RESEARCH

A minimal 3D model of mosquito flight behaviour around the human baited bed net

Jeff Jones, Gregory P D Murray and Philip J McCall*

Abstract

Background: Advances in digitized video-tracking and behavioural analysis have enabled accurate recording and quantification of mosquito flight and host-seeking behaviours, facilitating development of individual (agent) based models at much finer spatial scales than previously possible.

Methods: Quantified behavioural parameters were used to create a novel virtual testing model, capable of accurately simulating indoor flight behaviour by a virtual population of host-seeking mosquitoes as they interact with and respond to simulated stimuli from a human-occupied bed net. The model is described, including base mosquito behaviour, state transitions, environmental representation and host stimulus representation.

Results: In the absence of a bed net and human host bait, flight distribution of the model population was relatively uniform throughout the arena. Introducing an unbaited untreated bed net induced a change in distribution with an increase in landing events on the net surface, predominantly on the sides of the net. Adding the presence of a simulated human bait dramatically impacted flight distribution patterns, exploratory foraging and, the number and distribution of landing positions on the net, which were determined largely by the orientation of the human within. The model replicates experimental results with free-flying living mosquitoes at human-occupied bed nets, where contact occurs predominantly on the top surface of the net. This accuracy is important as it quantifies exposure to the lethal insecticide residues that may be unique to the net roof (or theoretically any other surface). Number of net contacts and height of contacts decreased with increasing attractant dispersal noise.

Conclusions: Results generated by the model are an accurate representation of actual mosquito behaviour recorded at and around a human-occupied bed net in untreated and insecticide-treated nets. This fine-grained model is highly flexible and has significant potential for in silico screening of novel bed net designs, potentially reducing time and cost and accelerating the deployment of new and more effective tools for protecting against malaria in sub-Saharan Africa.

Keywords: Mosquito, Malaria, Vector control, Host-seeking, Bed net, Behaviour, Computational model, in silico, tracking, Insecticide, Repellent, Anopheles, Attractants

Background

Whether to combat insecticide resistance, exploit new knowledge of vector biology, increase community acceptance or to accommodate changes in abiotic conditions, mosquito vector control methods are under constant pressure to improve. In sub-Saharan Africa, where 90% of malaria occurs, long-lasting insecticidal nets (LLINs) have been the main driving force in the reduction of malaria cases [1, 2], but widespread and growing insecticide resistance to the pyrethroid insecticides used on nets has stalled, and threatens to reverse, recent advances [3–6]. Novel insecticide compounds used either alone or in combination with existing pyrethroids, provide one solution [7, 8] but this raises significant questions regarding the optimal placement of compounds on the net walls or roof, both for maximizing safety and efficacy and...
minimizing cost. A detailed understanding of the spatio-temporal nature of mosquito responses to various insecticidal treatment(s) on the bed net interface is critical to balance these competing requirements for the numerous combinations possible.

The development and eventual implementation of a new bed net is an expensive and time-consuming process and the pipeline from early phase screening of chemicals with potential through the range of laboratory and field tests needed to generate the evidence required by regulatory authorities before the commodity finally reaches the affected communities, can take up to ten years.

The first studies on the behaviour of mosquitoes at the bed net interface demonstrated that significantly greater numbers of mosquitoes landed on the bed net roof [9, 10]. Later technological developments in imaging and computing enabled tracking of entire flight paths [11, 12], and 3D reconstructions of arrival patterns [13, 14] revealed complex but consistent behaviours, ultimately revealing how insecticide treatments on bed nets affect the behaviour of malaria vector mosquitoes. The growing body of data arising from these studies not only builds the evidence base required to accelerate the development process, but it also provides an excellent foundation for developing models of host-seeking behaviour with potential to validate experimentally the new tools at earlier stages. One such model is presented here: a fine-grained agent-based approach for modelling how indoor insecticide treatments deployed as residues on bed nets affect the behaviour and survival of mosquitoes.

To successfully find, select and feed on a human host, mosquitoes process input from multiple sensory modalities with the relative importance of particular cues differing at different stages of the approach. These include isolated carbon dioxide (CO2) concentration, olfactory [15–17], visual [18, 19], auditory [20, 21] and tactile sensory cues [22], or combinations thereof [23].

The interaction between host-seeking mosquitoes and the human host begins long before the mosquito enters the house but the quantity of sensory information is likely to be at its most intense in a room around a human-occupied insecticidal bed net. This is a unique context for a mosquito behaviour model as most modelling approaches to disease transmission by insect vectors either set the scale at a much higher level, emphasizing interactions at village scale (and above) [24, 25], or at the opposite end of the scale, modelling individual mosquitoes in experimentally controlled settings such as wind tunnels and olfactometers [26].

Until there is a more thorough understanding of the mechanisms guiding mosquito host location processes, the development of accurate fine-grained models of insect flight behaviour and host interactions will be restricted. Instead, agent-based modelling provides an alternative approach because, as noted in [27], it can represent stochasticity and heterogeneity at fine-grained scales and generate emergent behaviours not explicitly encoded into the model. Previous agent-based approaches to mosquito flight behaviour have included a 2D model of insect flight to study orientation and tracking of odour plumes [28], and a 2D model of mosquito interactions with insecticidal bed nets which was used to assess community scale protection [29].

The unresolved complexity of the roles and relative importance of different sensory stimuli were circumvented by developing a minimally simple agent-based model of 3D mosquito flight behaviour. The model is stimulus-agnostic and uses a single generic ‘attractant’ signal emanating and dispersing from a host. The model was used to study mosquito flight behaviour and landing distribution patterns on unoccupied and human-occupied bed nets. The eventual aim is to employ the model as a virtual experimental tool for accelerated exploration and evaluation of novel bed net treatments or designs. Hence, model parameters were validated with the results from previous experimental work on mosquito behaviour at the bed net interface.

Materials and methods

The Indoor Vector Control Testing System (InVeCTS) model creates a virtual environment in which to assess mosquito populations’ interactions with their host and environment. It uses an agent-based approach with fine-grained spatial representation in which a mosquito population can interact with a human host emanating the hypothetical spatially distributed attractant stimulus over time. Mosquito flight occurs in real time and all mosquito flight paths and interactions with the environment are recorded for subsequent analysis.

Environment and mosquito representation

A population of 25 mobile virtual mosquitoes is created. The flight tracking system data used to explore the parameter space of the model mosquitoes was generated from experiments with adult female Anopheles gambiae sensu stricto (s.s.), generally considered the most important vector of all, using the “Kisumu” strain, a widely used colony-raised insecticide susceptible strain. These individuals fly in a continuous 3D space representation inside a discrete spatial arena, representing an insectary whose dimensions directly correspond to an experimental insectary at Liverpool School of Tropical Medicine (LSTM) (5.6 m long × 3.6 m wide × 2.3 m high). This virtual insectary can contain a bed net and human host, as shown in Fig. 1.
The population is introduced at the release site (Fig. 1e) and begins exploration of the arena. A hypothetical attractant plume is projected with the size and shape of a human host of approximately 180 cm in height. To represent and store the attractant profile the arena is divided into a $56 \times 36 \times 23$ cubic lattice of $10 \text{ cm}^3$ cells. The host bait profile is configurable to represent hosts of different sizes (Fig. 1d) and the cells making up the bait profile are colour-coded to indicate regions where greater concentrations of attractant are emanated. As previously stated, a wide range of environmental cues (for example, CO$_2$, skin odours, sound, vision, temperature) are known to influence mosquito behaviour but their relative contribution and sequential importance in host location is still uncertain [30–32]. To circumvent this uncertainty, and to simplify the model, a generic representation of a single spatial attractant emanating and diffusing from the host was used. This is a simplifying assumption that the multimodal nature of the attractant profile can be represented in this manner. A simple cellular automata-based dispersion mechanism was used in order to enable real-time dispersal of the attractant profile.

**Projection and dispersal of human bait attractant**

At each model time step, attractant is projected at bait profile locations and isotropic dispersion is implemented in parallel for all cells in the 3D lattice volume by distributing the attractant between the current cell and the 26 nearest neighbour cells. Vertical dispersion (a very coarse approximation of convection) is implemented for all cells by dispersing a fraction of each cell to the cell immediately above the current cell in the arena. Such a simple scheme was used for two reasons. Firstly, this is because of the uncertainty of the relative importance of different environmental cues, and secondly, it enables the visualization of the dispersal in real-time which is not yet possible with more complex representations of environmental dispersion.

Real-time projection, diffusion and convection of bait profile attractant in the lattice is parameterized by independent weight parameters for diffusion ($W_d$) and convection ($W_c$). At every time step $t$ a generic chemottractant, weighted by the bait profile map (a simple representation of a human form resting within the bed net, see Fig. 1d), is projected into the 3D attractant lattice at locations directly corresponding to the bait map profile shape. The bait profile map values are weighted to account for regions projecting higher levels of attractant, for example, from the mouth.

**Attractant plume diffusion**

At every model time step $t$, volumetric diffusion of attractant is approximated by the following cellular automaton-based method.

For each cell in the attractant 3D lattice (axes $x,y,z$):

$$C_{xyz}^{(t+1)} = (C_{xyz}^t + \text{local neighbourhood attractant values (radius 1)})/27) \times W_d$$

To approximate non-uniform dispersal of attractant (for example, turbulent air currents) and their effects on host-seeking behaviour each cell can be modified by adding a value randomly sampled from a Gaussian distribution with mean 0 and standard deviation 1. Perturbation by Non-uniform dispersal can be increased by multiplying this noise value by a scaling parameter $D_s$. 

---

**Fig. 1** Arena and mosquito representation in the InVeCTS model. Left: Mosquito flight environment. The environment consists of the main insectary arena comprising **a** a cuboidal recording volume corresponding to that used in [11] **b** a bed net **c** containing a human host **d** Mosquito release location is indicated in **e**. Right: Schematic representation of virtual mosquito agent with its current directional vector (long arrow), its offset Left and Right sensors (L,R), Sensor Offset distance (SO), Sensor Angle (SA) and Rotation Angle (RA) parameters.
**Attractant plume convection**

Directly after the diffusion step volumetric convection upwards is approximated by copying of vertically stacked horizontal ‘sheets’ of cells containing attractant values to cells in the sheets above (Insectary ceiling is set to \( Y = 0 \), layers below are \( Y + 1 \) etc). Projection is weighted by convection weight parameter \( W_c \).

For each cell in the attractant lattice (axes \( x, y, z \)): 
\[
C_{xyzt}^{(t+1)} = (C_{x+y+1}^{(t+1)}) \cdot W_c
\]

A visualization of the effect of attractant plume dispersion is shown in Fig. 2. To reiterate, this is a simple approximation of attractant dispersion. However, it is sufficient to generate a stable 3D attractant field and avoids emphasizing any one particular (potentially erroneous) stimulus type or sensory modality over others.

**Mosquito sensory-motor Algorithm**

After the diffusion and convection stage each mosquito samples the attractant value in the insectary from two offset sensors (sensor offset distance may be adjusted with the parameter \( SO \). Sensor angle may be adjusted with the parameter \( SA \). If the sensor values are different the agent orients itself locally in space by rotating about its own axis towards the strongest value, by the amount in degrees given by the Rotation Angle (\( RA \)) parameter. Additionally, the agent orientation is subject to random modification by the \( pCD \) (probability of Change Direction) parameter. If a randomly sampled value from the uniform distribution is \(< pCD \) (default value 0.25), the agent will select a random orientation in 3D space. At each model step each agent moves forwards in its current orientation by a velocity given by the parameter \( V \) (default value 1 cm).

**Mosquito behaviour transition function**

Agent behaviour in response to environmental cues is determined by a set of states and the subsequent transition between these states. To avoid a fully deterministic response to population behaviour, transition between states is mediated by probabilistic sampling, implemented by individual parameters, corresponding to a Markov process of state transitions (see Table 1 for all model parameters). A schematic overview of the behavioural transitions can be found in Additional file 1.

The agents are initially in the **PRE-RELEASE** state and their state is updated every model step. A mosquito enters the **FLYING** state when a randomly sampled value from a uniform distribution is \(< 0.1 \). If an agent is at a wall, ceiling, bed net, or floor surface it will rest if it lands on or is resting on a treated bed net its start position, or mosquito release location, and experimental run time. Although measurements of mosquito flight speed vary in the literature (and indeed during different mosquito behaviours) a mosquito flight speed of 300 mm/s was selected to approximate the flight speed recorded experimentally under the same conditions [12].

Varying the \( SA \) and \( RA \) parameters affects flight tortuosity which subsequently affects arena occupancy and foraging behaviour. Examples of the effects of varying \( RA \) parameter can be seen in Additional file 2. After evaluating a range of \( SA \) and \( RA \) angle combinations, fixed values were selected, \( SA \) of 45° and \( RA \) of 22° (see Additional files 2 and 3 for the evaluation of sensory parameters and description of tortuosity calculation). This resulted in path tortuosity which approximated those of the image tracking experiments reported in [11].

### Table 1: Model parameters. Parameters are divided into groups reflecting the effect on the environment, agent sensory-motor behaviour, and agent behavioural transitions

| Parameter type | Name | Default value | Description |
|----------------|------|---------------|-------------|
| **Environment** | \( W_d \) | 0.7 | Diffusion damping factor |
| | \( W_c \) | 0.3 | Convection damping factor |
| | \( D_s \) | 0 | Dispersal noise value |
| | \( tx \) | 0 (untreated net) 0.1 (treated net) | Net contact toxicity value |
| **Mosquito Flight** | \( SA \) | 45 | Sensor angle (deg) |
| | \( RA \) | 22 | Rotation angle (deg) |
| | \( SO \) | 1 | Sensor offset distance (cm) |
| | \( V \) | 1 | Velocity (cm) per model step |
| **Mosquito State** | \( pCD \) | 0.25 | Probability of changing direction |
| | \( pRest \) | 0.1 | Probability of resting on surface |
| | \( pLeave \) | 0.05 | Probability of leaving surface |

in the current location until a randomly sampled value from a normal distribution is \(< pLeave \) parameter. It the agent lands on, or is resting on a treated bed net its starting health value (100) is decremented by a toxicity value \( tx \). If the health of an agent reaches zero it is removed from the arena.

### Parameter selection and model validation

Model parameters were set to match values obtained by previous image tracking experiments in the same LSTM insectary space [12]. The dimensions of the virtual arena were identical to the insectary, as were positions of the bed net, host, number of mosquitoes, mosquito release location, and experimental run time. Although measurements of mosquito flight speed vary in the literature (and indeed during different mosquito behaviours) a mosquito flight speed of 300 mm/s was selected to approximate the flight speed recorded experimentally under the same conditions [12].
**Fig. 2** Visualisation of attractant plume within the arena. Left side shows orientation of cross-sectional slice through attractant field within the arena and camera position. Right side shows attractant concentration scaled individually for each image to the look-up table (top). 

- **a** Gradient concentration of attractant plume (looking from above insectary, slice taken from coronal plane, 0.7m above floor).
- **b** Gradient concentration of attractant plume (looking along axial plane of human profile). Section is taken at the head region of the human profile.
- **c** Gradient concentration of attractant plume (looking at side-on human profile). Section is taken as a sagittal slice along the middle of the human profile.
Limiting assumptions of the model are described in the discussion section.

**Model scheduler and model experiment setup**

At the start of an experiment, the system creates the virtual insectary environment and the agent population. The population is placed at the release location and the model begins iterating through its main run-time loop as shown in Fig. 3.

Each model step comprises 1/30th second. The model halts after 1 h, or 108,000 model steps. After the model halts, log files pertaining to virtual mosquito activity (three-dimensional coordinate positions, agent state and agent health), bed net contact locations, and agent presence within spatial regions surrounding the bed net are saved to disk for further analysis.

The user can interact with the model during run-time, visualizing the current state of the model and population flight behaviour in 3D space using a mouse. Specific experimental configurations can be saved and re-loaded for repeated runs.

Five control experiments were performed in the virtual insectary where no bait or no bed net was present, as a baseline for the flight behaviour of the model. Twenty experiments were performed for each of the human bait conditions (no bait, head facing left, and head facing right). The head facing left profile was used assess the effect of noisy attractant dispersal, performing five experiments at each noise parameter value.

**Results**

**Spatio-temporal flight activity and foraging behaviour.**

Control experiments, where neither bait nor bed net were present, established baseline flight behaviour and distribution. Occupancy of all regions is plotted in Fig. 4 across the X and Z planes. The heatmap images indicate a top-down summary of occupancy results from above the entire insectary, demonstrating that in the absence of a host or bed net, flight activity was relatively uniform throughout the arena (Fig. 4a).

When an unbaited net is introduced to the arena, the flight distribution pattern in the arena is still relatively uniform except in those regions containing the empty bed net where the mosquito population cannot enter (Fig. 4b). When the stimulus profile of a human shape is placed on the bed, within the net, the flight distribution shows a marked change, with the population showing a strong preference to fly in the regions of the strongest diffusing and convective stimuli, corresponding to the orientation of the human bait profile (Fig. 4c and d).

As in nature [11], foraging in the InVECTS model population is strongly affected by the presence or absence of a human host. Figure 5 (top) demonstrates spatial foraging at 11 min into a 1-h experiment. In the presence of bait, a narrow field of the arena is explored, whereas the virtual population explores a much larger area of the arena when no host is present. Temporal exploration of arena occupancy over time (the total fraction of the arena explored by the population) confirms that the presence of bait results in less foraging (indicative of behaviour oriented towards a source). When bait is absent the population adopts an exploration strategy, with many casting flights as they seek recognizable host cues. Although it is difficult to compare foraging activity on a per-mosquito basis (due to the high individual variability in flight behaviour) the initial time taken to find and contact the net in each experiment was lower in baited conditions (mean 270 model steps, sd +− 42.23) than in unbaited conditions.
conditions (307 model steps, sd +− 101). These differing behavioural responses are induced solely by the presence or absence of dispersed attractant stimuli within the arena. Example video recordings of short model runs in unbaited and baited conditions are shown in Additional files 4 and 5 respectively and the mosquito trails indicate differences in foraging behaviour.

**Flight path tortuosity**

Flight path tortuosity was calculated using the rolling tortuosity metric described in Additional file 3. To summarize, a sliding window of 50 mosquito movements along the flight path was assessed for all points and only uninterrupted segments of flight paths were used (i.e. those not including landing and resting). Path tortuosity under different bait conditions are shown in Table 2. Tortuosity in unbaited arenas was slightly lower than measured previously in [11], however that study only included information from the recording volume captured by the cameras (where tortuosity is increased by the mosquito responses to the net). The model tortuosity includes the entire insectary space with more free flight space which reduces overall tortuosity. Under baited conditions (which attracts greater mosquito activity at the net) the model tortuosity closely matches experimental values.

**Net surface contact distribution**

Net contact locations in the unbaited condition occurred mainly along the sides of the bed net, particularly along the longest axis of the net (Fig. 6, top), with relatively little contact at the top surface of the net.

Under simulated human bait conditions, contact events showed a predilection for the top surface of the net and the X coordinates of the contact patterns were strongly influenced by the orientation of the human bait beneath (Fig. 6, bottom). An illustration comparing the distribution of net contact numbers and spatial distribution patterns in unbaited and baited experiments is shown in Fig. 7.

The total time spent in contact with the net surface itself is shown in Table 3. The mean contact time for an unbaited net (1.24 m for the model, compared to 2.4 m in the experimental results) is far less than the time spent at the baited net (31.51 m for the model versus 33.1 m in the experimental results). In response to a simulated treated net the model population spent a mean of 13.69 m in contact with the net, compared to 7.3 m in the experimental results. The difference in contact time at the treated net may partly be explained by the fact that the model does not incorporate a representation of internal mosquito energetics which may be responsible for reduced mosquito activity over time [11].

Furthermore, the range of contact times over multiple number of experiments in the model is also slightly less than the experimental findings for all bait and treatment conditions. This reflects the high variability of individual mosquitoes and suggests that stochasticity in the model could be increased in future research. This could be achieved by increasing individual virtual mosquito stochasticity (via the probabilistic mosquito state parameters) or by increasing the global stochasticity of the attractant signal via the dispersal noise parameter.

It has been suggested that strong air currents may affect host-seeking behaviour and, potentially, subsequent mosquito distribution on bed nets [17]. The effect of noise-contaminated attractant dispersal on virtual mosquito host-seeking and the number and spatial distribution of bed net contact locations was assessed. Figure 8 demonstrates the spatial effects showing, at low noise levels, a wider region of contact on the net surface (Fig. 8a–e). At higher noise levels, the mosquitoes increasingly contact the net at the sides, as opposed to the top surface (Fig. 8f–h). This change of net contact height with increasing noise (Fig. 9a) is accompanied by a reduction in total net contact (Fig. 9b).

**Peripheral bed net region activity**

To assess peripheral activity within the regions surrounding the bed net the recording volume was subdivided into polyhedral shapes similar to, but not exactly identical, to the polygonal regions used in an earlier experiment [11]. The reason for the differences is due to the 2D nature of the projection of the image tracking method and the slight difference in net orientation used in [11] (the net was tilted to allow better visualisation of the top surface of the net for the capture cameras). The outer regions of the bed net in the model are divided into 18 regions which allows differentiation between the top surface of the net (12 sub regions), the two end edges of the net (edges corresponding to those near the head and feet) and the two sides of the net (4 regions consisting of arms and legs for both sides of the net). A 3D visualization of the peripheral regions is shown...
in Additional file 6. Occupancy of the regions is recorded during experimental runs. Each time a mosquito is located within a particular region, a counter for that region is incremented. The total mean region distribution data for control and baited conditions is summarized in Additional file 7.

A visual summary of flight tracks and peripheral region occupancy of the model output compared to experimental data in response to unbaited, untreated and insecticide-treated nets is shown in Fig. 10 and the data in Additional file 7. In the absence of a human host, there is slightly greater activity in regions to the side and ends of the bed net than in regions just above the net. This is partly due to the elongated nature of the virtual insectary, but also because the two sides of the net present a larger surface area than the roof of the net. When a host is present in the net, however, the activity profile changes markedly. The attractant stimuli diffusing from the human host profile strongly attracts the mosquito population to the top regions surrounding the net. The presence of a baited net also affects flight activity in the peripheral regions.

Table 2 Model vs experimental flight tortuosity

| Tortuosity   | Unbaited net | Head facing left | Head facing right |
|--------------|--------------|------------------|-------------------|
| Model mean   | 1.120        | 1.635            | 1.640             |
| Model range  | 1.120–1.120  | 1.63–1.65        | 1.63–1.65         |
| Experimental mean [11] | 1.31          | 1.66 (both orientations combined) |
| Experimental range | 1.16–1.47     | 1.52–1.79        |                   |

Fig. 5 Effect of host presence on foraging and arena occupancy. Top: Snapshot of occupancy in a baited (left) and unbaited (right) arena after 20,000 model steps (approximately 11 min). Partially transparent voxels indicate regions of the virtual insectary which have been occupied. Bottom: Occupancy index (the fraction of the arena explored by the population) for an unbaited bed net (red) compared to baited conditions (green represents head facing left and blue is head facing right).
in total, resulting in approximately 8 times as much peri-bed net activity compared to an unbaited net. A baited net coated with insecticide ($tx = 0.1 \text{ 'dose' accumulation per contact, decrementing an initial health value of 100}$) exhibits the same strong attraction to the top surface of the net. Repeated contact, however, diminished the health of the mosquitoes until the population was entirely killed in a mean time of 36.34 m (s.d. 2.56 m). This corresponds closely with the findings from experimental tracking which found that activity after 30 m was negligible on LLIN treated nets [11].

**Discussion**

The human home is exploited by numerous parasitic arthropods, including vectors of malaria, dengue, yellow fever, lymphatic filariasis, leishmaniasis and Chagas disease. In Africa, most cases of malaria are transmitted indoors, despite the wide range of behavioural preferences shown by those Anopheles spp. that are vectors [33, 34]. In turn, humans have exploited this behaviour using a number of methods to target them at various locations inside the home. Of the methods used to date, insecticide-treated bed nets (ITNs/ LLINs) have been shown to be highly effective [1], and they remain the main intervention in malaria prevention in Africa. As a first stage in developing models that capture vector behaviour from house entry to exit, incorporating spatial movement and resting site preferences, this agent-based model of mosquito flight and host-seeking dynamics was developed, based on the actual conditions and their impacts on vector behaviour measured previously [11].

This is the first fine-grained model to simulate mosquito flight and distribution patterns in 3D space from quantification of actual tracked flight behaviours recorded in [11] and similar findings from earlier studies [9, 10]. Notably here, the patterns of net contact and flight distribution in the peri-bed net region were accurately reproduced by the model. Specifically, the differences in number of net contacts in no-host and host-present conditions, the preference for the top surface of the bed net

---

**Fig. 6** Distribution of bed net contact sites (shown from above, net shape indicated by dashed line). Spatial 2D heat map distribution of occupancy in the bed net XZ plane and 1D frequency distributions. There is relatively little contact at the top of the net in the unbaited condition where most contact occurs at the side net surfaces. In the baited conditions there are a large number of contacts on the top net surface at locations corresponding to the upper torso and head of the host. These contact sites correlate with bait orientation.
which is of critical importance relating to the design of next-generation bed nets and with respect to damaged nets \cite{35}, the orientation of net contacts depending on host orientation, and patterns of flight activity in regions surrounding the bed net.

It is the combined effect on this behaviour of the attractants emanating from the host within the net and the potentially repellent or irritant properties of the insecticide treatment on the net, that determine whether or not a mosquito makes sufficient net contact

![Fig. 7](mosquito_net_contact_distributions.jpg)

**Table 3** Model vs experimental Net Contact Time for all contacts

| Net Contact Time (m) | Unbaited Net | Baited Net | Treated Net |
|----------------------|--------------|------------|------------|
| Model mean           | 1.24         | 31.51      | 13.69      |
| Model range          | 0.99–1.49    | 29.45–32.65| 13.57–13.77|
| Experimental mean \cite{11} | 2.4          | 33.1       | 7.3        |
| Experimental range   | 2.1–6.8      | 24.3–41.2  | 3.9–10.7   |

(See figure on next page.)

**Fig. 8** Effect of noise contamination on attractant dispersal and mosquito bed net contact locations. Spatial heat map distribution of mosquito activity in the $XZ$ plane (i.e. looking from above, net shape indicated by dashed line) under noisy dispersal conditions with a head facing left host. Gaussian noise sigma multiplier from no noise contamination (a) up to sigma 10 (b–f). Increasing noise spreads mosquito distribution across the top surface of the net and at higher noise levels, increases net contact at the sides of the net.
to acquire a lethal dose. That this is accurately represented in the model is shown by Fig. 10. Without a host acting as attractant, activity around the bed net was relatively uniform (Fig. 10, top row), reflecting the shape of the arena and positioning of the bed net. When a human host was present, the behaviour of the mosquito population changed in response to different stimulus conditions, aggregating preferentially on the roof of the bed net (Fig. 10, middle row). The foraging behaviour of the virtual population measured by an occupancy metric based on the fraction of the arena explored, shifted from an 'exploration' behaviour (wide spatial casting throughout the arena) to an 'exploitation' behaviour (a narrower focused exploration as the population oriented to an

Fig. 9 Effect of noise contaminated attractant dispersal on total number of net contacts and contact height. **a** mean net contact height decreases as dispersal noise increases, moving from the top surface (orange) to the sides of the net. **b** The mean number of total net contacts also decreases as dispersal noise increases.
source of attractants). This difference in behaviour was entirely provoked by the diffusion of the attractant plume from a host. With an untreated net this foraging and attempted penetration of the net persists over the entire hour whereas LLIN treated nets, in experimental and simulation conditions, dramatically reduced the activity of the mosquito population (Fig. 10, bottom row).

Turbulent dispersal of the attractant field was approximated using a noise parameter which added Gaussian noise during the diffusion method. Increasing dispersal noise resulted in fewer net contacts, a wider distribution pattern and, at high noise values, a change in contact distribution from the top surface of the net to the sides of the net.

Limitations of the model
These results demonstrate that a fine-grained modelling approach has utility as an in silico method of performing virtual mosquito bio-assay experiments to explore 3D indoor flight behaviour. As with all modelling approaches, limiting assumptions have been made. Here, the most significant are associated with host attraction: visual stimuli are not represented and the multiple sensory modalities known to have a major role as attractant stimuli were greatly simplified. The latter, the spatially dispersed stimuli — CO₂, host odour (from the body and exhaled breath) and body temperature are known to be of major importance in host location and conveniently were suitable for grouping. Although other sensory modalities also contribute [31, 36], the interactions are complex and not yet fully elucidated, whereas the mosquito behaviours seen at insecticidal bed nets can be explained in terms of the these grouped stimuli alone. This simplification was also driven by the need for real-time simulation performance. Alternate methods of attractant dispersion and sampling, for example, computed by Computational Fluid Dynamics (CFD) methods, are a future possibility, although this would likely bias the model to particular mechanisms of stimulus dispersion.

The model mosquitoes do not incorporate any approximation of energy expenditure and therefore do not display a tail-off in flight activity over time, as previously reported in [11]. The tracking data used for the basis of

Fig. 10 Spatio-temporal flight tracks and peri-bed net regional occupancy. Simulation (Cols 1 & 3) and experimental (Col 2 & 4) data. Track colour indicates time (scale for flight tracks indicated below Cols 1 & 2). Region colour indicates mean occupancy time indicated below Cols 3 & 4. Conditions shown are for an unbaited net (top row), an untreated net (middle row) and LLIN treated net (bottom row). Model tracks show a single mosquito for clarity and circular regions denote landing sites. Experimental images reproduced courtesy of [11].
the model was based on the behaviour of a single mosquito species (An. gambiae s.s.). However, published data from tracking experiments shows that, in terms of spatial activity, the behaviour of Anopheles arabiensis [37] and Culex quinquefasciatus [12] at human-baited ITNs is remarkably similar to that of An. gambiae.

The relative simplicity of the mosquito behavioural stimulus-response transitions was chosen to attempt to reduce the wide range of potential parametric influences on model behaviour, a known issue with agent-based approaches. Nevertheless, despite these limiting assumptions, the model has demonstrated that it is able to reproduce the broad findings of [9, 10], most notably that the distribution of mosquito landing sites occurs predominantly on the top surface of the net and is affected by disturbances in dispersion by shifting this distribution to lower regions of the net. Furthermore, the model also accurately reproduces experimental tracking findings of [11], exhibiting similar flight tortuosity, bed net contact time, peri-bed net distribution and activity decay in response to simulated insecticide-treated bed nets.

Application of the model and scope for further work

By making adjustments to the model parameters, the model can accommodate far more variation than described here. Examples would include: additional vector mosquitoes with different arrival patterns at the bed net or different host species preferences (certain resistant strains of An. gambiae mosquitoes, for example, exhibit different activity levels during flight tracking than susceptible strains); the influence of a second host (human or other) in the room; effects of the new insecticide treatments on the bed net (level of pre-contact repellency or post-contact irritation), whether applied to all surfaces equally or to the roof alone; combining the insecticidal bed net with residual insecticide sprayed on the wall.

Ongoing experimental evaluation of the so-called next generation bed nets (bi-treated nets with pyrethroid and non-pyrethroid insecticides) will greatly expand the model’s power to predict how new net treatments perform, and determine whether they are best deployed alone or in combination.

Recently, an expanded version of the model was used to evaluate a range of novel bed net designs, and rapidly identified the best of seven candidate(s) [38]. The rapid process means the new design will be evaluated in initial trials in sub-Saharan Africa in early 2021. This research demonstrates that simple models of mosquito flight around bed nets can assist in enabling rapid assessment of novel vector control tools.
3. Churcher TS, Lissenden N, Griffin JT, Worrall E, Ranson H. The impact of pyrethroid resistance on the efficacy and effectiveness of bednets for malaria control in Africa. Elife. 2016;5:e16090.

4. Toé KH, N'Fale S, Dabiré RK, Ranson H, Jones CM. The recent escalation in strength of pyrethroid resistance in Anopholes coluzzii in West Africa is linked to increased expression of multiple gene families. BMC Genomics. 2015;16:1.

5. Ranson H, Lissenden N. Insecticide resistance in African Anopheles mosquitoes: a worsening situation that needs urgent action to maintain malaria control. Trends Parasitol. 2016;32:187–96.

6. Hemingway J, Ranson H, Magli A, Kolaczinski J, Fornadel C, Gimnig J, et al. Averting a malaria disaster: will insecticide resistance derail malaria control? Lancet. 2016;387:1785–8.

7. Baylii K, Ndo S, Namountougou M, Sanou R, Ouattara A, Dabiré RK, et al. Evaluation of efficacy of Interceptor-G2, a long-lasting insecticide net coated with a mixture of chlorfenapyr and alpha-cypermethrin, against pyrethroid resistant Anopheles gambiae s.l in Burkina Faso. Malar J. 2017;16:190.

8. Toé K, Müller P, Badolo A, Traoré A, Sagnon N, Dabiré R, et al. Do bednets including piperonyl butoxide offer additional protection against populations of Anopheles gambiae s.l that are highly resistant to pyrethroids? An experimental hut evaluation in Burkina Faso. Med Vet Entomol. 2018;32:407–16.

9. Lynd A, McCall PJ. Clustering of host-seeking activity of Anopheles gambiae mosquitoes at the top surface of a human-baited bed net. Malar J. 2013;12:1–8.

10. Sutcliffe JF, Yin S. Behavioural responses of females of two Anopheles mosquito species to human-occupied, insecticide-treated and untreated bed nets. Malar J. 2014;13:1–19.

11. Parker JE, Angarita-Jaimes N, Abe M, Townes CE, Townes D, McCall PJ. Infrared video-tracking of Anopheles gambiae at insecticide-treated bed nets reveals rapid decisive impact after brief localised net contact. Sci Rep. 2015;5:13392.

12. Angarita-Jaimes N, Parker J, Abe M, Mashauri F, Martine J, Townes C, et al. A novel video-tracking system to quantify the behaviour of nocturnal mosquitoes attacking human hosts in the field. J Roy Soc Interface. 2016;13:20150074.

13. Spitzer J, Spoor CW, Greco F, ter Braak C, Beeuwkes J, van Brugge SP, et al. A 3D analysis of flight behavior of Anopheles gambiae sensu stricto mosquitoes in response to human odor and heat: PLoS ONE. 2013;8:e62955.

14. Körner C, Townes CE, Angarita-Jaimes N, Parker JE, McCall P, Townes DP. 3D tracking of mosquitoes: A field compatible technique to understand malaria vector behaviour. In: 3D Image Acquisition and Display: Technology, Perception and Applications. J Opt Soc Am. 2016:TWSA–4.

15. Sutcliffe JF. Sensory bases of attractiveness: morphology of mosquito olfactory sensilla-a review. J Am Mosq Control Assoc. 1994;10:309–15.

16. Amer A, Melh犰m H. The sensilla of Aedes and Anopholes mosquitoes and their importance in repellency. Parasitol Res. 2006;99:491–9.

17. Dekker T, Cardé RT. Moment-to-moment flight manoeuvres of the female yellow fever mosquito (Aedes aegypti L) in response to plumes of carbon dioxide and human skin odour. J Exp Biol. 2011;214:3480–94.

18. Allan SA. Physics of mosquito vision—an overview. J Am Mosq Control Assoc. 1994;10:266–71.

19. Hawkes F, Gibson G. Seeing is believing: the nocturnal malarial mosquito Anopholes coluzzii responds to visual host-cues when odour indicates a host is nearby. Parasit Vectors. 2016;9:320.

20. Andrés M, Seifert M, Spalholz C, Warren B, Weiss L, Giraldo D, et al. Auditory efferent system modulates mosquito hearing. Curr Biol. 2016;26:2028–36.

21. Lapshin D. Mosquito bioacoustics: auditory processing in Culex pipiens L Males (Diptera, Culicidae) during flight simulation. Entomol Rev. 2012;92:605–21.

22. Ingham VA, Anthousi A, Douris V, Harding NJ, Lycett G, Morris M, et al. A sensory appendage protein protects malaria vectors from pyrethroids. Nature. 2019;779:376–80.

23. Montell C, Zwiebel L. Mosquito sensory systems. In: Adv In Insect Phys. vol. 2016;5:1:1–328.

24. Echhoff PA. A malaria transmission-directed model of mosquito life cycle and ecology. Malar J. 2011;10:303.

25. Gu W, Killeen GF, Mbogo CM, Regens JL, Githure JJ, Beier JC. An individual-based model of Plasmodium falciparum malaria transmission on the coast of Kenya. Trans R Soc Trop Med Hyg. 2003;97:43–50.

26. Ottenborou PE, Nikbakhtzadeh MR, Foster WA. Orientation of Anopheles gambiae (Diptera: Culicidae) to plant-host volatiles in a novel diffusion-cage olfactometer. J Med Entomol. 2015;53:237–40.

27. Smith NR, Trauer JM, Gambhir M, Richards JS, Maude RJ, Keith JM, et al. Agent-based models of malaria transmission: a systematic review. Malar J. 2018;17:299.

28. Cummins B, Cortez R, Foppa IM, Wallbeck J, Hyman JM. A spatial model of mosquito host-seeking behavior. PLoS Comput Biol. 2012;8:e1002500.

29. Scherbackheva A, Haano H, Killeen GF. Modeling host-seeking behavior of African malaria vector mosquitoes in the presence of long-lasting insecticidal nets. Math Biosci. 2018;295:36–47.

30. McMeniman CJ, Corfas RA, Matthews BJ, Ritchie SA, Vosshall LB. Multimodal integration of carbon dioxide and other sensory cues drives mosquito attraction to humans. Cell. 2014;156:1060–71.

31. van Breugel F, Riffell J, Fairhall A, Dickinson MH. Mosquitoes use vision to associate odor plumes with thermal targets. Curr Biol. 2015;25:2123–9.

32. De Moraes CM, Stanczyk NA, Betz HS, Pulido H, Sim DG, Read AF, et al. Malaria-induced changes in host odors enhance mosquito attraction. Proc Natl Acad Sci. 2014;111:11079–84.

33. Bayoh MN, Walker ED, Kosgei J, Ombok M, Olang GB, Githeko AK, et al. Persistently high estimates of late night, indoor exposure to malaria vectors despite high coverage of insecticide treated nets. Parasit Vectors. 2014;7:390.

34. Huho B, Briët O, Seyoum A, Sikaala C, Bayoh N, Gimnig J, et al. Consistently high estimates for the proportion of human exposure to malaria vector populations occurring indoors in rural Africa. Int J Epidemiol. 2013;42:235.

35. Sutcliffe J, Yin S. How many holes is too many? A prototype tool for estimating mosquito entry risk into damaged bed nets. Malar J. 2017;16:304.

36. Cardé RT. Multi-cue integration: how female mosquitoes locate a human host. Curr Biol. 2015;25:R793–R795795.

37. Parker JE, Jaimes NCA, Gleave K, Mashauri F, Abe M, Martine J, et al. Host-seeking activity of a Tanzanian population of Anopheles arabiensis at an insecticide treated bed net. Malar J. 2017;16:270.

38. Murray GPD, Lissenden N, Jones J, Voloshin V, Toe KH, Sherrard-Smith E, et al. Novel bednet design targets malaria vectors and expands the range of usable insecticides. Nat Microbiol. 2020;5:40–7.