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RESEARCH ARTICLE

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Colonization of *Anopheles coustani*, a neglected malaria vector in Madagascar

Tsarasoa M. Andrianinarivomanana^{1,*}, Fenomiaranjara T. Randrianaivo¹, Mandaniaina R. Andriamiarimanana¹, Mihary R. Razafimamonjy¹, Haja J.S. Velonirina¹, Nicolas Puchot², Romain Girod¹, and Catherine Bourgouin^{2,*}

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Abstract – *Anopheles coustani* has long been recognized as a secondary malaria vector in Africa. It has recently been involved in the transmission of both *Plasmodium falciparum* and *P. vivax* in Madagascar. As most secondary malaria vectors, *An. coustani* mainly bites outdoors, which renders the control of this mosquito species difficult using classical malaria control measures, such as the use of bed nets or indoor residual spraying of insecticides. For a better understanding of the biology and vector competence of a vector species, it is useful to rear the species in the laboratory. The absence of a colony hinders the assessment of the bionomics of a species and the development of adapted control strategies. Here, we report the first successful establishment of an *An. coustani* colony from mosquitoes collected in Madagascar. We used a forced copulation procedure as this mosquito species will not mate in cages. We describe our mosquito colonization procedure with detailed biological features concerning larval to adult development and survival, recorded over the first six critical generations. The procedure should be easily applicable to *An. coustani* from different African countries, facilitating local investigation of *An. coustani* vector competence and insecticide resistance using the colony as a reference.

Key words: Anopheles coustani, Colony, Forced mating, Rearing, Madagascar.

Résumé – Colonisation d'Anopheles coustani, vecteur négligé du paludisme à Madagascar. Anopheles coustani est reconnu depuis longtemps comme un vecteur secondaire du paludisme en Afrique. Il a récemment été impliqué dans la transmission de Plasmodium falciparum et de P. vivax à Madagascar. Comme la plupart des vecteurs secondaires du paludisme, An. coustani pique principalement à l'extérieur, ce qui rend difficile le contrôle de cette espèce de moustique par les mesures classiques de lutte contre le paludisme telles que l'utilisation de moustiquaires ou la pulvérisation intradomiciliaire d'insecticides à effet rémanent. Pour une meilleure compréhension de la biologie et de la compétence vectorielle d'une espèce vectrice, il est utile d'élever l'espèce en laboratoire. L'absence de colonie gêne l'évaluation de la bionomie d'une espèce et le développement de stratégies de contrôle adaptées. Nous rapportons ici le premier établissement réussi d'une colonie d' An. coustani issue de moustiques collectés à Madagascar. Nous avons utilisé une procédure de copulation forcée car cette espèce de moustique ne s'accouple pas en cage. Nous décrivons notre procédure de colonisation des moustiques avec des caractéristiques biologiques détaillées concernant le développement et la survie des stades larvaires aux adultes, enregistrées au cours des six premières générations critiques. La procédure devrait être facilement applicable aux An. coustani de différents pays africains, facilitant les enquêtes locales sur la compétence vectorielle d'An. coustani et sa résistance aux insecticides, en utilisant une colonie comme référence.

Introduction

Malaria is still a public health concern in many countries, mainly in Africa [38]. Malaria is transmitted by mosquitoes belonging to the *Anopheles* genus. The *Anopheles* genus is further subdivided into eight subgenera of which three (*Anopheles*,

*Corresponding authors: cabourg@pasteur.fr (Catherine Bourgouin); andrianina.tsarasu@gmail.com (Tsarasoa M. Andrianinarivomanana) Cellia, and Nyssorhynchus) contain all known dominant vector species (DVS) of human malaria [34]. For countries close to malaria elimination, residual malaria is of high concern. In this regard, the role of the so-called secondary malaria vectors, often outdoor biters, in maintaining malaria transmission has been considered [39]. However, for decades most laboratory and field studies focused on the primary or dominant malaria vectors [34].

With the stagnation in worldwide malaria incidence over the last decade, it is more and more important to better

¹ Institut Pasteur de Madagascar, Medical Entomology Unit, Antananarivo, Madagascar

² Institut Pasteur, Université de Paris Cité, Biology of Host-Parasite Interactions, Paris, France

understand the bionomics of secondary malaria vectors for which limited data are available. This is especially crucial for secondary malaria vectors in Madagascar, a country that has recorded a worrying increase in malaria cases despite indoor residual spraying and distribution campaigns for insecticidal-treated nets (WHO 2022).

Anopheles coustani is an Afrotropical anopheline species belonging to the Anopheles subgenus. In several African countries, including Madagascar, An. coustani is now widely recognized as a secondary vector of human malaria parasites [15, 17, 20, 24, 26]. Although it was reported as being both anthropophilic and exophagic (feeding outdoors on humans) in South Africa [10], An. coustani is often referred to as a zoophilic mosquito [35]. Nevertheless, because of its abundance it can be a locally major malaria vector [15, 28]. Indeed, in Madagascar, An. coustani was shown to be the major vector of both Plasmodium vivax and Plasmodium falciparum in a village where it showed high anthropophilic behavior, associated with both endophilic and exophilic behavior [18]. Furthermore, An. coustani has been implicated as a potential vector of Rift Valley Fever virus in Madagascar [31] and of Chikungunya virus in Senegal [13].

Anopheles coustani Laveran was first described in 1900 by Alphonse Laveran from samples received from Madagascar [23]. Anopheles mauritianus described at about the same time from mosquitoes collected in the Mascarene Islands was later considered as identical to An. coustani Laveran [14, 32]; details are presented by M. Coetzee [11]. Using cytogenetics on larval polytene chromosomes, M. Coetzee reported the existence of two cryptic species within An. coustani collected in South Africa and Eswatini (former Swaziland) [10]. When crossed, these species named species A and species B produced infertile male F1. More recently, using An. coustani samples from Madagascar and morphology as well as cytogenetics, the same author confirmed that the Malagasy mosquitoes belong to the former species A, and conclude that species B was a different species named An. crypticus [11]. Of note, Malagasy specimens were then referred to as neotypes as the original type specimens were lost. In 2020, sampling the genetic diversity of An. coustani populations in Zambia and the Democratic Republic of Congo revealed two distinct phylogenetic groups [8].

To our knowledge, no colony of *An. coustani* has ever been established. As for other species, having at hand a colony will be instrumental for various types of studies: insecticide assays with the colony as a reference, behavioral study for refining vector control strategies, assessing whether the bacteria *Asaia* or *Wolbachia* could be considered as additional tools for controlling mosquito populations or their vectorial competence toward *Plasmodium* and viruses [6, 7, 33].

Establishing a novel mosquito colony from field collected samples can be very tricky. The main challenge is mating behavior. Indeed, many anopheline mosquitoes are eurygamous, mating in open space often in swarms, and anopheline females are also known for their refractoriness to multiple mating [9, 36]. *Anopheles coustani* is no exception, as evidenced in preliminary assays while working with large numbers of F1 progeny obtained by in-tube forced oviposition of gravid wild females [27]. Even though F1 females would readily feed on blood, none would lay eggs over more than a week post blood

feeding. Observation of a subset of the spermatheca of F1 females revealed that none had been fertilized (Jean Jose Nepomichene and Bourgouin, unpublished).

To overcome the absence of mating in a constrained environment (rearing cages), a "forced mating" technology was developed [4], and successfully used for establishing anopheline colonies [1, 29], even though the technique is highly demanding in human resources and skill. However, forced mating does not equal female fertilization. Indeed, it was reported that males from major anopheline vectors from the *Nyssorhynchus* subgenus, including *Anopheles darlingi*, would readily copulate by the forced mating technique but would fail to inseminate the females ([22]; Puchot, personal observation). On the contrary, several colonies of anophelines from both the *Anopheles* and *Cellia* subgenus were created and maintained using the forced mating technique [1, 3, 29].

Based on successful published records on anopheline from the *Anopheles* subgenus and our preliminary observations, we initiated the establishment of the first *An. coustani* colony using the forced mating technique. Here, we provide details on the major initial steps until we obtained the eighth generation and discuss ways of improvement toward the establishment of a free-mating colony of *An. coustani*.

Material and methods

Producing the F1 generation from wild *An.* coustani

Anopheles coustani females were captured in Maroharona (17°36'44.33"S, 46°56'2.11"E), municipality of Andriba (Maevatanana district) in the Northwest fringe of the Central Highlands of Madagascar, in September 2019. The mosquitoes were collected in zebu parks using mouth aspirators, as previously described [27]. Engorged females were transferred into 30 cm³ cages, provided with a 10% sucrose solution on cotton bowls. The cages were covered with a wet black fabric and transported over the next day to the Medical Entomology Unit at the Institut Pasteur de Madagascar in Antananarivo, located a 6 h drive away from Andriba. Morphological identification on live mosquitoes was confirmed one by one using a dichotomous identification key adapted from Gjrebine [19], Fontenille [16], and Coetzee [12]. Females were transferred to a rearing cage $(32.5 \times 32.5 \times 32.5 \text{ cm BugDorm-4}^{\otimes} \text{-Bioquip, Rancho})$ Dominguez, CA, USA).

A beaker half-filled with dechlorinated water was placed inside the cage for egg collection and surveyed every day for 5 days. Each batch of eggs was treated for 10 min with a 0.01% formaldehyde solution and extensively rinsed with dechlorinated water for limiting potential contamination with micro-organisms such as microsporidia. Eggs were then transferred to larval rearing pans ($26 \times 26 \times 11$ cm (Gilac®), filled with 1L of dechlorinated tap water (water height ~ 1.5 cm). L1 and L2 larvae were fed on Tetramin Baby fish food (Tetra®), whereas L3 and L4 were fed on ground cat food. When pupae started appearing, they were counted every day and transferred to a small beaker containing dechlorinated water and placed in a novel rearing cage. A 10% sucrose solution-containing cotton

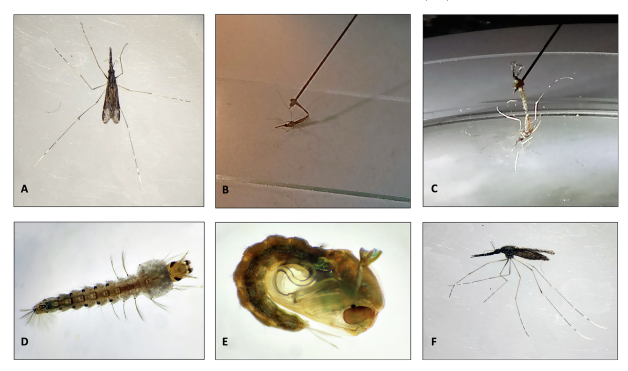


Figure 1. Illustrations of *An. coustani* forced copulation procedure and developmental stages. Female *An. coustani* (A and F). Forced copulation: the male is hooked on a dissecting needle and presented to a cold anesthetized female; decapitated male (B); non-decapitated male (C). Fourth instar larvae (D). Pupa (E).

bowl was placed on top of the rearing cage and changed every other day.

Insectary conditions

Larvae and adults were reared at 26.9 ± 0.7 °C, $77 \pm 7.7\%$ RH, with a 12:12 night and day light regimen.

Establishing the An. coustani colony

From the F1 adults obtained from wild F0 mosquitoes, the artificial mating (forced mating also called forced copulation) strategy of males and females was used. This followed the procedure initially described by Baker et al. [4], refined by Ow Yang et al. [29] and nicely documented by the MR4 staff [25]. Females were given a blood meal 24 h before forced copulation. Males and females were cold anesthetized just before the process as illustrated in Figure 1 panels B&C. Each male (3-5 days old) was used for fertilizing 2 females (5-10 days old). We found it unnecessary to decapitate males for copulation stimulation. After forced copulation, females were transferred to a novel rearing cage and provided a blood meal every other day to stimulate egg production. From egg production to adult emergence, the procedure described above was used and led to the successful production of eight generations between September 2019 and May 2020. Because of the general movement restriction imposed in Madagascar during the Covid-19 outbreak, the colony could not be maintained further than the F8.

Recorded rearing parameters

At each generation, the following parameters were recorded: number of females successfully fertilized and their egg production, L1 to pupae growing success, as well as larvae to adult developmental time.

Results and discussion

Establishing the first six generations of *An. coustani*

Starting with 31 gravid F0 wild females free to lay eggs in cage, we obtained 260 eggs, which is quite a low and challenging number (8.4 eggs/female). This number is actually very similar to previous results [27]. The 260 eggs were obtained from a unique gonotrophic cycle of the females as they refused to feed on blood provided through the Hemotek membrane feeding device. For the sake of the project, the females from each subsequent generation were fed on a rabbit multiple times.

As previous attempts revealed that *An. coustani* would not mate in cages, the production of each generation was performed by artificial copulation at the ratio of 1 male for 2 females. It was not necessary to decapitate the male to obtain efficient copulation and insemination of the females. Table 1 summarizes the number of males and females used for forced mating at each generation, as well as the number of blood meals that the mated females received, and the mean number of eggs laid per female at each generation. The mean number of eggs produced per female increased slightly between F1 and F5, despite a reduced

Table 1. Number of males and females used at each generation in the forced mating procedure and mean egg production by each mated female

Generation	Number of mosquitoes used for copulation		Number of blood meals after forced	Mean number of eggs per female	
	Males	Females	— mating		
F1	19	37	10	13.19	
F2	15	29	7	8.48	
F3	23	48	10	15.13	
F4	26	57	8	11.90	
F5	20	38	7	25.18	

number of blood meals. This is possibly due to the acquisition of better skills for performing the forced copulation technique. Accordingly, this led to an increase of total adults from F1 to F6 (Figure 2). The production of F7 and F8 was achieved; however, no data on egg and adult numbers were recorded due to the Covid-19 lockdown.

Bionomic features

Egg to adult production

As depicted in Table 2, each generation suffered a drastic reduction, with a mean survival rate from egg to adult of 36.3%, with a broad range from 24.2 to 62.2%. This latter value happened at generation F3, but no clue of this result could be identified. Nevertheless, the egg to adult survival rate increased slightly across the generation and was higher than previous results obtained by Jean Jose Nepomichene in 2017, on F1 only [21]. Except for the F3 generation, a high mortality rate occurred during the larval stages. Normalization of larval density and food amount might contribute to limit this rather high mortality rate. It is also striking that the mortality rate of pupae was quite high (18.7–44.7%). These mortality rates might be a consequence of poor larval development linked to restricted food availability and larval competition.

Larval to adult development

Previous work at a field station (Andriba, Madagascar) revealed that the larval to adult development of An. coustani F1 was quite asynchronous and long. Indeed, larval development lengthened more than 25 days, while An. arabiensis F1 population produced side by side would reach the adult stage in 10-15 days, as usually observed in insectary with tightly controlled parameters (Bourgouin et al., unpublished, 2016). Under the rearing conditions reported here, the larval to adult stage of An. coustani decreased across the generations and exhibited a different pattern of adult emergence (Figure 3). Indeed, F1 adults emerged in two peaks. The first peak occured between day 16 and day 18 from the day of egg laying and the second peak at day 20. From F2 to F4, adult emergence followed a 2-peak pattern, but each peak occurred earlier, except for the second peak of F2 that was at day 20, as for F1. Interestingly, F5 exhibited a broad single peak of adult emergence between day 11 and

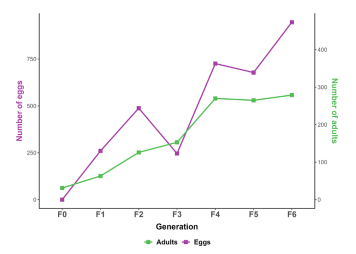


Figure 2. Yield in eggs and adults from F1 to F6. The graph represents the number of eggs obtained at each generation (Purple line and left scale) and the subsequent number of adults (green line and right scale).

day 17, while for F6, a sharp single peak of adult emergence centered at day 13. These changes in adult emergence patterns possibly reveal a trait of adaptation to colonization.

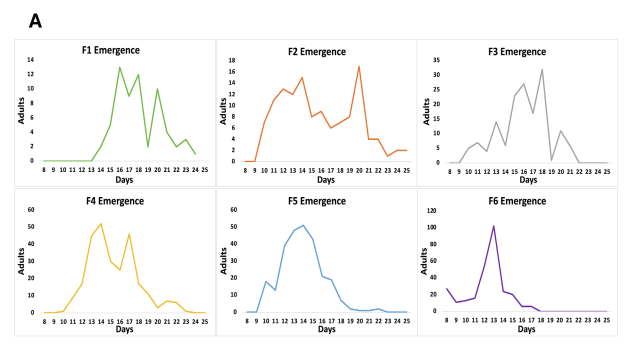
Concluding remarks

We report here the first successful estabishment of a colony of An. coustani. As this mosquito species would not mate in cages, each of the 8 generations produced resulted from forced mating. This is comparable to the Anopheles cracens colony recently established to the 6th generation by forced mating [1]. On the contrary, a novel Anopheles atroparvus colony was established to the 10th generation without forced mating [5]. Given that Anopheles coustani and An. atroparvus are from the Anopheles subgenus, while An. cracens belongs to the Cellia subgenus, no conclusions can be drawn as to whether species from the Anopheles subgenus would require forced copulation for colony establishment. Indeed, even a reference colony of Anopheles dirus (Cellia subgenus) is maintained by forced copulation at the MR4 repository (WRAIR2 strain, https://www.beiresources.org/Catalog/BEIVectors/MRA-700. aspx)

As maintaining a colony by forced copulation is highly demanding in human resources and skill, the biggest refinement would be to obtain a free-mating colony. Towards this goal, it would be of great interest to test whether the tools developed for establishing *An. darlingi* colonies would be beneficial, notably the use of light flashes to stimulate free copulation and shift in night temperature [2, 30, 37]. As discussed previously, additional refinements in the larval feeding regimen and larval density would possibly contribute to a decrease in the mortality rate during both larval and pupal stages. Overall, having at hand an *An. coustani* colony, whether free mating or maintained by forced copulation, as *An. dirus*, will provide an unvaluable tool to better understand the biology of this secondary malaria vector and assessing relevant malaria transmission control strategies.

Table 2. Summary of the raw data obtained across the production of the first six generation of *An. coustani* starting with 31 wild gravid females producing 260 eggs.

Parental generation	Number of mated females	Number of eggs	Number of pupae	Number of adults	Egg to pupa mortality (%)	Pupa to adult mortality (%)	Egg to adult survival (%)
F0	31						
F1	37	260	114	63	56.2	44.7	24.2
F2	29	488	155	126	68.2	18.7	25.8
F3	48	246	221	153	10.2	30.8	62.2
F4	57	726	420	270	42.1	40.0	37.2
F5	38	678	nd	265	nd	nd	39.1
F6	nd	957	nd	279	nd	nd	29.2



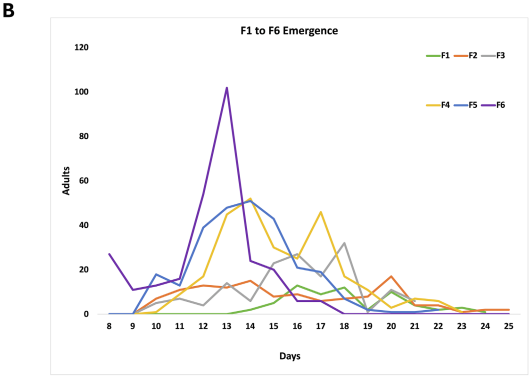


Figure 3. Graphic representation of *An. coustani* adult emergence at each generation. (A) Individual graphs showing the pattern of emergence of F1 to F6. (B) Aggregated graphs highlighting the clear changing patterns of adult emergence from F1 to F6 both in mode and date of the peak of emergence. The scale of the abscissa axis corresponds to days till the first appearance of L1.

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Conflict of interest

The authors declare that there are no competing interests.

Author contribution

Conceived the project: TMA, CB and RG. Designed the experiments: TMA, NP, CB. Performed the experiments: TMA, FTR, MRA, MRR and HJVS. Analyzed the data: TMA and CB. Drafted the paper: TMA and CB. Reviewed and edited the manuscript: TMA, RG and CB. All authors contributed to the article and approved the submitted version.

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