Bulk-up synchronisation of successive larval cohorts of Anopheles gambiae sensu stricto and Anopheles coluzzii through temperature reduction at early larval stages: effect on emergence rates, body size and mating success

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Research

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Abstract

Background: Malaria persists as a huge medical and economic burden. Although the number of cases and death rates have reduced in recent years, novel interventions are a necessity if such gains are to be maintained. Alternative methods to target mosquito vector populations that involve the release of large numbers genetically modified mosquitoes are in development. However, their successful introduction will require innovative strategies to bulk-up mosquito numbers and improve mass rearing protocols for Anopheles mosquitoes.

Methods: Here we aimed to exploit the relationship between mosquito aquatic stage development and temperature so that multiple cohorts of mosquitoes, from separate egg batches, could be synchronised for a 'bulk-up' of the number of mosquitoes released. First instar larvae were separated into two cohorts: the first, maintained under standard insectary conditions at 27 °C, the second subjected to an initial 5-day cooling period at 19 °C.

Results: Cooling of 1st instars slowed the mean emergence times of An. coluzzii and An. gambiae by 2.4 and 3.5 days, respectively, compared to their 27 °C counterparts. Pupation and emergence rates were good (> 85%) in all conditions. Temperature adjustment had no effect on mosquito sex ratio and adult fitness parameters such as body size and mating success were also comparable with the control cohort.

Conclusions: Bulk-up larval synchronization is a simple method allowing more operational flexibility in mosquito production towards mark-release-recapture studies and mass release interventions.

Background

Malaria is a persistent public health issue. Despite over 50 years of sustained effort to control the disease through the use of antimalarial drugs and vector control, transmission has been interrupted in only a limited number of countries. The World Health Organization reported 228 million cases and 405,000 deaths in 2018 (1). Most of these deaths occurred in children under the age of 5 living in sub-Saharan Africa. In recent years, the introduction of insecticide treated bednets (ITNs), mainly long lasting insecticidal nets (LLINs) and indoor residual spraying (IRS) combined with Artemisinin-based combination therapies has resulted in a decline in malaria incidence, thus providing renewed hope for elimination goals (2, 3). However, such gains are beginning to diminish, once again threatened by the development and spread of resistance to all antimalarials and insecticides introduced (4, 5). Furthermore, behavioural changes in mosquito vectors, whereby they bite at dawn or early evening rather than through the night, when people are under bednet protection, diminishes the effectiveness of current intra-domiciliary control measures (6-9). Therefore, if reductions in malaria burden are to be at least sustained, alternative complementary approaches are necessary (10).

Following recent advances in genetic engineering, genetic vector control strategies for malaria mosquitoes are now at the forefront of research and development goals (11-13). These include a range of different approaches that are either self-limiting or self-sustaining. Self-limiting strategies involve the use of
genetically modified sterile males or mosquitoes modified with a gene drive mechanism that is spatially or temporally self-limited (14, 15). These methods bear similarities with the traditional sterile insect technique (SIT). Their impact depends on effective mating between released mosquitoes and the target population, and require repeated, inundative mass release of mosquitoes (16, 17). Self-sustaining strategies employ a gene drive mechanism which means that a desirable trait such as male biased sex ratio (18, 19), reduced female fertility (20) or an antiparasitic effector gene (21) is inherited at a higher rate than mendelian inheritance. The spread of such self-propagating genes can lead to population suppression, reducing the number of biting females or population replacement with mosquitoes that are refractory to the malaria parasite (22, 23). The self-sustaining strategies are a longer-term goal that would ideally require relatively smaller initial releases of mosquitoes thereby making them more cost efficient (13, 15, 16). However, the deployment of such genetic tools on a broader scale will still necessitate the production and release of much larger numbers of mosquitoes (13, 15, 16). In addition to mosquito release interventions per se, ecological studies that focus on mosquito survival, dispersal or estimation of population sizes such as Mark-Release-Recapture studies also rely on the punctual release of mosquitoes reared at a much smaller scale (24).

One major challenge in rearing Anopheline mosquito vectors for release studies and interventions is that their eggs hatch shortly after being laid and can only survive a limited number of days without water, hence, egg-to-adult rearing needs to be continuous (25). This imposes constraints on rearing protocols and infrastructures and means that the release cohort largely depends on the number of adults in the preceding generation. There have been efforts towards the optimisation of Anopheline egg storage. Through elaborate drying and cooling methods, it is now possible to increase egg storage times by up to 4-6 days, however beyond that point, hatch rate and larval development are negatively impacted (26-28). Therefore, other avenues to bulk-up Anopheles mosquitoes for mass release, without affecting their phenotypic quality, should be explored.

The development rate of insects is mainly temperature dependent and offers the potential opportunity to slow or accelerate development (29). In Anopheles gambiae, the relationship between mosquito aquatic stage development and temperature has been well studied (30-34). Within a minimum and maximum threshold, development rate increases linearly with an increase in temperature. Indeed, Barreaux et al. (2018) reported a 1.4-day difference in time to pupation between larvae maintained at 21°C and 29°C (31). similarly, Christiansen-Jucht and colleagues reported a linear increase in development rate from 23°C to 31°C, but at 35°C all larvae died before emergence (34). Bayoh and Lindsay (2003) showed that development rate increased linearly with temperatures from 22°C to 28°C resulting in a ~10-day shift in egg to adult development time. No adults emerged at temperatures below 18°C or above 34°C (32).

In this study, we aimed to exploit this relationship to mimic synchronisation of successive egg batches obtained from repeated blood-feeding of a single female cohort, without impacting negatively on mosquito survival. The rearing temperature of 1st instar larvae of An. gambiae s.s. and An. coluzzii laboratory strains was reduced with the aim of slowing down development by approximately 3 days, the
time required for one gonotrophic cycle by females at 27°C (35-37). The impact of the temperature alteration on the pupation and emergence rates, developmental times, adult phenotypic quality and mating success was evaluated. The ability to slow down a larval cohort by 3 days, hence to synchronize the emergence of adult progeny resulting from multiple blood feeds and successive egg batches from the same pool of females, has important implications for the optimisation of mass production and release methods for *Anopheles* s.l.

**Methods**

**Mosquito maintenance**

*Anopheles gambiae* s.s, Kisumu strain; an old strain colonized originally from Kisumu, Kenya, East Africa, and *Anopheles coluzzii*, VK3 strain; a strain colonized in 2018 from Vallee du Khou, Burkina Faso, West Africa, were maintained in the Manson Insectaries at the Centre for Applied Entomology and Parasitology, Keele University, UK. The strains were kept under our standard Manson insectary conditions: 27 ± 2°C, 12/12-hour light/dark cycle at 70 ± 5% relative humidity unless otherwise stated (38). Adults had a constant supply of 10% glucose and were blood fed on defibrinated (fibrin removed to prevent clotting) horse blood (TCS Biosciences) using the Hemotek membrane feeding system (Blackburn, United Kingdom). Polystyrene cups, lined with Whatman filter paper, containing 50:50 deionized: mineral water was provided for oviposition. After hatching, 200 first instar larvae / 500ml mixed water (250ml deionized water + 250ml mineral water) were placed in trays (34cm X 24cm) and supplemented with 2 drops of Liquifry. Feeding with solid food commenced after 24h, and all trays were provided with an additional 500ml water on day 5. Larvae were fed with an optimised feeding regime using ground TetraMin fish food (Tetramin, Tetra, Melle, Germany) and transferred to adult cages (5l plastic, 20.5cm height x 20cm diameter), upon pupation as described elsewhere (38, 39).

**Manipulation of larval temperature**

For each strain, 1<sup>st</sup> instar larvae from one egg batch were split into two groups: control and temperature manipulated (Figure 1). The larvae in the control group were trayed in accordance with standard insectary protocol as described above (200 larvae/tray), 8 trays in total. The larvae in the temperature manipulated group were trayed at 2000 larvae/tray and placed in a climate chamber at 19°C 12/12-hour light/dark cycle at 70 ± 5% relative humidity (Panasonic MLR Climatic Test Chambers 352H-PE Kadoma, Osaka, Japan). The temperature manipulated larvae remained at 19°C for 5 days and fed, first with Liquifry (as described previously), and then *ad libitum* with ground TetraMin fish food. On day 5, larvae kept at 19°C were transferred to standard insectary conditions (27 ± 2°C) and re-trayed at 200 larvae/tray (500ml of mixed water was added, with an additional 500ml of tap water) 8 trays total. Larvae in the 27°C control group were reared according to the standard insectary protocol for the duration of the experiment.
Adult development and mating

Adult emergence and pupae failing to emerge were recorded daily, as well as dead adults. For each experimental group, pupae were collected each day, sexed and placed into separate cages for males and females. Male and female mosquitoes aged 3-5 days old were combined (40 males + 40 females) into mating cages (6 cages total) and allowed to mate overnight. The following morning, mosquitoes were transferred to -20°C and stored in 75% ethanol. Spermathecae from female mosquitoes were dissected and burst open in a drop of water. The presence of a coagulated sperm bundle provided confirmation of a successful mating event. Wing length was recorded for all females and a subsample of 15-30 males/condition as a proxy for adult size. In brief, a binocular microscope, calibrated using a stage micrometer (1mm= 10 eye piece units at x1 magnification) was used to measure one wing from each adult. Wings were measured from the distil end of the allula to the apical margin (radius veins) as described previously (40).

Statistical analysis

Binomial variables such as pupation rates, emergence rates, sex ratio, and insemination rates were analysed via logistic regression. Emergence times were analysed via proportional hazard analysis. Likelihood odds ratios were used for post-hoc comparisons following logistic regression and proportional hazard analysis. Continuous data, such as wing length, (body size) was checked for normality and parametric and non-parametric tests were used where appropriate. In all multivariate analyses, interactions between independent variables were tested but removed from models if not significant. All analyses were carried out using the JMP 14 statistical software (SAS Institute, North Carolina)

Results

Effect of temperature manipulation on pupation and emergence rates

Logistic regression analysis indicated that the reduction in temperature, to 19°C, during early larval development had no overall impact on pupation rates of An. coluzzii or An. gambiae (Likelihood ratio Chi-square = 3.63, df =1 P = 0.057). There was a significant difference in pupation rates between the two species (LR = 144.23, df =1, P< 0.001). Higher pupation rates were observed for An. coluzzii at both 19°C (98%) and 27°C (97.5%), however, An. gambiae also achieved high pupation rates at both temperatures; 85% (19°C) and 89% (27°C) (Figure 2a).

High overall emergence rates (> 85%) were observed for both species, however the effect of temperature depended on species (Table 1). Indeed, higher emergence rates were observed for An. coluzzii at 19°C, whereas for An. gambiae emergence was higher at 27°C (Figure 2b).
Table 1 Logistic regression (Likelihood ratios) of the effect of temperature and species on emergence rates

| Source                | DF | LR ChiSquare | Prob>ChiSq |
|-----------------------|----|--------------|------------|
| Species               | 1  | 30.61        | < 0.001    |
| Temperature           | 1  | 5.79         | 0.016      |
| Temperature*Species   | 1  | 24.15        | < 0.001    |

DF Degrees of freedom, LR Likelihood Ratio

**Effect of temperature manipulation on sex ratio**

None of the *An. coluzzii* and *An. gambiae* treatment groups differed from a male:female ratio of 1:1. For *An. coluzzii* at 19°C the proportion of males was 0.52 (95% CI 0.48-0.56) and at 27°C the proportion of males was 0.54 (CI 0.50-0.58). For *An. gambiae* the male:female ratio for both temperatures was 0.51:0.49 (CIs +/- 0.04). Logistic regression analysis indicated that sex-ratios did not significantly differ between species (LR Chi-square= 2.14, df =1, P =0.144) nor by temperature condition (LR Chi-square =0.002, df =1, P =0.965).

**Effect of temperature manipulation on emergence times**

Proportional hazards analysis revealed that the emergence time of both *An. coluzzii* and *An. gambiae* was significantly affected by the 5-day cooling period (Table 2). *An. coluzzii* took on average 2.4 and *An. gambiae* 3.5 days longer to emerge compared with those maintained at 27°C (Figure 3). There were also significant differences in emergence times between species and sex. The interactions between species, sex and temperature were also found to be significant (Table 2).

Table 2 Logistic regression (Likelihood Ratios) for the effect of species, temperature and sex on emergence time

| Source                | DF | LR ChiSquare | Prob>ChiSq |
|-----------------------|----|--------------|------------|
| Species               | 1  | 259.71       | < 0.001    |
| Temperature           | 1  | 2025.21      | < 0.001    |
| Sex                   | 1  | 19.51        | < 0.001    |
| Temperature*Species   | 1  | 46.01        | < 0.001    |
| Temperature*Sex       | 1  | 5.60         | 0.018      |

**Effect of temperature manipulation on adult fitness parameters**
Multivariate analysis showed that mosquito wing length was significantly affected by species, temperature and sex (Table 3). *An. coluzzii* individuals were significantly smaller than *An. gambiae* and male mosquitoes significantly smaller than females. Those exposed to the 19°C 5-day cooling period were significantly smaller than their counterparts maintained at 27°C (Table 3, Figure 4a).

**Table 3 General linear model effect data for the effect of species, temperature and sex on mosquito wing lengths**

| Source   | DF  | Sum of Squares | F Ratio | Prob > F |
|----------|-----|----------------|---------|----------|
| Species  | 1   | 1.20           | 70.86   | < 0.001  |
| Temperature | 1 | 0.51           | 30.29   | < 0.001  |
| Sex      | 1   | 0.89           | 52.09   | < 0.001  |

Insemination rates were similar at both temperatures for both species (Figure 4b). When female size was also considered, differences were apparent between species, between temperatures and at different wing lengths (Table 4). Overall, inseminated females were larger in all conditions (Figure 4c).

**Table 4 Logistic regression (Likelihood Ratios) for the effect of species, temperature and wing length on mosquito inseminations rates**

| Source       | DF | L-R ChiSquare | Prob>ChiSq |
|--------------|----|---------------|------------|
| Species      | 1  | 6.63          | 0.010      |
| Temperature  | 1  | 4.16          | 0.041      |
| Wing length (mm) | 1 | 94.00        | < 0.001    |

**Discussion**

Here we show that through temperature manipulation it is possible to delay emergence of mosquitoes by up to 3 days; the approximate length of the gonotrophic cycle of *Anopheline* females. These finding are important for ecological studies that require small punctual releases and for interventions requiring mass releases focussing on *Anopheline* vector species. Currently the logistics and planning for *Anopheline* production revolve around the assumption that achieved mosquito numbers, at a particular time point, directly depend on the quantity of eggs produced by a single gonotrophic cycle. The findings of this study offer the potential to effectively double the progeny produced from one female cohort, thereby bringing much needed flexibility to *Anopheline* rearing practices.

The 3-day delay was achieved by subjecting first instar larvae to a 5-day cooling period at 19°C. The alteration in temperature had no effect on pupation rates although there was a difference in the rate of pupation between *An. coluzzii* and *An. gambiae*. We also found that cooling had a minimal effect on
emergence rates, that were > 85%, but affected the two species conversely. In An. coluzzii, it resulted in an increase in emergence rate but in An. gambiae it resulted in small decrease in emergence rates. Overall, pupation and emergence rates were high and in line with reports elsewhere for laboratory reared Anopheles (41, 42).

We found no effect of temperature reduction on sex ratio, which was equivalent to a 1:1 male to female ratio in both species. Any evidence of female bias would have important consequences for male-focused mass release programmes. Imbalances have been reported following temperature and diet alterations for Aedes mosquitoes (43, 44). However, for Anopheles mosquitoes no such differences have been found (45, 46).

Adult phenotypic quality and mating competitiveness are crucial to the success of release programmes (39, 47, 48). Several studies have reported negative carry-over effects on the phenotypic quality of adult mosquitoes following experimental manipulations of larval conditions such as temperature, density and food availability (31, 38, 49). Here we found that male and female adults reared at 19°C were smaller than those reared at 27°C, but the 0.05 mm (1.5%) reduction in size observed was unlikely to be biologically important. Indeed, the negligible size differences found did not translate to a negative impact on insemination rates. In the natural setting, An. gambiae s.l. mate in swarms that are typically composed of males and females visit to choose a mate and leave in copula (17, 50). Smaller males have reduced spermatogenesis and are less competitive in terms of mating than medium-to-large sized mosquitoes, making them poor candidates for release programmes (51, 52). Compared to the size distribution from those reports (2.48-3.12), males produced in this study at either temperature, were relatively large (2.98-3.08 mm) and consistent with the optimal size of 3mm for mating found in field studies (17).

Smaller females have reduced fecundity, have been shown to require multiple blood feeds before completion of a gonotrophic cycle and may be less attractive to males (34, 39, 53, 54). Although we found no difference in overall insemination rates in relation to larval cooling, inseminated An. coluzzii and An. gambiae females were 0.08 mm (2.7%) and 0.09 mm (2.9%) larger than non-inseminated ones, respectively. Although, this is again a very small size difference, the finding that larger females were more likely to mate is consistent with results from insectary and field swarm studies that suggest that males might prefer to mate with larger females (17).

The current study opted to slow down larval development rate by lowering the temperature rather than speed it up by increasing the temperature. Studies elsewhere have shown that at temperatures >34°C there are negative, irreversible carry-over effects on surviving adult mosquitoes and overall survival is lower (32, 34, 55). Indeed, although adults develop quicker, they are smaller (31, 34, 56) possibly because food consumption cannot sustain the rate of metabolism (57). Therefore, our work corroborated previous studies which found that cooling temperatures serve as a reversible inhibitor to mosquito development with negligible impacts on mosquito phenotypic quality, provided they are not maintained throughout their entire development (32, 58). We opted for a relatively short 5-day cooling period of 1st instars, which
allowed rearing at 10-fold higher density and *ad libitum* feeding. In preliminary studies we attempted to also maintain 1\textsuperscript{st} instar larvae at comparable densities at 27°C but found that larval competition negatively affected development rates and success. Hence, keeping 1\textsuperscript{st} instars at high densities was only possible for larvae kept at cooler temperature which reduced their metabolism and food consumption (57, 59). The optimised protocol presented here therefore exploits the relationship between development rate, temperature, density and food availability to adjust emergence time by ~3 days. As an incubator/fridge will be required for the cooled temperature condition, the 10-fold higher density culture at 19°C make the method both practical and scalable whilst minimizing pressure in terms of insectary space.

**Conclusions**

In conclusion, we present an optimised translatable methodology to increase *Anopheles* mosquito numbers for release studies and programmes. The optimised regime including a 5-day reduction in temperature (from 27°C to 19°C), adapted feeding and increased density represents a practical and scalable addition to mosquito production protocols. Here we achieved a 2.4 and 3.5-day delay for *An. coluzzii* and *An. gambiae* emergence times, respectively with no or negligible impacts on mosquito numbers, adult body size and mating rates. Using 18°C to slowdown larval development will ensure that a 3-day delay is achieved under all circumstances. As the 3-day delay spans the duration of one gonotrophic cycle the inclusion of a cooling period into mosquito mass rearing protocols offers the potential to synchronise successive larvae batches from a single pool of females. This is a modest but much needed step towards the optimisation of rearing techniques geared specifically for *Anopheles* mosquitoes, one of the most important groups of disease vectors.

**Declarations**

**Ethics approval and consent to participate**

Not applicable

**Consent for publication**

Not applicable

**Availability of data and materials**

All data generated or analysed during this study are included in this published article.

**Competing interests**

The authors declare that they have no competing interests

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**Authors’ contributions**

Experiments were designed by FT and HM, SS and SA. Experiments we conducted by QZ, FM and SA. Data was analysed by FT and QZ. HM and FT wrote the manuscript with input from FAA. All authors read and approved the final manuscript.

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**Figures**
Figure 4

The effect of temperature manipulation on adult fitness parameters. (a) The effect of temperature on adult size of male and female An. coluzzii and An. gambiae mosquitoes. (b) The effect of temperature on mating status of An. coluzzii and An. gambiae females. (c) The effect of temperature on mating status with regard to mosquito size. Error bars represent 95% confidence intervals.