1	Revisiting the paradigm of anhematophagy in male mosquitoes
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32 Abstract

33 Female mosquitoes are reproductively obligate bloodfeeders which feed on vertebrate 34 blood to obtain nutrients required for egg production (driving transmission of vectorborne pathogens in the process), and which rely on plant sugars for their non-35 36 reproductive energy requirements. Male mosquitoes, on the other hand, are thought to 37 rely exclusively on plant sugars for their energetic needs; indeed, this dichotomy is one 38 of the central tenets of medical entomology. Here, we show that male *Culex tarsalis* and 39 Aedes aegypti mosquitoes will readily take blood from a membrane feeder when reared 40 under dehydration conditions with no toxic effects. Mosquitoes with impaired humidity 41 detection do not increase their bloodfeeding rates when dehydrated compared to wild-42 type controls. While conventionally reared males ignore a human host, dehydrated 43 males are attracted to and attempt to probe, with some success, although they cannot 44 access host capillaries. However, they will take blood from a vertebrate host wound. 45 When fed a blood meal containing West Nile virus, male mosquitoes can become 46 infected with and orally transmit the pathogen at rates and titers equivalent to females. 47 These data suggest that under some circumstances male mosquitoes may be able to 48 probe and/or ingest blood and transmit pathogens to vertebrate hosts, and that their role 49 in maintaining pathogen transmission cycles should be re-examined.

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51 **Key words:** *Culex tarsalis, Aedes aegypti,* mosquito, bloodfeeding, sugar feeding, 52 vector-borne disease

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63 Introduction

64 Female mosquitoes (except those which are autogenous) are reproductively obligate 65 bloodfeeders, feeding on vertebrate blood to obtain nutrients required for egg 66 production, and relying on plant sugars for their non-reproductive energy requirements. 67 Male mosquitoes, on the other hand, rely exclusively on plant sugars for their energetic needs [1]. Indeed, this difference is one of the central tenants of medical entomology: 68 69 female mosquitoes bloodfeed, males do not. Female bloodfeeding allows transmission 70 of blood-borne pathogens, such as viruses or parasites, between vertebrate hosts, 71 which is why the majority of mosquito research is performed on females rather than 72 males; even when males are studied, it is usually within the context of how they affect 73 females (mating behavior and fertility, pathogen transmission modulation etc...) [2-5].

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75 Female mosquitoes are adapted to feed on blood, and this adaptation is reflected in the 76 biology of their midgut, where transcripts related to blood digestion are enriched in the 77 female compared to the male [6]; one would expect that male mosquitoes should not be 78 attracted to blood as a nutrition source as they are thought to lack the proper physiology 79 to digest and process it. However, there is one interesting report in the literature where 80 male mosquitoes were attracted to and fed on blood. Nikbakhtzadeh and colleagues [7] 81 documented bloodfeeding behavior in a laboratory colony of the mosquito Culex 82 quinquefasciatus. When presented with defibrinated sheep blood on a cotton pledget 83 (and to a much less efficient extent, a Parafilm membrane), male mosquitoes took a 84 bloodmeal. However, blood was toxic to male mosquitoes, which died in a dose-85 dependent manner when blood was mixed with sugar [7], consistent with physiological 86 adaptations to sugar vs bloodfeeding in males vs. females [6]. Interestingly, males did 87 not show a preference for sugar compared to blood in a dual-choice assay [7], and the 88 reason they fed on blood at all, particularly as it was toxic, remains an open question. 89 As this is (to our knowledge) the only observation of male mosquito bloodfeeding 90 behavior, it is difficult to speculate. However, there are a multitude of observations that 91 male mosquitoes are attracted to human host odors and this behavior is suppressed by 92 mosquito repellants [8], which includes species from arguably the three most important 93 mosquito genera that act as disease vectors to humans (Anopheles, Culex, and Aedes).

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95 Here, we present studies on male bloodfeeding behavior in the mosquitoes Cx. tarsalis 96 and Ae. aegypti. Cx. tarsalis is one of the major West Nile virus (WNV) vectors in North 97 America, where it is widely distributed across the Western United States [9]. It is 98 genetically diverse, generally feeds on birds in the wild, and can be facultatively 99 autogenous [9-10]. After becoming infected with WNV during a bloodfeeding event, it 100 can also transmit the virus vertically to offspring at relatively high rates [11-12]. Ae. 101 aegypti is one of the major invasive arbovirus vectors in the world [13]. We 102 opportunistically observed Cx. tarsalis and Ae. aegypti males taking blood during 103 unrelated laboratory studies, and undertook experiments to document and understand 104 the behavior. We found that when dehydrated, male Cx. tarsalis and Ae. aegypti will 105 predictably take human blood from a membrane feeder, determined the mechanism 106 driving male bloodfeeding behavior, and present results of experiments examining 107 potential for male mosquitoes to be involved in pathogen transmission cycles. These 108 results are a paradigm shift in our understanding of male mosquito biology and suggest 109 they may be more directly involved in pathogen transmission cycles than previously 110 recognized.

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112 Methods

Human subjects: All experiments with a human volunteer used the senior author (JLR)
 under PSU IRB Exempt Protocol STUDY00024284.

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<u>Mosquitoes:</u> *Cx. tarsalis* strain (KNWR) and *Aedes aegypti* (Liverpool) were maintained at 25°C, 16:8 h light:dark diurnal cycle with 80% relative humidity, with 10% sucrose solution provided at all times through a cotton wick. For general rearing, mosquitoes were provided with expired anonymous human blood (BioIVT) through a water-jacketed glass membrane (Parafilm) feeder or a Hemotek feeder for egg development.

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122 <u>Male dehydration:</u> To stimulate bloodfeeding, male mosquitoes were held at 25°C, 75%

123 RH without sugar or water for 24 hours [14].

<u>Survival analysis:</u> Bloodfed male mosquitoes were isolated and placed into cup cages, held at the previously described standard insectary conditions, and provided with a cotton ball soaked in 10% sucrose solution. Control males were non-bloodfed and were maintained under the same conditions. Dead mosquitoes were counted every day and removed from the cages. Significant differences in survival between mosquito groups was determined with Kaplan-Meier analysis using GraphPad Prism version 9.0.4.

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<u>Dehydration and bloodfeeding behavior in male *Cx. tarsalis*: *Cx. tarsalis* males were reared conventionally (80% RH, with free access to 10% sucrose solution in water), or under dehydrating conditions (75% RH, 25°C, with no access to water or sugar for 24 hours) [14], then were offered a bloodmeal through a membrane feeder for 30 minutes. The number of fed and unfed mosquitoes at the end of the feeding period were counted. Data were analyzed by Fishers Exact test.</u>

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139 Ionotropic receptor 93a (Ir93a) mutant mosquito assays: CRISPR protocols have not yet 140 been developed for Cx. tarsalis, but we noted during experiments that males of the 141 species Ae. aegypti (where CRISPR mutagenesis is routine) would also take blood from 142 a membrane feeder, so we obtained an Ae. aegypti line that was a CRISPR knockout 143 mutant for the Ir93a gene, which inhibits its ability to sense humidity [15]. The mutation 144 was introgressed into the wild-type Liverpool background for comparison with Liverpool 145 controls, and both lines reared as described above. For experiments, at 5-6 days post-146 emergence, males of each strain were transferred to 10 x 10 x 10 cm cages and 147 deprived of sucrose and water (or held at normal conditions as controls) for 24 hours 148 before being offered an anonymous human bloodmeal using an artificial feeding system 149 (Hemotek). Bloodfeeding rates for each genotype and condition were recorded. Data 150 were analyzed by Fishers Exact tests and confidence intervals calculated from the 151 binomial distribution.

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Landing and probing experiments: Cages of 50 *Cx. tarsalis* males (reared under standard or dehydrating insectary conditions) were allowed to probe on the hand of the senior author for five minutes. Mosquito landings (defined as a mosquito alighting on the

volunteer hand for any period of time) and probing behavior (defined as exploring and
probing with mouthparts) were counted during the 5 minute interval. The experiment
was repeated 6 times. Data were analyzed by Mann-Whitney U test.

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Host bloodfeeding by male mosquitoes: The senior author had an unrelated small (3mm) scratch on their hand from obtained from a pet cat a day earlier. A sterile razor blade was used to pick the scab off the scratch allowing a minimal amount of blood to be exposed. The wounded hand was placed in a cage of 20 dehydrated male *Cx. tarsalis* mosquitoes and their behavior recorded.

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166 WNV feeds: Dehydrated Cx. tarsalis males and female controls were allowed access to 167 an infectious blood meal consisting of a 1:1 mix of anonymous human blood (BioIVT) and 5.0 x 10⁷ FFU/mI (focus-forming units/mI) of WN02-1956 (GenBank: AY590222). A 168 169 subset of male and females were processed immediately after feeding ("day zero") to 170 check for virus viability. Mosquito virus infection and transmission assays were 171 performed at 7 and 14 days post-blood feeding. Fully engorged mosquitoes were sorted 172 from non-fed ones for analysis. Mosquitoes were anesthetized with triethylamine 173 (Sigma, St. Louis, MO), legs/wings from each mosquito were removed and placed 174 separately in a 2-mL tube filled with 0.5 mL mosquito diluent (MD: 20% heat-inactivated 175 fetal bovine serum (FBS) in Dulbecco's phosphate-buffered saline, 50 µg/mL 176 penicillin/streptomycin, 50 ug/mL gentamicin, and 2.5 µg/mL fungizone, with a sterile 2.0 177 mm stainless steel bead (Next Advance, Inc. Innovative Lab Products for the Life 178 Sciences). The proboscis of each mosquito was positioned in a tapered capillary tube 179 containing approximately 10 µL of a 1:1 solution of 50% sucrose and FBS to induce 180 salivation. After 30 min, the tube contents were expelled into 0.3 mL MD, and bodies 181 were placed individually into a 2-mL tube filled with 0.5 mL MD and a stainless steel 182 bead as described above. Mosquito bodies and legs/wings were homogenized for 30 183 sec with TissueLyser (QIAGEN, Hilden, Germany) at 24 cycles/sec, followed by 184 centrifugation for 1 min. Mosquito bodies, legs/wings, and salivary secretion samples 185 were tested for live, infectious WNV using focus-forming assays (FFAs; see below).

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187 WNV FFAs: WNV titers were quantified by FFA, which detects live, infectious virus. 188 C6/36 cells were seeded into 96-well plates at a density of 1X10⁵ cells/well and 189 incubated overnight at 28°C in complete RPMI medium without CO₂. The next day, 190 medium was removed from the wells. Samples from male and female bodies or 191 legs/wings were serially diluted in a serum-free RPMI medium; saliva samples were 192 undiluted. 30 µl of each sample was added in duplicate to the prepared C6/36 cells. 193 Cells were incubated for 1 hour at 28°C without CO₂, after which the inoculum was 194 removed. 100 µl of RPMI containing 0.8% methylcellulose was added to limit viral 195 spread. Infected cells were incubated for 48 hours at 28°C without CO₂. At 48 hours 196 post-infection, infected C6/36 cells were fixed with 50 µl 4% formaldehyde for 30 197 minutes at room temperature (RT). Cells were washed, permeabilized with 0.2% triton-X, and blocked with 3% BSA. 30 µl monoclonal flavivirus antibody (Clone D1-4G2-4-15, 198 199 BEI-resources, NR-50327) was added and incubated overnight at 4°C. After washing, 200 30µl of fluorescent secondary antibody (1:1000 dilution: Goat anti-Mouse IgG (H+L) 201 Highly Cross-Adsorbed Secondary Antibody, Alexa Fluor 488, Invitrogen/Thermo 202 Fischer, A-11029) was added and incubated overnight at 4°C. Cells were maintained in 203 100 µI PBS to prevent drying. West Nile virus foci were imaged using a FITC filter on an 204 Olympus BX41 microscope with a UPIanFI 4x objective and counted. Infection rate (IR) 205 was defined as the proportion of mosquitoes exposed to virus that had WNV-positive 206 bodies. Dissemination rate (DR) was defined as the proportion of mosquitoes with 207 WNV-positive bodies that had WNV-positive legs/wings. Transmission rate (TR) was 208 defined as the proportion of mosquitoes with WNV-positive legs/wings that had WNV-209 positive saliva. IR, DR, and TR were analyzed with Fisher's exact tests. Viral titers were 210 analyzed using Mann-Whitney U tests.

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213 **Results**

<u>Bloodfeeding is not toxic to *Cx. tarsalis* males:</u> While we were bloodfeeding during an experiment related to relative humidity (e.g. [16]), we noted incidentally that in addition to females, male mosquitoes were probing the membrane and were taking blood (Figure 1A,B). As this was a spontaneous occurrence, the total number of males in the

218 cage was not recorded but was on the order of 50-70 based on standard rearing 219 practices in our lab. Out of this total, we isolated seven blood-engorged males. These 220 males were placed into a survival cup and a survival experiment conducted, comparing 221 their survival to 10 non-bloodfed males from the same initial cage. Although 222 Nikbakhtzadeh et al. [7] demonstrated than blood was highly toxic to male Cx. 223 quinquefasciatus, we did not observe any acute toxicity to blood in male Cx. tarsalis; 224 indeed, survival in bloodfed males was marginally (although not statistically) higher than 225 non-bloodfed (Figure 1C).

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227 <u>*Cx. tarsalis* male bloodfeeding is driven by dehydration:</u> As we previously demonstrated 228 that dehydration stimulates elevated bloodfeeding behavior in females [14, 16], we 229 tested the hypothesis that dehydration was driving bloodfeeding behavior in males. 230 Cages of male mosquitoes were reared under conventional insectary conditions or 231 under dehydrating conditions [14]. No conventionally reared male mosquito (N = 64) 232 took a bloodmeal from the membrane feeder, while 44/163 dehydrated males took a 233 bloodmeal (P < 0.00001).

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235 Male mosquito bloodfeeding behavior is dependent on their ability to sense humidity: 236 Ae. aegypti and Anopheles gambiae mosquitoes sense humidity through ionotropic 237 receptor Ir93a, by which they locate oviposition sites, and CRISPR Ir93a knock-out 238 mutants are impaired in this behavior [15]. We noted anecdotally that in our lab, male 239 Ae. aegypti mosquitoes would also take blood from a membrane feeder, and as 240 CRISPR protocols have not yet been developed for Cx. tarsalis, we used an Ir93a Ae. 241 aegypti KO mutant for these assays [15]. When reared under standard insectary 242 conditions, bloodfeeding rates did not differ statistically between wild-type and mutant 243 mosquitoes. However, when reared under dehydrating conditions, bloodfeeding rates 244 for the mutant did not increase, while wild-type mosquitoes had significantly elevated 245 bloodfeeding behavior (P = 0.0284) (Figure 2).

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247 <u>Dehydrated male mosquitoes will probe the hand of a human volunteer:</u> The hand of a 248 human volunteer was exposed to cages of conventionally reared or dehydrated male

249 Cx. tarsalis. Conventionally reared males showed little interest in the host, with 250 infrequent landings that lasted less than 5 seconds. None demonstrated probing 251 behavior. In contrast, dehydrated males landed significantly more often on the hand of 252 the volunteer (P = 0.0065), most landings lasted until the end of the time period, and 253 probing behavior was observed in the majority of landings (P = 0.0022) (Figure 3, Supplementary Videos 1 and 2). One dehydrated male mosquito (out of 6 separate 254 255 trials) was able to lightly pierce the skin of the volunteer at the base of the wrist, 256 although it was unable to reach the capillaries and acquire a bloodmeal (Supplementary 257 Video 3). The bite resulted in a mild immunogenic reaction that disappeared after 258 approximately 10 minutes (Supplementary Figure 1). To confirm that only males were in 259 the cage, after the study was concluded the entire cage was killed by freezing and every 260 mosquito visually examined for the presence of a female or a gynandromorph; only 261 males were identified. While this is only a single observation and thus definitive 262 conclusions cannot be drawn, to our knowledge, this is the first documented case of a 263 male mosquito biting a vertebrate host.

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265 Dehydrated male mosquitoes will take blood from a vertebrate host wound: Dehydrated 266 male Cx. tarsalis show keen interest in probing a human host, but were not able to acquire blood, even from the single observed "successful" probing attempt. We 267 268 hypothesized that if blood was made more accessible, male mosquitoes would take a 269 bloodmeal. The senior author serendipitously had a small scratch on their hand 270 (acquired from a pet cat a day earlier). The scab was peeled back using a sterile razor 271 blade, exposing a small amount of blood. The volunteer placed their hand in a cage of 272 20 dehydrated male mosquitoes. Males were attracted to the wound, and wound 273 probing behavior was observed by 5 males (see Supplementary Video 4 for example). 274 One male out of the 5 that probed fed and took a bloodmeal from the wound 275 (Supplementary Video 5, Figure 4). At the conclusion of the experiment, the fed male 276 was dissected to confirm the presence of blood in the gut (Figure 4).

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278 <u>Male Cx. tarsalis mosquitoes are competent vectors for West Nile virus:</u> Since we 279 determined that male Cx. tarsalis will probe a human hand or ingest blood from a

280 wound, allowing ingestion of a blood meal from a vertebrate host, we asked the 281 guestion: can male mosquitoes become infected with and transmit arboviruses? We 282 offered dehydrated male mosquitoes a bloodmeal spiked with WNV and assayed their 283 vector competence at day 7 and day 14 post-exposure. Female Cx. tarsalis were 284 exposed to virus at the same time as a control. We found that both female and male Cx. 285 tarsalis were able to become infected with, disseminate, and orally transmit virus; males 286 transmitted at both day 7 and 14, while females only had detectable virus in their saliva 287 at day 14. After adjusting for multiple comparisons, infection rates (IR), dissemination 288 rates (DR), and transmission rates (TR) did not differ statistically between males and 289 females at either timepoint (Table 1).

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291 We quantitated all viral titers using an infectious virus assay. First, a subsample of 292 males and females were assayed immediately after feeding ("day zero") to confirm virus 293 viability. All fed males and females had detectable live infectious virus in their bodies, 294 although females had statistically higher viral titers (P = 0.005), likely because they 295 could physically ingest a larger volume of blood. At day 7 post-exposure, viral titers 296 were not statistically different between males and females in the bodies, the legs/wings, 297 or the saliva (Figure 5A). At day 14 post-exposure, females had higher viral titers in their 298 bodies (P = 0.001) and legs/wings (P = 0.0083) compared to males, suggesting either 299 greater viral replication rates, or simply more tissue available for virus replication due to 300 the larger size of the females. However, viral titers in saliva between males and females 301 were statistically similar (Figure 5B).

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304 **Discussion**

Previous work showed that blood was toxic to male *Cx. quinquefasciatus* mosquitoes [7], suggesting that in this species male bloodfeeding seems to be a maladaptive trait, perhaps a laboratory artifact. In our study, we demonstrate that males of other species (*Cx. tarsalis* and *Ae. aegypti*) can tolerate bloodfeeding, and that male bloodfeeding behavior is driven by water homeostasis during dehydration conditions. When mosquitoes cannot sense humidity due to Ir93a mutagenesis, dehydration does not

increase blood seeking behavior. These results are consistent with the role of dehydration on bloodfeeding behavior in female mosquitoes, where dehydration can stimulate females to increase their bloodfeeding rates as well [14, 16-17] and thus may reflect an adaptive trait where mosquitoes (female or male) can maximize their water intake during drought or periods of low relative humidity if other sources (nectar or free water) are not available.

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318 The mouthparts of male mosquitoes are thought to be physically incapable of 319 penetrating vertebrate skin; however, in our experiments they were proven adequate to 320 pierce a Parafilm membrane. Dehydrated Cx. tarsalis males were significantly attracted 321 to and actively probed the hand of a human host, and one individual was even able to 322 slightly penetrate the outer epidermis, leading to a transitory immune reaction 323 (Supplementary Video 3 and Supplementary Figure 1). As the saliva of males differs from that of females, lacking various proteins needed for immunomodulation and 324 325 bloodmeal acquisition [18], and it is likely that very little saliva was transferred compared 326 to the bite of a female mosquito, it is not surprising that the host immune reaction was 327 mild and rapidly resolving. When allowed access to a wound, dehydrated male 328 mosquitoes readily probed the wound and one took a bloodmeal. As this experiment 329 was facilitated by the fortuitous presence of a pre-existing wound on the hand of the 330 senior author, it could not be deliberately repeated (as we were not allowed to make a 331 deliberate wound due to IRB concerns). Still, it does suggest that male mosquitoes have 332 the ability to take blood under specific rare circumstances that require dry periods and, 333 likely, a host with a wound. According to fossil evidence, male mosquitoes are thought 334 to once have had the ability to feed on vertebrate blood, and to have lost this ability over 335 evolutionary time [19]. It is possible that the neural circuitry regulating host seeking and 336 bloodfeeding behavior may still be conserved among male mosquitoes, or alternatively 337 that this is simply a unique response to dehydration conditions in the lab.

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Interestingly our data demonstrate that, if male *Cx. tarsalis* orally acquire a WNV infection, they are competent vectors and transmit the virus at similar rates and titers compared to females. In our experiments we explicitly used an assay that quantified

342 live, infectious viral particles rather than quantitative PCR to rule out results that might 343 be due to carryover of non-infectious viral RNA. Our results suggest that male *Cx.* 344 *tarsalis* retain the receptors necessary for viral infection on their midgut, salivary glands, 345 and other body tissues.

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347 Finally, there is the question "is bloodfeeding behavior by male mosquitoes 348 epidemiologically significant"? It is already known that male mosquitoes can be 349 indirectly important for vector-borne disease transmission dynamics. For example, 350 mating can affect key physiological parameters in females related to pathogen infection 351 and transmission [2]. More directly, in some species, including Cx. tarsalis and WNV, 352 male mosquitoes can be infected with arboviruses by vertical transmission from infected 353 mothers [11, 20-21]. Infected males can also transmit some viruses venereally to 354 females during mating where they can be transmitted to vertebrate hosts during feeding 355 [20-21]. Consistent collection of males using host-derived attractants suggest that males 356 are commonly found to move toward hosts [8], increasing the potential of male feeding 357 on host-derived fluids under specific conditions (dry periods with a lack of sugar and 358 water resources). Our study suggests the rare possibility of edge cases where male 359 mosquitoes could be more directly implicated in virus transmission, where males 360 undergoing dehydrating conditions (for example, during drought) acquire virus through 361 vertical transmission from infected mothers or by feeding on an open wound of an 362 infected vertebrate host, then transmit to a naïve host through feeding on an open 363 wound or by probing the skin, as mosquitoes often transmit the bulk of virus when 364 probing skin prior to actually taking a bloodmeal [22].

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We must emphasize that while compelling, the results presented in this research are laboratory-based, and there is no peer-reviewed evidence of male mosquito bloodfeeding or pathogen transmission in nature (although we suspect that researchers have not rigorously looked for these phenomena). However, while arbovirus transmission by males is unlikely to be a major factor in driving disease dynamics, these data suggest that their canonical role as non-bloodfeeders needs to be reexamined and their contribution to pathogen transmission explicitly quantified,

- 373 particularly in light of recent vector-borne disease control strategies that rely on the
- mass release of male mosquitoes into natural populations [23-26].
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503 **Declarations**

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505 Ethics approval and consent to participate: All experiments with a human volunteer 506 used the senior author (JLR) under PSU IRB Exempt Protocol STUDY00024284.

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515 Authors' contributions: JB, REJ, RSK, JBB, and JLR conducted the research, JBB 516 contributed materials and reagents, JLR analyzed the data, JB, REJ, JBB, and JLR 517 wrote the manuscript.

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- 529 530 531 532 533 **Figure legends** 534 Figure 1. Male Cx. tarsalis bloodfeeding behavior and survival. A) mosquitoes 535 congregating at and bloodfeeding from a paraffin membrane. Arrow points to male 536 orienting toward the membrane. B) Blood-engorged male mosquito. Un-engorged male 537 can be seen in-frame. C) Survival curve of bloodfed vs. non-bloodfed male Cx. tarsalis 538 mosquitoes. No significant difference was observed between treatments. 539 540 Figure 2. CRISPR deletion of Ir93a ablates male mosquito bloodfeeding behavior 541 under dehydration conditions. When reared conventionally, both wild-type and 542 mutant Ae. aegypti exhibit baseline levels of bloodfeeding behavior. When reared under 543 dehydration conditions, wild-type males significantly increase bloodfeeding behavior but 544 humidity-insensitive mutant mosquitoes do not. Confidence intervals were calculated 545 from the binomial distribution. WT = wild-type. 546 547 Figure 3. Host probing behavior of dehydrated male mosquitoes. A) Control cage
- of conventionally reared male *Cx. tarsalis*. Mosquitoes ignore the human host. B) Cage of dehydrated male *Cx. tarsalis*. Dehydrated males land on and probe the human host. C, D) Stills from Supplementary Videos 1 and 2 showing dehydrated male mosquito probing behavior. See videos for complete behavioral responses. E) Landing responses for dehydrated vs. conventionally reared ("standard") male *Cx. tarsalis*. F) Probing behavior for dehydrated vs. conventionally reared ("standard") male *Cx. tarsalis*. Error bars = SEM. ** = P < 0.01

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556 **Figure 4. Male** *Cx. tarsalis* **taking a human bloodmeal from a wound.** A) Male *Cx. tarsalis* feeding on an open wound. B) Close-up of feeding behavior. C) Blood is observable in the male mosquito gut. D) Blood in the male mosquito gut was confirmed by dissection. See Supplementary Video 3 for complete behavioral response.

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Figure 5. West Nile virus vector competence for male and female *Cx. tarsalis.* A) 7 days post-exposure. B) 14 days post-exposure. Red = bodies (infection); blue = legs/wings (dissemination); black = saliva (transmission). Males = closed circles, females = open circles. Zero values had 0.01 added purely for log-scale plotting purposes (10^{-2} = uninfected); analysis was performed on untransformed data. ** = *P* < 0.01.

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Table 1. WNV Infection rate (IR), dissemination rate (DR) and transmission rate (TR) of dehydrated males and females at 7 and 14 days post-infection. No comparisons were statistically significant after correcting for multiple comparisons.

	Ν	IR	DR	TR
Day 7				
Males	43	0.53	0.70	0.56
Females	10	0.50	0.60	0.00
Day 14				
Males	14	0.50	0.86	0.50
Females	12	0.92	0.91	0.30















Supplementary material

Supplementary Figure 1. Host immune reaction to probing of dehydrated male *Culex tarsalis* mosquito. A) Bite reaction 2 minutes post-probing (arrow). B) Immune reaction resolved by 10 minutes post-probing.



Supplementary Video 1. Probing behavior of dehydrated male *Cx. tarsalis* mosquito on the thumb of a human volunteer.

Supplementary Video 2. Probing behavior of dehydrated male *Cx. tarsalis* mosquito on the index finger of a human volunteer.

Supplementary Video 3. Probing behavior of dehydrated *Cx. tarsalis* male mosquito on the wrist of a human volunteer. This mosquito succeeded in slightly penetrating the outer epidermis (see Supplementary Figure 1).

Supplementary Video 4. Male *Cx. tarsalis* probing a human host wound.

Supplementary Video 5. Male Cx. tarsalis feeding from a human host wound.