Identification of *Spiroplasma insolitum* symbionts in *Anopheles gambiae* [version 1; peer review: 2 approved, 1 not approved]

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Abstract

**Background:** Insect symbionts have the potential to block the transmission of vector-borne diseases by their hosts. The advancement of a symbiont-based transmission blocking strategy for malaria requires the identification and study of *Anopheles* symbionts.

**Methods:** High throughput 16S amplicon sequencing was used to profile the bacteria associated with *Anopheles gambiae sensu lato* and identify potential symbionts. The polymerase chain reaction (PCR) with specific primers were subsequently used to monitor symbiont prevalence in field populations, as well as symbiont transmission patterns.

**Results:** We report the discovery of the bacterial symbiont, *Spiroplasma*, in *Anopheles gambiae* in Kenya. We determine that geographically dispersed *Anopheles gambiae* populations in Kenya are infected with *Spiroplasma* at low prevalence levels. Molecular phylogenetics indicates that this *Anopheles gambiae* associated *Spiroplasma* is a member of the *insolitum* clade. We demonstrate that this symbiont is stably maternally transmitted across at least two generations and does not significantly affect the fecundity or egg to adult survival of its host.

**Conclusions:** In diverse insect species, *Spiroplasma* has been found to render their host resistant to infection by pathogens. The identification of a maternally transmitted strain of *Spiroplasma* in *Anopheles gambiae* may therefore open new lines of investigation for the development of symbiont-based strategies for blocking malaria transmission.

**Keywords**

symbiont, malaria, mosquito, Anopheles, Spiroplasma, Plasmodium, vector borne disease
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Introduction
Malaria remains a major health problem in many developing countries, particularly in Sub-Saharan Africa (WHO, 2015). Malaria transmission dynamics are dependent on aspects of the physiology and ecology of their vectors, Anopheles mosquitoes. Historically, the most successful malaria interventions have been aimed at controlling the vector to break the disease transmission cycle (Raghavendra et al., 2011). The wide distribution of insecticide treated bednets (ITNs) has had a significant impact on reducing the number of malaria cases over the past 15 years, accounting for more than 50% of the malaria deaths averted in this period (Bhatt et al., 2015). However, vector resistance to insecticides used in ITNs is spreading rapidly and there are clear signs of behavioral resistance; mosquitoes that formerly bit indoors are now biting outdoors where nets offer no protection (Kaballe et al., 2013). This may reverse significant reductions in malaria disease burden and therefore new strategies are desperately needed to control mosquito populations or their capacity to transmit parasites. One of the most promising new tools for controlling vector borne diseases involves bacterial symbionts that decrease the vectorial capacity of their insect hosts (Iturbe-Ormaetxe et al., 2011). These symbionts are maintained in host populations through maternal transmission and can spread through insect populations. These features render them potentially a much more sustainable and cost-effective strategy for the control of vector-borne disease transmission than conventional methods (McGraw & O’Neill, 2013).

In the last decade, there have been many significant advances in the development of symbiont-based strategies for arboviral disease control, primarily centred on the bacterial symbiont, Wolbachia (Jeffries & Walker, 2016). Wolbachia can be transinfected into Ae. aegypti and Ae. albopictus where it blocks the transmission of arboviruses including Dengue, Chikungunya, Yellow Fever and Zika (Bian et al., 2010; Blagrove et al., 2013; Blagrove et al., 2012; Dutra et al., 2016; Ferguson et al., 2015; Moreira et al., 2009; van den hurk et al., 2012; Walker et al., 2011; Ye et al., 2015). In addition, Wolbachia-induced reproductive manipulation (cytoplasmic incompatibility) can drive the rapid spread of this endosymbiont through wild Ae. aegypti populations (Dutra et al., 2016; Hoffmann et al., 2011). While there is much interest in using a similar strategy to control malaria, there has been limited progress in identifying suitable, maternally transmitted symbionts in Anopheles mosquitoes. Numerous studies failed to identify Wolbachia from Anopheles species (Ricci et al., 2002), and although transinfestation of Anopheles stephensi has been achieved (Bian et al., 2013), attempts to generate stable transinfected lines of An. gambiae have remained unsuccessful. Wolbachia has more recently been reported at low frequency, and very low apparent density, from certain field populations of Anopheles coluzzii and An. gambiae (Baldini et al., 2014). The natural Wolbachia – Anopheles gambiae system reported seems unlikely to have the characteristics required for development as a transmission blocking strategy since it has not been possible to select lines with high density and stable maternal transmission (Shaw et al., 2016).

To advance the prospect of a symbiont-based strategy for malaria control it will be important to continue to identify, generate and study a broad range of Anopheles – symbiont systems. Spiroplasmas are members of the Mollicutes, a bacterial group that split from a Gram-positive clostridial lineage of the eubacteria around 600–800 mya and has undergone degenerative evolution. Spiroplasmas are arthropod ‘specialists’ and all known species have some form of interaction with this clade (Gasparich et al., 2004). Members of this genus are functionally diverse, exhibiting a broad array of infection and transmission strategies: they can be pathogens, commensals or mutualists and rely on vertical or horizontal transmission (Regassa & Gasparich, 2006). In addition, Spiroplasma can confer a variety of insect hosts with resistance to a range of eukaryotic parasites, including nematodes, parasitoids and fungal pathogens (Jaenike et al., 2010; Lukasik et al., 2013; Xie et al., 2010), and they are therefore a good candidate for a symbiont that could be useful for control of Plasmodium.

Several Anopheles mosquito microbiome surveys have identified Spiroplasma from pathogenic clades (Lindh et al., 2005; Segata et al., 2016). In this study, we detected the presence of a novel strain of Spiroplasma in Anopheles gambiae mosquitoes. We sampled Anopheles gambiae sensu lato (s.l.) populations from geographically dispersed study sites in Kenya and found that the strain was present at low frequencies across both regions. We have also demonstrated that this Anopheles associated Spiroplasma is maternally transmitted.

Methods
Sampling sites and mosquito collection
Mosquitoes were collected in Karima (0° 41.373’S; 37°19.742’E) and Mbui-Njeru (0° 41.911’S; 37° 20.929’E) villages in Central Kenya region (Mwea) and Kirindo (0° 26’33.1’S 34° 14’58.9’E), Nyawiya (0° 26.7547’S; 34°15.0548’E) and Mageta Island (0° 07.1468’S and 34° 01.018’E) in Western Kenya between April 2016 and July 2017. Karima and Mbui-Njeru villages are located in Mwea, a rice producing region, where rice paddies and associated irrigation canals surrounding the villages provide suitable breeding habitats for Anopheles mosquitoes, resulting in very high Anopheles mosquito density (Mwangangi et al., 2010). The annual rainfall varies from a maximum of 1,626 mm to a minimum of 356 mm, with an average of 950 mm per year. The average temperatures are 21.3°C (range: 16.0 to 26.5°C) and the relative humidity averages 59.5% (range: 52 to 67%). The Western Kenya region lies within the Kenyan part of the Lake Victoria basin. The main socio-economic activities are small scale fishing and farming. Small sun-lit pools are the main larval habitats for Anopheles, mosquito densities are significantly lower than the Mwea region and highly seasonal. The region receives between 250mm and 1200mm of rainfall annually, with the average annual rainfall estimated at 1,100mm. The average temperatures are 22.3°C (range: 15.0 to 28.5°C) and the relative humidity averages 60.5% (range: 51 to 68%). Mosquitoes were collected by manual aspiration in houses and livestock sheds and collection of Anopheles mosquito larvae. All collected mosquitoes used in this study were identified as Anopheles gambiae s.l.
prior to analysis. Mosquito rearing was done in accordance with centre-wide approved standard operating procedures and occupational health and safety guidelines. The study protocol (NON-KEMRI 545) was approved by the Ethical Review Committee of the Kenya Medical Research Institute (KEMRI/RES/7/3/1).

High-throughput 16S rRNA amplicon sequencing

To maximize our chances of detecting potential symbionts we pooled 10 mosquitoes from each location (Central Kenya and Western Kenya). The pools were comprised of DNA from mosquito ovaries (5 mosquitoes) and whole mosquitoes (5 mosquito samples), since endosymbionts are generally at highest density in ovaries but can also be found in high densities in other tissues. The DNA samples were sent to the Research and Testing Laboratory (Lubbock, Texas, USA) for PCR amplification with ‘universal’ 16S rDNA primers (Lane et al., 1991; Lane et al., 1985), followed by MiSeq illumina sequencing. Samples were amplified in a two-step process that involved 25 µl reaction using Qiagen Hotstart Taq mastermix (Qiagen Inc, Valencia, California, USA), 1 µl of each 5 µM primer, and 1 µl of template. Reactions were performed on ABI Veriti thermocyclers (Applied Biosystems, Carlsbad, California). The PCR cycling conditions were 95°C for 5 min, then 25 cycles of 94°C for 30 s, 54°C for 40 s, 72°C for 1 min, followed by one cycle of 72°C for 10 min and 4°C hold. Amplified products were visualized with eGels (Life Technologies, Grand Island, New York, USA). Products were then pooled in equimolar concentrations and each pool was selected using Agencourt AMPure XP (BeckmanCoulter, Indianapolis, Indiana, USA) in ratios of 0.75. The selected pools were then quantified using the Qubit 2.0 fluorometer (Life Technologies, Grand Island, New York, USA) and loaded on a MiSeq Illumina (Illumina, Inc. San Diego, California, USA) 2x300 flow cell at 10 pM. The High-throughput 16S rRNA amplicon sequences reported in this study have been deposited in NCBI under Bioproject number PRJNA399254, Biosample Accession number SAMN07528657 and SAMN07528758.

_Spiroplasma_ screening by PCR and DNA sequencing

DNA was extracted from individual whole mosquitoes and tissues of _An. gambiae_ s.l. using a previously described protein precipitation method (Herren & Lemaître, 2011). To screen samples for the presence of _Spiroplasma insolitum_ GAMB we used the primers RPOB3044F_ALL and RPOB3284R_INS targeting the 350 bp region of the _rpoB_ gene. For molecular phylogenetic analyses, we used the primers SINSFTSZ294F and SINSFTSZ727R (see Supplementary Table 1 for the full list of primers used). The reactions were performed in a 10 µl reaction volume that included 5X HOT FIREPol Blend Master Mix (Solis BioDyne, Tartu, Estonia) and 2 µl of DNA template. The cycling conditions included initial enzyme activation at 95°C for 15 min, followed by 35 cycles of denaturation at 95°C for 30 s, annealing at 58°C for 30 s, elongation at 72°C for 30 s, then a hold temperature of 72°C for 7 min. Once samples were found positive for _Spiroplasma_, sequencing of amplicons was performed on a number of samples for validation (Macrogen, Amsterdam, Netherlands). PCR products were cleaned prior to sequencing using the ExoSap-IT purification protocol (USB Corporation, Cleveland, Ohio, USA). The gene sequences generated in this study have been deposited into GenBank under accession numbers MF695842, MF695843, MF695844 and MF695845.

Molecular phylogenetic analysis

Sequence alignments were performed using Clustal W in Geneious 8.1.9 software (www.geneious.com, Kearse et al., 2012). The trees were constructed by the maximum-likelihood method with a Tamura-Nei model in Geneious 8.1.9 software. Support for tree topology assessed by bootstrap resampling. To determine the phylogenetic position of _Spiroplasmas_ identified in this study relative to previously identified _Spiroplasmas_, we compared sequence of 16S rRNA, _rpoB_ and _ftsZ_ genes. Nucleotide sequences of the other _Spiroplasma_ species were derived from GenBank database (accession numbers shown in Figure 3). The length of the compared sequences was 301 bp for 16S rRNA, 210 bp for _rpoB_ and 260 bp for _ftsZ_.

Establishment of iso-female lineages

_Anopheles gambiae_ larvae collected from Central Kenya region were reared in the _icipe_ mosquito insectary in Mbita, Kenya. Female mosquitoes that successfully reached adult stage were placed in standard 30cm x 30cm x 30cm rearing cages at a density of 30–100 mosquitoes per cage, ensuring a minimum of 30% males. Mosquitoes were then blood fed on _Plasmodium_ uninfected human blood, as previously described (Gouagna et al., 2003) and allowed to individually oviposit. After oviposition, eggs were counted and each female was screened for the presence of _Spiroplasma_. Adult progeny from infected mothers were counted for egg to adult survival rates and maintained to investigate transmission across multiple generations using the same experimental design. The eggs and adult progeny from some uninfected mothers were also counted to determine egg to adult survival rates. The effects of _Spiroplasma_ on female fecundity were determined using female mosquitoes collected as larvae from Central Kenya using the same strategy described above. To reduce a potential bias from non-mated females, only broods consisting of more than 10 eggs were used to evaluate fecundity.

Mitochondrial DNA analysis

To determine the diversity of mosquito mtDNA, the 655bp _ND5_ gene was amplified using the primers described by Besansky (Besansky et al., 1997). Single PCR reactions were performed on the Veriti Thermal Cycler (Applied Biosystems, Carlsbad, CA, USA). PCR cycling conditions were initial denaturation at 95°C for 15 min, followed by 35 cycles of denaturation at 95°C for 30 s, annealing at 55°C for 30 s, elongation at 72°C for 30 s, then a hold temperature of 72°C for 7 min. PCR products were visualized on a 1% agarose gels. The PCR products were purified using ExoSap-IT purification protocol (USB Corporation Cleveland, OH). The DNA Sequences were cleaned and aligned using the MUSCLE algorithm (Edgar, 2004) in Geneious 8.1.9 software. Minimum spanning haplotype network (Bandelt et al., 1999) was constructed using Popart (http://popart.otago.ac.nz).

Results

_Spiroplasma_ sequences isolated from _An. gambiae_

Two pools of whole mosquitoes and ovaries were used to generate DNA for High-throughput sequencing of 16S rDNA,
which resulted in 195,592 and 18,921 high-quality 16S rRNA sequences, from the Central Kenya pool (CK) and the Western Kenya pool (WK), respectively. Enterobacteriacea was most predominant group in CK with approximately 79% of the sequences, whereas both Propionibacteriaceae and Enterobacteriacea dominated in WK with 26% and 22%, respectively (see Figure 1). In the CK sample, a relatively small fraction of the 16S sequence reads (0.02%) were from Spiroplasmataceae. The Spiroplasma 16S sequence reads matched the 16S rDNA gene of Spiroplasma insolitum strain M55 with 100% identity. Spiroplasma insolitum M55 was originally isolated from a flower in Maryland, USA (Hackett et al., 1993).

** Spiroplasma insolitum prevalence in field populations of An. gambiae s.l. **

We developed a set of primers to target the rpoB gene of Spiroplasma insolitum. These primers were designed based on several Spiroplasma insolitum rpoB sequences from previous studies (Watanabe et al., 2013). The specificity of these primers for Spiroplasma insolitum was investigated on a panel of diverse Spiroplasma species. These primers were then used to determine the population-level prevalence of *Anopheles gambiae* associated Spiroplasma insolitum in mosquito samples obtained from Western Kenya (Kirindo, Nyawiya and Mageta Island) and Central Kenya (Mwea), see Figure 2. In all sites, mosquitoes were collected by mouth aspiration in houses across one rainy season (October–December 2016 or April–June 2017). In Mwea, we also collected *Anopheles gambiae* larvae, which were allowed to eclose before being screened for Spiroplasma as 21 day old adults. In Mwea, approximately 8% (n=490) of *An. gambiae s.l* harbored *Spiroplasma*. When collected as larvae, the rate of infection was higher, at 14% (n=163). In Western Kenya, the prevalence of *Spiroplasma* was generally lower and absent from one site. In Kirindo, the rate of *Spiroplasma* prevalence was 4% (n=173) and in Mageta the prevalence was 3% (n=66), whereas no infections were found in mosquitoes obtained from Nyawiya (n=222).

**Molecular phylogenetic analyses of Spiroplasma**

To determine the phylogenetic position of *Anopheles* associated *Spiroplasma insolitum* relative to other members of this clade, and to determine if multiple *Spiroplasma* strains are present in these populations, we developed primers to specifically target and amplify a region of the *Spiroplasma insolitum* ftsZ gene (Supplementary Table 1). In addition, we sequenced the region of rpoB amplified by our *Insolitum*-specific primers. The high-throughput sequencing that we carried out to investigate microbial diversity enabled us to obtain the sequence of a fragment of 16S rDNA. These sequences were used for the construction of phylogenetic trees, which indicate the strain of *Spiroplasma* from *An. gambiae s.l* can be classified into the citri-clade and confirm it clusters with *Spiroplasma insolitum* (see Figure 3). The 16S rDNA

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**Figure 1. Relative abundance of Bacterial families in An. gambiae from Kenya.** The relative abundance of bacterial sequences obtained from mosquitoes and mosquitoes from Central (CK) and Western Kenya (WK) Regions as determined by high throughput sequencing of 16s rRNA gene. Notably, 0.02% of the CK reads corresponded to *Spiroplasmataceae.*
Figure 2. Spiroplasma prevalence in *An. gambiae* populations. The prevalence of *Anopheles* associated *Spiroplasma* Western Kenya (blue) and Central Kenya (red). In central Kenya, mosquitoes were collected as both adults and larvae, whereas only adults were collected in Western Kenya. For each site, the prevalence is given as an average of several collections across the course of one rainy season. In central Kenya, 8% (n=490) of *An. gambiae* s.l. harbored *Spiroplasma*. When collected as larvae, the observed prevalence of infection was higher, at 14% (n=490). Western Kenya sites tended to have a lower prevalence of *Spiroplasma*; Magenta 3% (n=66), Kirindo 4% (n=173) and no infections were found in Nyawiya (n=222).

fragment sequence was found to be identical to that of three previously described strains of *Spiroplasma insolitum*, M55, TU-14 and NBRC. The sequenced region of the *rpoB* gene from *Anopheles* associated *Spiroplasma* was also found to be identical to M55, TU-14 and NBRC and two strains of *S. insolitum* that are endosymbionts of flower bugs of the genus *Orius*, SpOriA/B (Watanabe et al., 2013). The region we sequenced of the *ftsZ* gene from *Anopheles* associated *Spiroplasma* was identical to M55, TU-14 and NBRC. Notably, *ftsZ* sequence data is not available for SpOriA/B. These results indicate that this *Anopheles* associated *Spiroplasma* strain is *Spiroplasma insolitum*, henceforth referred to as *S. insolitum* GAMB. Since all the *S. insolitum* GAMB genes (*16S rDNA, rpoB* and *ftsZ*) we sequenced were identical, we find no evidence for multiple strains of *S. insolitum* co-existing in the populations of *Anopheles gambiae* s.l. studied.

*Spiroplasma insolitum* GAMB is maternally transmitted in *Anopheles gambiae* s.l but does not bias sex ratio or affect egg to adult survival

To determine if *Spiroplasma insolitum* GAMB is maternally transmitted, we collected mosquitoes from the field and established iso-female mosquito lineages. We collected larvae from Mwea (where *Spiroplasma insolitum* GAMB prevalence was highest) and maintained them until they eclosed as G₀ adults, at which point they were blood fed then allowed to oviposit prior to screening for *Spiroplasma* infection. Three G₀ females carried *Spiroplasma* and from these, individual F₁ female offspring were maintained to enable further screening for *Spiroplasma*. Most F₁ did not produce viable offspring; in only one instance we obtained F₂ females (see Figure 4). This is not altogether surprising, as field caught *Anopheles gambiae* s.l. are known to perform poorly prior to becoming ‘acclimatized’ to laboratory conditions (Diop et al., 1998). We found that *Spiroplasma insolitum* GAMB is maternally transmitted with very high efficiency, but that transmission efficiency did vary slightly between iso-female lineages. In two cases we observed perfect maternal transmission, whereas the rest had transmission efficiencies between 43% and 87%. The iso-female lineage that produced F₂s showed 100% transmission from G₀ to F₁ and 83% transmission from F₁ to F₂.

Maternally transmitted symbionts are known to manipulate the sex ratio of their hosts to gain a transmission advantage (Hurst & Majerus, 1993). To determine if *Spiroplasma insolitum* GAMB affects the sex ratio of *Anopheles* hosts, we monitored the sex ratio of offspring in *Spiroplasma* infected isofemale lineages. The sex ratio did not differ substantially from the expected 50% female/male in the two lineages producing greater than 10 progeny (see Figure 4), and therefore we conclude that *Spiroplasma insolitum* GAMB is not a male-killer.

We also monitored the fecundity and egg to adult survival rate for *Spiroplasma* infected and uninfected iso-female lineages (see Figure 5). We did not observe any significant difference between the fecundity and survival rate of *Spiroplasma* infected and uninfected individuals, indicating that *Spiroplasma insolitum* GAMB is not pathogenic.
Figure 3. Molecular phylogenetic analysis of Spiroplasma. Phylogenies are based on (a) 16S rRNA (b) rpoB (c) ftsZ. The trees were constructed by the maximum-likelihood method with a Tamura-Nei model using unambiguously aligned sites (total sites are 301, 210, 260 bp, respectively). The numbers above branches indicate a bootstrap value for 1000 replicates. Branch lengths indicate substitutions per site (based on scale bars).
Figure 4. Vertical transmission and sex ratio in *Spiroplasma* infected iso-female lineages. *Anopheles gambiae* mosquitoes collected from Mwea as larvae were used to establish iso-females lineages. Offspring from *Spiroplasma*-infected iso-female lineages were screened for the presence of *Spiroplasma* at adult stage. The number of progeny screened are shown for each female and in the subsequent generation (G₀, F₁, and F₂). At each generation the infection levels and sex ratio were monitored. The bars represent the % sex ratio of the total number of offspring from each iso-female lineage. The presence of between 40%–60% males in all but one family demonstrates that *Spiroplasma* is not a male killer. The observed maternal transmission efficiency (TE), ranged between 43% and 100% with an average of 82.6%.

Figure 5. Fecundity and egg to adult survival rates are not significantly affected by *Spiroplasma*. *Anopheles gambiae s.l.* mosquitoes collected as larvae from mwea oviposited individually. The fecundity and the survival of eggs into adulthood was monitored. The fecundity (A) and egg to adult survival rate (B) for *Spiroplasma*-infected mosquitoes was not significantly different from that of uninfected mosquitoes in unpaired t-tests. For fecundity P=0.84, t=1.976, df=62. For egg to adult survival P=0.32, t=1.012, df=50. The data shown is pooled from 3 independent experiments. Shown on the graph are the means with 95% confidence interval.

Association between *Spiroplasma insolitum* GAMB and mtDNA haplotypes

We investigated possible associations between mtDNA haplotypes and *Spiroplasma insolitum* GAMB infection. Symbionts that have recently infected an insect population and are maintained by high efficiency maternal transmission can be expected to be associated with one or a few mitochondrial DNA haplotypes. In contrast, if a symbiont infection is associated with many or most mtDNA haplotypes, this suggests an older infection, paternal transmission or an appreciable level of horizontal transmission in addition to maternal transmission. We sequenced the ND5 mtDNA gene, which has been widely used for haplotyping *Anopheles gambiae* mosquitoes (Besansky et al., 1997). Of 21 *Anopheles gambiae* specimens collected in Central Kenya (Mwea), 11 were shown to be *Spiroplasma insolitum* GAMB positive based on PCR based screening. We identified a total of 6 haplotypes, four of these are identical to haplotypes reported previously (Aboud et al., 2014; Besansky et al., 1997), while two were novel (M1 and M2, see Figure 6). In the
GAMB is maternally transmitted. We (Hurst & Majerus, 1993). Male is known to achieve trans-ovarial (Mwea). Other microorganisms (e.g., Spiroplasma) are known to persist in the intestinal tract of Anopheles gambiae s.l. mosquitoes (Goto et al., 2006) and are likely transmitted to the offspring via the fecal-oral route. However, the transmission efficiency of GAMB from mother to offspring is not well understood.

Figure 6. Minimum spanning haplotype network for mitochondrial ND5 gene for Spiroplasma-infected and uninfected mosquitoes from Mwea. The population genetics of Spiroplasma-infected and uninfected mosquitoes based on ND5 mitochondrial DNA loci. Minimum spanning haplotype network reflects loci from 21 mosquitoes, 11 of these were Spiroplasma-infected. The circle size corresponds to the haplotype frequencies. The shading reflects the proportion of each haplotype that is Spiroplasma-infected. Notably, the majority of haplotypes (B1, H47, M2 and B41) had both Spiroplasma-infected and uninfected individuals.

4 most common haplotypes, we found both Spiroplasma insolitum GAMB infected and uninfected individuals, suggesting that this symbiont is either an ancient infection or exhibits appreciable levels of horizontal transmission.

Discussion

We have identified a strain of Spiroplasma that is associated with Anopheles gambiae mosquitoes in Kenya. This Spiroplasma was initially identified from 16S rDNA high throughput sequencing reads from a pool of Anopheles gambiae mosquitoes from Central Kenya (Mwea). 16S rDNA sequence revealed that this Anopheles associated Spiroplasma strain was a member of the citri clade and grouped with Spiroplasma insolitum. Most of the known insect endosymbiotic Spiroplasmas are found in this clade, for example, S. poulsonii, S. citri and S. insolitum are all species which have been studied as insect endosymbionts (Haselkorn, 2010; Watanabe et al., 2014). It is notable that Spiroplasma insolitum GAMB is very closely related to SsPori/A/B, a strain of Spiroplasma insolitum that is an endosymbiont of flower bugs in the genus Orius (Watanabe et al., 2014). We are aware of two other studies that identified Spiroplasma sequence associated with Anopheles mosquitoes. An investigation on midgut bacteria in Anopheles gambiae and Anopheles funestus from Western Kenya detected 16S sequence corresponding to Spiroplasma in Anopheles funestus (Lindh et al., 2005). Another study investigated the microbiome of the reproductive tracts of Anopheles gambiae and Anopheles coluzzi in Burkina Faso and found evidence for the presence of Spiroplasma in both species (Segata et al., 2016). In both studies, the Spiroplasma identified appears to be closely related to Spiroplasma ixodetis, which was initially discovered as a pathogen associated with ticks (Tully et al., 1995), and is thus quite different from the Spiroplasma identified in our study.

We demonstrate that Spiroplasma insolitum GAMB is found at relatively low frequencies in Anopheles gambiae s.l. mosquito populations. Frequencies tended to be higher in the Central Kenya Region (Mwea) than in the Western Kenya Region (Magenta, Kirindo and Nyawinya). These two regions have quite different mosquito habitats; in western Kenya most mosquitoes emerge from isolated puddles whereas Mwea (Central Kenya) is a rice growing region where Anopheles larvae are abundant in rice paddies and irrigation canals. While both sites experience an increase in mosquito abundance during the rainy season, the difference is less pronounced in Mwea due to year round irrigation providing more permanent larval habitats. We also noted that a higher infection rate was observed in Mwea when we collected larvae instead of adults. A possible explanation for this is that a greater number of these samples had Spiroplasma levels that were above the detection limit, or above levels required for the bacteria to be maintained through pupal morphological re-organization into the adult stage. This could be due the favorable laboratory larval and adult rearing conditions, which would likely result in more nutrients available to host and symbiont. Additionally, the mosquitoes collected as larvae (aged to 21 days) were likely to be older than field caught mosquitoes (unknown age). Spiroplasma densities in insects are known to significantly increase over the life of the host (Goto et al., 2006; Herren et al., 2014) and this could also have caused the observed increase in number of Spiroplasma positives.

Since insect associated Spiroplasma are known to exhibit a variety of different transmission patterns it was important to determine if Spiroplasma insolitum GAMB is maternally transmitted. We established iso-female lineages from infected field collected larvae and demonstrated that maternal transmission does occur with a high level of efficiency. We note that transmission efficiency does appear to vary slightly between the iso-female lineages tested. Two families exhibited very high maternal transmission to F1s, whereas transmission efficiency for the third was about 50%, however the third only generated 6 female offspring. While the spatial localization and mechanistic basis of Spiroplasma insolitum GAMB maternal transmission were not investigated, the closely related Spiroplasma poulsonii is known to achieve trans-ovarial maternal transmission by subversion of the yolk uptake pathway in Drosophila melanogaster (Herren et al., 2013). Other microbial symbionts that persist in the intestinal tract are more likely to achieve maternal transmission by the fecal-oral route. While trans-ovarial maternal transmission tends to be higher efficiency, there are reports of very high efficiency transmission via the fecal-oral route (Hosokawa et al., 2015).

Many maternally transmitted insect symbionts have evolved strategies to bias sex ratio towards females to gain a transmission advantage (Werren & O’Neill, 1997). The most common manifestation of this is male-killing in which the endosymbiont confers male-specific embryonic lethality (Hurst & Majerus, 1993). Male
killing has been observed in numerous strains of endosymbiotic Spiroplasma (Anbutsu & Fukatsu, 2011). We monitored the sex ratio of the Spiroplasma-carrying lineages and found very close to 50% male offspring in the two lineages where more than ten offspring could be examined, suggesting that Spiroplasma insolitum GAMB is not a male killer.

A number of Spiroplasma are known to be pathogenic to their arthropod hosts (Clark et al., 1985; Nunan et al., 2005). By monitoring the fecundity and egg to adult survival rate we determined that Spiroplasma insolitum GAMB was not pathogenic. This finding when coupled with its phylogenetic position suggests that Spiroplasma insolitum GAMB is likely to either be a commensal or mutualist, although adult fitness assays are also needed. From the standpoint of developing Spiroplasma insolitum GAMB as part of a future microbe based transmission blocking strategy this is advantageous, as a pathogenic phenotype could limit the capacity for Spiroplasma to spread through the host population.

We did not observe a clear correlation between mtDNA haplotype and Spiroplasma infection. This could be due to two major possibilities. The first, that Spiroplasma infection has been maintained in this species for a very long period of time enabling diversification of mtDNA within the infected lineage (as is the case for obligate symbionts, Moran, 2006). A second possibility is that there is significant horizontal transmission of Spiroplasma between the An. gambiae mosquitoes and/or that paternal as well as maternal transmission occurs, resulting in the wide and almost even distribution of Spiroplasma infection between mitochondrial haplotypes. Given that Spiroplasma insolitum GAMB is not an obligate symbiont (as it is not found in all Anopheles gambiae s.l. individuals), it seems most likely that Spiroplasma insolitum GAMB is both horizontally and vertically transmitted. The phylogeny of Spiroplasma also suggests a high frequency of horizontal as well as vertical transmission (Haselkorn et al., 2009) and many Spiroplasma likely utilize both forms of transmission. From the standpoint of using Spiroplasma as a tool for blocking VBD transmission, the prospect of strains being both vertically and horizontally transmitted is of considerable interest and could render them easier to spread through host populations.

We have reported the identification of novel strain of Spiroplasma in Anopheles gambiae s.l. mosquitoes. Questions that now need to be addressed, once stable infected colonies have been successfully created, include the effects of these Spiroplasma on Plasmodium transmission, effects on adult lifespan, where it localizes within the mosquito, and mechanisms of vertical and horizontal transmission. Spiroplasmas are known to protect a variety of insect hosts from diverse parasites (Jaenike et al., 2010; Lukasik et al., 2013; Xie et al., 2010) and therefore the discovery of Spiroplasma insolitum GAMB could provide a step towards the development of novel malaria control strategies.

Data availability
Figshare: Identification of Spiroplasma insolitum symbionts in Anopheles gambiae

Dataset 1: Spiroplasma prevalence in Anopheles gambiae s.l. field populations. (Herren, 2017a)
Dataset DOI: http://dx.doi.org/10.6084/m9.figshare.5384089.v4

Dataset 2: Spiroplasma transmission and effects on survival and fecundity. (Herren, 2017b)
Dataset DOI: http://dx.doi.org/10.6084/m9.figshare.5384101.v3

The High-throughput 16S rRNA amplicon sequences reported in this study have been deposited in NCBI under Bioproject number PRJNA399254, Biosample Accession number SAMN07528657 and SAMN07528758.

The gene sequence data has been deposited in Genbank under accession numbers MF695842, MF695843, MF695844 and MF695845.

Competing interests
No competing interests were disclosed.

Grant information
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The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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Supplementary material
Supplementary Table 1: Primer sequences used to investigate Spiroplasma and host mtDNA haplotype in Anopheles gambiae s.l.
Click here to access the data.


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In this paper the authors analyze the microbiota of Anopheles gambiae mosquitoes from 3 geographically distinct regions of Kenya, with an emphasis on the structure (and to a lesser extent, function) of Spiroplasma populations within infected hosts. This analysis is performed by 1) deep sequencing of 16s rRNA sequences from whole mosquitoes and mosquito ovaries, 2) PCR screening for Spiroplasma specifically, and 3) observing the effect of Spiroplasma infection on female reproductive biology (maternal transmission, fecundity, offspring sex ratio and egg to adult survival). This study, which provides further confirmation that Spiroplasma is present in An. gambiae, is an important contribution to our collective knowledge of this bacterium's population structure within this prolific mosquito disease vector.

I have a few questions/comments regarding the manuscript in its current form:

The high-throughput experimental design and data representation is ambiguous in places.

- In the Methods, I believe the authors are trying to say that they made 5 samples of ovary and whole organism DNA, each of which contained material from 10 mosquitoes. However, as written in the paper, this is difficult to understand.

- The authors indicate that they used whole mosquitoes "since endosymbionts are generally at highest density in ovaries but can also be found in high densities in other tissues." If they were interested in characterizing the microbiota found outside of the ovaries, then it might have been a better idea to sequence the 16s rRNA from mosquito carcasses that had the ovaries removed.

- The Results indicate that only 2 of these 5 pools were sequenced, and only one from each location. This sample size is low (and the Western Kenya pool gave rise to low number of reads). I suspect this analysis was performed as a preliminary indicator of whether or not symbionts are present, and thus whether or not the authors should proceed with further analyses.

- Missing from the Methods is a description of how the 16s data analysis was performed.
A final ambiguity involves the lack of information regarding microbiota tissue distribution. DNA was extracted from whole mosquitoes and ovaries. Were the whole mosquito and/or ovary specific libraries sequenced separately? Is the 16s data provided from whole mosquito or ovary samples? If libraries from both sample types were sequenced, it would be interesting if the authors presented the data in a tissue specific manner so as to correlate bacterial diversity and abundance with the distinct tissues used to make the libraries.

For all PCR assays, it would be nice to have more information regarding negative controls. To do so, the authors could have used mosquitoes that they know are Spiroplasma negative. I say this because BLAST analysis of the Rpob and FtsZ primers used in this study revealed that they exhibit very high identity with these genes in other bacteria. Can the authors conclusively rule out non-specific amplification from bacterial members of the microbiota that are not Spiroplasma?

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Partly

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
There is great interest in using host-associated microbes to control insect disease vectors. Maternally transmitted microbial symbionts are especially promising because many have evolved sophisticated strategies to spread through host populations and/or suppress interfere with insect parasites and pathogens. In this study, the authors report the discovery and initial characterization of a strain of Spiroplasma insolitum that infects Anopheles gambiae mosquitoes. It occurs at low frequency in western and central Kenya. The authors establish infected isofemale lines in the lab, rear them for two generations, and show that offspring are infected as well (although transmission efficiency is often low). Mitochondrial sequence analysis suggests that horizontal transmission is pervasive. This is an important contribution to the study of mosquito symbiosis and it will be interesting to characterize this symbiont in greater detail.

It would be useful to have more information about the establishment and rearing of the iso-female lines (this is not always clear), because it is a bit difficult to interpret the results on transmission to offspring. Were some or all offspring screened as eggs? Were some or all offspring reared individually? If offspring were reared in the same container as larvae, then it is possible that there was horizontal transmission from adult females, and then between larvae. It would be useful to rear infected and uninfected individuals together. If uninfected mosquitoes do not pick up the infection from infected ones, then vertical transmission may predominate.

Some other comments:

- It would be useful to have a sentence explaining why Spiroplasma-infected mosquitoes were only kept in the lab for 2 generations.

- The fact that the strain is identical at all genes sequenced in the study to S. insolitum from insects from other orders also suggests predominantly horizontal transmission. It would also be useful to mention more clearly that a number of cultivable/horizontally transmitted Spiroplasma have been isolated from (non-anopheline) mosquitoes.

- In Figure 4, it would be useful to add arrows to indicate which F1 lines are the mothers of the F2.

- It would be useful to have more information about Figure 5. Was transmission to offspring measured in these Spiroplasma-infected females? Which mothers are these (connect to Figure 4? F1)?

- Why were mosquitoes collected as larvae older than field-caught mosquitoes (second last page)?

Is the work clearly and accurately presented and does it cite the current literature? Yes

Is the study design appropriate and is the work technically sound? Yes

Are sufficient details of methods and analysis provided to allow replication by others? Partly

If applicable, is the statistical analysis and its interpretation appropriate? Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Partly

**Competing Interests:** Collaborate with former supervisor of two of the coauthors.

**Reviewer Expertise:** Insect parasites and symbionts

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 06 October 2017

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The authors show the relative abundance bacteria families found in *Anopheles gambiae* mosquitoes collected in West Kenya (3 areas) and Central Kenya (2 areas). Thereafter, the authors focus the entire paper on *Spiroplasma, insolitum*, and claim that this species is different from *Spiroplasma* already described in African *Anopheles* mosquitoes.

1. The paper should be revised and written more carefully. There are incomplete sentences and also errors in graph (Figure 2, pie graph Mwea does not appear to be 14%).

2. The methodology is poorly described. One example is that the rearing procedures after the mosquitoes were brought from field to laboratory were not described, and in this case are crucial. Another important point: How were the field mosquitoed treated during sample preparation for sequencing? Were the males used to cross with field collected females laboratory sterile mosquitoes, or they were infected too?

3. Moreover the experimental design could be improved considerably:

   - The number of mosquitoes used in different analysis is still too low! Sample size needs to be higher.
   - Although authors say that they had pools with samples of 5 whole mosquitoes and samples of 5 mosquito ovaries, nothing more is done or said about the differences among the two pools. It would be interesting to also include male in the analysis, as well as compare different tissues (including midgut, ovaries, testis and accessory glands, for example).
- Include sterile laboratory mosquitoes as controls.
- Authors should compare results using cultivable and culture-independent methods.
- Experiments performed to verify vertical transmission and sex ratio determination were performed with low number of mosquitoes, two F1 had really low number of offspring and most of F1 produced no offspring. So, the conclusions to address those questions (maternal transmission and sex ratios) were made based on very few mosquitoes, that came from the low number of parental mosquitoes that were able to produce viable offspring.
- Fecundity assays performed with low number of mosquitoes. Were those assays made with G0? How those results correlate with the data of Figure 4?
- In Figure 5, instead of means, this type of data requires medians.
- It is important to keep in mind that the relative abundance of Spiroplasma, based on data that the authors had shown (with a low number of pools), is quite low, and the possibility that other bacteria play a role in fecundity and survival rate for example, cannot be ruled out. This was not addressed or discussed.

4. In my opinion data provided and sample size are not enough to support the conclusions.

5. Not clear why authors focused on Spiroplasma, that has low relative abundance and prevalence in the areas and claim they might be good symbiont to protect mosquitoes from Plasmodium. A lot needs to be done and better addressed.

6. Discussion is still poor.

Is the work clearly and accurately presented and does it cite the current literature?
Partly

Is the study design appropriate and is the work technically sound?
No

Are sufficient details of methods and analysis provided to allow replication by others?
Partly

If applicable, is the statistical analysis and its interpretation appropriate?
Partly

Are all the source data underlying the results available to ensure full reproducibility?
Partly

Are the conclusions drawn adequately supported by the results?
No

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Biochemistry, vector biology, vector-parasite interaction

We have read this submission. We believe that we have an appropriate level of expertise to state that we do not consider it to be of an acceptable scientific standard, for reasons outlined above.