A conserved female-specific larval requirement for MtnB function facilitates sex separation in multiple species of disease vector mosquitoes

Keshava Mysore1,2, Longhua Sun1,2, Joseph B. Roethele1,2, Ping Li1,2, Jessica Igiede1,2, Joi K. Misenti1,2 and Molly Duman-Scheel1,2*

Abstract

Background: Clusters of sex-specific loci are predicted to shape the boundaries of the M/m sex-determination locus of the dengue vector mosquito Aedes aegypti, but the identities of these genes are not known. Identification and characterization of these loci could promote a better understanding of mosquito sex chromosome evolution and lead to the elucidation of new strategies for male mosquito sex separation, a requirement for several emerging mosquito population control strategies that are dependent on the mass rearing and release of male mosquitoes. This investigation revealed that the methylthioribulose-1-phosphate dehydratase (MtnB) gene, which resides adjacent to the M/m locus and encodes an evolutionarily conserved component of the methionine salvage pathway, is required for survival of female larvae.

Results: Larval consumption of Saccharomyces cerevisiae (yeast) strains engineered to express interfering RNA corresponding to MtnB resulted in target gene silencing and significant female death, yet had no impact on A. aegypti male survival or fitness. Integration of the yeast larvicides into mass culturing protocols permitted scaled production of fit adult male mosquitoes. Moreover, silencing MtnB orthologs in Aedes albopictus, Anopheles gambiae, and Culex quinquefasciatus revealed a conserved female-specific larval requirement for MtnB among different species of mosquitoes.

Conclusions: The results of this investigation, which may have important implications for the study of mosquito sex chromosome evolution, indicate that silencing MtnB can facilitate sex separation in multiple species of disease vector insects.

Keywords: Aedes aegypti, Aedes albopictus, Anopheles gambiae, Culex quinquefasciatus, Sex, Larvae, Methionine, Saccharomyces cerevisiae, Yeast, Larvicide, RNAi

Background

Blood-feeding vector mosquitoes transmit disease-causing pathogens that lead to hundreds of thousands of human deaths each year. For example, Zika, yellow fever, chikungunya, and dengue result from infections with arboviruses transmitted through the bites of Aedes mosquitoes, including Aedes aegypti and Aedes albopictus [1].
Given poor progress in vaccine development and distribution, mosquito control is the primary mechanism for prevention of these and other mosquito-borne illnesses. Unfortunately, the emergence of insecticide resistance, concerns for the negative impacts of pesticides on the environment, and a lack of support for mosquito control programs compromise current strategies for managing mosquitoes [1].

Female mosquitoes possess sex-specific morphological, physiological, and behavioral traits, such as blood-feeding behavior, that promote the spread of disease-causing pathogens. Although genes that regulate female-specific mosquito development may represent novel targets for vector control, a majority of these genes have not yet been identified [2]. Discovery and functional characterization of female-specific genes in multiple mosquito species could reveal mechanisms that regulate female mosquito development, provide insight into the evolution of these mechanisms, and promote the development of novel mosquito control interventions. For example, gene drive strategies that involve the manipulation of sex-specific genes are being developed and characterized [3–5]. Moreover, renewed interest in the sterile insect technique (SIT), which involves the release of sterile male insects to reduce insect pest populations [6], has recently emerged in the mosquito control community [7, 8]. Broad use of SIT for mosquito control has been hampered by the need to remove female mosquitoes, which is required to reduce health and nuisance biting risks posed by the release of female mosquitoes. Removal of females is also believed to increase the efficacy of this intervention, while also reducing costs [7, 8]. Likewise, the Wolbachia-based incompatible insect technique (IIT), which is proving useful for mosquito population suppression [9] and can be paired with SIT [10, 11], also requires mass rearing and release of adult male mosquitoes [7, 8]. Transgenic-based population suppression approaches involving the release of insects carrying a dominant lethal (RIDL) [12, 13], as well as various emerging gene drive technologies to block pathogen transmission or to suppress mosquito populations [14], also involve the release of male mosquitoes [5]. Thus, the development and implementation of effective and affordable sex-separation technologies that can be deployed worldwide are rate-limiting steps in the global introduction of numerous population-based mosquito control methods [8]. The identification of genes with sex-specific functions could permit the elucidation of novel male sex-sorting technologies that could support these interventions.

*Aedes aegypti* sex determination is regulated by a non-recombining Y-chromosome-like male-determining region, the M locus, which is present on chromosome 1 [15, 16] and contains the male-determining factor *NIX* [17]. Males, which possess one copy of the chromosome bearing the M locus and one which lacks it, have an M/m genotype, while *A. aegypti* females, which lack the male-determining locus, have an m/m genotype [18]. Clusters of loci that cause sex-specific lethal effects are believed to reside in the vicinity of the sex-determining region [19, 20]. However, the identities of these genes are not known. Characterization and sequencing of the M/m locus region had been thwarted by the repetitive nature of DNA located in this centromeric region of chromosome 1. However, recent innovations in sequencing technology generated an improved and re-annotated genome assembly that facilitated better estimation of the M/m locus, providing opportunities to study the evolution of the sex-determining chromosomes and prospects for evaluating the functional contributions of genes linked to the M/m locus [21]. In this investigation, methylthioribulose-1-phosphate dehydratase (*MtnB*), which resides adjacent to the M/m locus, was identified as a female-specific larval lethal locus.

*MtnB*, an evolutionarily conserved component of the methionine salvage pathway, functions to convert 5-methylthiadenosine (MTA) to methionine. This process occurs through six enzymatic reactions, the third of which, the dehydration of 5-methylthioribulose-1-phosphate (MTRu-1-P) to 2,3-diketo-5-methylthiopentyl-1-phosphate (DK-MTP-1-P), is catalyzed by *MtnB* [22–25]. This pathway facilitates recycling of MTA, a by-product of polyamine synthesis, back to methionine, an essential amino acid, while also serving as a useful means of recycling the sulfur present in MTA [26]. The methionine salvage pathway has also been implicated in cell death and inflammation [26–29]. In this investigation, RNAi-based silencing of *MtnB* in *A. aegypti* revealed a female-specific requirement for this gene in larvae that was exploited for the development of a scalable yeast-based oral RNAi method for production of fit adult males. Characterization of *MtnB* in several additional species of disease vector mosquitoes, including *Aedes albopictus* (vector of multiple arboviruses), *Anopheles gambiae* (malaria parasite vector), and *Culex quinquefasciatus* (lymphyatic filariasis and West Nile viral vector) [1], suggested that targeting *MtnB* during larval development could facilitate male sex sorting in multiple species of disease vector mosquitoes.

**Methods**

**Mosquito rearing**

The *A. aegypti* LVP-IB12, *A. albopictus* Gainesville, *A. gambiae* G3, and *C. quinquefasciatus* JHB (BEI Resources, NIAID, NIH) strains were reared as described [30], except that sheep blood (purchased from HemoStat Resources, NIAID, NIH) strains were reared as described [30].
Laboratories, Dixon, CA, USA) was provided with an artificial membrane feeding system (Hemotek Limited, Blackburn, UK).

**Larval soaking**

Custom small interfering RNAs (siRNAs) corresponding to mosquito *MtnB* genes (Table 1) were selected using the Integrated DNA Technologies (IDT) Custom Dicer-Substrate siRNA (DsiRNA) tool [31]. Custom siRNAs (see Table 1 for further information), as well as a control siRNA lacking a known target in mosquitoes [32], were purchased from IDT (Coralville, IA, USA) for use in larval soaking screening experiments that were performed in duplicate using the method of Singh et al. [33]. For each soaking experiment, 20 first instar larvae (L1) were soaked in 0.5 µg/µl control or experimental siRNA (Table 1) for 4 h and then reared as described [34]. Soaking data were statistically evaluated using the chi-squared test to identify statistically significant differences between the observed and expected survival of female and male mosquitoes, assuming expected 1 male:1 female ratios on the basis of laboratory observations of the mosquito-strains when reared in the laboratory insectary.

**Production of yeast interfering RNA larvicides**

Yeast strains were prepared as described [35] through cloning of custom shRNA expression cassettes corresponding to target sequences of interest (Table 1; synthesized by Invitrogen, Carlsbad, CA, USA) into the pRS426 GPD shuttle vector [36]. After confirming the clones through restriction digestion and sequencing, the resulting plasmids were used to transform *S. cerevisiae* CEN. PK yeast (genotype MATa/α ura3-52/ura3-52 trp1-289/trp1-289 leu2-3_112/leu2-3_112 his3 Δ1/his3 Δ1 MAL2-8C/MAL2-8C SUC2/SUC2 [37]. Transformants were selected based on growth on minimal media lacking uracil. shRNA expression was confirmed through PCR conducted as described [38] using the following primers: forward 5′-CAG GAT ATT CTT TCA TAG CAT GTT C-3′ (specific to the 3′ end of the *MtnB* hairpin) and reverse primer 5′-TCC TCT TCT TCT CGGG TAG AGC-3′ (which corresponds to the terminator sequence). An image of the PCR products visualized with ethidium bromide staining (Additional file 1: Fig. S1) was edited using Adobe Photoshop 2021 software (to remove non-relevant lanes and to label the figure). Following strain confirmation, dried inactivated yeast interfering RNA was prepared for use in larvicide assays as described [35].

**Yeast interfering RNA larvicide screening**

Yeast interfering RNA larvicides were evaluated through larvicide assays that conformed to the World Health Organization (WHO) [39] guidelines, which were performed as previously described [35]. In summary, 20 L1 larvae were placed in 500 ml plastic cups containing 50 ml of distilled water and 50 mg of dried inactivated yeast (either experimental larvalid yeast or yeast prepared from a control interfering RNA strain [34]). The emergence rates and sexes of the resulting adults were recorded in each of nine replicate trials that were performed for each control and larvicalid yeast treatment. Data were evaluated using the chi-squared test to identify statistically significant differences between the observed and expected 1 male:1 female ratios, female survival, and male survival for each treatment.

**qRT-PCR assays**

*MtnB* silencing was confirmed through qRT-PCR assays that were performed as described [40]. In summary, TRIzol Reagent (Invitrogen, Carlsbad, CA, USA) was used according to the manufacturer’s instructions to prepare total RNA from 20 pooled third instar larvae that had fed on either control or *MtnB*.496 yeast throughout larval development. cDNA was prepared from total RNA extracts using the High Capacity RNA to cDNA Kit (Applied Biosystems, Foster City, CA, USA) according to the manufacturer’s instructions. Real-time PCR quantification assays were performed using an Applied

---

**Table 1** siRNA target sequences, corresponding genes, and resulting male:female sex ratios following treatments

| siRNA | Target sequence | Corresponding genes | Species | siRNA soaking treatment | Male mortality | Female mortality |
|-------|-----------------|---------------------|--------|-------------------------|---------------|-----------------|
| 496   | GAA CAU GUA AAG AAU AUC CUG | AAEL011830 | A. aegypti | 23:12 (40) | 1.0000 | 0.0033 |
| 529   | GCAUCAAGGCUUAGA AAG AAA UUA | GAPW0103631.1, Aa-53178 mRNA | A. albopictus | 10:4 (20) | 1.0000 | 0.0108 |
| 523   | CGUGGAUGCAUAAUCAUAGA AAG | AGAP000470 | A. gambiae | 28:12 (40) | 1.0000 | 0.0033 |
| 534   | AGAAAUUGUGUAGAUGAUGUGCA | CPU011357 | C. quinquefasciatus | 25:7 (40) | 1.0000 | 0.0001 |

siRNAs and corresponding target sequences/genomes in the indicated species, and the altered male:female ratios resulting from siRNA soaking treatments are indicated (P values correspond to Fisher’s exact tests)
Two hundred L1 LVP-IB12 larvae were placed in 34 × 25 × 4 cm trays (1426B, Bioquip, Rancho Dominguez, CA, USA) bearing 1 L of distilled water. Per Zhang et al. [42], larvae were cultured using a mass-rearing diet that consisted of a slurry of 250 mg bovine liver powder (MP Biomedicals, Santa Ana, CA, USA) and 150 mg of ground shrimp (Tetra GMBH, Melle, Germany) mixed with 10 ml water, but with the brewer's yeast component of the diet [42] replaced with yeast interfering RNA. The yeast interfering RNA component (control or Mtn.496) was combined with the liver powder and shrimp slurry and fed to larvae as follows: 100 mg at L1, 100 mg at L2, 100 mg at L3, and 200 mg at L4. Each day, larval trays were examined for pupae, which were removed, sorted by sex, and counted. Mortality and sex ratio distortions were evaluated in three biological replicate trials, each with two replicate trays of larvae fed using diets prepared with control interfering RNA or MtnB.496 yeast. Statistically significant differences between the observed and 1:1 expected survival levels for males and females were evaluated using the chi-squared test. The fitness of surviving males was ascertained through comparisons of wing lengths, which were measured as described [43]. Wing length data were evaluated with Student's t-test.

Results
Silencing MtnB during A. aegypti larval development selectively kills female mosquitoes

Given that the M/m locus is believed to be tightly linked to developmental genes that confer sex-specific effects in A. aegypti, MtnB, which is located on chromosome 1 and flanks the M/m locus [21, 44], was hypothesized to function as a potential sex-specific lethal gene. MtnB was silenced through larval soaking with siRNA #496, which corresponds to the MtnB transcript. Treatment with siRNA #496 resulted in significant female mortality yet had no significant impact on male survival (further details, including P values, are provided in Table 1). Given the results of this brief siRNA soaking-mediated transient gene silencing assay, S. cerevisiae, which was recently developed as a system for expression and sustained delivery of interfering RNA to mosquito larvae, was used to express a short hairpin RNA (shRNA) corresponding to siRNA #496. shRNA expression in this yeast strain, hereafter referred to as MtnB.496, was confirmed through PCR amplification of cDNA corresponding to the shRNA transcript (Additional file 1: Fig. S1). Dried inactivated MtnB.496 yeast was fed to larvae throughout larval development. Larvae were also reared on yeast that expresses an shRNA with no known target in mosquitoes [34] for comparison. MtnB transcripts, which are detected during larval and pupal development (Fig. 1a), are silenced in larvae that consume MtnB.496 yeast (Fig. 1b; 88 ± 4% reduction in transcript levels in comparison to larvae reared on control interfering RNA yeast; t-test, P < 0.001). Larval consumption of MtnB.496 yeast,
but not control interfering RNA yeast, resulted in significantly higher male:female ratios among the surviving mosquitoes (Fig. 1c; $\chi^2 = 31.387, P < 0.0001$). Although no significant impact on male survival was observed following consumption of MtnB.496 yeast ($\chi^2 = 3.214, P > 0.05$), only 13 ± 2% of expected adult female mosquitoes survived, with a majority of animals dying as late larvae or early pupae ($\chi^2 = 128.571, P < 0.0001$).

Scaled production of fit adult *A. aegypti* male mosquitoes

To confirm that the impacts of MtnB.496 larvicide are specific to female larvae, life history traits were assessed in adult male mosquitoes that had consumed the yeast throughout larval development. The percentage of treated males that mated with wild-type females was not significantly different than adult males that were reared on control yeast interfering RNA larvicides (Fig. 2a, $t = -1.925, P > 0.05$) or fecundity were observed (Fig. 2b, $t = 0.698, P > 0.05$). These findings indicated that MtnB.496 yeast larvicide treatment did not produce any detectable effects on male fitness in the assays conducted.

To assess whether the RNAi-based yeast larvicides could also facilitate scaled production of males, a larval diet and feeding regimen employed at *Aedes* mass-rearing facilities [42] was modified through replacement of the nutritional yeast component of the diet with dried inactivated yeast larvicide MtnB.496. The modified diet was tested on *A. aegypti* larvae grown in mass-rearing trays. Under these conditions, the larvicides continued to induce significant *A. aegypti* female mortality (Fig. 2c, $\chi^2 = 1275.764, P < 0.0001$) with no impacts on male survival (Fig. 2c, $\chi^2 = 0.5777, P > 0.05$). The fitness of male survivors, which was estimated through measurement of wing lengths, a correlate of body size and
fitness, was not significantly different than that of males reared on the standard mass-rearing diet (Fig. 2d, \(t = 1.031, P > 0.05\)).

**MtnB orthologs are female-specific lethal genes in multiple species of disease vector mosquitoes**

*MtnB* orthologs have been identified in *A. gambiae* and *C. quinquefasciatus*, and BLAST searches identified an ortholog in *A. albopictus* (Table 1). RNAi-mediated targeting of these genes through brief soaking treatments in siRNAs #529 (corresponds to *A. albopictus* *MtnB*), #523 (corresponds to *A. gambiae* *MtnB*), and #534 (corresponds to *C. quinquefasciatus* *MtnB*) resulted in significant female-specific larval death, yet did not significantly impact male survival (Fig. 2c, \(P > 0.05\) vs. mass-rearing diet, chi-squared test; data were compiled from four replicate containers per condition, each bearing 500 larvae) or wing size (Fig. 2d, \(P > 0.05\) vs. mass-rearing diet, t-test; \(n = 50\) wings/treatment), which correlates with fitness. Error bars represent SEM in all panels. NSD: Not significantly different

Larval consumption of *MtnB* by *A. albopictus* larvae resulted in significantly higher than expected male:female ratios (Fig. 3a; \(\chi^2 = 24.202, P < 0.0001\)). Only 24±1% of expected *A. albopictus* female mosquitoes survived (Fig. 3a, \(\chi^2 = 101.829, P < 0.00001\)), but no significant impact on male survival was observed (Fig. 3a, \(\chi^2 = 0.004, P > 0.05\)). Likewise, *MtnB* resulted in significantly higher male:female ratios among surviving *A. gambiae* adults (Fig. 3b; \(\chi^2 = 7.453, P < 0.01\)). Although *MtnB* had no significant impact on the survival of males through adulthood (\(\chi^2 = 0.004; P > 0.05\)), only 43±2% of expected *A. gambiae* female mosquitoes survived (Fig. 3b, \(\chi^2 = 37.551, P < 0.0001\)). Finally,
significantly higher than expected male:female ratios were also observed among *C. quinquefasciatus* adult survivors following larval consumption of MtnB.534 (Fig. 3c; \(\chi^2 = 28.751, P < 0.0001\)), a difference that coincided with only 21 ± 1% of expected *C. quinquefasciatus* adult emergence (Fig. 3c; \(\chi^2 = 95.890; P < 0.00001\)), with no significant corresponding changes in male survival (Fig. 3c; \(\chi^2 = 3.051, P > 0.05\)).

**Discussion**

**Implications for understanding the evolution of sex chromosomes in mosquitoes**

Sex chromosome evolution is believed to occur in several stages [45–47], starting with a homologous pair of autosomes that acquire sex-determining loci. A proto-Y chromosome, which has a male fertility locus (M) and a dominant female suppressor locus (Su\(^y\)), as well as a proto-X chromosome carrying a female fertility locus (F) and a male sterility locus (m) form. Suppressed recombination in the sex-determining region evolves, eventually spreading along the proto-sex chromosomes, which accumulate transposable elements and non-coding sequences, undergo chromosomal rearrangements, and accumulate sexually antagonistic genes with different alleles that differentially benefit either males or females. The proto-sex chromosomes eventually evolve into heteromorphic X and Y chromosomes, with the Y chromosome eventually degenerating and reducing in size [45–47].

Sex chromosomes in different mosquito species have progressed to varying stages of this evolution. Culicines, including *Aedes* and *Culex* mosquitoes, possess homomorphic sex chromosomes in which the sex-determining alleles are linked to the smallest homomorphic first chromosome [48]. In *A. aegypti*, the homomorphic sex chromosomes have evolved into proto-sex chromosomes bearing a sex determining M/m locus [15, 16]. The M locus at the proto-Y chromosome contains the male-determining factor, *Nix* [17], which directs male-specific splicing of the sex chromosome gene *doublesex (dsx)*. In females, which have two copies of the proto-X (m) chromosome, the female-specific Dsx\(^x\) form regulates ovary development and fertility [43]. Although a sex-differentiated region of suppressed recombination extends ~100 Mb beyond the M/m locus [21, 49, 50], heteromorphic X and Y chromosomes have not yet evolved in *A. aegypti*. Likewise, *A. albopictus* also possesses homomorphic sex chromosomes, and the *A. albopictus* *Nix* gene was identified as a strong candidate for a male-determining factor in this species [17, 51, 52]. In contrast, *A. gambiae* has already evolved separate X and Y chromosomes,
with the Yob gene functioning as a male-determining factor on the Y chromosome [53]. Despite different mosquito species having advanced to various stages of sex chromosome evolution, this investigation functionally verified that MtnB is required for survival of female larvae in A. aegypti (Figs. 1c, 2c), A. albopictus (Fig. 3a), A. gambiæ (Fig. 3b), and C. quinquefasciatus (Fig. 3c) mosquitoes. These findings suggest that loci which cause sex-specific lethal effects and shape the boundaries of the sex-determination locus were accumulating on the sex chromosomes prior to separation of the culicine and anopheline mosquitoes. Future studies will be directed toward understanding the nature of this female-specific requirement during larval development.

Krzywinska et al. [20] discuss possible mechanisms underlying the sex-specific lethality they observed in A. aegypti that had inherited sex chromosomes produced through male meiotic recombination events that occurred between the sex-determination locus and a linked enhanced green fluorescent protein (EGFP)-marked transgene. An explanation which is consistent with the MtnB silencing results presented here proposed that such recombination events could lead to the loss of a haploinsufficient sex chromosome gene for which two copies are required in m/m females. If MtnB is haploinsufficient in females, any female that inherits a recombinant m chromosome lacking a copy of the MtnB gene will die, an event that is sometimes referred to as “suppression of recombination” because resulting female offspring are not observed in adult populations. Likewise, as observed in this investigation, silencing MtnB using RNAi-based yeast larvicides can also result in female death. These silencing results suggest that the presence of haploinsufficient genes adjacent to the sex-determination locus may have helped to shape the boundaries of the sex-determination region during mosquito sex chromosome evolution, as previously suggested [20].

Sex-separation applications
Papathanos et al. [54] recommend that sex-sorting methodology should be simple, stable, and produce competitive males, ideally through female elimination during development. The yeast female-specific larvicide system described herein meets these criteria (Fig. 2). It is also recommended that the system be cost-effective, scalable, and suitable for implementation in remote or resource-limited regions of the world [54], criteria that would be met through the use of yeast, which is routinely produced at commercial scale by the food industry for worldwide distribution. Moreover, the demonstrated ability to seamlessly incorporate yeast into a standard mass-rearing diet (Fig. 2c) makes this system simple to implement.

Sex-sorting technology should meet regulatory standards, ideally requiring no modification of existing regulatory protocols that may have already been acquired for the mosquito technology under development. It is also important that the technology for elimination of females be conditional, so as not to interfere with strain maintenance [54]. The yeast sex-sorting system meets these criteria, as it induces no permanent changes to the mosquito genome, but acts through conditional gene silencing during female development. The use of yeast RNAi-based larvicides would therefore circumvent the need to further genetically manipulate existing mosquito strains developed for population control strategies. Furthermore, the yeast would be used only during the larval stages and would be heat-killed prior to use indoors at mass-rearing facilities. This may mitigate a need for modification of existing regulatory permits previously obtained for male mosquito releases, a prospect that can be further discussed with the appropriate regulatory bodies. Thus, sex-specific yeast interfering RNA technology meets many of the criteria for a useful sex-separation system [54]. Future studies will be directed toward scaled industrial-sized production of yeast interfering RNA larvicides for distribution of yeast to mass-rearing mosquito facilities worldwide.

In this investigation, replacement of the nutritional yeast component of a larval mass-rearing diet [42] resulted in female-specific larval killing, facilitating production of fit male mosquitoes at a ratio of 5 male:1 female mosquitoes (Fig. 2d). Silencing MtnB did not eliminate all females (Fig. 2b, c) and this yeast strain could not likely be used as a stand-alone strategy for male mosquito production. It would be interesting to explore methods for further improving the female-killing capacities of the yeast strains (i.e. increased shRNA expression levels, stability, or uptake). We hope to explore this in future studies, as a stand-alone strategy that generates >99% male separation, as observed through use of a sophisticated camera system recently developed for A. aegypti [9], would be preferable. Although the automated camera system is highly effective, it is not yet clear if such technology can be deployed worldwide or adapted for use is all mosquito species. In such instances, and particularly when resources are limited, replacing nutritional yeast with larvicial yeast during the mass-rearing process could be used in combination with existing sex-separation technologies, particularly manual separation strategies, which are highly labor-intensive [8].

Female-specific yeast larvicide technology could particularly benefit the development and implementation of population-based control strategies for C. quinquefasciatus (Fig. 3c), in which sex-sorting methods other
than physical separation have yet to be established [7, 8]. The female-specific mortality levels observed in the current investigation were relatively lower in *A. gambiae* with respect to the other mosquitoes evaluated (Fig. 3). However, given the conservation of *MtnB* in many species of *Anopheles* mosquitoes [44], it may nevertheless still be worth exploring the potential for using *MtnB* silencing to sex other *Anopheles* species. For example, the importance of developing sex-separation strategies that could be implemented in malaria vectors such as *Anopheles arabiensis* [55] and *Anopheles stephensi* [56] has been discussed [44], and *MtnB* is conserved in both species, as well as several other anophelines. It may therefore be helpful to pair a yeast larvicide diet with the manual or visual sorting technologies that capitalize on the conserved female-specific larval requirement for *MtnB* for using several species, including female lethal strains or transgenic sex-sorting strains that have been generated for *Anopheles* mosquitoes [7]. One could also envision pairing the RNAi yeast diet sex-separation approach with various transgenic sex-sorting strains that have been generated for several species, including female lethal strains or transgenic fluorescent marker strains [57, 58] that facilitate automated sorting. Likewise, yeast feeding could be combined with imaging-based sorting strategies [9] and automated pupal size estimators [59], or paired with feeding of toxicant-infused blood meals to females [60]. Such combined approaches may better permit the >99% removal of females that is generally accepted as a minimal requirement for male-only release control strategies [7].

**Conclusions**

These studies have uncovered a conserved female-specific larval requirement for *MtnB*, which encodes a component of the methionine salvage pathway, in distantly related mosquitoes. The results of this investigation may have important implications for the study of mosquito sex chromosome evolution, and it will be interesting to further investigate this topic in future studies. It is anticipated that female-specific yeast interfering RNA technologies that capitalize on the conserved female-specific requirement for *MtnB* function in mosquito larvae could permit mass production of fit males, thereby facilitating emerging population-based strategies for control of multiple disease vector mosquito species.

**Abbreviations**

DK‑MTP‑1‑P: 2,3‑Diketo‑5‑methylthiopentyl‑1‑phosphate; MTA: 5‑Methylthioadenosine; MTRu‑1‑P: 5‑Methylthioribulose‑1‑phosphate; Dxx: doublesex; L1: First instar larvae; IDT: Integrated DNA Technologies; LVP‑IB12: Liverpool‑IB12; MtnB: Methylthioribulose‑1‑phosphate dehydratase; shRNA: Short hairpin RNA; siRNA: Small interfering RNA.

**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1186/s13071-021-04844-w.

**Additional file 1: Fig. S1.** Confirmation of shRNA expression in recombinant yeast strain MtnB.496. PCR reactions performed with primers corresponding to the MtnB.496 shRNA transcript generated a ~100 bp amplicon (see DNA standard at far left; cDNA template was prepared from MtnB.496 yeast total RNA). Negative control PCR reactions included an amplification with cDNA prepared from non-transformed yeast (marked by — sign) and a reaction with no cDNA added (marked by knot symbol). A representative ethidium bromide‑stained agarose gel from one of two comparable biological replicate experiments is shown.

**Acknowledgements**

We thank Na Wei and Chien‑Wei Wang for their suggestions on yeast strain production and Zhiyong Xi for his recommendations on preparation of the larval mass-rearing diet. We thank members of the lab for useful discussions.

**Authors’ contributions**

MDS conceived of the study. KM, PL, JR, and MDS developed methodology. KM, LS, PL, JR, JI, JKM and MDS performed the investigation. KM and MDS supervised the research and administered the project. MDS acquired funding for this research. All authors read and approved the final manuscript.

**Funding**

This work was funded by NIH‑NIAID Awards 1 R21 AI144256‑01 and R21AI156170‑01 to MDS. The funders were not involved in study design, data collection and interpretation, or the decision to submit the work for publication.

**Availability of data and materials**

All data generated or analyzed during this investigation are included in this manuscript. Yeast strains and corresponding plasmids generated during the course of this study will be made available upon satisfactory completion of a material transfer agreement with Indiana University and following procurement of required import permits by the requesting party.

**Declarations**

**Ethics approval and consent to participate**

Not applicable. No vertebrate animals or human subjects were used in this research investigation.

**Consent for publication**

Not applicable.

**Competing interests**

MDS was named as the inventor on U.S. patent application number 62/751,052. The pending application did not impact her data interpretation or decision to publish this work. All other authors declare that they lack competing interests.

**Author details**

1Department of Medical and Molecular Genetics, Indiana University School of Medicine, Raclin‑Carmichael Hall, 1234 Notre Dame Ave., South Bend, IN 46617, USA. 2University of Notre Dame Eck Institute for Global Health, Notre Dame, IN, USA.

Received: 31 March 2021 Accepted: 11 June 2021

Published online: 26 June 2021
References

1. Centers for Disease Control CDC. Mosquitoes and diseases. A-Z. https://www.cdc.gov/mosquitoes/about/diseases.html. Accessed Jan 2020.

2. Duman-Scheel M, Syed Z. Developmental neurogenetics of sexual dimorphism in Aedes aegypti. Front Ecol Evol. 2015;3:61.

3. Kyrou K, Hammond AM, Galizi R, Kranjc N, Burt A, Beaghton AK, et al. A CRISPR-Cas9 gene drive targeting doublesex causes complete population suppression in caged Anopheles gambiae mosquitoes. Nat Biotechnol. 2018;36(11):1062–6.

4. Simon J, Hammond AM, Beaghton AK, Galizi R, Taxarchi C, Kyrou K, et al. A male-biased sex-distorter gene drive for the human malaria vector Anopheles gambiae. Nat Biotechnol. 2020;38(9):1054–60.

5. Adelman ZN, Tu Z. Control of mosquito-borne infectious diseases: sex and gene drive. Trends Parasitol. 2016;32(3):219–29.

6. Knipling EF. Possibilities of insect control or eradication through the use of sexually sterile males. J Econ Entomol. 1955;48:459–62.

7. Lutrat C, Giesbrecht D, Marois E, Whyard S, Baldet T, Bouyer J. Sex determination. A male-determining factor in the mosquito Aedes aegypti (L.). Genetica. 1978;48(2):137–43.

8. Papathanos PA, Bourtzis K, Tripet F, Bossin H, Virginio JF, Capurro ML, et al. Lethal genes on the sex chromosomes concealed in a population of Aedes aegypti. Front Ecol Evol. 2015;9(7):e0003864.

9. Crawford JE, Clarke DW, Criswell V, Desnoyer M, Cornel D, Deegan B, et al. Improved reference genome of Aedes aegypti L. Genetica. 1990;46(1):49–66.

10. Zheng X, Zhang D, Li Y, Yang C, Wu Y, Liang X, et al. Incompatible and sterile insect techniques combined eliminate mosquitoes. Nature. 2019;572(7767):56–61.

11. Kittayapong P, Ninphanomchai S, Limohpasmanee W, Chansang C, et al. Sex sorting for pest control: It’s raining men! Trends Parasitol. 2010;26(10):485–92.

12. Carvalho DO, McKemey AR, Garziera L, Lacroix R, Donnelly CA, Alphey L. A male-biased sex-distorter gene drive for the human malaria vector Anopheles gambiae. Nat Biotechnol. 2020;38(9):1054–60.

13. Hall AB, Basu S, Jiang X, Qi Y, Timoshevskiy VA, Biedler JK, et al. Sex determination. A male-determining factor in the mosquito Aedes aegypti (L.). Genetica. 1978;48(2):137–43.

14. Newton ME, Wood RJ, Southern DI. Cytological mapping of the M and D loci in the mosquito, Aedes aegypti (L.). Genetica. 1978;48(2):137–43.

15. Krzywinski J. The sex locus is tightly linked to factors conferring sex-specific lethal effects in the mosquito Aedes aegypti. Heredity. 2010;105(1):46–56.

16. Albers E. Metabolic characteristics and importance of the universal methionine salvage pathway compound 4-methylthio-2-oxobutanate causes apoptosis independent of down-regulation of ornithine decarboxylase. Biochem Pharmacol. 2006;72(7):806–15.

17. Hill CL, Sharma A, Shouche Y, Severson DW. Dynamics of midgut microbial communities in mosquito larvae. Methods Mol Biol. 2019;1858:213–31.

18. Mumberg D, Muller R, Funk M. Yeast vectors for the controlled expression of heterologous proteins in different genetic backgrounds. Gene. 1995;156(1):119–22.

19. van Dijken JP, Bauer J, Brambilla L, Duboc P, Francois JM, Gancedo C, et al. An interlaboratory comparison of physiological and genetic properties of four Saccharomyces cerevisiae strains. Enzyme Microb Technol. 2000;26(9–10):706–14.

20. Mysore K, Hapairai L, Sun L, Li P, Emrich SJ, Severson DW, et al. Preparation and use of a yeast shRNA delivery system for gene silencing in mosquito larvae. Methods Mol Biol. 2019;1958:213–31.

21. Hill CL, Sharma A, Shouche Y, Severson DW. Dynamics of midgut microflora and dengue virus impact on life history traits in Aedes aegypti. Acta Trop. 2014;140:151–7.

22. Zhang D, Zhang M, Wu Y, Giles JL, Yamada H, Wu Z, et al. Establishment of a medium-scale mosquito facility: optimization of the larval mass-rearing unit for Aedes albopictus (Diptera: Culicidae). Parasit Vectors. 2017;10(1):569.

23. Hill CL, Sharma A, Shouche Y, Severson DW. A targeted approach to the identification of candidate genes determining susceptibility to Plasmodium gallinaceum in Aedes aegypti. Mol Genet Genomics. 2003;269(6):735–64.

24. Pirkov I, Norbeck J, Gustafsson L, Albers E. A complete inventory of all enzymes in the eukaryotic methionine salvage pathway. FEBS J. 2008;275(16):4111–20.

25. Mary C, Duek P, Salleor L, Tienz P, Burmann D, Bairoch A, et al. Functional identification of APIP as human mtrn8, a key enzyme in the methionine salvage pathway. PLoS One. 2012;7(12):e28877.

26. Albers E. Metabolic characteristics and importance of the universal methionine salvage pathway recycling methionine from 5'-methylthioadenosine. IUBMB Life. 2009;61(12):1132–42.

27. Avila MA, Garcia-Trevijano ER, Lu SC, Corrales FJ, Mato JM. Methylthioadenosine. Int J Biochem Cell Biol. 2004;36(11):2125–30.

28. Wang X, Baxi V, Murphy M, Kruger ME, Kruger ME. The methionine salvage pathway compound 4-methylthio-2-oxobutanate causes apoptosis independent of down-regulation of ornithine decarboxylase. Biochem Pharmacol. 2006;72(7):806–15.

29. Li C, Li Z, Stetten E, Arnesano F, Losacco M, Natiello G, et al. Methionine can favor DNA platination by trans-coordinated platinum antitumor drugs. Angew Chem Int Ed Engl. 2009;48(45):8497–500.

30. Clemons A, Mori A, Haugen M, Severson DW, Duman-Scheel M. Culturing and egg collection of Aedes aegypti. Cold Spring Harb Protoc. 2010;2010(10):pdbprot5507.

31. IDT. Custom Dicer-Substrate siRNA (DsRNA). https://www.idtdna.com/site/order/designtool/index/DSIRNA_CUSTOM (2017). Accessed Apr 2010.

32. Tomchaney M, Mysore K, Sun L, Li P, Emrich SJ, Severson DW, et al. Examination of the genetic basis for sexual dimorphism in the Aedes aegypti (dengue vector mosquito) pupal brain. Biol Sex Differ. 2014;5:10.

33. Singh AD, Wong S, Ryan CP, Whyard S. Oral delivery of double-stranded RNA in the yellow fever mosquito, Aedes aegypti: implications for pest mosquito control. J Insect Sci. 2013;13:69.

34. Mysore K, Hapairai L, Wu N, Realey JS, Scheel ND, Severson DW, et al. Preparation and use of a yeast shRNA delivery system for gene silencing in mosquito larvae. Methods Mol Biol. 2019;1958:213–31.

35. Mumberg D, Muller R, Funk M. Yeast vectors for the controlled expression of heterologous proteins in different genetic backgrounds. Gene. 1995;156(1):119–22.

36. van Dijken JP, Bauer J, Brambilla L, Duboc P, Francois JM, Gancedo C, et al. An interlaboratory comparison of physiological and genetic properties of four Saccharomyces cerevisiae strains. Enzyme Microb Technol. 2000;26(9–10):706–14.

37. Mysore K, Hapairai L, Sun L, Li P, Wang CW, Scheel ND, et al. Characterization of a dual-action adulticidal and larvicidal interfering RNA pesticide targeting the Shaker gene of multiple disease vector mosquitoes. PLoS Negl Trop Dis. 2020;14(7):e008479.

38. WHO. Guidelines for laboratory and field testing of mosquito larvicides. www.cdc.gov/mosquitoes/about/diseases.html. Accessed Jan 2020.
46. Charlesworth B. The evolution of sex chromosomes. Science. 1991;251(4997):1030–3.

47. Charlesworth D, Charlesworth B, Marais G. Steps in the evolution of heteromorphic sex chromosomes. Heredity. 2005;95(2):118–28.

48. Sinkins SP, Gould F. Gene drive systems for insect disease vectors. Nat Rev Genet. 2006;7(6):427–35.

49. Juneja P, Osei-Poku J, Ho YS, Ariani CV, Palmer WJ, Pain A, et al. Assembly of the genome of the disease vector Aedes aegypti onto a genetic linkage map allows mapping of genes affecting disease transmission. PLoS Negl Trop Dis. 2014;8(1):e2652.

50. Fontaine A, Filipovic I, Fansini T, Hoffmann AA, Cheng C, Kirkpatrick M, et al. Extensive genetic differentiation between homomorphic sex chromosomes in the mosquito vector, Aedes aegypti. Genome Biol Evol. 2017;9(9):2322–35.

51. Miller JR, Koren S, Dilley KA, Puri V, Brown DM, Harkins DM, et al. Analysis of the Aedes albopictus C6/36 genome provides insight into cell line utility for viral propagation. Gigascience. 2018;7(3):1–13.

52. Gomulski LM, Mariconti M, Di Cosimo A, Scolari F, Manni M, Savini G, et al. The Nix locus on the male-specific homologue of chromosome 1 in Aedes albopictus is a strong candidate for a male-determining factor. Parasit Vectors. 2018;11(Suppl 2):647.

53. Krzywinska E, Krzywinski J. Effects of stable ectopic expression of the primary sex determination gene Yob in the mosquito Anopheles gambiae. Parasit Vectors. 2018;11(Suppl 2):648.

54. Papathanos PA, Bossin HC, Benedict MQ, Catteruccia F, Malcolm CA, Alphey L, et al. Sex separation strategies: past experience and new approaches. Malar J. 2009;8(Suppl 2):55.

55. Mashatola T, Ndo C, Koekemoer LL, Dandalo LC, Wood OR, Malakoane L, et al. A review on the progress of sex-separation techniques for sterile insect technique applications against Anopheles arabiensis. Parasit Vectors. 2018;11(Suppl 2):646.

56. Wu Y, Hu W, Biedler JK, Chen XG, Tu ZJ. Pure early zygotic genes in the Asian malaria mosquito Anopheles stephensi. Parasit Vectors. 2018;11(Suppl 2):652.

57. Catteruccia F, Benton JP, Crisanti A. An Anopheles transgenic sexing strain for vector control. Nat Biotechnol. 2005;23(11):1414–7.

58. Magnusson K, Mendes AM, Windbichler N, Papathanos PA, Nolan T, Dottorini T, et al. Transcription regulation of sex-biased genes during ontogeny in the malaria vector Anopheles gambiae. PLoS ONE. 2011;6(6):e21572.

59. Zacares M, Salvador-Herranz G, Almenar D, Tur C, Argiles R, Bourtzis K, et al. Exploring the potential of computer vision analysis of pupae size dimorphism for adaptive sex sorting systems of various vector mosquito species. Parasit Vectors. 2018;11(Suppl 2):656.

60. Mukiama TK. Y-autosome genetic sexing strain of Anopheles albimanus (Diptera: Culicidae). Int J Trop Insect Sci. 1985;6:649–52.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.