The microbiome and mosquito vectorial capacity: rich potential for discovery and translation

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Abstract

Microbiome research has gained considerable interest due to the emerging evidence of its impact on human and animal health. Similar to higher organisms, the gut-associated microbiota of mosquitoes affect host fitness and other phenotypes. It is now well established that microbes can alter pathogen transmission in mosquitoes, either positively or negatively, and avenues are being explored to exploit microbes for vector control. However, less attention has been paid to how microbiota affect phenotypes that impact vectorial capacity. Several mosquito and pathogen components, such as vector density, biting rate, survival, vector competence and pathogen extrinsic incubation period all influence pathogen transmission. Interestingly, the mosquito gut-associated microbes can impact each of these components, and therefore ultimately modulate vectorial capacity. Promisingly, this expands the options available to exploit microbes for vector control by also targeting parameters that affect vectorial capacity. However, there are still many knowledge gaps in the biology of the mosquito – microbe symbiosis that need to be addressed in order to understand these interactions more thoroughly and exploit them efficiently. Here, we review current evidence of the impacts of the microbiome on aspects of vectorial capacity highlighting opportunities for novel vector control strategies and areas where further studies are required.

Keywords: microbiome, vectorial capacity, density, competence, biting, extrinsic incubation period, longevity, mosquito, symbiosis, pathogen transmission
Background

The microbiome is a collection of microorganisms within or on an organism. In mosquitoes, the microbiome, which consists of bacteria, viruses and fungi, profoundly alters host phenotypes. Acquisition and the composition of the microbiome is influenced by several abiotic and biotic factors, including host and microbial genetics [1–4] and the environment [5–7]. Therefore microbiomes of mosquitoes can vary substantially between individuals, life stages, species, and over geographical space [8,9], and this variation likely contributes to differences in host phenotypes. Similarly, the horizontal and vertical transmission routes microbes exploit to colonize their host means that mosquitoes reared in a laboratory setting have a vastly different microbiome compared to their field counterparts [10–12]. As such, undertaking studies with a field relevant microbiome has been challenging. Within the mosquito, microbes colonize different tissues, perhaps by intracellular routes [13], however the reproductive organs [14,15] and salivary glands [16] appear to have the greatest diversity of microbes. The latter, in addition to the gut are relevant tissues for pathogen transmission. However, microbes that reside in the gut [17], Malpighian tubules [18] or other tissues may have relevance for other life history traits which influence vectorial capacity.

Vectorial capacity describes the ability of a population of vectors to transmit pathogens to a host and has been represented by the vectorial capacity equation (Fig. 1). This was created by Garret-Jones in 1964 and represents the number of secondary cases of vector infection per unit of time given the introduction of an infectious individual into a naïve population [19,20]. Pathogen transmission is modelled by the vectorial capacity equation, which is a vector-centric adaptation of the basic reproductive number (\(R_0\)) equation [21]. The components of the vectorial capacity equation are the following: vector biting rate (a), vector density (m), probability of vector daily survival (p), vector competence (b) and pathogen extrinsic incubation period (N). An infected person gets bitten by \(ma\) vectors each day. Of
these $ma$ bites, only a proportion $b$ is infectious to the vector, giving a total of $mab$ vectors infected by the primary case. The proportion of vectors surviving the extrinsic incubation period is $p^N$, so $mabp^N$ vectors become infectious. Each of these infectious vectors then survives for an average time of $1/-\ln(p)$, and during this time, it bites people at the rate of $a$ bites per day, making a total of $a/-\ln(p)$ bites. Thus, there are $mahp^N$ infectious vectors arising from the primary case making $a/-\ln(p)$ infectious bites on susceptible hosts, resulting in the following vectorial capacity: $ma^2hp^N/-\ln(p)$. Therefore, each component of the equation will have a certain impact on the ability of mosquitoes to transmit pathogens. As such, targeting any of these components could result in a reduction of pathogen transmission.

Some components of the vectorial capacity equation have traditionally received more attention than others by mosquito control efforts. Probability of daily survival and density have been targeted by insecticides and larvicides respectively, achieving significant reduction of vector-borne diseases, but the emergence of insecticide resistance is compromising these strategies. Vector competence has been the main focus of the design of novel vector control methods, such as release of Wolbachia-infected mosquitoes for population replacement, which has showed unprecedented success in dengue control [22]. However, little attention has been paid to other aspects of mosquito biology which can have equal or potentially greater effect on pathogen transmission. In that sense, the great diversity of mosquito gut-associated microbes could offer new tools to target different components of vectorial capacity [23,24]. In order to leverage the microbiome for vector control, it is imperative to understand how microbial symbionts modulate vector biology. In this review we compile evidence of the impact of the mosquito gut-associated microbiome on each component of the vectorial capacity equation. We also discuss other vector systems, what we can infer from other insect models and what is known about Wolbachia when there is a
lack of evidence on how gut-associated microbes influence traits relevant for vectorial capacity in mosquitoes.

Influence of microbiota on vector competence (b)

Microbes that associate with vectors, including bacteria [25], virus [26], fungi [27] and microsporidia [28] can modulate vector competence. Vector competence is fundamental to vectorial capacity since it determines the susceptibility of the mosquito to become infected by a pathogen, and the higher the vector competence, the higher the vectorial capacity. Gut associated microbiota can interfere directly with pathogens through mechanisms such as lysis and biofilm formation [25] or indirectly by affecting intrinsic aspects of the vector that determine its vector competence, like midgut and salivary gland barriers [29–31] and the immune system [1,32]. In addition, microbiota can potentially have other functions in pathogen transmission, since it may be transmitted to the mammalian host during feeding on the host [33]. The role of the gut microbiome in modulating vector competence for several pathogens has been well studied and reviewed extensively in mosquitoes [34–41] and other vectors [42–48], so we have focused our attention on the other components of the vectorial capacity equation.

Influence of microbiota on pathogen extrinsic incubation period (N)

The extrinsic incubation period (EIP) is the time that it takes for pathogens to develop in the vector and little is known regarding how microbes influence this time. The EIP affects vectorial capacity since it influences the number of infected mosquitoes that live long enough to become infectious and can vary depending on host and pathogen genetic factors and environmental conditions. There is evidence that Wolbachia infection can extend the EIP for DENV in Aedes aegypti [49,50] and the authors argue that this may be due to the antiviral properties of Wolbachia, which delay the time that the virus titres reach an infectious
threshold. Given gut-associated microbiota modulate pathogens, it would be interesting to explore how the microbiota could be exploited to delay the EIP. Alternatively, microbiota that enhance the EIP could be potentially targeted to prevent a positive effect on pathogen transmission.

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**Influence of microbiota on vector density (m)**

Vector density is the number of vectors per host, and there is increasing evidence suggesting that the mosquito gut microbiota can modulate this facet of vectorial capacity. A reduction in vector density leads to progressive population reduction in successive generations, resulting in reduced vectorial capacity. This principle has been the cornerstone of many vector control strategies, whereby larvicides or the release of *Wolbachia*-infected male mosquitoes are used to reduce the number of vectors in a population [51,52]. Gut-associated microbes can influence vector density through the modulation of development, reproductive outputs and resistance to abiotic stress.

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1. **Growth and development**

Recent work has elucidated the importance of microbes as a factor influencing growth and moulting of mosquito larvae into adults by regulating growth signalling and serving as a food source. Axenic (microbe-free) larvae fail to moult under normal rearing conditions [53], and exhibit differential expression of genes relating to amino acid transport, hormone signalling, and metabolism compared to normal larvae [54]. Although some studies have produced larvae that developed without bacteria [55–57], the addition of living microbes appear to induce gut hypoxia and activation of growth-related signalling pathways that larvae require to achieve the critical size necessary for moulting [58–60]. In addition, gut hypoxia depends on bacterial density, as shown by *Ae. aegypti* larvae showing higher growth rates [60] and *Aedes albopictus* exhibiting enhanced adult emergence [61]. This indicates that the
mechanisms responsible for regulation of host development occur via bacterial metabolism. Without bacteria-induced hypoxia [62], the larva fails to make adequate nutrient stores, so the mosquito is under microbial influence for accumulation of nutrient reserves that will take it into adulthood. Most mosquito species are detritivore as larvae, using bacteria as a food source [63], but predaceous species also access bacteria as food when prey are not available, so bacteria can contribute to nutritional supply when food availability is a limiting factor [64,65]. Reliance on microbe-induced hypoxia for signalling appears to be conserved across mosquito lineages, including detritovore larvae from the Culicinae and Anophelinae subfamilies, and predaceous larvae of Toxorhynchites amboinensis [66], indicating that the role of larval gut microbiota on mosquito development is not limited to detritivory. Another condition that relies on the nutritional stocks built during larval development is autogeny, which is the ability of some mosquito species to produce their first clutch of eggs without taking a blood meal. Although both anautogenous and autogenous mosquito species rely on the larval microbiota for development, the autogenous Aedes atropalpus shows limited rescue of development by some bacterial taxa when reared in a monoculture, in contrast to its anautogenous relative Ae. aegypti [67]. This suggests that autogenous species may have more specific requirements for microbiota composition due to their reliance on larval nutrition. Gut microbes simultaneously regulate signalling and serve as a food source, and further study is required to identify potential interactions of these dual functions and their impacts on vector life history.

Characterization of microbiota effects on vector development begins with tracing impacts of individual microbial taxa and continues with the study of bacterial communities and their diversity. Although multiple microbial taxa individually support normal development [53], outcomes may differ according to nutrient conditions: Ae. aegypti reared on E. coli, Saccharomyces cerevisiae, or Chlamydomonas reinhardtii, vary in their survival to
adulthood and longevity as a function of the diet they are reared on [60]. *Culex pipiens* reared on the human pathogen *Cryptococcus gattii* show detrimental development compared to individuals reared on *S. cerevisiae* or yeasts isolated from wild *Cx. pipiens* and *Cx. theileri* [68]. Furthermore, colonization by a single strain may manifest as more or less supportive of host fitness in different life history traits. Naturally occurring bacterial strains *Klebsiella sp.* and *Aeromonas sp.* are able to support *Cx. pipiens* larval development from first to second instar and are the most attractive to ovipositing females, but fail to produce surviving adults [69]. Particularly impactful microbes may also alter development even when they are not the sole occupant of the larval gut. For example, supplementation of conventionally reared larvae with a culture of *Asaia* increases the rate of *Anopheles gambiae* development, however it is unknown whether this effect results from *Asaia* metabolism specifically, or merely from the increased bacterial density [70]. Diversity and community composition of the microbiota also impact development. Larvae reared on combined microbial isolates have higher pupation and survival rates than those reared on monocultures, indicating that a combination of cells of differing nutrient compositions and/or metabolic processes may have additive effects for larval nutrition [71]. In addition, antibiotic treatment which decreases diversity and abundance of the gut microbiota delays larval development by 2-4 days in *An. stephensi* [72]. However, supplementation of the disturbed microbiota with antibiotic resistant *Asaia* restores development, suggesting that the roles of density and diversity in the gut microbiota's modulation of host phenotype is complex and requires further testing.

2. Reproductive output

The microbiome influences the mosquito reproductive output, which is the culmination of several physiological processes and population dynamics, beginning with sex ratio and mating behaviour, and resulting in egg production and hatching. Sex ratio is the number of
males or females relative to the total number of emerged adults. The female-biased sex ratio of *Ae. aegypti* can be shifted towards a male-biased sex ratio when larvae are fed with bacteria or yeast symbionts [73]. This may be the result of differences in larval metabolism and development between males and females, so further investigation is needed to understand the mechanisms underpinning this phenotype.

Mating starts with an effective encounter between individuals which requires a certain population density and coordinated locomotor behaviour. There is evidence these traits can be influenced by the gut microbiome. For example, studies in *Drosophila* indicate larvae congregate in response to acetoin produced by the gut microbiome [74] and the absence of gut microbiota leads to a hyperactive adult behaviour [75]. Normal behaviour can be restored with the addition of *Lactobacillus*, bacteria that produces xylose isomerase and influences a neuronal pathway [75]. Some mosquitoes mate in swarms, and variation in microbiota between swarms has been observed [76], however further work is required to determine the cause of these differences. Mating behaviour continues with the mating choice. In *Drosophila*, a greater number of matings were observed when males and female were reared on diet containing the same microbial consortia as opposed to diets with different microbial communities [77,78]. Changes in the levels of sex pheromones and production of volatile compounds seem to be two factors responsible for this phenotype [79]. First, the production of hydrocarbons is regulated by the insulin signalling pathway, which is enhanced by the microbiome [80]. Second, changes in the ratios of cuticular hydrocarbons affect mating recognition and sexual attractiveness [81–83]. Further investigation is required in order to disentangle the effects of the microbiota on host mating behaviour since this could affect genetic control strategies in vectors, and there is evidence that similar traits occur in mosquitoes. For example, transgenic mosquitoes with enhanced immunity have a modified microbiome and these mosquitoes have a mating fitness advantage compared to their wild
Egg production, oviposition, and hatching in insects are all affected by microbiota, and this impact on fecundity translates to changes in vector density. In general, fecundity in mosquitoes is governed by nutritional availability acquired during blood feeding, so blood digestion by adult females is necessary for egg production. A significant increase of microbe levels occurs after mosquitoes take a bloodmeal [85–87]; and treatment of Ae. aegypti with antibiotics impedes digestion of blood proteins and consequently reduces egg production [88], suggesting that the microbiome contributes to blood digestion. Recently it has been shown that sequential bloodmeals promote pathogen infection [89,90], and it would be intriguing to determine the role of the microbiome in this phenotype. Additionally, when Ae. aegypti eggs are laid in water containing bacteria, they hatch at a higher rate than those laid in sterile water [91]. Taking together, it is evident that microbes enhance mosquito fecundity and therefore the mechanisms that facilitate these phenotypes could be targeted to reduce vector density.

3. Resistance to abiotic stress

Some vector species can survive (or are adapted to live) under adverse conditions, such as desiccation, brackish water or competitive environments, which permits colonisation of a broader range of environments. Resistance of mosquito eggs to desiccation is variable among species, and three main factors drive this variability: chitin content, egg volume and shell density [92]. The evidence that the gut associated microbiota regulates two enzymes involved in chitin synthesis (GFAT and CHS2) in An. gambiae [31] suggests the potential for
the microbiome to influence resistance to desiccation. Once eggs have hatched, larvae have to persist in their aquatic environment. While most mosquito species breed in fresh water, *Culex sitiens* and *An. sundaicus* survive in brackish water [93]. In general, the microbiome can confer resistance to salinity in plants and animals [94,95], suggesting similar advantages could be conferred by gut-associated microbes to their mosquito hosts. Finally, mosquito larvae need to survive in competitive environments surrounded by their conspecifics, other species, and even predaceous larvae. The influence of the microbiome on larval competition is still to be determined, but *Wolbachia* infection has been shown to cause a density-dependent effect on larva survival [96]. Microbes that protect mosquitoes against abiotic stresses would be good candidates for paratransgenesis as this trait would likely facilitate their spread and persistence in the mosquito population.

**Influence of microbiota on probability of vector daily survival (p)**

The probability of daily survival is the chance that a vector survives each day, and the microbiome has the potential to affect this by altering adult nutrition and fitness, interacting with other microbes and modulating insecticide resistance.

**1. Adult nutrition and fitness**

The microbiome can impact insect survival by affecting host fitness, nutrition, homeostasis and metabolism of their host. One indicator of mosquito fitness (among many others) is body size, and in general microbiota enhance development and size of mosquitoes. For example, *An. gambiae* and *An. stephensi* supplemented with *Asaia* have increased growth rates [70]. Similarly, when *An. coluzzi* mosquitoes were reared on three distinct diets larger mosquitoes where seen to harbour a greater bacterial load [97]. Mosquito larvae fed solely with either bacteria and yeast still developed, although were smaller than their counterparts fed on food sources [73], suggesting that microbes alone provide some sustenance for the insect.
Smaller mosquitoes are more susceptible to environmental stressors and thus have a reduced chance of survival [98], therefore microbe stimulation of nutrition can influence vector populations dynamics.

Adult mosquitoes obtain their nutrients from two food sources, sugar and blood, and the microbiomes play an important role at food digestion and nutrient provision. *Enterobacteriaceae* is the most active family of the gut microbiota of *Ae. albopictus* at assimilating fructose, the main sugar component of nectar [99] and this sugar will be used by bacteria as an energy source to produce other nutrients for the mosquito host. The impact of the gut associated microbiota on nutrition has also been studied in model insects, and results in these systems could shed insights into mechanisms occurring in mosquitoes. Examples include complementation of vitamins missing from the diet in other hematophagous insects [100] and *Drosophila* [101], and altering expression of genes involved in energy storage and regulation of differentiation in *Riptortus pedestris* [102].

In *Ae. aegypti* [103] and *An. arabiensis* [104] disturbance of gut homeostasis resulted in a shortened lifespan, so inducing microbiome dysbiosis in vectors may be explored as a novel control strategy. There is precedence for microbial-based life shortening approaches with the modelling and empirical evidence suggesting the wMelPop strain of *Wolbachia* reduces transmission [105–107], however this strategy was not perused after it was apparent that *Wolbachia* interfered with pathogens meaning population replacement could be undertaken. Microbiome-mediated alterations in metabolites in the host can lead to different survival outcomes. A recent study demonstrated that bacteria which lowered methionine content of food extended *Drosophila* host lifespan [108]. Although this was tested in flies, methionine acts as a larvicide against several mosquito species such as *An. quadrimaculatus, Ae. albopictus* and *Cx. tarsalis* [109], suggesting that similar processes could occur in
mosquitoes. Another study in *Drosophila* showed that the production of lithocholic acid by the gut microbiota elongated host survival through upregulation of genes involved in glucose homeostasis [110], offering a potential target in the host to shorten lifespan. The insulin growth factor signalling pathway is central to regulation of lifespan [111–113], and it can be impacted by bacterial metabolism in mosquitoes [62], although the mechanisms are unknown.

Host-microbe symbioses are complex and are influenced by host physiology, microbial composition, and the timing of infection. The lifespan of *An. coluzzii* is extended with exposure to doxycycline but shortened with azithromycin [114], suggesting that changes in microbiome composition are driving this phenotype, although direct effects from the antibiotic need to be considered. Similarly, axenically reared or antibiotic-treated *Drosophila* had reduced lifespans, but if flies were exposed to bacteria in their first week as adults, their lifespan was similar to their conventionally reared counterparts [115]. In contrast, a study that compared axenic *D. melanogaster* with gnotobiotic flies infected with *Acetobacter pomorum* found no differences in survival. However, axenic flies had greater glucose levels and lower oxygen consumption, suggesting a potential overall slowing of respiration [116]. These findings indicate that the host changes caused by microbiota may manifest as intermediate phenotypes rather than detectable changes in lifespan and thus studies that measure overall fitness outcomes may miss subtle effects of the microbiota. Further work is needed to identify which affected host functions contribute to longevity, and whether similar longevity phenotypes may obscure other trait differences. Host-microbe interactions become even more complex when some members of the microbiome shift from a commensal to a pathogen status and vice versa [117]. This can happen due to temperature, presence of pathogens, and other unknown factors [118,119]. The possibility for this transition of symbiotic status and the general complexities of the host-microbe interactions
should not be ignored when considering basic research questions and ultimately when considering microbiome control strategies.

2. Microbe – microbe interactions

The diverse microbes that reside within insects may interact with pathogens that are detrimental to the host, making the vector either more resistant, tolerant or susceptible to infection and thus impacting lifespan. For example *Rickettsia*, an endosymbiont of whiteflies reduces the density of pathogenic *Pseudomonas* resulting in an extended lifespan [120]. In contrast, the infection of mosquitoes with the pathogenic fungus *Beauveria bassiana* causes microbiome dysbiosis and over proliferation and translocation of *Serratia marcescens* from the gut to the hemocoel subsequently killing the insect [121]. Microbes can benefit from each other, like symbiotic bacteria and yeast in *Drosophila* [122], but they can also exclude one another, like *Enterobacteriaceae* and *Serratia* [123] or *Asaia* and *Wolbachia* [124] in mosquitoes. These complex interactions determine microbiome composition and colonisation of the host [10,125], influencing host physiology and lifespan [126] and therefore the effectiveness of microbial control of mosquitoes [127].

3. Insecticide resistance

Gut-associated microbiota may indirectly affect mosquito lifespan by mediating resistance to insecticides. Evidence is emerging that mosquitoes with differing resistance status have distinct microbiomes [128,129], but further work is required to investigate the causality and the mechanisms underpinning these interactions. *Streptococcus, Pseudomonas, Klebsiella* and *Pantoea* correlated with insecticide resistance in *An. arabiensis* [130], *An. albimanus* [131,132] and *An. stephensi* [133]. *Wolbachia* has also been associated with insecticide resistance in *Culex pipiens* [134]. Detoxifying symbionts in the gut microbiome have been shown to confer insecticide resistance in other insects like wasps [135], honeybees [136]...
and other insect pests [137]. Although the mechanisms have not been described in mosquitoes, the ability of some of these bacteria to degrade insecticides [131] provides a possible explanation. Additionally, bacteria present in the soil may become resistant to insecticides due to chronic exposure [138] and these bacteria may colonize insects, either transiently or stably. A more complete understanding of the role of the microbiome on insecticide resistance will enable the development of strategies to mitigate the emergence of resistance and extend the longevity of currently used formulations.

**Influence of microbiota on vector biting rate (a)**

Vector biting rate is the average number of times that a vector bites per unit of time and can be modulated by the microbiome by impacting feeding behaviour and host preference. An increased biting rate leads to a higher vectorial capacity, since the vector has more opportunities to acquire and transmit pathogens. Feeding behaviour is disrupted in Ae. aegypti by Serratia [139] and in Anopheles mosquitoes when exposed to heat-killed E. coli [140] or Chromobacterium [141]. Microbiota also have the potential to affect host seeking behaviour through modulation of their chemosensory system. In D. melanogaster, symbionts determine larval pheromone preference [74] and affect the adult olfactory system, influencing food choice [142–144]. Additionally, gut bacteria are known to modulate expression levels of vitellogenin genes in the true bug, Riptortus pedestris [102], and in Ae. albopictus, vitellogenin expression regulates host-seeking behaviour [145]. Therefore, the ability of the microbiome to impact host seeking behaviour through modulation of vitellogenesis should be further investigated.

**Conclusions and future perspectives**

There is emerging evidence that the microbiome of vectors can influence many traits important for vectorial capacity. At the same time, many studies highlight the complexities
of microbial communities and variability of the microbiome in mosquitoes. Attempts to disentangle this complexity often examine the effect of a specific microbe on the host, such as those that exploit mono-axenic gnotobiotic infections, however it is unclear if these findings translate to mosquitoes with a complete microbiome consisting of many microbes. Additionally, applied strategies need to be effective in mosquitoes with divergent microbiomes which mosquitoes possess in the field. Other challenges for the scientific community to solve include moving beyond simply describing the microbiome of distinct mosquito cohorts or mosquitoes with differing treatments and validate microbes or microbial consortia that are the causal agent of phenotypes in the host followed by investigating the mechanisms responsible for these interactions. Additionally, a more complete understanding of microbiome acquisition and factors that affect composition and abundance will be essential to develop strategies to introduce microbes into field mosquito populations. Progress in these areas will help unleash the full potential of the microbiome for vector control which will include strategies that modulate vectorial capacity.

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Authors’ contributions

CCU and SZ performed the literature review and wrote the manuscript. GLH and KLC assisted in reviewing literature and provided critical and intellectual input to the manuscript. PJM provided useful comments and edits on the manuscript. All authors read and approved the final manuscript. The authors declare that they have no competing interests.
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Figure. 1. Vectorial capacity (VC) equation and the effects of the microbiome on mosquito vectorial capacity. The mosquito microbiome can modulate the five components of vectorial capacity. It can affect the probability of vector daily survival (p) by interacting with the host and the pathogen or by affecting insecticide resistance. It can also modulate biting rate (a) pathogen extrinsic incubation periods (N) and vector competence (b). Vector density (m) can also be modulated by the microbiome through effects on growth and development, reproductive output and their life under adverse conditions.