Insecticide resistance selection and reversal in two strains of *Aedes aegypti* [version 1; peer review: 1 approved with reservations]

Jonathan Thornton1*, Bruno Gomes1,2*, Constância Ayres3, Lisa Reimer1

1Vector Biology Department, Liverpool School of Tropical Medicine, Liverpool, L35QA, UK
2Laboratório de Bioquímica e Fisiologia de Insetos, Oswaldo Cruz Institute (IOC-FIOCRUZ), Rio de Janeiro, 21040-360, Brazil
3Aggeu Magalhães Institute, Oswaldo Cruz Foundation (IAM-FIOCRUZ), Recife, Brazil

Abstract

Background: Laboratory reared mosquito colonies are essential tools to understand insecticide action. However, they differ considerably from wild populations and from each other depending on their origin and rearing conditions, which makes studying the effects of specific resistance mechanisms difficult. This paper describes our methods for establishing multiple resistant strains of *Aedes aegypti* from two colonies as a new resource for further research on metabolic and target site resistance.

Methods: Two resistant colonies of *Aedes aegypti*, from Cayman and Recife, were selected through 10 generations of exposure to insecticides including permethrin, malathion and temephos, to yield eight strains with different profiles of resistance due to either target site or metabolic resistance. Resistance ratios for each insecticide were calculated for the selected and unselected strains. The frequency of *kdr* alleles in the Cayman strains was determined using TaqMan assays. A comparative gene expression analysis among Recife strains was conducted using qPCR in larvae (CCae3A, CYP6N12, CYP6F3, CYP9M9) and adults (CCae3A, CYP6N12, CYP6BB2, CYP9J28a).

Results: In the selected strain of Cayman, mortality against permethrin reduced almost to 0% and *kdr* became fixated by 5 generations. A similar phenotype was seen in the unselected homozygous resistant colony, whilst mortality in the susceptible homozygous colony rose to 82.9%. The Recife strains showed different responses between exposure to adulticide and larvicide, with detoxification genes in the temephos selected strain staying similar to the baseline, but a reduction in detoxification genes displayed in the other strains.

Conclusions: These selected strains, with a range of insecticide resistance phenotypes and genotypes, will support further research.
on the effects of target-site and/or metabolic resistance mechanisms on various life-history traits, behaviours and vector competence of this important arbovirus vector.

**Keywords**
Aedes aegypti, insecticide resistance, mosquito, pyrethroid, permethrin, temephos, malathion, kdr,
**Introduction**

*Aedes aegypti* is one of the most significant mosquito species of public health concern due to its role as a vector of several arboviruses, including dengue, yellow fever, chikungunya and Zika. Over 4 million disability-adjusted life years worldwide were attributed to mosquito-borne viruses in 2013 (Moyes et al., 2017). Dengue virus, the most ubiquitous arbovirus, is found in 128 countries across temperate and tropical regions, and 3.9 billion people currently live at risk of infection (Guzman & Harris, 2015; Pollet et al., 2018). Yellow fever has re-emerged as an important disease in Africa and the Americas, with outbreaks occurring in regions that previously had low vaccination coverage and low-to-zero yellow fever incidence (Douam & Ploss, 2018; Monath & Vasconcelos, 2015). Large outbreaks of chikungunya virus have also been described since 2000 (Burt et al., 2017). Meanwhile, the World Health Organization declared a “public health emergency of international concern” during the Zika virus epidemic of 2015 and following the discovery of its association with microcephaly (Kindhauser et al., 2016).

Vector control is the primary strategy to prevent transmission of arboviruses due to the absence of prophylactic drugs or vaccines for most diseases (Silva et al., 2018). Chemical insecticides, biological agents, and habitat management (George et al., 2015; Horstick et al., 2010) are three common methods of controlling *Aedes spp.* However, insecticide resistant *Ae. aegypti* are commonly reported in Latin America and southern Asia, and have been reported in Africa (Moyes et al., 2017; Vontas et al., 2012), threatening the efficacy of vector-borne disease control programs (Corbel et al., 2017; Moyes et al., 2017; Ranson et al., 2010).

Experimental studies comparing the attributes of susceptible and resistant mosquito colonies are crucial to elucidate resistance mechanisms (Davies et al., 2008; Feyereisen et al., 2015; Ingham et al., 2018; Li et al., 2007; Melo-Santos et al., 2010; Moyes et al., 2017; Poupardin et al., 2014; Ranson et al., 2010; Rinkevich et al., 2013; Stevenson et al., 2012; Strode et al., 2012; Vontas et al., 2012; Weetman et al., 2018), insecticide mode of action (Ffrench-Constant et al., 2013; Yunta et al., 2016), and the fitness costs of resistance (Brito et al., 2013; Diniz et al., 2015). This information is necessary to develop new insecticide formulations and alternative control methods that avoid cross-resistance (de Lourdes Macoris et al., 2018; Jankowska et al., 2017). However, differences between susceptible and resistant colonies due to differing genetic backgrounds – caused by bottlenecks or genetic drift – may influence study outcomes (Ffrench-Constant et al., 2013). The ability to select sub-strains from a single parent colony which exhibit phenotypically distinct insecticide susceptibility profiles could help address the limitations of using disparate colonies. The aim of our study was to select multiple strains of *Ae. aegypti*, through exposure to a range of insecticides, that vary in resistance phenotype. Here we present the results after ten generations of insecticide exposure using two parent colonies: Cayman (CAY), a pyrethroid-resistant colony conferred by two target-site mutations in the sodium channel gene (V1016I and F1534C) (Harris et al., 2010b), and Recife (REC), a temephos-resistant colony conferred by overexpression of multiple detoxification genes (Diniz et al., 2015).

**Methods**

**Summary of the study design**

We exposed mosquitoes from one of two colonies (CAY and REC) to different insecticide selection pressures and monitored key indicators of resistance over time (Figure 1). We established the following eight strains of *Ae. aegypti*: CAY-P exposed to permethrin, CAY-RR unexposed and homozygous for resistance alleles (V1016I and F1534C), CAY-SS unexposed and heterozygous for resistance alleles, CAY-SS exposed and homozygous for susceptible alleles, REC-R exposed to temephos, REC-M exposed to malathion, REC-P exposed to permethrin, and REC-U unexposed.

First, discriminating concentration assays were conducted to identify a suitable concentration of permethrin, malathion and temephos for the selection procedure. Second, mosquitoes were exposed to an insecticide selection regime as outlined in Figure 1. Every three generations during the selection process, we monitored phenotypic resistance according to WHO diagnostic concentrations [30] and we assessed the presence of *kdr* alleles (CAY) or the upregulation of a select panel of detox genes (REC). Finally, resistance ratios for the eight strains were compared against a fully susceptible colony (New Orleans). Details of these procedures are described below.

**Initial Discriminating Concentration assays in larvae and adults**

Discriminating concentration assays using WHO tubes were conducted to determine an appropriate concentration of insecticide to expose each colony. The criteria is based on LC₅₀ to provide insecticide pressure, but within concentrations that mosquitoes are likely to encounter in the field. Papers impregnated with insecticide were prepared according to the standard WHO protocol (WHO, 2013). Filter papers of 12 cm x 15 cm were impregnated with a 1:1:1 volume mix of insecticide, acetone and corning oil and left to dry in a fume hood for 24 hours. Papers were stored wrapped in aluminium foil and placed in plastic bags at -20°C and used in up to five assays or within six months. The adult mosquitoes were selected with either permethrin or malathion and larvae were selected with temephos (see “Mosquito selection regime”). Groups of 25 L3 larvae were exposed to different concentrations of temephos in 200 mL of water to confirm the current LC₅₀. The LC50 identified for each colony-insecticide combination was used in subsequent selection procedures described below. However, we could not achieve an LC50 in the Cayman/permitrin combination that was relevant to concentrations that mosquitoes would typically encounter. We therefore decided to expose Cayman to a high concentration (3%) that is similar to doses received in the wild.

**Mosquito selection regime**

*Cayman colony*. We exposed 2–5 day old adult female mosquitoes in one strain, CAY-P, to 3% permethrin every generation
according to the standard WHO protocol (WHO, 2013). After a one-hour exposure and 24 hours recovery time, mortality was recorded and the surviving adults were allowed to mate and bloodfeed to create the next generation. We maintained another strain without insecticide exposure for ten generations (CAY-U). After five generations without insecticide exposure (generation 18), we used molecular tools to manually split CAY-U (see “Manual selection of kdr alleles”) across generations 6 and 7 due to the high frequency of kdr alleles (I1016 and C1534). Four unselected strains were established: i) homozygous susceptible individuals (CAY-SS: V1016 and F1534), ii) homozygous resistant individuals (CAY-RR: I1016 and C1534), and iii) two heterozygote strains created by crossing resistant females with susceptible males (CAY-RS) or susceptible females with resistant males (CAY-SR) (see Figure 1).

Recife colony. We selected larvae with temephos every three generations to create REC-R (Melo-Santos et al., 2010). Groups of L3 larvae were exposed to 0.5 mg/L temephos in plastic trays for 24 hours. At 24 hours, the mortality was recorded and

**Figure 1. Experiment outline.** Strains used were Cayman, which has target-site resistance to pyrethroids/DDT, and Recife, which has metabolic resistance to temephos. LC50s were determined by discriminating dose assays, then the LC50 was used to select the strain. Cayman was split into a strain selected with permethrin and an unselected strain, which was manually selected into R and S homozygotes and then crossed with reciprocal homozygotes to produce heterozygote strains. Recife was selected with malathion and permethrin, one strain was maintained with temephos exposure every three generations and one strain was left unselected. Each strain was subjected to bioassays using the WHO tube assay at the WHO recommended concentrations every three generations, and nucleic acid extraction and analysis of kdr alleles or detoxification genes was performed. At the end of 10 generations, each strain underwent bioassays to determine the LC50 and LC95 compared to the susceptible strain.
surviving larvae were transferred to fresh water, provided with food, and allowed to pupate and emerge as usual. Adult females were allowed to mate and bloodfeed to create the next generation. Two additional strains were established by exposing 2–5 day old adult female mosquitoes to malathion (REC-M: 1% WHO papers for 6 generations and 1.5% WHO papers for 3 generations) or permethrin (REC-P: 0.4% WHO papers for 6 generations and 0.75% WHO papers for 3 generations) every generation according to the standard WHO protocol (WHO, 2013) (see Figure 1). The exposure concentrations were determined by the initial discriminating dose. Survivors were allowed to mate and bloodfeed to create the next generation. The concentration was increased at generation 43 (7th generation of insecticide exposure) following a decrease in mortality to 25% after exposure in generation 42 (see Extended data: Table S1 (Reimer, 2020)).

**Evaluating phenotypic resistance**

We performed WHO tube assays (WHO, 2013) pre-selection and every three generations (G15, G18 and G21 for Cayman; G39, G42 and G45 for Recife) during the selection regime for permethrin or malathion. Assays were performed with standard WHO papers at diagnostic concentration (0.75% for permethrin, 5% for malathion), ordered from the Vector Control Research Unit, School of Biological Sciences, Universiti Sains Malaysia. Four exposure tubes and one negative control tube were filled with up to 25 mosquitoes each. Mosquitoes were exposed for one hour, returned to the holding tubes and provided with 10% sucrose solution. Mortality was recorded after 24 hours.

**Detection of kdr alleles**

For DNA analysis, 50 female mosquitoes were analysed every third generation of selection. Mosquitoes were killed at -20°C and stored on silica gel. We extracted DNA from individual mosquitoes using the Livak method (Livak, 1984). TaqMan® SNP Genotyping Assays for Vgsc-1016 and Vgsc-1534 alleles in *Aedes* (Extended data: Table S2 (Reimer, 2020)) were used to screen for *kdr* in the Cayman strains. The lack of resistant *kdr* alleles at Vgsc-1016 and Vgsc-1534 was confirmed in a small subsample of Recife pre-selection (G37). TaqMan reactions were performed in 10 μl volumes containing 1X TaqMan® Gene Expression Master Mix (Thermo Fisher Scientific, MA, USA), 800 nM of each primer, and 200 nM of each probe on an Mx3005P qPCR thermal cycler (Agilent Technologies, CA, USA) with initial denaturation of 10 min at 95°C followed by 40 cycles of 15 s at 92°C and 1 min at 60°C.

**Detoxification gene expression**

For RNA analysis, total RNA was extracted from pools of five female mosquitoes using Quick-RNA™ Miniprep (Zymo Research, CA, USA) and the purity and quantity of RNA were individually determined using a Nanodrop spectrophotometer (Nanodrop Technologies, DE, USA). SuperScript® III First-Strand Synthesis System performed cDNA synthesis from total RNA using oligo-dT20 primer (Thermo Fisher Scientific). Four genes associated with insecticide resistance were selected to screen expression profiles in larvae (CCae3A, CYP6N12, CYP6F3 and CYP9M9) and adults (CCae3A, CYP6N12, CYP6BB2 and CYP9J28a). cDNA was diluted ten-fold and qPCR reactions were performed in 20 μl volumes containing 2 μl of cDNA, 1x PowerUp™ SYBR® Green Master Mix (Thermo Fisher Scientific) and 300 nM of each primer on an Mx3005P qPCR thermal cycler (Agilent Technologies) with initial denaturation of 10 min at 95°C followed by 40 cycles of 15 s at 92°C and 30 s at specific TA (Extended data: Table S3 (Reimer, 2020)). The specificity of the primers was verified by melting curve analyses.

Relative fold gene expression was calculated using the comparative CT method (2^ΔΔCt method), taking into account PCR efficiency (Pfaffl, 2001). The genes coding for the 60S ribosomal protein L8 (RPL8) and the 40S ribosomal protein S7 (RPS7) were defined as reference genes. Between three and five biological replicates were performed for each strain and REC as baseline, respectively. All samples were run in duplicate. Results were expressed as mean transcription ratio in each strain and life stage ± SD relative to the mean transcription ratio of the specific life-stage of REC. Mann–Whitney U tests from the R package “stats” (R version 3.5.2: Copyright (C) 2020 The R Foundation for Statistical Computing) were used to compare transcription ratios between the selected strains and REC. The sequential Holm-Bonferroni procedure (Holm, 1979) was used to adjust α to account for multiple comparisons.

**Manual selection of kdr alleles**

CAY-U was maintained without insecticide pressure for five generations (G13-G18) before starting the selection process to establish homozygous strains for *Vgsc*-1016 and *Vgsc*-1534, which over the course of the process showed complete linkage in all individuals. These strains were established by separating pupae by sex and removing a leg from each adult for genotyping, as described above, before returning the adults to a cage for mating. The process was divided into two steps: 1) in generation G18, all the isoleucine homozygote mosquitoes for *Vgsc*-1016 were removed, and only heterozygotes and valine homozygotes were allowed to mate; 2) in generation G20, two homozygous strains were established for *Vgsc*-1534 (CAY-SS: Phe/Phe; CAY-RR: Cys/Cys), and all the heterozygotes were removed. Both CAY-SS and CAY-RR were screened by TaqMan assays for *Vgsc*-1016 and *Vgsc*-1534 in generation G21 to confirm the genotype of each strain. An extra selection was repeated in G21 for CAY-SS to remove the few individuals with *kdr* alleles.

**Determination of resistance ratio based on LC10 and LC50**

Standardised larval trays were prepared with 200 L1 larvae and provided a yeast tablet every other day until pupation. Adult two to five day old female mosquitoes were exposed to insecticide papers of a range of concentrations of malathion and permethrin, as described previously. Mortality was calculated 24 hours after exposure, and at least three replicates of each assay were performed. For larval bioassays, stock concentrations of temephos were prepared at 0.05 – 1 mg/ml. 1 ml of stock was added to 750 ml of water mixed in a 1:1 ratio from distilled water and larval water from the trays. The water was mixed and
aliquoted into five pupae pots of 150 ml each and groups of 25 L3 larvae in 50 ml water were added to each. This process was repeated for each concentration in the assay. Larval mortality was recorded at 24 hours. LC$_{50}$ and LC$_{95}$ estimates were generated from the data using the statistical software R-2.15.2. Resistance ratios were calculated based on comparison to the reference colony New Orleans, which is fully susceptible to all three insecticides.

Mosquito colony maintenance
All mosquitoes were maintained in the insectary of the Liverpool School of Tropical Medicine under controlled temperature (26 ± 2°C), relative humidity (75 ± 20%), and photoperiod (12:12 L:D). Adult mosquitoes were housed in BugDorm cages (MegaView Science Co, Ltd, Taiwan) and provided with constant access to 10% sucrose solution on a cotton pad, which was changed weekly. Eggs were obtained by feeding mated adult females on blood using a Hemotek feeder (Hemotek Ltd, Blackburn, UK). Due to issues with our supplier, the blood source was changed from human to horse at the beginning of the selection procedures. However, issues with mosquito egg-laying performance forced us to switch back to human blood mixed from separate bags of red blood cells and plasma in a 50:50 ratio from supplier overstock. Larvae were reared in plastic trays and fed Brewer’s Yeast tablets (Nature’s Aid ®).

Results
Cayman
In the unexposed strain CAY-U, we observed a slight increase in mortality over time for the standard WHO bioassay (0.75% permethrin) (Figure 2). In the permethrin selected CAY-P strain, mortality decreased to nearly 0% compared to the baseline of 4.6% mortality after only three generations of insecticide exposure at 3% permethrin. The kdr resistant alleles were still present in CAY-U at a high frequency (Table 1).

Kdr allele frequency. Both kdr alleles (V1016I and F1534C) showed complete linkage in all 230 individuals. The allele frequency for both kdr alleles in the Cayman colony was 93% at baseline with a high frequency of resistant homozygotes (86%). Selection with 3% permethrin (CAY-P) lead to kdr fixation within five generations. In CAY-U, kdr allele frequency was 68% split between a similar proportion of heterozygotes (46.4%) and resistant homozygotes (44.6%) (Table 1).

Only 1.27% of CAY-RR died in standard WHO tube bioassays with permethrin whereas CAY-SS had a mortality rate of 82.9%.

Resistance ratio to permethrin. CAY-P displayed a resistance ratio over 32x that of CAY-SS when compared to the reference colony New Orleans (Table 2). CAY-P and the unexposed CAY-RR had similar resistance ratios of over 200x resistance
compared to New Orleans. Resistance ratios in the heterozygote strains CAY-RS and CAY-SR were similar to each other, 2x higher than CAY-SS, and 14x lower than CAY-RR.

Recife
In the Recife colony, for both the malathion and permethrin-selected strains, a sharp drop in mortality to the WHO tube assay was observed, followed by a recovery then a shallower drop, rather than a gradual increase in resistance over selected generations (Figure 3). However, the mortality of REC-P for standard WHO bioassay (0.75% permethrin) on later generations (generations 42 and 45) is around 75%, while REC-M present mortality values for standard WHO bioassay (5% malathion) above 90% on later generations (Figure 3).

Differential expression of detoxification genes. The detoxification genes with differential expression among REC strains varied between larval and adults (Table 3 and Table 4). In REC-R, we observed no changes in gene expression from baseline in larval stages (Table 3), but expression of CYP9J28 and CYP6BB2 in adults increased significantly. Some genes were significantly downregulated at this stage in other Recife strains: i) REC-M (CCae3A, CYP6F3, and CYP9M9; MW: $p <0.000666$); ii) REC-U (CCae3A; MW: $p <0.002664$); iii) REC-P (CYP6F3; MW:

### Table 1. Frequency of Kdr alleles V1016I and F1534C in the Cayman strains across generations.

<table>
<thead>
<tr>
<th>Strain</th>
<th>G</th>
<th>Details</th>
<th>V1016I N</th>
<th>Val/Val</th>
<th>Val/Ile</th>
<th>Ile/Ile</th>
<th>V1534C N</th>
<th>Phe/Phe</th>
<th>Phe/Cys</th>
<th>Cys/Cys</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAY</td>
<td>G13</td>
<td>Baseline</td>
<td>43</td>
<td>-</td>
<td>0.140</td>
<td>0.860</td>
<td>42</td>
<td>-</td>
<td>0.143</td>
<td>0.857</td>
</tr>
<tr>
<td>CAY</td>
<td>G15</td>
<td>unexposed</td>
<td>48</td>
<td>-</td>
<td>0.208</td>
<td>0.792</td>
<td>48</td>
<td>-</td>
<td>0.208</td>
<td>0.792</td>
</tr>
<tr>
<td>CAY</td>
<td>G15</td>
<td>PERM 3%</td>
<td>47</td>
<td>-</td>
<td>0.085</td>
<td>0.915</td>
<td>48</td>
<td>-</td>
<td>0.083</td>
<td>0.917</td>
</tr>
<tr>
<td>CAY</td>
<td>G18</td>
<td>unexposed</td>
<td>56</td>
<td>0.089</td>
<td>0.464</td>
<td>0.446</td>
<td>52</td>
<td>0.077</td>
<td>0.462</td>
<td>0.462</td>
</tr>
<tr>
<td>CAY-P</td>
<td>G18</td>
<td>PERM 3%</td>
<td>49</td>
<td>-</td>
<td>-</td>
<td>1.000</td>
<td>40</td>
<td>-</td>
<td>-</td>
<td>1.000</td>
</tr>
<tr>
<td>CAY</td>
<td>G18</td>
<td>1st separation</td>
<td>351</td>
<td>0.046</td>
<td>0.313</td>
<td>0.641</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>CAY-U</td>
<td>G20</td>
<td>2nd separation</td>
<td>N/A</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>416</td>
<td>0.269</td>
<td>0.510</td>
<td>0.221</td>
</tr>
<tr>
<td>CAY-RR</td>
<td>G22</td>
<td>screen</td>
<td>23</td>
<td>-</td>
<td>-</td>
<td>1.000</td>
<td>23</td>
<td>-</td>
<td>-</td>
<td>1.000</td>
</tr>
<tr>
<td>CAY-RR</td>
<td>G22</td>
<td>screen</td>
<td>23</td>
<td>0.978</td>
<td>-</td>
<td>0.022</td>
<td>23</td>
<td>0.978</td>
<td>-</td>
<td>0.022</td>
</tr>
<tr>
<td>CAY-RR</td>
<td>G22</td>
<td>cleaning</td>
<td>189</td>
<td>1.000</td>
<td>-</td>
<td>138</td>
<td>0.957</td>
<td>-</td>
<td>0.043</td>
<td></td>
</tr>
</tbody>
</table>

CAY-U: strain without insecticide exposure; CAY-P: strain selected with 3% permethrin; CAY-SS: homozygous for susceptible alleles without insecticide exposure; CAY-RR: homozygous for resistant alleles without insecticide exposure; CAY-RS: heterozygous by the cross between CAY-RR females and CAY-SS males; CAY-SR: heterozygous by the cross between CAY-SS females and CAY-RR males.

### Table 2. Lethal concentrations and resistance ratios of Cayman strains for permethrin.

<table>
<thead>
<tr>
<th>Insecticide</th>
<th>Strain</th>
<th>LC$_{50}$</th>
<th>RR$_{50}$</th>
<th>LC$_{95}$</th>
<th>RR$_{95}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Permethrin</td>
<td>New Orleans</td>
<td>0.066 (0.053-0.083)</td>
<td>N/A</td>
<td>0.327 (0.211-0.507)</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>CAY-SS</td>
<td>0.432 (0.354-0.528)</td>
<td>6.54</td>
<td>1.232 (0.897-1.169)</td>
<td>3.77</td>
</tr>
<tr>
<td></td>
<td>CAY-RS</td>
<td>1.904 (0.904-1.116)</td>
<td>15.19</td>
<td>2.244 (1.755-2.870)</td>
<td>6.86</td>
</tr>
<tr>
<td></td>
<td>CAY-SR</td>
<td>0.973 (0.863-1.097)</td>
<td>14.72</td>
<td>2.529 (1.947-3.245)</td>
<td>7.73</td>
</tr>
</tbody>
</table>

LC$_{50}$: Lethal concentration for 50% mortality; RR$_{50}$: resistance ratio for LC$_{50}$; LC$_{95}$: Lethal concentration for 95% mortality; RR$_{95}$: resistance ratio for LC$_{95}$; CAY-SS: homozygous for susceptible alleles without insecticide exposure; CAY-RR: homozygous for resistant alleles without insecticide exposure; CAY-RS: heterozygous by the cross between CAY-RR females and CAY-SS males; CAY-SR: heterozygous by the cross between CAY-SS females and CAY-RR males; CAY-P: strain selected with 3% permethrin.
Figure 3. Mortality of Recife strains against WHO standard malathion 5.0% (top) and permethrin 0.75% (below).
p <0.002664) (Table 3). In adults of the REC-R strain, two genes were significantly upregulated in generation G45 (CYP9J28 and CYP6BB2) (MW: p <0.004329). A significant upregulation in these genes was observed in REC-U (CYP9J28; MW: p <0.002498) and REC-M (CYP9J28 and CYP6BB2; MW: p <0.002165). Moreover, the CCae3A gene was upregulated in REC-M (MW: p <0.000250) in adults (Table 4).

### Table 3. Mean fold change in gene expression for larvae, Recife colony.

<table>
<thead>
<tr>
<th>Strain</th>
<th>G</th>
<th>CCae3A</th>
<th>CYP6F3</th>
<th>CYP6N12</th>
<th>CYP9M9</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MF</td>
<td>sd</td>
<td>MF</td>
<td>sd</td>
<td>MF</td>
</tr>
<tr>
<td>REC-Baseline</td>
<td>G37</td>
<td>1.06</td>
<td>0.39</td>
<td>1.13</td>
<td>0.54</td>
</tr>
<tr>
<td>REC-U</td>
<td>G45</td>
<td>0.34</td>
<td>0.28</td>
<td>0.65</td>
<td>0.34</td>
</tr>
<tr>
<td>REC-R</td>
<td>G45</td>
<td>0.73</td>
<td>0.23</td>
<td>0.92</td>
<td>0.38</td>
</tr>
<tr>
<td>REC-M</td>
<td>G45</td>
<td>0.38</td>
<td>0.12</td>
<td>0.41</td>
<td>0.07</td>
</tr>
<tr>
<td>REC-P</td>
<td>G45</td>
<td>1.77</td>
<td>0.39</td>
<td>0.41</td>
<td>0.13</td>
</tr>
</tbody>
</table>

G: Generation; MF: mean fold; sd: standard deviation. REC-U: strain without insecticide exposure; REC-R: strain with temephos exposure; REC-M: strain selected for malathion; REC-P: strain selected for permethrin. Bold: Expression values significantly different from Baseline (Holm–Bonferroni method).

### Table 4. Mean fold change in gene expression for adults, Recife colony.

<table>
<thead>
<tr>
<th>Strain</th>
<th>G</th>
<th>CCae3A</th>
<th>CYP9J28</th>
<th>CYP6N12</th>
<th>CYP6BB2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MF</td>
<td>sd</td>
<td>MF</td>
<td>sd</td>
<td>MF</td>
</tr>
<tr>
<td>REC-Baseline</td>
<td>G37</td>
<td>1.35</td>
<td>0.79</td>
<td>1.04</td>
<td>0.29</td>
</tr>
<tr>
<td>REC-U</td>
<td>G45</td>
<td>1.75</td>
<td>0.43</td>
<td>2.63</td>
<td>0.47</td>
</tr>
<tr>
<td>REC-R</td>
<td>G45</td>
<td>1.7</td>
<td>0.52</td>
<td>2.56</td>
<td>0.31</td>
</tr>
<tr>
<td>REC-M</td>
<td>G45</td>
<td>2.89</td>
<td>0.37</td>
<td>3.37</td>
<td>2.11</td>
</tr>
<tr>
<td>REC-P</td>
<td>G45</td>
<td>1.56</td>
<td>1.00</td>
<td>0.72</td>
<td>0.38</td>
</tr>
</tbody>
</table>

G: Generation; MF: mean fold; sd: standard deviation. REC-U: strain without insecticide exposure; REC-R: strain with temephos exposure; REC-M: strain selected for malathion; REC-P: strain selected for permethrin. Bold: Expression values significantly different from Baseline (Holm–Bonferroni method).

Resistance ratio to permethrin, malathion and temephos.

Nearly all strains were more resistant to all insecticides than New Orleans. REC-P was 5x more resistant to permethrin than REC-U, REC-M, and REC-R (Table 5). REC-R and REC-M were slightly more resistant to malathion (~2x) than REC-U or REC-P. REC-R, REC-M and REC-P were more resistant to temephos (>2x) than REC-U.

**Discussion**

Inference of insecticide resistance in adults of *Ae. aegypti* is rarely performed by dose-response curves. A recent review (Moyes et al., 2017) highlights the lack of literature that calculate resistance ratios based on dose-response or lethal time (see S2 file in (Moyes et al., 2017)). This practice limits comparative analysis of our results with other resistance studies. Resistance ratios for permethrin in CAY with homozygous resistance alleles (I/CC; CAY-RR and CAY-P) were of a similar magnitude to homozygous resistant *Ae. aegypti* from Cayman Islands populations (Harris et al., 2010a). In other field populations, a significant positive correlation between the frequency of IICC individuals and resistance ratio for permethrin was also observed, which indicates a higher resistance for these double homozygotes (Estep et al., 2018). The lack of variation in permethrin resistance ratio between IICC strains regardless of selection pressure indicates that this allele is primarily responsible for the phenotype observed. Moreover, the other CAY strains presented lower resistance ratios to permethrin. The susceptible double homozygous strain (VV/FF; CAY-SS) had a resistance ratio similar to susceptible homozygous field populations (RR: 0.8 – 7.0) in Asia (Brengues et al., 2003) and lower than REC-P (the selected strain for permethrin in REC). This intermediate level of resistance in heterozygotes is consistent with the recessive nature of kdr alleles in mosquitoes and other dipterans (Gomes et al., 2017; Huang et al., 2004; Saavedra-Rodriguez et al., 2007). However, the combination of multiple heterozygote kdr alleles in *Ae. aegypti* can present a stronger resistance phenotype in the future, as is observed in Thailand where triple heterozygotes (S/P989 + V/G1016 + F/C1534) had a higher resistance ratio to deltamethrin than kdr homozygotes at F1534C (Plernsub et al., 2016).
Changes in temephos resistance in our REC strains differed from previous studies in two primary ways. First, the resistance level in our REC-R temephos selected strain was lower than the resistance level reported in previous studies where REC was put under similar selection pressure (Diniz et al., 2015; Melo-Santos et al., 2010). This is likely because we used the New Orleans colony as the denominator in calculating the resistance ratio, while the Rockefeller colony was used in other studies. Rockefeller has a lower LC50 on average than New Orleans (see S1 file in (Moyes et al., 2017)). Second, in this study, temephos resistance in the REC-U unselected strain was not completely reversed while previous studies have documented reversal in a similar number of generations. This may be because our starting material had been under temephos selection pressure for longer prior to starting the experiments.

In the Recife strains, the response to the adulticides malathion and permethrin was different to the response to the larvicide temephos. REC P and REC M showed a similar LC50 to temephos as the REC-R colony and a slightly lower LC95. The gene expression within REC-R showed no significant variation compared to the baseline colony. However, REC-P, REC-M and REC-U showed downregulation of detoxification genes compared to the baseline, consistent with the lower tolerance to temephos displayed in the LC95s.

In contrast to larvicide exposure, resistance ratios for REC-P and REC-M showed a different pattern. Substantial differences between REC-P compared to either REC-R or REC-U suggest that exposure to permethrin increased the tolerance of this insecticide in the Recife colony. REC-M and REC-R present similar resistance ratios to Malathion, while the unexposed strain

**Table 5. Lethal concentrations and resistance ratios of Recife strains for three insecticides (i.e. permethrin, malathion and temephos).**

<table>
<thead>
<tr>
<th>Insecticide</th>
<th>Strain</th>
<th>LC50</th>
<th>RR50</th>
<th>LC95</th>
<th>RR95</th>
</tr>
</thead>
<tbody>
<tr>
<td>Permethrin</td>
<td>New Orleans</td>
<td>0.066 (0.053-0.083)</td>
<td>N/A</td>
<td>0.327 (0.211-0.507)</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>REC-U</td>
<td>0.131 (0.106-0.162)</td>
<td>1.98</td>
<td>0.404 (0.259-0.631)</td>
<td>1.24</td>
</tr>
<tr>
<td></td>
<td>REC-R</td>
<td>0.155 (0.121-0.198)</td>
<td>2.35</td>
<td>0.629 (0.390-1.014)</td>
<td>1.92</td>
</tr>
<tr>
<td></td>
<td>REC-M</td>
<td>0.111 (0.0922-0.134)</td>
<td>1.68</td>
<td>0.451 (0.283-0.717)</td>
<td>1.38</td>
</tr>
<tr>
<td></td>
<td>REC-P</td>
<td>0.657 (0.585-0.738)</td>
<td>9.94</td>
<td>2.876 (2.292-3.608)</td>
<td>8.80</td>
</tr>
<tr>
<td>Malathion</td>
<td>New Orleans</td>
<td>0.329 (0.274-0.394)</td>
<td>N/A</td>
<td>1.423 (1.109-1.825)</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>REC-U</td>
<td>0.566 (0.490-0.654)</td>
<td>1.72</td>
<td>2.093 (1.694-2.586)</td>
<td>1.47</td>
</tr>
<tr>
<td></td>
<td>REC-R</td>
<td>0.898 (0.741-1.087)</td>
<td>2.73</td>
<td>7.709 (4.302-13.812)</td>
<td>5.42</td>
</tr>
<tr>
<td></td>
<td>REC-M</td>
<td>1.006 (0.876-1.155)</td>
<td>3.06</td>
<td>4.583 (3.430-6.124)</td>
<td>3.22</td>
</tr>
<tr>
<td></td>
<td>REC-P</td>
<td>0.614 (0.566-0.666)</td>
<td>1.87</td>
<td>1.091 (0.957-1.245)</td>
<td>0.77</td>
</tr>
<tr>
<td>Temephos</td>
<td>New Orleans</td>
<td>0.011 (0.010-0.011)</td>
<td>N/A</td>
<td>0.032 (0.028-0.036)</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>REC-U</td>
<td>0.145 (0.141-0.149)</td>
<td>13.81</td>
<td>0.304 (0.284-0.326)</td>
<td>9.53</td>
</tr>
<tr>
<td></td>
<td>REC-R</td>
<td>0.342 (0.328-0.356)</td>
<td>32.57</td>
<td>1.163 (1.065-1.269)</td>
<td>36.46</td>
</tr>
<tr>
<td></td>
<td>REC-M</td>
<td>0.376 (0.357-0.396)</td>
<td>35.81</td>
<td>0.845 (0.798-0.938)</td>
<td>26.49</td>
</tr>
<tr>
<td></td>
<td>REC-P</td>
<td>0.355 (0.339-0.372)</td>
<td>33.81</td>
<td>0.810 (0.750-0.873)</td>
<td>25.39</td>
</tr>
</tbody>
</table>

LC50: Lethal concentration for 50% mortality; RR50: resistance ratio for LC50; LC95: Lethal concentration for 95% mortality; RR95: resistance ratio for LC95. REC-U: strain without insecticide exposure; REC-R: strain with temephos exposure; REC-M: strain selected for malathion; REC-P: strain selected for permethrin.
REC-U presents a lower resistance ratio than REC-M/REC-R. Malathion exposure over more generations will be required to increase the divergence between REC-R and REC-M phenotypes. We have experienced three main limitations in the selection of these strains: i) inconsistent blood sources, ii) time required to reverse the resistance mechanism (Diniz et al., 2015), iii) potential diversity loss associated with bottlenecks and/or high mortality due to aggressive artificial selection. We experienced difficulties in maintaining Ae. aegypti after a few generations using horse blood. Unfortunately, this was the only blood source available for our lab after the source of human blood was interrupted. Moreover, REC-M exhibited a drastic increase in mortality (over 90%) when we adjusted the malathion exposure concentration to 1.5% at generation 43. Mortality levels in malathion selection remained higher than 50% after this adjustment, and we had difficulties maintaining this strain post-generation 45. Future malathion selection will require a longer build-up to create a more viable resistant strain.

Conclusions
We generated strains of Ae. aegypti which differ in phenotypic resistance to permethrin, malathion and temephos. The selected CAY and REC strains will allow for further research on the effects of target-site and metabolic resistance, respectively, on the life-history traits, behaviour and vector competence of this important arbovirus vector. The strains can also be used to compare the efficacy of novel insecticide formulations in strains with similar genetic backgrounds and different mechanisms of resistance.

Data availability
Underlying data

This project contains the following underlying data:
- Raw values for insecticide selection mortality per generation
- Raw values for mortality in WHO bioassay
- CT values for detoxification genes in larvae
- CT values for detoxification genes in adults
- Raw values for mortality to a range of insecticide concentrations, used to calculate LC50, LC95 and resistance ratios

Extended data

This project contains the following extended data:
- Table S1. Insecticide selection mortality (%) for Cayman and Recife strains against standard WHO tube concentrations.

Data are available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY 4.0).

References

George L, Lenhart A, Toledo J, et al.: Community-Effectiveness of Temephos for


Open Peer Review

Current Peer Review Status: ?

Version 1

Reviewer Report 17 August 2020

https://doi.org/10.21956/wellcomeopenres.17519.r39800

© 2020 Smith L. This is an open access peer review report distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Leticia Smith
National Institutes of Health, Rockville, MD, USA

This study used two insecticide resistant parental colonies of Aedes aegypti to develop multiple strains with resistance to different insecticides. The aim was to create these strains as a tool so that the mechanisms of resistance can be studied without interference from varying genetic backgrounds. Understanding insecticide resistance in mosquitoes is as important now as ever and because insecticides will continue to be used to control disease vectors in the foreseeable future, understanding the mechanisms underlying insecticide resistance is critical for vector control. For this reason, having multiple strains with the same genetic background but different resistance mechanisms and resistance to different insecticides is a great value to the insecticide resistance research community.

Below are my suggestions and specific points related to the questions above that I hope will improve this publication.

The rationale for developing the new method wasn't clearly explained, but perhaps part of my confusion is that there didn't seem to be anything new about the method. The rationale for developing the strains was clearly articulated, but the method of generating strains with the same genetic background through exposing versus not exposing sub-samples of the population to insecticides has been done numerous times already. Also selecting strains based on resistance genotype has been done before. I may have simply missed it, but perhaps the novelty of this method needs to be elaborated and/or more clearly articulated. As it is now, it seems like the goal was to simply select strains from a parental colony.

The description of the selection method was technically sound, but the evaluation of resistance part of the method could use some improvement. For the detection of kdr mutations, is there a reason why V410L was not checked? This is a common mutation in Ae. aegypti in the Americas and was only first detected after the initial study that collected the parental strain was published. How were the metabolic genes selected? Are they known to be responsible for resistance in the Recife region where the colony came from? These genes can vary across populations and by insecticide selection, so some clear explanation of how these particular ones were tested is needed. Also, why
was this assay done only on the REC strains and not CAY? Some of these genes are also associated with pyrethroid resistance and are likely involved in the resistance observed in the CAY strain. Perhaps insecticide bioassays using CYP, carboxylesterase, or GST synergists would be a better evaluation of metabolic resistance than gene expression? For determining the resistance ratios, why use the WHO assay?

As far as details provided to allow replication of the method development and its use by others, some important information is missing or needs clarification. More information on the parental colonies would be helpful. Although the citations provide information on where the strains originated from, they do not say how many generations the strains have been in the lab prior to this study, or whether they were previously selected for insecticide resistance in the lab or are the original field collected colony. The source and stereochemistry (when applicable) for the insecticides used needs to be provided. The generation numbers used are a bit confusing. It would be helpful if only the generations from this study were used and not a mix between generations selected and what might be total number of generations in the lab. Also, in the mosquito selection regime section, the number of generations don’t seem to add up. After 5 generations without exposure, wouldn’t that be generation 5? If the authors mean after five additional generations without exposure, wouldn’t that be generation 15 (10 + 5)? Figure 1 shows the selection process, but that doesn’t seem to match the selection process described in the text. CAY-P was selected at every generation, not every third generation. REC-R was selected as larvae, so not using the WHO bioassay, and REC-M and REC-P seems to have been selected a total of nine generations (the figure shown only three selections). What was the calculation or method used to determine $LC_{50}$ and $LC_{95}$? R is a programming language and the software is only an environment to perform computations, this gives no information of how the $LC_{50}$ was calculated. Maybe a link to a GitHub project or profile with the code used? Or a citation for the calculation methodology used? The authors also need to be more specific about how the resistance ratio was calculated. How were they compared? How was significance determined? How was control mortality accounted for? For figures 2 and 3, change “time point” to “generations” and to make it clearer and add a description of what the bars represent (error, st. dev., 95%CI?).

The authors provide a great statement in the beginning for the discussion about the need to perform more dose-response curves in insecticide resistance studies, but sadly this study also didn’t exactly provide a dose-response curve. The issue with using a concentration, such as in the WHO assay, is that the exact amount of insecticide the individuals were exposed to remains unknown. Some animals may be more active and therefore be exposed to more by walking around the tube more than others. Also, an individual that is being affected may fall on its side more frequently and in the process get a higher exposure than a healthy animal that is standing straight. While LC assays are great for certain purposes, they are not optimal for determining resistance ratios and dose-response curves. Would it be possible to do topical assays on the final strains to get a dose response RR instead of a concentration response RR?

In regard to using the New Orleans colony as the denominator in calculating the resistance ratio, a recent study showed that the NO strain has multiple $kdr$ mutations\(^1\), just something to keep in mind. In regard to the potential loss of diversity associated with bottlenecks limitations mentioned by the authors in the discussion section, perhaps backcrossing back to the parental colony after each selection could reduce this problem. With enough selections and backcrossing you should retain only the resistance mutations (if they were present in the original population) and any genes that are linked to the resistance alleles (this one is harder to separate out).
On a very minor note, the citation for Harris et al. 2010 was listed twice.

References

Is the rationale for developing the new method (or application) clearly explained?
No

Is the description of the method technically sound?
Partly

Are sufficient details provided to allow replication of the method development and its use by others?
Partly

If any results are presented, are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions about the method and its performance adequately supported by the findings presented in the article?
Yes

*Competing Interests*: No competing interests were disclosed.

*Reviewer Expertise*: Insecticide resistance, mosquito biology, insect physiology.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 02 Dec 2020

Lisa Reimer, Liverpool School of Tropical Medicine, Liverpool, UK

Thank you to Reviewer 1 for the helpful and insightful comments. We have made additions to the manuscript to clarify the scope, re-analysed the data and provided a few additional methodological details and rationale. There were quite a few suggestions for additional research or using different methods than what we have (such as tarsal assays, synergist assays, sequencing). While these are all valid suggestions they were beyond the scope of this methods article. We have hopefully clarified the aims of this paper, which is to describe the methods underlying the selection and creation of the 8 mosquito strains. These strains have been provided to other researchers to address some of the key questions about insecticide resistance and the purpose of this article is to provide the complete background
on the process. Our primary outcome was the resistance ratio (compared to a susceptible lab strain). Where we chose a few markers (kdr alleles, metabolic resistance genes, knockdown in WHO assays to select insecticides) the purpose was to track the impact the process of selection was having not to provide a complete characterisation of all possible mechanisms. We have provided a response to the below comments and have uploaded a new version of the manuscript to reflect the changes.

Comment 1
The rationale for developing the new method wasn't clearly explained, but perhaps part of my confusion is that there didn't seem to be anything new about the method. The rationale for developing the strains was clearly articulated, but the method of generating strains with the same genetic background through exposing versus not exposing sub-samples of the population to insecticides has been done numerous times already. Also selecting strains based on resistance genotype has been done before. I may have simply missed it, but perhaps the novelty of this method needs to be elaborated and/or more clearly articulated. As it is now, it seems like the goal was to simply select strains from a parental colony.

This is correct, what we present here is not completely novel methodology but it is a record of the methods used to develop these mosquito resources and it provides an initial characterisation of resistance status. These strains have been shared with other researchers who are currently investigating different aspects of insecticide resistance and we wanted to have a published record of how the resources were established. While the methods are known amongst the scientific community, these selected strains are still a unique resource which allows for the study of multiple resistant phenotypes. We request editorial advice on the most appropriate classification for this manuscript.

Comment 2
The description of the selection method was technically sound, but the evaluation of resistance part of the method could use some improvement. For the detection of kdr mutations, is there a reason why V410L was not checked? This is a common mutation in Ae. aegypti in the Americas and was only first detected after the initial study that collected the parental strain was published. How were the metabolic genes selected? Are they known to be responsible for resistance in the Recife region where the colony came from? These genes can vary across populations and by insecticide selection, so some clear explanation of how these particular ones were tested is needed. Also, why was this assay done only on the REC strains and not CAY? Some of these genes are also associated with pyrethroid resistance and are likely involved in the resistance observed in the CAY strain. Perhaps insecticide bioassays using CYP, carboxylesterase, or GST synergists would be a better evaluation of metabolic resistance than gene expression? For determining the resistance ratios, why use the WHO assay?

CAYMAN has been maintained in laboratory since 2008 by LITE and is screened for two kdr alleles (F1534C and V1016I). Moyes et al. (2017) was an important reference in our rationale for the choice to continue screening F1534C and V1016I since they are considered key mutations associated with pyrethroid resistance in Aedes aegypti when compared with other kdr alleles. We understand that V410L is among the many kdr alleles of Aedes aegypti in the Americas, and was detected approximately 10 years after our colony was established.
though was found in historic samples). We chose to continue the tracking of F1534C and V1016I to perform our selection process. Our results indicate that these two resistance alleles are linked in CAYMAN and it is possible that other \textit{kdr} alleles exist in this strain. The information about alternative resistant alleles which may be contributing further to the resistant phenotype can be explored through sequencing studies which are currently underway by other researchers. We did not screen detox genes in CAYMAN for characterization, since resistant phenotypes suggest similar resistance for both homozygotic resistant strains (CAY-P, CAY-RR).

**RECIFE**: We selected a battery of detox genes associated with metabolic resistance in \textit{Aedes aegypti} from Americas (based on the literature) to perform an initial screen in our strains. The purpose of the initial screen was to determine whether the selection process was having an impact and to supplement the primary outcome measure which were the resistance ratios. However, as above, further research studies based on RNAseq analysis will be conducted by other researchers. We agree that transcriptomic analyses will be very valuable but they are beyond the scope of this methods article.

We have added some information in the text to clarify genes used and the rationale:

- Line 75-81: “Here we present the resistant phenotypes after ten generations of insecticide exposure... Additionally, we present differences in a select number of resistant alleles and metabolic genes which can be used to inform further research.”
- Line 188-192: “Four genes associated with insecticide resistance were selected to screen expression profiles in larvae (CCae3A, CYP6N12, CYP6F3 and CYP9M9: all associated with temephos resistance in multiple studies) and adults (CCae3A and CYP6N12 associated with temephos resistance; and CYP6BB2 and CYP9J28a associated with pyrethroid resistance in multiple studies, Moyes et al. 2017).”

We also updated the abstract to clarify which alleles were tracked during the selection.

**Bioassays with synergist**: We agree that studies with synergists and antagonists are interesting for determining metabolic resistance, but this was not a priority during our selection process.

**Quantitative approach with WHO assays**: we address this concern under comment 4 below

**Comment 3**
As far as details provided to allow replication of the method development and its use by others, some important information is missing or needs clarification. More information on the parental colonies would be helpful. Although the citations provide information on where the strains originated from, they do not say how many generations the strains have been in the lab prior to this study, or whether they were previously selected for insecticide resistance in the lab or are the original field collected colony. The source and stereochemistry (when applicable) for the insecticides used needs to be provided. The generation numbers used are a bit confusing. It would be helpful if only the generations from this study were used and not a mix between generations selected and what might be total number of generations in the lab. Also, in the mosquito selection regime section, the
number of generations don't seem to add up. After 5 generations without exposure, wouldn't that be generation 5? If the authors mean after five additional generations without exposure, wouldn't that be generation 15 (10 + 5)? Figure 1 shows the selection process, but that doesn't seem to match the selection process described in the text. CAY-P was selected at every generation, not third generation. REC-R was selected as larvae, so not using the WHO bioassay, and REC-M and REC-P seems to have been selected a total of nine generations (the figure shown only three selections). What was the calculation or method used to determine LC50 and LC95? R is a programming language and the software is only an environment to perform computations, this gives no information of how the LC50 was calculated. Maybe a link to a GitHub project or profile with the code used? Or a citation for the calculation methodology used? The authors also need to be more specific about how the resistance ratio was calculated. How were they compared? How was significance determined? How was control mortality accounted for? For figures 2 and 3, change “time point” to “generations” and to make it clearer and add a description of what the bars represent (error, st. dev., 95%CI?).

Colony: We include description for all three colonies used in this study (lines 85-100).

“A Total of three colonies from *Aedes aegypti* were used in this study: Colony CAYMAN (CAY) was originally established in 2008 with *Aedes aegypti* collected in Grand Cayman island (Caribbean). This colony is highly resistant to pyrethroids and DDT, attributed to kdr alleles (F1534C and V1016I), and has been routinely selected with 0.75 % permethrin for 1 hour in the Liverpool Insect Testing Establishment (LITE: https://lite.lstmed.ac.uk/lite-facilities/lite-insectaries/aedes-aegypti-cayman). Colony RECIFE (REC) was originally established in 2004 with *Aedes aegypti* collected in Araripina, Brazil (7° 32’ S and 40° 34’ W; Melo-Santos et al. 2010). This colony is resistant to temephos (OP), and biochemical assays indicate a higher activity of multiple detox enzyme families. Larvae from this colony have been routinely selected with 0.5 mg/L temephos for 24 hours. Colony NEW ORLEANS is a susceptible colony established at LSTM in the 1970s. This colony originated in New Orleans (USA) and it is maintained in laboratory without insecticide exposure. Routine screening for target-site mutations associated with resistance indicates a lack of kdr alleles in the sodium channel and resistant alleles in Ace-1 (LITE: https://lite.lstmed.ac.uk/lite-facilities/lite-insectaries/aedes-aegypti-new-orleans).”

Stereochemistry: We include more information about the insecticides (lines 129-132)

“The adult mosquitoes were selected with either permethrin (Sigma Aldrich, Pestanal®, CAS # 52645-53-1, >95.0% sum of cis+trans 97.8%) or malathion (Sigma Aldrich, Pestanal®, CAS # 121-75-5) and larvae were selected with temephos (CHEM SERVICE INC., CAS # 3383-96-8) (see “Mosquito selection regime”).”

Generations: We agree with the reviewer about the generation information in the text. To avoid confusion, we use the generation number starting in the beginning of our experiment for both Recife (G1-G10) and Cayman (G1-G11). The information about colony age at the beginning of our experiment was only included in the figure 1.
We updated:
- The text of the manuscript (mainly methods);
- Tables 1, 3, and 4;
- Figures 1 to 3;
- supplementary information: CT values adults.csv; Mortality after insecticide selection.csv; WHO bioassay results.csv; Table S1. Insecticide selection total mortality.csv.

Information about analysis:
Thank for your comment. We recognize that the information about analysis was incomplete.
- We include more details about the analysis and the R script.

Lines 235-244: “This process was repeated for each concentration in the assay. Larval mortality was recorded at 24 hours and Abbot's formula (Abbot 1925) was used to adjust mortality when necessary. The data from bioassays was organized by concentration in each biological replicate. These data were used to create general linear models (G.L.M.) with “logit” models for each strain using “glm” function in statistical software R-2.15.2. Lethal concentration and 95% confidence intervals were calculated using the r-package “MASS”. Resistance ratios were calculated based on comparison of LC$_{50}$ and LC$_{95}$ from each strain against the reference colony New Orleans, which is fully susceptible to all three insecticides. Confidence intervals for ratios were calculated using the method MOVER-R (Newcombe 2016) presented in the R-package “pairwiseCI.”
- We update the Table 2 and 5
- We update the information about resistance ratios in Results (lines 273-277, and lines 297-300).
- We also include new references:

Figure 2 and 3:
We include “Bars represent 95% CI.”

Comment 4
The authors provide a great statement in the beginning for the discussion about the need to perform more dose-response curves in insecticide resistance studies, but sadly this study also didn’t exactly provide a dose-response curve. The issue with using a concentration, such as in the WHO assay, is that the exact amount of insecticide the individuals were exposed to remains unknown. Some animals may be more active and therefore be exposed to more by walking around the tube more than others. Also, an individual that is being affected may fall on its side more frequently and in the process get a higher exposure than a healthy animal that is standing straight. While LC assays are great for certain purposes, they are not optimal for determining resistance ratios and dose-response curves. Would it be possible to do topical assays on the final strains to get a dose response RR instead of a concentration response RR?
We understand the critique on using WHO assays to calculate lethal concentrations and resistance ratios. The calculation of doses based on mosquito weight may refine a resistance ratio calculation reducing certain limitations of WHO assays (or similar assays such as CDC bottle assay). However, topical assays also present disadvantages to study insecticide resistance since they require immobilization (e.g. using cold temperature, CO₂) and insecticide is normally apply in thorax or head (not the most common area of entry in the field). For this reason, we believe that WHO assays are a valid method for this manuscript, and they are a clear improvement over qualitative assays with reference concentrations.

We have included a discussion on the limitations of WHO assays for quantitative approach:

Lines 359-364: “Our method to calculate resistance ratios using WHO tubes is not able to define the individual dose received per mosquito, since mosquito weight and individual activity against the treated surface could not be measured. Topical assays, which don't mimic natural exposure routes, do allow for controlled application and would further reduce the individual variation observed within each strain.”

We also added to the discussion:

Lines 303-306: “Inference of insecticide resistance in adults of *Ae. aegypti* is rarely performed by quantitative methods. A recent review (Moyes *et al.*, 2017) highlights the lack of literature that calculate resistance ratios based on dose-response, lethal concentration or lethal time (see S2 file in (Moyes *et al.*, 2017)).”

**Comment 5**

In regard to using the New Orleans colony as the denominator in calculating the resistance ratio, a recent study showed that the NO strain has multiple kdr mutations1, just something to keep in mind.


Thank you for the concern. However, our susceptible New Orleans colony is an old colony without *Kdr* alleles such as “Liverpool”, and “Rockefeller” (see comment 3). Moreover, Fan et al. is focused on recent collections from Americas and Africa that includes recent field collection from New Orleans.

We include the description of all colonies used in the study (Lines 88-105).

In regard to the potential loss of diversity associated with bottlenecks limitations mentioned by the authors in the discussion section, perhaps backcrossing back to the parental colony after each selection could reduce this problem. With enough selections and backcrossing you should retain only the resistance mutations (if they were present in the original population) and any genes that are linked to the resistance alleles (this one is harder to separate out).
Thank you for this suggestion.

On a very minor note, the citation for Harris et al. 2010 was listed twice.

References were corrected to remove this duplication.

*Competing Interests:* No competing interests were disclosed.