses are computed for each cluster independently. Hypotheses within a large cluster (more than 10 measurements) are limited to a
scanback [27] at each step, we choose the hypothesis with the highest probability to reduce to one the number of hypotheses at the previous step. Child hypotheses resulting from a pruned parent hypotheses are also removed.

New targets are automatically initialized from unassigned measurements and confirmed if they are tracked for more than three frames. New target distributions are sampled from a normal distribution with a low standard deviation in position (5 mm) about the triangulated point, and a large standard deviation in velocity (500 mm s\(^{-1}\)) about zero. The combined likelihood function resamples the distribution to equally favour particles getting projected on either side of the streak in the next time step.

Occlusions are not directly addressed as part of any data-association strategy, because existing strategies assume that motion coherence will automatically associate the right tracks in a future time step. In our case, occlusions undermine the velocity estimate, making future associations less reliable. An occlusion is detected if (i) two measurement pairs within a hypothesis consist of the same measurement from a single camera, or (ii) multiple hypotheses assign the same measurement to two or more targets. We interpret an occlusion as a combination of individual streaks, which are then used to extract velocity information as described in §3.1.

In order to cluster the pixels in an occlusion blob, we use the information about the number of mosquitoes hypothesized in the occlusion as well as their position and velocity estimates to model the blob as a mixture of Gaussians. An expectation-maximization algorithm [39] uses position estimates for initial means and velocity estimates for covariance in each dimension to hard-cluster the pixels into individual streaks. This set of individual streaks is used as an initial guess to soft-cluster\(^3\) the pixels into more accurate overlapping streaks. Using the shortest distance of a pixel from the line that passes through the split streak, we allow multiple assignments of each pixel to individual streaks.

\(^3\)Soft clustering allows a single pixel to be assigned to more than one cluster, whereas hard-clustering assigns each pixel to exactly one cluster.

4. DATA COLLECTION AND TRACKING RESULTS

To validate the mosquito-tracking system and film mosquito swarms in the field, we used a pair of phase-locked Hitachi KP-F120CL cameras in a stereo configuration. Each camera captured 10-bit images at 25 frames per second and 1392 × 1040 pixel resolution. Figure 6 shows a schematic of the data collection system. The video streams were recorded using a 2.8 GHz quad core laptop, an Imperx FrameLink Express frame grabber (Imperx Inc., Boca Raton, FL USA), and STREAMPIX v. 5 software (Norpix Inc, Quebec, Canada). Each camera was calibrated onsite using a checkerboard and the MATLAB Calibration Toolbox [40]. Reprojection error, which is a measure of calibration accuracy, was in sub-pixels for each camera. Relative camera orientation and position were determined by extrinsic calibration by taking multiple pictures of a stationary checkerboard with both cameras. During filming, the camera height, azimuth and elevation were recorded to create a ground-fixed reference frame. We used a Kestrel 4500 portable weather station (Nielsen-Kellerman, Boothwyn, PA, USA) to sample other environmental factors such as wind velocity and humidity at 0.1 Hz.

Filming was done in the village of Donéguebogou, Mali in Western Africa. Donéguebogou is 29 km north of Bamako and has been the site of previous research on *A. gambiae* mosquitoes [5,10]. Swarms formed approximately 20 min after sunset, initially with only one or two males then increasing in numbers, and lasted for 20 min. Most couples were seen 5–10 min after the swarm was first observed. Couples formed only for a few minutes during this period, then were no longer observed, though the males continued to swarm for many minutes after the last couple had formed. We filmed swarms of *A. gambiae* that formed over bare ground or markers.

Female mosquitoes are difficult to detect and track because they fly faster than the average male (see §4.3) and appear as a faint streak much of the time. However,
a mosquito couple is distinguishable to the human eye owing to its distinct flying pattern and darker appearance against the sky. Upon spotting a couple, we noted the frame number displayed on the laptop screen. The couples were located after filming by manually reviewing the video footage at the designated frame. Out of the two mating mosquitoes, the female mosquito was identified as the mosquito that entered the swarm last. We tracked the pair, first as a couple and then individually, by playing the sequence backwards. Parameter values used for data collection and tracking are described below and in table 2. The validation and evaluation of tracking performance follow.

4.1. Parameters used for data collection and tracking

The camera baseline \( b \), i.e. the distance between cameras, affects the disparity \( \Delta u \) in pixel positions of an object in a stereo camera set-up [34]. A large disparity reduces uncertainty along the camera axis, which in turn improves accuracy as well as the ability to resolve occlusions. For a stereo-camera configuration with focal length \( f \) and no vertical offset between centres, the baseline and disparity are related according to \( \Delta u = (bf)/z \) [34], where \( z \) is the distance along the camera axis of the target from the stereo set-up. The overlap between camera views is \( (I_w/\Delta u)/I_w \), where \( I_w \) is the image width resolution in pixels. A large overlap is desirable for maximum coverage. Since the majority of swarms were filmed with \( 1.5 \leq z \leq 2.5 \text{ m} \), we selected a baseline of 20 cm to achieve 80–90% overlap and three to five pixel separation between two mosquitoes that are 3 cm apart (approx. two body lengths) along the camera axis.

In addition to the intensity threshold \( t_{\text{int}} \) described in §3.1, foreground segmentation requires setting the sliding window \( t_{\text{win}} \) and a threshold on area of the blobs \( t_{\text{area}} \). We selected \( t_{\text{win}} = 7 \) frames centred on the current frame, although swarms filmed at short ranges required a sliding window in the range of 3–5 frames. We computed the area-threshold limits \( 20 \leq t_{\text{area}} \leq 150 \) from several different swarms to achieve the best rejection of noise as well as large insects.
The covariance $\Sigma_{ep} = \text{diag}\{4, 4\}$ pixels$^2$ for endpoints was computed by manually selecting the endpoints of streaks in a random sampling of frames and comparing with the calculated value. The disturbance $w$ for the constant-velocity model was sampled from $N(0, 100 \text{m}^2 \text{s}^{-4})$, whose covariance was found by fitting a normal distribution to the acceleration values of manually generated tracks.

### 4.2. Validation and performance of tracking system

We tested the position accuracy of our tracking system using a calibration checkerboard with squares of known dimensions by manually clicking pairs of points whose separation distance was in the range 3–40 cm. This method yielded an error of $5 \pm 5$ mm for 50 pairs. Average position error by tracking an artificial mosquito projected on the stereo images as in §3.2 over multiple runs was $5 \pm 4$ mm. We also reconstructed tracks from a single swarming event on Aug. 29 using two independent stereo camera rigs. We created a common reference frame by measuring the height, azimuth and elevation of the cameras. The videos were time-synced using a laser pointer flashed at the end of the sequence. The mean distance between independent tracks of the same mosquito (200 data points) was $4.4 \pm 1.3$ cm, although up to 3 cm error can be attributed to the inter-frame time difference between the camera systems (caused by a possible mismatch in the pair of frames that contained the laser flash). A mosquito flying at an average speed of $1.5 \text{ m s}^{-1}$ will cover 3 cm in 1/50th of a second.

Figure 7 shows the results of using the OSPA metric (see the electronic supplementary material, equation (3.3)) to compare tracks from the multi-target tracking system to the manually generated ground-truth. We tested two swarms with 10 and 20 mosquitoes, respectively. The order parameter and the cut-off parameter for computing OSPA values were set 2 and 50 mm, respectively. Decomposing OSPA into position and cardinality errors shows that the average root mean square (RMS) position errors in the 10- and 20-mosquito swarms were $2.17 \pm 0.58$ and $2.3 \pm 0.46$ cm, respectively. Correspondingly, average absolute position errors for the 10- and 20-mosquito swarms were $1.74 \pm 0.56$ and $2.03 \pm 0.47$ cm, respectively. A low cardinality error was often accompanied by relatively high position error during periods when the swarm was dense, because of occlusions and false tracks. As would be expected for a stereo set-up, position error was highest (44%) in the range measurement (along the camera optical axis) when compared with either of the other two dimensions. OSPA was larger for the 20-mosquito swarm, mainly owing to cardinality errors. The position error is likely a consequence of image noise, which resulted in partially segmented streaks. (We mitigate this problem by filtering trajectory data using a Kalman smoother.) Average reprojection error on the images was less than two pixels.

The labelling error, which captures track continuity and identity swaps, was computed separately. An identity swap results in a labelling error of 2 before or after the swap in the sequence. Track fragmentation results in a labelling error of 1 after the disconnect occurs. We randomly selected 100 instances of 25 continuous frames in a swarm of 10 mosquitoes. The average labelling error (most of which was due to track fragmentation) was $2.1 \pm 1.4$ tracks. A simple average of track lengths across six swarms ranged between 15 and 25 frames corresponding to 0.6–1 s. Track fragmentation occurs owing to early terminations, which can be caused by the following:

- partially segmented streaks owing to noise, cloudy background and clutter. Partially segmented streaks
in one frame often violate the epipolar constraint. Decreasing the intensity threshold to get full streaks adds noise to the measurements. (A possible solution that we are exploring in ongoing work is to reconstruct the streak using velocity estimates.); and
— occlusion between a tracked and untracked target. Occlusions between a tracked (known) target and an as yet uninitialized target are not detected. The success rate of surviving such an occlusion depends on the motion of the tracked target after the occlusion. A manoeuvre or successive occlusion may terminate the track.

4.3. Tracking results

This section presents a subset of the three-dimensional trajectory data generated using the mosquito tracking system. We filmed 21 swarms and 13 mating events between 17 August 2010 and 3 September 2010. Out of the 21 swarms, 18 formed over bare ground and three formed over natural markers. (A natural marker is an area of high contrast with the rest of the ground such as a patch of grass.) Anopheles gambiae can be divided into two incipient species namely the M and S molecular forms [42]. In Diabaté et al. [10], a strong association between the swarming marker type and molecular form has been found. The M form was found to swarm over natural markers, whereas the S form swarms over bare ground [10]. We collected a few mosquitoes from each swarm and performed a polymerase chain reaction (PCR) test to determine the molecular form. All sequences presented in this paper were of type S. Each day two teams of three to five people with identical camera rigs selected separate swarming sites for filming. The swarming sites were usually within a few hundred metres of each other. Swarming sites were surveyed the day before to record average swarm size and location. Filming locations spread throughout the village (figure 8) were chosen based on swarm size (less than about 100 mosquitoes for tractability in tracking) and the presence of few trees or houses in the background (i.e. in the direction of the setting sun). Once filming began, 60–90 s stereo video sequences were recorded as 10-bit synchronized tiff images on separate solid-state drives. (The drives were backed up daily on two separate discs.) A filming session typically produced five to eight video sequences before it became too dark to film.

To create representative trajectory dataset, we selected six video sequences that contain a mating event. We call these the mating sequences. We refer to the mating mosquito as the female and the focal male. We also selected six other video sequences with no female present, called the male-only sequences, to produce full-length trajectories of swarming behaviour. Trajectory data presented here are from swarms filmed on 20, 21, 25, 26, 28 and 29 August and 1 September. Male-only sequences last between 20 and 35 s, whereas mating sequences start a few seconds prior to the detection of female within the field of view and end when the couple flies out of field of view (0–5 s).

Figure 9 shows the position and velocity of a randomly selected male A. gambiae in the Aug. 29 male-only sequence, which was a swarm that formed over bare ground (S molecular form). The swarm consisted of 20 mosquitoes at the beginning of the
sequence and dropped to 19 after 10 s. The mosquito movement is characterized by quasi-periodic motion in each of the three spatial dimensions. The instantaneous mean position of the mosquitoes in the swarm, i.e. the swarm centroid, is also shown. The origin of the inertial frame is located at ground level under the camera rig; the inertial frame is oriented along east–west, north–south and vertical directions. The 3σ bounds for position and velocity of all of mosquitoes in this swarm are shown in grey. Swarm size (twice the 3σ bounds) averaged 1.17 m in the horizontal plane and 0.56 m in the vertical. The average swarm size across all planes ranged between 0.52 and 1.86 m. The average height of the swarm was 1.89 m. The average velocity along each dimension is close to zero with highest standard deviation in the east–west direction (0.514 m s⁻¹).
m s\(^{-1}\)) followed by the north–south (0.332 m s\(^{-1}\)) and the vertical (0.281 m s\(^{-1}\)).

Figure 10 shows the ratio between horizontal and vertical speed for each swarm. The Aug. 28 sequence was filmed on a day with relatively high wind (approx. 0.8 m s\(^{-1}\)) when compared with other sequences. The mosquito movement for that swarm was characterized by a rolling motion in the direction of the wind and relatively higher vertical velocities. In five out of the six swarms that we used to generate male-only sequences, we witnessed mating events at a later time. The horizontal and vertical speeds of female mosquito that formed couples are also plotted in figure 10. Non-parametric Kruskal–Wallis tests on each dataset show that the average male and female speeds in the same sequence are significantly different for each sequence. The maximum p-value among all mating sequences was 0.0003. (In contrast, the maximum p-value for male speeds during the same mating sequence was 0.051.)

Figure 11 shows the position and velocity of a female mosquito that formed a successful couple in the Aug. 29 sequence. The mating sequence was filmed about a minute after the male-only sequence on the same date. The female appeared in the field of view 5 s prior to coupling. The movement of the female crosses the 3σ boundaries of the swarm in the north–south direction.
The average speed of the female was higher than the male mosquito until just before the couple forms, when the focal male speeds up. The vertical movement shows that the female stayed predominantly at heights corresponding to the lower half of the swarm. A three-dimensional reconstruction of the mating mosquitoes in six mating sequences is shown in figure 12. Across all mating sequences, the female mosquito covered an average 59 per cent more distance than the focal male during the same time interval.

Figure 13 shows the separation distance and speeds from six mating sequences. The amount of time we observed the females in the swarm before forming a couple was up to 5 s. In each mating sequence that lasted longer than 0.5 s, the number of close encounters (moments when the separation distance between the mating mosquitoes dropped below three body lengths, or 4 cm) with the successful male mosquito was in the range 3–6.

5. CONCLUSIONS AND ONGOING WORK

We describe a tracking system to reconstruct the three-dimensional trajectories of wild mosquito swarms. We address noisy images by adaptively seeking missing measurements and exploit streak orientation and length to extract velocity information. A probabilistic data association method that uses multiple hypotheses is modified to address occlusions. We evaluate the system using an established multi-target tracking metric and validate using independent measurements of the same swarm. Tracking results are presented in the form of three-dimensional trajectories of swarming and mating mosquitoes. To date, the data produced from the tracking system described in this paper are an order of magnitude larger (97 trajectories and 55 000 position points) than the last published result on reconstruction of wild mosquito swarms, and the first to contain three-dimensional trajectories rather than three-dimensional positions. In ongoing work, we are investigating these trajectories to characterize swarming and mating behaviour.

As part of ongoing work on the tracking system, we are working to include the streak intensity in the image as part of the likelihood function. This will help predict the appearance of a mosquito on the image plane as a function of its velocity, thereby allowing the possibility of streak retrieval. Such an approach would, for example, reduce track terminations and create longer tracklets. Another aspect of the tracking system that we are investigating is the automatic detection of mating events. In order to avoid sifting through video streams to locate mating events, the distinct flying pattern and appearance of the mating couple...
can be used for automatic detection and backwards tracking of the female. With enough mating events, a higher order motion model (that depends on more than one previous time step) will automatically predict and detect mating events.

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