Exploring malaria vector diversity on the Amazon Frontier

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Abstract

Background: Deforestation in the Amazon and the social vulnerability of its settler communities has been associated with increased malaria incidence. The feeding biology of the most important malaria vectors in the region, notably *Nyssorhynchus darlingi*, compounds efforts to control vectors and reduce transmission of what has become known as “Frontier Malaria”. Exploring Anophelinae mosquito diversity is fundamental to understanding the species responsible for transmission and developing appropriate management and intervention strategies for malaria control in the Amazon River basin.

Methods: This study describes Anophelinae mosquito diversity from settler communities affected by Frontier Malaria in the states of Acre, Amazonas and Rondônia by analysing COI gene data using cluster and tree-based species delimitation approaches.

Results: In total, 270 specimens from collection sites were sequenced and these were combined with 151 reference (GenBank) sequences in the analysis to assist in species identification. Conservative estimates found that the number of species collected at these sites was between 23 (mPTP partition) and 27 (strict ABGD partition) species, up to 13 of which appeared to be new. *Nyssorhynchus triannulatus* and *Nyssorhynchus braziliensis* displayed exceptional levels of intraspecific genetic diversity but there was little to no support for putative species complex status.

Conclusions: This study demonstrates that Anophelinae mosquito diversity continues to be underestimated in poorly sampled areas where frontier malaria is a major public health concern. The findings will help shape future studies of vector incrimination and transmission dynamics in these areas and support efforts to develop more effective vector control and transmission reduction strategies in settler communities in the Amazon River basin.

Keywords: Malaria, Species discovery, Mosquito, Anophelinae, Amazon, Deforestation

Background

Deforestation is the permanent destruction of forests in order to harvest timber, develop farms and pasture, and build roads and urban areas. Many of the world’s most biologically diverse regions are now subject to the highest rates of deforestation and one of the most severely affected of these is the Amazon tropical rainforest. In Brazil over the past 50 years, deforestation in the Amazon has reached unprecedented levels, with estimated losses of between 4 and 29 thousand km² annually [1]. During the same period, the human population in the region has increased from approximately 4 to 24 million [2]. Additional effects of regional deforestation include a dramatic loss of endemic species [3], creation of social conflict, contribution to global climate change, and elevated human vulnerability to socio-environmental conditions [4]. It is also having a negative impact on public health [5–7], and increases vector-borne diseases in the Amazon River basin [8]. One of the most important public health impacts is the increased transmission of *Plasmodium* species responsible for human malaria [9–13].

The increased incidence of *Plasmodium* infection in humans in Amazonian villages resulting from...
deforestation has been described as “Frontier Malaria”. This has been characterized as a temporal process comprised of epidemic, transition, and endemic phases [14, 15]. The epidemic phase involves changes in the natural forest landscape, including the biotic and abiotic conditions of forest larval habitats, that can favour species that are actively involved in the transmission of the *Plasmodium* to humans. Ongoing anthropogenic changes in the environment, which alter the abiotic characteristics and ecology of larval habitats [16], lead to an increase in abundance of the most important vector species and, ultimately, higher biting rates and *Plasmodium* infection in settler communities with poor housing conditions, lack of access to health services and low or no immunity to the pathogens [17]. The transition phase occurs several years after the initial settlements and sees gradual declines of transmission as a result of improved housing, infrastructure and public health services. The final endemic phase is reached when transmission reaches low and stable levels, normally within 10 years of the initial settlement. A developed public health service can achieve effective diagnosis and treatment of *Plasmodium falciparum* infection, due to symptoms occurring before gametocyte production. *Plasmodium vivax* infection, however, is more difficult to diagnose and treat, as the pathogen can exist at much lower densities [18] and persist as hypnozoites in the liver [19]. The only licensed treatments of the liver stage of *P. vivax* are primaquine and tafenoquine, but they are of limited use in some populations due to the adverse effect of acute haemolysis in people with G6PD deficiency [20, 21]. *Plasmodium falciparum* transmission emerges in the early stages of settlement and is joined by *P. vivax* in later stages, with the latter generally becoming more prevalent. An emerging public health consensus is that the dynamics of malaria in the Amazon River basin is unstable, with waves of disease emergence accompanied by explosive epidemics in many localities. These are usually associated with changes to natural environments and ecologies, waves of economic development, and migratory influxes between endemic and non-endemic areas. This dynamic process has challenged control programmes developed to mitigate the burden of malaria in the Amazon [22].

There are several vectors of malaria in the Neotropics that vary in importance at the local, regional and continental scale [23]. Although the highest rates of experimental *Plasmodium* infection have been found in *Nyssorhynchus aquasalis* and *Nyssorhynchus albitalis* [24], with the latter presenting higher biting indices [25], *Nyssorhynchus darlingi* has the highest rates of natural infection and is seen as the most important malaria vector through much of South America [25, 26] (herein *Nyssorhynchus* is elevated from subgenus to genus rank, as proposed by Foster et al. [27]). This species is the most anthropophilic, endophilic and opportunistic vector, and is relatively abundant in the more populated localities of the Amazon [16]. It is also strongly associated with human environments in deforested areas in the Amazon [28, 29].

A great variety of potential vectors have been recorded in the deforested areas of the Amazon [30–32], but the frequent occurrence of morphologically indistinguishable sibling species, which form species complexes [33], means that vectors are often misidentified or cryptic species boundaries go undetected [34–36]. Some vectors, such as *Nyssorhynchus triannulatus* and *Anopheles peryassui*, inhabit the forest edge, away from domestic environments and conventional vector control activities, but may be associated with human activity such as deforestation [37]. Failure to properly elucidate the diversity present in vector species complexes at locations where forest gives way to frontier settlements will likely obstruct the understanding of vector ecology and the *Plasmodium* transmission cycle associated with deforestation in the Amazon and impede efforts to develop strategies to control frontier malaria [10, 38]. It is, therefore, essential to develop and employ a range of tools to discover potential vectors in malaria endemic areas.

Molecular tools are commonly employed to identify known species and delimit new species [39]. DNA barcoding is now seen as an important first step in biodiversity assessment and sorting specimens into tentative species, and COI the marker of choice because of the availability of universal primers that amplify across a diverse range of species, and the relatively low intraspecific and high interspecific divergence that occurs at this locus. It is clear that barcoding approaches are helping to resolve morphologically indistinguishable species within Anophelineae [31, 34, 40, 41].

Although barcoding studies frequently employ phylogenetic (and normally NJ) trees to delimit species, resolving intraspecific from interspecific phylogenetic structure is frequently subject to observer bias [42, 43]. Approaches explicitly designed for empirically and objectively delimiting species boundaries have been developed, among which are clustering and tree-based approaches, which do not require a priori definitions of taxa [44] or even a threshold for intraspecific diversity [45]. The use of such approaches can therefore allow species richness and delimitation to be independently estimated and tested against a priori species morphological definition.

The current study aims to describe Anophelineae species diversity in rural settlements affected by frontier malaria in the Amazon River basin, employing clustering-based
(Automatic Barcode Gap Discovery; [44]) and tree-based (multi-rate Poisson tree processes; [45]) species delimitation approaches. The ability to describe diversity and delimit species in this setting is fundamental to understanding the vectors driving malaria transmission and allows for the development of more effective vector control and management strategies in frontier settlements in the Amazon River basin. In the study we pay particular attention to vector complexes and the potential for these to harbour new morphologically similar and potentially medically important species.

**Methods**

**Mosquito collection**

In total, 270 mosquito specimens from 47 collection sites in the Brazilian states of Acre, Amazonas and Rondônia were included in this study (Fig. 1). Collection sites were selected based on a high annual malaria parasite index (annual parasite index, API $\geq 50$) in 2015 or 2016 and levels of forest cover depletion greater than 10%, assuming this was representative of a human colonization process in the Amazon Forest. Field females were collected at the forest edge by either Shannon trap or human landing interception, using a small, manual aspirator. Females were euthanized with ethyl acetate vapors inside a plastic...
chamber in the field, and kept in silica gel and separated hourly. Collections at each site were made on a single day from 06:00 p.m. to 06:00 a.m., during the months of January, April, May, August, and October 2015. Specimens were identified from the Genus Anopheles and the recently elevated Genus Nyssorhynchus [27]. After species identification, females were kept individually frozen at −80 °C. One or two legs were removed with small, pointed scissors and used for sequencing the 658 base pair COI barcode region of the mitochondrial genome. One hundred and fifty-nine sequences from thirty species were downloaded from GenBank and used in the phylogenetic analysis as references (Additional files 1, 2).

DNA extraction
Genomic DNA extraction was performed from 1 or 2 legs of each Anophelinæe specimen (Additional file 1). Each extraction was carried out as follows: legs were macerated in 10 µl of NaCl 0.9% with an autoclaved pistil; 20 µl of Chelex-100 5% was added then, vortexed; the solution was incubated at 99 °C for 10 min and then centrifuged at 13,000 rpm for 5 min at 25 °C; the supernatant was recovered and an aliquot was frozen at −20 °C and the remaining DNA at −70 °C in the entomological frozen collection of the Faculdade de Saúde Pública, São Paulo, Brazil.

DNA amplification
Primers described by Folmer et al. [46] were used to amplify the barcode region of the cytochrome c oxidase gene (COI). Each reaction was performed with final volume of 25 µl containing 2 µl of DNA extracted by Chelex, 1× PCR Buffer (Invitrogen), 1.5 mM MgCl2 (Invitrogen), 0.2 mM each dNTPs (Amresco), 0.1 µM each primer (LCO1490 5’-GGTCAACAAATCATATAAGATATTG G-3’ and HCO2198 5’-TAAACTTCAGGGTGACCA AAAATCTA-3’), 0.625 U Taq Platinum polymerase (Invitrogen) and the remaining volume of ultra-pure water. The thermocycler conditions were 94 °C for 3 min, 5 cycles of 94 °C for 30 s, 45 °C for 30 s, 68 °C for 60 s, followed by 35 cycles of 94 °C for 30 s, 51 °C for 30 s, 68 °C for 60 s and a final extension at 68 °C for 10 min [47]. PCR products were purified using PEG precipitation (20% polyethylene glycol 8000/2.5 M NaCl) and then analyzed in an Applied Biosystems 3130 DNA Analyser (PE Applied Biosystems). Sequences were edited in Sequencher v.4.9 software (Genes Codes Corporation, Ann Arbor, MI, USA), and the primer regions removed. In addition to these novel sequences, 162 sequences from GenBank were also included, to serve as references in the phylogenetic and species delimitation analysis. The COI gene sequences were aligned first by nucleotides using the Muscle algorithm [48] implemented in SeaView [49], and then by amino acid using TransactorX [50].

Phylogenetic analysis
A Bayesian phylogenetic analysis was applied to sequences using a codon partitioning scheme to allow different partitions to have their own model characteristics (composition, rate matrix and among-site variation) and to allow for among-partition rate variation. Optimal evolutionary models for the sequences were determined for each partition using the corrected Akaike Information Criterion (AICc) in jModelTest 2 [51]. With the exception of 2 singletons, codon position 2 was invariable and so was excluded from phylogenetic analyses. The optimal models for codon positions 1 and 3 were GTR+G and GTR+G+I, respectively. All Bayesian phylogenetic analyses were performed using MrBayes 3.2 [52] at the CIPRES Science Gateway [53] and each analysis consisted of two simultaneous runs, which were then repeated to provide confirmation of convergence of posterior probability distribution. Each Bayesian phylogenetic analysis was run for fifty million generations, with the first twenty-five million generations being discarded as burn-in. The Metropolis-coupled Markov chain Monte Carlo strategy was used with four heated chains; adequate mixing was achieved by setting the chain temperature to 0.03. Convergence of topology between the two simultaneous runs was monitored using the average standard deviation of split frequencies—this index consistently fell to below 0.01 in the post-burn-in samples. Convergence was also monitored by noting that the potential scale reduction factor values were all approximately 1.0 in the post-burn-in samples. Consensus trees were constructed containing nodes with posterior probability support of at least 70% using SumTrees [54, 55]. Pairwise Kimura-two-parameter (K2P) [56] distances were calculated from the COI alignment using APE [57].

Species delimitation
The first species delimitation method employed was the Automatic Barcode Gap Discovery—ABGD [44], which is a distance-based method that assigns sequences into potential species based on the detection of a barcoding gap (interspecific variation > intraspecific variation). It does not require a priori species designation but does

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require the setting of an intraspecific threshold (set to the range $P_{min}=0.005–P_{max}=0.1$), below which interspecific diversity will not be detected.

The second species delimitation method employed was the multi-rate Poisson tree process—mPPT (multi-rate Poisson tree processes; [45]), which is a “phylogenetics-aware” method that uses differences in mutation rate in a phylogenetic tree to resolve interspecific from intraspecific diversity. Unlike the previous method, it does not rely on a priori distance thresholds but does require a bifurcating non-ultrametric phylogenetic tree for input. A maximum likelihood (ML) tree was generated using RAxML-HPC BlackBox 8.2.10 [58] on the CIPRES Science Gateway [53] using gene partitioning by codon (with the 2nd codon position excluded due to invariability) under the model GTR+$\Gamma$, based on 1000 bootstrap replicates.

**Results**

**Phylogenetics**

A total of 421 sequences (270 newly sequenced from Amazonian rural settlement collections, and 151 downloaded from GenBank databases) from the genus *Anopheles* (in the Arribalzaga series from subgenus *Anopheles*) and genus *Nyssorhynchus* (in the Albimanus and Argyritarsis Series) were included in the analysis (Additional file 1). After alignment these yielded 359 COI haplotypes of 658 base pairs in length. *Chagasia bonneae* (GenBank accession no. KF671010) was also included to serve as an outgroup taxon.

Results of the Bayesian analysis found no evidence for reciprocal monophyly between the subgenera *Anopheles* and *Nyssorhynchus* (Figs. 2a and 3a). The subgenus *Anopheles* is paraphyletic with respect to *Nyssorhynchus*, with two of its clades forming a three-way polytomy with the latter (BPP 85%; Fig. 2a).

**Genus Anopheles (see Fig. 2a)**

*Anopheles peryassui* was recovered as a strongly supported but highly diverse monophyletic clade (99% BPP). It contained a haplotype (Sample id: RO38_2; Accession number: MF381690) that is at least 7.4% different (K2P distance) from its conspecifics. Although *Anopheles intermedius* from the state of São Paulo was included in the analysis, it appeared from the phylogenetic tree (and was confirmed in subsequent species delimitation analysis) that this sequence was the sole representative of *An. intermedius* in this study.

*Anopheles punctimacula* formed two strongly supported monophyletic clades (both with BPP of 100%) that appear to have very different evolutionary histories. The first clade was sister (96% BPP) to a clade containing all but one (*An. peryassui*) of the ingroup species. This first clade is also highly diverse, containing a haplotype (*Anopheles punctimacula* AC22_64) that is at least 5.9% different (K2P distance) from other haplotypes within the clade. The second *An. punctimacula* clade contains haplotypes from *An. punctimacula* sensu stricto/lineage A and *An. punctimacula* lineage B detailed in Loaiza et al. [36]. It is recovered in a disparate part of tree from the first *An. punctimacula* clade.

*Anopheles fluminensis* was recovered as two phylogenetically distant clades (both with BPP of 100%). The first clade includes *An. fluminensis* from the type locality in Rio de Janeiro, and herein considered sensu stricto. It is found on a three-way polytomy with the *Nyssorhynchus* clade and a clade containing *Anopheles costai*, *Anopheles malefactor*, *An. punctimacula* s.s./lineage A, *An. punctimacula* lineage B and the second *An. fluminensis* clade. This second clade was found to be sister (100% BPP) to a highly distinct lineage of *An. malefactor*. There is a difference of 8.0–9.3% (K2P distance) between these two *An. fluminensis* clades. The highly distinct *An. malefactor* lineage is herein denoted *An. near malefactor* and is clearly resolved from *An. malefactor* sensu lato by a difference of 8.4–9.0% (K2P distance).

*Anopheles costai* was resolved (100% BPP) as a highly diverse clade (0.0–7.8% K2P distance). The *An. costai* haplotype relationships within this clade are suggestive of between three and six lineages, but the lineage composition of this clade will be more clearly described in the results from species delimitation below.

**Genus Nyssorhynchus (see Fig. 3a)**

Members of the Oswaldoi Subgroup included in this study are the Oswaldoi Complex, the Konderi Complex and the Nuneztovari Complex. Members of the Oswaldoi Complex in this study are *Nyssorhynchus oswaldoi* sensu stricto, *Ny. oswaldoi* A, *Ny. oswaldoi* B, and *Nyssorhynchus rangeli*. One of these, *Ny. oswaldoi* A (100% BPP), forms a monophyletic clade within the three-way polytomy (97% BPP) with *Nyssorhynchus konderi* and *Ny. near konderi*. *Nyssorhynchus rangeli* (100% BPP) is a monophyletic clade on a five-way polytomy (97% BPP) that also contains a clade (96% BPP) comprised of *Ny. oswaldoi* s.s. (87% BPP) and *Ny. oswaldoi* B (100% BPP), which are reciprocally monophyletic. The two members of the Konderi Complex, *Ny. konderi* (100% BPP) and *Ny. near konderi* (100% BPP), each form monophyletic clades within the three-way polytomy described above. The Nuneztovari Complex (100% BPP) is composed of *Nyssorhynchus dunhami* and several lineages from *Nyssorhynchus goeldii* and *Nyssorhynchus nuneztovari*. Of the five *Ny. goeldii* and *Ny. nuneztovari* clades (Groups I–V) detailed in Scarpassa et al. [40], *Ny. goeldii* Group I (100% BPP), *Ny. nuneztovari* Group II (100% BPP) and Group
IV (100% BPP) are found to be monophyletic and Ny. goeldii Group III are Ny. goeldii Group V are paraphyletic. Nyssorhynchus dunhami is also paraphyletic with respect to other members of the Nuneztovari Complex.

Sequences from six members of the Strodei Subgroup are included in this study. Nyssorhynchus benarrochi is resolved into two quite distinct clades. The first is identified from Acre and Rondônia (100% BPP) and is found as
a sister (97% BPP) to the Ny. nuneztovari complex. The second is a distantly related Ny. benarrochi B clade (100% BPP) found to be sister (87% BPP) to the Nyssorhynchus strodei subgroup (100% BPP). Nyssorhynchus arthuri A (100% BPP) and Ny. arthuri C (98% BPP) formed distinct clades on a three-way polytomy (100% BPP) with the Ny. strodei/Nyssorhynchus albertoi clade (100% BPP), which are unresolved.

The Albitarsis Series is comprised of the Albitarsis Group, made up of the Albitarsis Complex, and the Braziliensis Group, made up of Nyssorhynchus braziliensis. In this study, the Albitarsis Complex is resolved as a monophyletic clade (100% BPP), and all (Ny. albitarsis, 100% BPP; Nyssorhynchus deaneorum, 100% BPP; Nyssorhynchus marajoara, 97% BPP; Nyssorhynchus oryzalimentes, 100% BPP) but one (Ny. albitarsis H) of its multiple species form monophyletic clades. Nyssorhynchus braziliensis (100% BPP) is resolved from all other species in Nyssorhynchus (100% BPP) and shows high levels of haplotype diversity, up to 3.3% (K2P distance).

Nyssorhynchus darlingi (100% BPP) is resolved as a sister species (99% BPP) to the highly diverse Ny. triangulatus clade (100% BPP), with intraspecific diversity of up to 3.8% (K2P distance).

Species identification
Collection and reference specimens are delimited into 30 “species” groups by the mPTP analysis (Additional file 3; Figs. 2b, 3b). The ABGD analysis provided three possible partitions for these specimens before grouping them together as a whole (Fig. 4). The most conservative partitioning produces 36 groups (Additional file 4; Figs. 2c, 3c) and is herein referred to as the “strict” partition. The most fragmented partitioning produces 44 groups (Additional file 5) and is herein referred to as the “relaxed” partition.

Genus Anopheles (see Fig. 2b, c)
Under the mPTP (phylogeny-aware) analysis, 3 groups are delineated for An. costai. The first group, herein denoted An. costai G1, has a large distribution, and is identified in the states of Acre and Amazonas and the Colombian departments of Putumayo (Accession number: JX205128) and Amazonas (Accession number: KF698865). This group has good support in the Bayesian phylogenetic tree (89% BPP). The second delineated group, herein denoted An. costai G2, is only identified from rural settlements in Acre and is well supported in the Bayesian phylogenetic analysis (100% BPP). Both An. costai G1 and An. costai G2 are consistently delineated in the strict and relaxed ABGD species partition. The third group includes An. costai sensu stricto haplotypes from the state of São Paulo (the type locality; Field Specimen ID: SP02_17_3) as well as haplotypes from other locations in Acre, Amazonas, Bahia, Rondônia, and the Colombian departments of Meta (Accession numbers: HM022403 and HM022404) and Putumayo (Accession number: JX205127). This group is further partitioned into four groups in both the strict and relaxed ABGD (K2P distance-based) species partitions. These groups are herein denoted An. costai sensu stricto (identified from Bahia and São Paulo), An. costai G3 (identified from Acre, Amazonas and Rondônia), An. costai G4 (identified from Acre, Amazonas and Colombia) and An. costai G5 (identified from Acre only).

Anopheles fluminensis is consistently partitioned into two groups across all species delimitation analyses, and these groups represent the two polyphyletic clades identified in the Bayesian phylogenetic analysis. The first group includes specimens from five rural settlements in Acre in addition to specimens from Rio de Janeiro (the type locality—Reserva Biológica do Tinguá; Accession number: MF381677), São Paulo and Ecuador, but denoted An. nr fluminensis in Linton et al. [59]. This group is tentatively denoted An. fluminensis sensu stricto, although there is considerable intra-group diversity that ranges from 0.6 to 3.3% (K2P distances), with the São Paulo and Rio de Janeiro haplotypes (0.6% different) having distances of between 2.0 and 3.3% with the other members of the group. The second group is comprised of specimens collected from eight rural settlements in Acre, and is herein denoted An. fluminensis G1. The two groups differ by between 7.7 and 9.3% (K2P distances). Anopheles
malefactor is also partitioned into two groups across all species delimitation analyses and is again consistent with the polyphyly observed to the phylogenetic analysis. The first group is *An. malefactor*, of González et al. [41] and Gómez et al. [31], whereas the second is herein denoted *An. near malefactor*, and they differ by between 8.4 and 9.0% (K2P distance).

Both mPTP and ABGD species delimitation approaches found three groups in *An. punctimacula*. Two of these groups are found in the highly diverse *An. punctimacula* clade and herein are denoted *An. punctimacula G1* from Acre and *An. punctimacula G2* from Acre and the Colombian department of Putumayo (Accession number: JX205122). The third group is made up of the *An. punctimacula ss, An. punctimacula* lineage A and *An. punctimacula* lineage B (from [36]).

The mPTP species delimitation approach identified two groups in *An. peryassui*. The first group, herein denoted *An. peryassui G1*, is represented by a single haplotype from Rondônia (Sample id: RO38_2; Accession number: MF381690), and identified as highly divergent (7.4% K2P distance) in the phylogenetic analysis. The second group is made up of haplotypes from Acre, Para, the Colombian departments of Amazonas (Accession numbers: KF698875–KF698877; [31]) and Meta (HM022405), and the state of Amapá, with this latter haplotype delineated

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### Delimitation of Collection Specimens

| mPTP       | ABGD - strict                      | ABGD - relaxec                      |
|------------|------------------------------------|-------------------------------------|
| An. punctimacula G1 | An. punctimacula G1                | An. punctimacula G1                |
| An. punctimacula G2 | An. punctimacula G2                | An. punctimacula G2                |
| An. malefactor | An. malefactor | An. malefactor |
| An. near malefactor | An. near malefactor | An. near malefactor |
| An. simoni G1 | An. simoni G1 | An. simoni G1 |
| An. simoni G2 | An. simoni G2 | An. simoni G2 |
| An. simoni G1, G2 & G3 | An. simoni G1, G2 & G3 | An. simoni G1, G2 & G3 |
| Ny. brasilensis | Ny. brasilensis | Ny. brasilensis |
| Ny. triangulata | Ny. triangulata | Ny. triangulata |
| Ny. darlingi | Ny. darlingi | Ny. darlingi |
| Ny. abithia H, Ny. eisenkon & Ny. marmorata | Ny. abithia H, Ny. eisenkon & Ny. marmorata | Ny. abithia H, Ny. eisenkon & Ny. marmorata |
| Ny. ringeli | Ny. ringeli | Ny. ringeli |
| Ny. covoeldi ss | Ny. covoeldi ss | Ny. covoeldi ss |
| Ny. covoeldi A | Ny. covoeldi A | Ny. covoeldi A |
| Ny. near konderi | Ny. near konderi | Ny. near konderi |
| Ny. konderi | Ny. konderi | Ny. konderi |
| Ny. covoeldi, Ny. guttii & Ny. nuciflavus | Ny. covoeldi, Ny. guttii & Ny. nuciflavus | Ny. covoeldi, Ny. guttii & Ny. nuciflavus |
| Ny. hentzii G1 & G2 | Ny. hentzii G1 & G2 | Ny. hentzii G1 & G2 |
| Ny. hentzii B | Ny. hentzii B | Ny. hentzii B |
| Ny. sabeth C | Ny. sabeth C | Ny. sabeth C |

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**Fig. 5** Summary of all species delimitation analysis of collection specimens from Acre, Amazonas and Rondônia
as a third group in both the strict and relaxed partitions of the ABGD species delimitation. The *An. intermedius* singleton from São Paulo was separately delimited across all species delimitation analyses.

**Genus Nyssorhynchus** (see Fig. 3b, c)

Within the Strodei Subgroup, *Ny. albortoi* and *Ny. strodei* are not delineated in either mPTP or ABGD analyses. *Ny. arthuri A* and *Ny. arthuri C* are delineated across all analyses but only *Ny. arthuri C* was collected in this study at a rural settlement in Rondônia. Two groups are identified from the Benarrochi Complex. The first is *Ny. benarrochi B*, which is delineated in the mPTP analysis and from the ABGD strict partition (but splits into 4 groups in ABGD relaxed partition). *Nyssorhynchus benarrochi B* was collected from five rural settlement in Acre. The second group is delimited in the mPTP analysis, but splits into two further groups, herein denoted *Ny. benarrochi G1* and *Ny. benarrochi G2*, in both the strict and relaxed partition from the ABGD analyses. *Nyssorhynchus benarrochi G1* was collected from a rural settlement in Acre, while *Ny. benarrochi G2* was collected from three rural settlements in Rondônia.

The sole representative from the Argyritarsis Series included in this study is *Ny. darlingi*. This species is clearly delineated across all analyses, and identified from six rural settlements in Acre and Amazonas. With respect to the Albitarsis Series, *Ny. braziliensis* is delimited in mPTP analysis and in the ABGD strict partition. However, it decomposes into five separate groups (of which 3 were singletons) in the ABGD relaxed partition. Within the Albitarsis Complex, only *Ny. albitarsis* and *Ny. oryzalimnetes* are delineated in analyses (ABGD strict and relaxed partitions). *Nyssorhynchus albitarsis*, *Ny. deaneorum* and *Ny. marajoara* are consistently grouped together in all species delimitation analyses.

The Oswaldoi Subgroup contains the Konderi, Oswaldoi and Nuneztovari Complexes, all of which are represented in this study. Species from this subgroup that can be clearly delimited across all species delimitation analyses are *Ny. rangeli* (rural settlements in Acre and Rondônia), *Ny. oswaldoi A* (rural settlements in Acre, Amazonas, Rondônia), *Ny. konderi* and *Ny. near konderi*.

In the case of *Ny. konderi*, the “An. konderi of Amapá” (GenBank accession: JF437967 and JF437968) in Ruíz-Lopez et al. [60], the “An. konderi of Sallum” (GenBank accession: KF809126 and KF809128) in Motoki et al. [60] belong to this lineage. This lineage was collected from rural settlements in Acre and Amazonas. *Nyssorhynchus oswaldoi* s.s. and *Ny. oswaldoi B* are delimited in the mPTP analysis and the ABGD relaxed partition, but not in the ABGD strict partition. This species delimitation (and the phylogenetic analysis) suggests the “An. konderi of Paraná and Rondônia” (GenBank accession: JF437969–JF437974) in Motoki et al. [60] belongs to *Ny. oswaldoi* s.s. (GenBank accession: KF809126 and KF809128). *Nyssorhynchus oswaldoi* s.s. was collected from a single rural settlement in Amazonas, whereas *Ny. oswaldoi B* was absent from all rural settlements in our study. None of the members of the *Ny. nuneztovari* complex are resolved in any of the species delimitation analyses. Finally, *Ny. triannulatus* is delineated as a single group across all analyses, and identified from a single rural settlement in Acre.

**Summary of species delimitation for collection specimens**

The most conservative species delimitation approach used in this study (mPTP) delimits at least 23 putative species from these collection sites (Fig. 5), and 30 from the study as a whole i.e. including reference sequences. The strict clustering-based (ABGD) partition delimits 27 putative species (Fig. 5) and 36 from the study as a whole, while the relaxed clustering-based (ABGD) partition delimits 33 putative species (Fig. 5) and 44 from the study as a whole.

**Discussion**

The pattern of deforestation and anthropogenic changes in natural environments, especially in the Amazon tropical rainforest, and the insidious poverty of settlers living in precarious conditions, have been associated with increases in the incidence of malaria [62]. Compounding this problem in areas of the Amazon River basin with active malaria is the primary vector *Ny. darlingi*, which is a generalist and opportunistic species that can blood-feed both indoors and outdoors complicating the dynamics of transmission [63]. There are also other important vectors involved in malaria transmission in the region, depending on the local ecological conditions. Considering the urgent need for studies focusing on practical malaria that includes accurate identification of malaria vectors, this study demonstrates the capability of the barcode region of the *COI* mitochondrial gene and commonly used species delineation approaches to delimit a diverse range of *Anopheles* species found in rural settlements affected by frontier malaria in the Amazon River basin. After assigning collection specimens to 19 morphospecies, conservative estimates delimit between 23 and 27 potential species, up to 13 of which appear to be new species.
Several species are well resolved in our phylogenetic tree and consistently delineated as coherent groups across analyses. These include Ny. darlingi, Ny. rangeli and An. intermedius. Ny. darlingi is the primary vector of malaria in the Amazon River basin [16]. It is a highly anthropophilic species and can vary from endophagy to exophagy, and endophily to exophily [64]. As such, vector control can be extremely challenging and the presence of this species at six rural settlements in Acre and Amazonas is significant for malaria transmission and vector control. In addition, Ny. rangeli has been found to be important in local malaria transmission in Colombia [65] and its larval habitat is associated with human-modified environments [66]. Its presence in rural settlements in Acre and Rondônia may indicate a potential role in frontier malaria at these localities. Anopheles intermedius sequence data (a singleton from São Paulo state) was included in the analysis as a reference as it is a potential malaria vector in the region, with natural infection detected in Amapa [67], Pará [68], and French Guiana [69]. However, this species was not detected at any of the study’s collection sites.

Although Ny. braziliensis was recovered as a monophyletic species and singularly grouped in both species delineation approaches, the species appeared to be highly diverse and partitioned into multiple groups in the relaxed ABGD partition. Insufficient sampling meant that we could not attribute any significance to this structure. The local abundance of this species in the Amazon River basin varies considerably [37, 67, 70] but it appears to be common in human-modified environments [71]. The species is highly anthropophilic [37] and it has been found naturally infected with P. malariae [70], P. vivax [25, 72] and P. falciparum [25, 73]. Its presence in rural settlements may therefore have implications for frontier malaria transmission.

To date, there is little genetic data available for An. periyassui, whose range covers most of northern South America. There is some evidence to suggest that the species may play a role in malaria transmission, with natural infection by P. falciparum and P. vivax reported in the state of Amazonas [74]. The current study included specimen reference data from Colombia in Gómez et al. [31], the only other source of An. periyassui genetic data available in GenBank. Results of the analyses demonstrated that An. periyassui appears to exist as a species complex with at least two (and possibly three) lineages identified, with the first found in Rondônia and the second found in Acre, Amapa, Colombia and Para (with the Amapa haplotype possibly splitting into a third lineage). However, the systematics of this complex remains poorly understood, because of the lack of material from the type locality in Rio Bonito in the state of Minas Gerais.

The genetic diversity of An. fluminensis has also been poorly explored. This lack of interest may be due to the understanding that An. fluminensis is not an important malaria vector [75], although specimens identified as An. near fluminensis have been incriminated as a malaria vector in the department of Junin, Peru [76] and more recently the species was found naturally infected by Plasmodium malariae in the Atlantic Forest of São Paulo, Brazil [77]. A study of mosquito diversity in the Ecuadorian Amazon found some genetic support for An. near fluminensis, demonstrating a 2.75–3.77% difference between sequences from the province of Orellana in Ecuador and São Paulo state, Brazil. However, despite the identification of similar levels of diversity between the two regions in this study, this is considered to be intraspecific, according to the approaches used, and grouped with the topotype specimen (Accession number: MF381677) from Rio de Janeiro. However, An. fluminensis was split into two sympatric lineages (with Ecuadorian specimens and sensu stricto specimens grouped together), which, given their distinct evolutionary histories, may be of varying epidemiological importance. Clearly, An. fluminensis has, to date, been understudied and overlooked but, given the species diversity that has been identified in this study from specimens morphologically identified as An. fluminensis and the limited sampling effort (only collected from Acre), a more thorough evaluation of lineage diversity and relationships across its considerable South American range is to be encouraged.

Like An. periyassui and An. fluminensis, there are scant genetic data available for An. costai. The species appears to be found mainly in forest habitat [78] and is not believed to be an important malaria vector. However, An. costai has been frequently misidentified as An. mediopunctatus [79], which has been found naturally infected with Plasmodium vivax in the state of Amazonas [74] and it may yet be recognized as epidemiologically important. This study reveals that An. costai exists as a diverse species complex (between 3 and 6 lineages), and several of its lineages have geographical distributions that span the Amazon River basin and beyond. More extensive sampling will be required to determine whether An. costai s.s. lineage identified in the clustering-based delimitation occurs in the Amazon River basin, while the ecologies of each species complex member should be explored to determine their potential importance in rural settlements affected by frontier malaria.

Anopheles punctimacula ranges from Mexico to Argentina and the Caribbean and has previously been implicated as a malaria vector in Panama [80] and Colombia [81], although this occurred prior to An. malefactor and An. calderoni emerging from this taxon [82, 83]. Anopheles punctimacula has previously been reported to exist
as a species complex, with at least two lineages from Panama supported across multiple genes [36]. The two lineages identified in this study appear to be distinctly related to those identified in Loaiza et al. [36], which included the sensu stricto form collected near the type locality in Colón, Panama (and consistently grouped with the Lineage A and B form denoted in that study). One of these lineages exists over quite a considerable range (~1000 km; Acre, Brazil—Putumayo, Colombia) and further studies of the phylogeny and ecology of these taxa necessitate employing a range of molecular and morphological markers and more comprehensive sampling in order to more clearly define evolutionary relationships, geographical boundaries, ecological niches and, ultimately, their role in malaria transmission in the region. Anopheles malefactor, previously elevated from synonymy with An. punctimacula [82], collected from a rural settlement in Acre, was clearly distinct from the An. malefactor from the Panama (type locality) and Colombia, and appears to be a new species and sister to a newly delineated An. flavimensis G1 lineage.

The close relationships observed among the Ny. konderi and Ny. oswaldoi complexes indicate that these should be considered a unique species complex named Oswaldoi–Konderi [34]. There are now considered to be five species in this Oswaldoi–Konderi Complex, which include Ny. oswaldoi s.s., Ny. oswaldoi A, Ny. oswaldoi B, Ny. konderi and Nyssorhynchus nr. konderi [34, 61]. Although Ny. konderi and Ny. oswaldoi are generally not considered vectors of human malaria, Quiñones et al. [65] found Ny. oswaldoi B (although denoted Ny. oswaldoi in their study) infected with P. vivax in the state of Putumayo, Colombia. The complex may therefore be of epidemiological importance. Although all five species could be effectively delineated in the current study, only four were identified from the collection sites in Acre, Amazonas and Rondônia. Ny. oswaldoi s.s. was identified from a site south of Amazonas and (with the inclusion of other data from GenBank) it can also be found Espírito Santo state (type locality; Accession number: JF923721), Paraná, Rondônia and São Paulo. This range covers both the Paraná and Amazon river basins, which are connected by the Parapetí River alluvial fan and may allow fluvial species relatively uninhibited expansion between basins [84]. Whereas Ny. oswaldoi A appears to be found throughout much of the Amazon River basin (Amazonas, Acre, Pará, Rondônia), Ny. oswaldoi B was not identified at the collection sites, although it has previously been identified from a site in the northern reaches of the Brazilian Amazon (Santana, Amapá; [34]). A Ny. oswaldoi specimen (Field Specimen ID: SP22_9) from Paráquera-Mirim, Paráquera-Açu, São Paulo was included in this study to aid in delineating geographic distribution. However, this specimen was consistently resolved from Ny. oswaldoi s.s. and is likely a new species within the Oswaldoi–Konderi Complex. The Ny. konderi specimens collected in this study were confined to sites in the state of Acre but this species has also been collected from sites in Amapá (previously denoted An. konderi Amapa [60]). Saraíva et al. [34] also identified this species from Amazonas, Pará and Rondônia. With respect to Ny. nr. konderi, previous studies have found this species in Colombia, Ecuador, Peru [61] and the Brazilian states of Amazonas and Rondônia [34]. In this study, this species was collected from Amazonas but also from a large number of rural settlements in Acre.

Recent studies have confirmed Ny. benarrochi as a species complex, with the identification of Ny. benarrochi B in Colombia [85] and Peru [86]. Although Ny. benarrochi is predominantly zoophilic [87] and therefore not believed to be an important malaria vector, highly anthropophilic behaviour does occur in some areas and it may play a role in malaria transmission in the Peruvian Amazon [85, 88]. This study confirms the existence of Ny. benarrochi B, using data from Conn et al. [86], in rural settlements in the state of Acre. It also identifies at least one other Ny. benarrochi lineage from Acre and Rondônia (and perhaps representing two). Unfortunately, this lineage cannot be confirmed as Ny. benarrochi s.s. as material from the type locality in the municipality of La Ceiba, Trujillo State, Venezuela is unavailable. It is, therefore, essential that collections be obtained from this type locality in order to undertake a meaningful systematic review of the Ny. benarrochi complex and determine whether species diversity within this complex is associated with variation in anthropophilia and vector competence.

Nyssorhynchus triannulatus is found throughout Central and South America, and has been incriminated in the transmission of human malaria in the states of Amapá [67], Amazonas [74] and Pará [68], Brazil. Recent studies have shown that Ny. triannulatus forms a species complex with two other recently identified species; Ny. halophylus [89] and Ny. triannulatus C [90]. Nyssorhynchus triannulatus sensu lato has been found to form paraphyletic (at a combined IT2S, white and COI gene tree; [91]) and monophyletic sister [using RAPD markers; 92] relationships with an Ny. halophylus and Ny. triannulatus C clade. Within Ny. triannulatus s.l. two further clades have been recovered with the COI gene, one from Central America and northern Colombia and the other primarily from the Amazon river basin [91], and several additional biologically meaningful subclades from within the Amazonian clade were also proposed, based on multiple gene (COI, white and ITS2) and haplotype network analysis. The haplotypes included in this study come from the Amazonian states of Acre and Amapá, as well as
from the states of Minas Gerais to the east, Espírito Santo on the Atlantic coast, São Paulo in the south east of Brazil, and neighbouring Colombia. Despite a considerable geographic distribution and habitat range [91, 92], results from the current analysis show that, although the *Ny. triannulatus* s.l. clade is extremely diverse, relationships among its COI haplotypes cannot be meaningfully partitioned into separate species i.e., all branch structure and pairwise distances within *Ny. triannulatus* are considered intraspecific. It appears that relationships may be better explained by complex population histories [92], where elevated levels of genetic diversity have been maintained perhaps by historical fragmentation, secondary contact and gene-flow followed by more recent divergence due to geographical isolation. Further phylogenetic and species delimitation analyses at multiple loci may reveal more compelling support for species designation within *Ny. triannulatus*, as phylogenetic analysis of COI frequently fails to resolve well supported species of Anophelinae mosquitoes, e.g., *Ny. strodei* and *Ny. albitors*. Conflict among gene trees due to incomplete lineage sorting and horizontal gene transfer (gene-flow) is pervasive in very closely related species, making molecular identification of such species difficult. High density codominant markers such as single nucleotide polymorphisms (SNPs) offer some opportunities to overcome such phylogenetic challenges [93–96] and an exploration of the hierarchy of structure within *Ny. triannulatus* (and *Ny. braziliensis*) may be better achieved using such markers in combination with population genetic approaches [97].

Several of the morphospecies collected in this study are highly diverse complexes, the diversity of which is supported by a range of ecological, morphological and genetic data. However, the use of species delimitation approaches with the COI gene failed to detect several of these species boundaries. Currently, 10 species have been identified within the *Ny. albitorsis* complex, five of which (*Ny. albitorsis, Ny. albitorsis H, An. deaneorum, Ny. marajoara* and *Ny. oryzalimnetes*) were collected in this study. A range of morphological and molecular data has been employed to resolve and support species in this complex [98–100]. Nevertheless, the close relationships observed among *Ny. albitorsis H, Ny. deaneorum* and *Ny. marajoara* at the COI gene in the present study were consistent with patterns of intraspecific variation when employing cluster- and tree-based species delimitation. It appears that species delimitation of this complex at the COI gene will fail to detect important species boundaries supported by other data and further exploration and discovery of species diversity in the Albitorsis Complex will require a multilocus approach, possibly using loci such as ITS2 and the *white* gene, which have been used to resolve many of the species within the complex [101, 102].

Similar issues were encountered when dealing with the Nuneztorvari Complex and Strodei Subgroup. The former is comprised of *Ny. dunnhami, Ny. goeldii, Ny. nuneztorvari* [103], supported by a range of morphological and molecular data [104–106]. Its geographical distribution runs from the Isthmus of Panama to northern South America and *Ny. nuneztorvari* sensu lato is considered one of the most important malaria vectors in the region [26]. A recent study by Scarpasa et al. [40] found *Ny. nuneztorvari* s.l. haplotypes were variously identified as *Ny. dunnhami, Ny. goeldii, Ny. nuneztorvari* and an additional unknown clade. They found that population genetic and phylogenetic analysis were in some agreement, although lineage delineation and relationships among COI trees that differed in sampling effort were to some degree incongruent. Although the current study combined the sequences from Scarpasa et al. [40] with those from specimens collected from the Brazilian Amazonian states of Acre, Amapá, Amazonas, Pará, Rondônia, it failed to delineate any species within the *Nyssorhynchus nuneztorvari* complex. The Strodei Subgroup is made up of *Ny. albertoi, Ny. strodei* [107], *Ny. striatius* [108], *Ny. rondoni* and finally *Ny. arthuri*, which is comprised of four species A - D [35]. Given the systematic complexity of this group and the recent emergence of many of its species, the importance of these species in malaria transmission is unknown. However, *Ny. strodei* has previously been found naturally infected with *P. vivax* in Ariquemes, Rondônia [73], although this case may refer to *Ny. arthuri C* [35]. *Nyssorhynchus albertoi* and *Ny. strodei*, despite having considerable support as separate species within the Strodei Subgroup [107], are known to be unresolved with the COI gene [35] and, unsurprisingly, species delimitation in the current study using this same gene failed to identify the respective species. The other members of the subgroup included in the study were resolved and clearly delineated. As in Bourke et al. [35], *Ny. arthuri A* was not detected from the Amazon region and *Ny. arthuri C* has so far only been identified from the state of Rondônia. Further analysis of this species range and habitat is required before its epidemiological importance in rural settlements can be determined.

The work presented here demonstrates a clear advantage to employing species delimitation approaches when the objective is to explore species diversity and discover new species in Anophelinae where cryptic species boundaries are common. Due to the relative speed at which COI can be sequenced and analysed, the study demonstrates the power of single-locus species delimitation approaches to establish a baseline of species diversity in Anophelinae in remote and unexplored regions of the Amazon River basin. However, it must be clearly stated that single-locus data alone should only be used to
provide a preliminary description of species boundaries (possible gene tree-species tree discordance) and the new species delimited in this study remain putative. Rather, it allows for the establishment of viable species hypotheses, to be tested against a range of independent data sources that may be, for example, morphological, molecular and/or ecological in nature. In particular, there have been few or no classical studies of Anophelinae taxonomy conducted in the Amazon River basin in recent years, and further studies of species diversity in Anophelinae are, therefore, encouraged to better explore morphological variation among these species. Future work on species exploration and discovery in the Amazon River basin should also employ a multilocus delimitation approach [43, 109] to better enable the resolution of all recognized species within important complexes, such as Albitarsis and Nuneztovari. In addition, the study demonstrates that exceptional diversity detected in morphospecies can be consistent with a model of intraspecific diversity and is suggestive of a complex evolutionary history that may be better explored using high-density markers and population genetic approaches.

This study has been successful in revealing a large number of unknown Anophelinae species that are likely to be new to science and occur in areas with endemic malaria transmission. One of the great challenges for malaria control in the Amazon River basin is the transmission that occurs outside of rural dwellings and the detection of new species that belong to groups containing important vectors will, therefore, have an important impact on the development of effective vector management and control strategies in the region. The World Health Organization recommends that vector management and control interventions should take account of potential impacts on the environment and biodiversity and should be focused on avoiding unintended impacts on non-vector species. The findings from this study will assist in refining such strategies, help build capacity in public health entomology and provide an important contribution to effective malaria control in the region.

Conclusion
Anophelinae mosquito diversity continues to be underestimated in poorly sampled tropical rain forest areas where frontier malaria is a major public health concern. Delimitation approaches used to explore Anophelinae mosquito diversity in these areas reveal a large number of unknown species that are likely to be new to science. These findings will help shape future studies of vector incrimination and transmission dynamics and support efforts to develop more effective vector control and transmission reduction strategies in settled communities in the Amazon River basin.

Additional files

**Additional file 1.** Sample information of collection specimens and reference sequences.

**Additional file 2.** COI sequence data for all collection specimens and references used in this study.

**Additional file 3.** Output from mPTP analysis of collection specimens and reference sequences.

**Additional file 4.** “Strict” partition from ABGD analysis of collection specimens and reference sequences.

**Additional file 5.** “Relaxed” partition from ABGD analysis of collection specimens and reference sequences.

Authors’ contributions
Study design: MAMS, BPB. Data analysis and illustrations: BPB. Interpretation of results: BPB, MAMS. Scientific writing: BPB, MAMS, JEC. Field collections: MAMS, ESB, LS, MC, GZL. Sequence data: TMPO, MAMS. All authors read and approved the final manuscript.

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Competing interests
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