Mitochondrial genome sequence diversity of Indian Plasmodium falciparum isolates

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We have analysed the whole mitochondrial (mt) genome sequences (each ~6 kilo nucleotide base pairs in length) of four field isolates of the malaria parasite Plasmodium falciparum collected from different locations in India. Comparative genomic analyses of mt genome sequences revealed three novel India-specific single nucleotide polymorphisms. In general, high mt genome diversity was found in Indian P. falciparum, at a level comparable to African isolates. A population phylogenetic tree placed the presently sequenced Indian P. falciparum with the global isolates, while a previously sequenced Indian isolate was an outlier. Although this preliminary study is limited to a few numbers of isolates, the data have provided fundamental evidence of the mt genome diversity and evolutionary relationships of Indian P. falciparum with that of global isolates.

Key words: malaria - Plasmodium falciparum - mitochondrial genome - evolution - India

Analyses of whole mitochondrial (mt) genome sequences have proven to be highly relevant in inferring both inter and intra-specific evolutionary histories of many model and non-model organisms (Hedges 2000, Hurst & Jiggins 2005, Haag-Liautard et al. 2008) including the malaria parasites which have shown promising evolutionary outcomes (Joy et al. 2003, 2006, Mu et al. 2005). For example, analyses of whole mt genomes in the two most common human malaria parasites, Plasmodium falciparum and Plasmodium vivax, have provided evidence of the origin and evolutionary histories of these species (Joy et al. 2003, 2006, Mu et al. 2005). In addition, useful inferences on the probable host-switching mechanism of P. falciparum, although much debated (Prugnolle et al. 2011), have been revealed to a certain extent.

Malaria is highly endemic in India, with P. falciparum causing malaria havoc (Singh et al. 2009). However, the mt genome sequence of Indian P. falciparum has not yet been analysed and compared to global mt genome sequences of this species. Therefore, the evolutionary history of global P. falciparum isolates remains incomplete without Indian mt genome sequence data, although two mt genome sequences of Indian origin are available in the public domain (Sharma et al. 2001, Tyagi et al. 2014). In order to fill this gap, we have utilised whole mt genomes of four P. falciparum field isolates [3 new to this study and 1 previously reported (Tyagi et al. 2014)] from low, mild and high endemic localities of India and compared them with the global data. We have also included the already published mt genome sequence data of a single P. falciparum isolate of unknown Indian origin (Sharma et al. 2001) in this study to obtain first-hand information on the extant diversity of Indian P. falciparum in comparison to global isolates.

We collected finger-prick blood samples from P. falciparum-infected malaria patients attending malaria clinics in three geographically distant localities of India with variable P. falciparum malaria epidemiology: Betul (state of Madhya Pradesh, high endemic), Goa (state of Goa, low endemic) and Mangalore (state of Karnataka, mild endemic). The samples were spotted (2, 3 spots) on Whatman filter paper, dried and transported to the laboratory in New Delhi. While the sample from Betul was collected by us, the samples from Goa and Mangalore were obtained from the parasite bank of the National Institute of Malaria Research (NIMR). The fourth sample came from Bilaspur [state of Chhattisgarh, high endemic (Tyagi et al. 2014)] (Fig. 1). This study was approved by NIMR, New Delhi, India, and written informed consents were obtained from all patients who participated in the study before the samples were collected. The locations of the sample collection sites in India are shown in Fig. 1.

For each collected sample, genomic DNA was isolated using the QIAamp mini DNA isolation kit (Qiagen, Germany) according to the manufacturer’s instructions. Because both P. falciparum and P. vivax occur in India in almost equal proportion (Singh et al. 2009) and the mt genome is conserved among different species of human malaria parasites (Hikosaka et al. 2011), we first tested each sample for incidences of mixed malaria parasite infections using nested polymerase chain reaction (PCR) detection assays with genus and species-specific primers based on the 18S rRNA gene (Gupta et al. 2010). In order to PCR amplify and sequence the whole mt genome of 5,967 nucleotide base pairs, we used 19 recently designed primer pairs [for details of the primer sequences and PCR conditions, see Tyagi et al. (2014)]. For each individual DNA fragment, the PCR protocols suggested by Tyagi et al. (2014) were followed. To avoid
infections with multiple clones of *P. falciparum* in a single patient, we sequenced all 19 DNA fragments of the *mt* genome from both directions (2X coverage, see below) and isolated clones showing double peaks in any single nucleotide position (Gupta et al. 2012). For each single-clone *P. falciparum* isolate, the 19 PCR-amplified products were separately purified with exonuclease and shrimp alkaline-phosphatase following standard protocols (Tyagi et al. 2014) and DNA sequencing was performed at an NIMR in-house facility following Sanger sequencing technology using an ABI 3730XL DNA Analyser (Applied Biosystems). For each *P. falciparum* isolate, all 19 DNA fragments downstream of the sequence were individually edited using the SeqMan and EditSeq modules of the Lasergene v.7 computer program (DNASTAR, Madison, USA), respectively and the sequences were manually assembled to form one complete *mt* genome. The newly generated whole *mt* genome sequences of the three isolates were submitted to GenBank with accession numbers KJ418723, KJ418724 and KJ418725.

To estimate *mt* genome diversity in Indian *P. falciparum* and infer evolutionary patterns by comparing to global isolates, we have estimated several population genetic parameters. Specifically, the number of segregating sites, number of haplotypes, haplotype diversity (Hd) and a measure of nucleotide diversity (π) (Nei 1987), were calculated and compared with data from other parts of the globe using the DnaSP v.5 computer program (Librado & Rozas 2009). Furthermore, to perform comparative and evolutionary studies of global *P. falciparum* isolates using whole *mt* genome sequences, five Indian whole *mt* genome sequences were analysed (of which 3 were newly generated in the present study and 2, Blsp1 (Tyagi et al. 2014) (GenBank accession KJ144901) and PfPH10 (Sharma et al. 2001) (GenBank accession AJ298788) were previously reported). In these analyses, one consensus sequence from each continent [Africa (n = 29), South America (n = 28) and Asia (n = 29)] and country [Papua New Guinea (PNG) (n = 10)] (Joy et al. 2003) were generated by alignment of the whole *mt* genome sequences from the respective places using the MegAlign module of Lasergene v.7 computer program (DNASTAR, Madison, USA). The 100 *mt* genome sequences of *P. falciparum* utilised here for comparative and evolutionary studies bear the GenBank accessions AY282924-AY283019 and AJ276844-AJ276847. The whole *mt* genome sequence of the reference 3D7 isolate (GenBank accession AY282930) was also included in this study. Therefore, 10 sequences were utilised in this study [5 Indian, 1 African (consensus), 1 South American (consensus), 1 PNG (consensus), 1 Asian (consensus) and the single 3D7 strain]. All 10 sequences were aligned following the CLUSTALW algorithm and a phylogenetic tree was constructed using the neighbour-joining (NJ) method implemented in MEGA v.5 computer program (Tamura et al. 2011). In order to estimate the strength of each internal node of the NJ phylogenetic tree, the tree topologies were simulated 1,000 times.

Following the PCR diagnostic approach to detect malaria parasites and based on the peaks of the DNA sequence chromatogram, we were able to isolate three pure single-clone *P. falciparum* isolates (Bet12, Goa2 and Mang2) and successfully sequence their complete *mt* genomes. Because complete *mt* genome sequences from two other Indian isolates (Blsp1 and PfPH10) were available in the public domain (National Center for Biotechnology Information, available from: ncbi.nlm.nih.gov/), we also utilised the data from these two isolates (altogether 5 Indian *mt* genome sequences) to estimate *mt* genome sequence diversity and to infer first-hand evolutionary relationships of Indian *P. falciparum* with that of global isolates. The whole *mt* genome sequence alignment of five Indian isolates (Bet12, Goa2, Mang2, Blsp1 and PfPH10) with the reference sequence from the 3D7 strain revealed 26 variable nucleotide sites in Indian *P. falciparum* (Table I). Considering that the *mt* genome is fairly conserved across populations and species in *Plasmodium* (Hikosaka et al. 2011), the observed high incidences of nucleotide substitutions in Indian isolates was quite surprising. However, a closer look at the alignment revealed that as many as 21 unique nucleotide substitutions were present in the PfPH10 isolate (Table I). A very similar pattern was also observed when the PfPH10 isolate was compared with the Blsp1 and 3D7 isolates (Tyagi et al. 2014). Because there were such unusual patterns of single nucleotide mutations in the PfPH10 isolate, which might bias the overall outcome of the *mt* genome diversity, we did not consider the data of PfPH10 in further population genetic analyses. Therefore, we restricted the dataset to four *mt* genome sequences of Indian *P. falciparum*. Multiple alignments of *mt* genome sequences of four Indian *P. falciparum* isolates (Bet12, Goa2, Mang2 and Blsp1) with the reference 3D7 isolate revealed five nucleotide substitutions (Table II). Out of these five single nucleotide polymorphisms (SNPs), four were found in the four Indian isolates and one in the 3D7 isolate. Out of the four SNPs found in Indian isolates, two were present in the intergenic regions and two were in the *cox I* gene (Table II). The two SNPs present in
the *cox I* gene, however, were found to be synonymous. Further multiple sequence alignment of the four Indian *mt* genome sequences with the 100 published sequences (Conway et al. 2000, Joy et al. 2003) indicated that three out of the four SNPs found in Indian *P. falciparum* (at positions 276, 725 and 2763) were novel and India-specific (data not shown). The fourth SNP (at the 2175th position) seems to be Asian-specific in nature, as this SNP was never detected in any global samples and was only detected in some Asian isolates (Joy et al. 2003). In comparison to the reference 3D7, one nucleotide substitution was detected in the Blsp1 isolate and two were found in each of the Goa2 and Mang2 isolates (Table II). The Bet12 isolate, however, contained four nucleotide substitutions, indicating that this isolate is highly diverged from the rest of the three Indian isolates (Table II). The four SNPs segregating in the four Indian isolates produced three haplotypes: GCTC (Blsp1), ATTT (Bet12) and GCCC (Goa2 and Mang2).

| Nucleotide positions | 276 | 725 | 2175 | 2763 | 4952 |
|----------------------|-----|-----|------|------|------|
| 3D7                 | G   | C   | T    | C    | T    |
| Blsp1                | .   | .   | .    | .    | .    |
| Bet12                | A   | T   | .    | T    | C    |
| Goa2                 | .   | .   | C    | .    | C    |
| Mang2                | .   | .   | C    | .    | C    |

*a*: Joy et al. (2003); *b*: Tyagi et al. (2014); *c*: present paper; Only sites showing nucleotide variations in the alignment were shown. To be noted that due to occurrence of four single nucleotide polymorphisms in the four Indian isolates, three haplotypes are formed: GCTC (Blsp1), ATTT (Bet12) and GCCC (Goa2 and Mang2).
limited to only four isolates, the high mt genome diversity observed was comparable to that of Africa. These results corroborate earlier observations on high genetic diversity in different genes in Indian *P. falciparum* (Singh et al. 2009) and therefore justify further in-depth analyses with a larger number of isolates from different eco-climatic and malaria endemic localities in India.

With the whole mt genome sequence information of Indian *P. falciparum*, we were interested to know the evolutionary relationships of global isolates using phylogenetic approaches. To do this, we used single consensus sequences of whole mt genomes from Africa, Asia, South America, PNG, the reference 3D7 and five mt genomes of Indian *P. falciparum* (Bet12, Goa2, Mang2, Blsp1 and PfPH10). The NJ phylogenetic tree (Fig. 2) showed congregation of global *P. falciparum* isolates (Africa, Asia, South America, PNG, 3D7) with two Indian sequences (Bet12 and Blsp1) to form a large single clade with high bootstrapped values of the internal node (Fig. 2). However, two other Indian isolates (Goa2 and Mang2) were found in a small separate clade with very weak bootstrapped values (Fig. 2). This might be because both Goa2 and Mang2 isolates bear a similar haplotype (GCC) (Table II) that is different from the other two isolates (Bet12 and Blsp1). Moreover, Goa and Mangalore are from the south-western part of India and are in close geographic proximity to each other (Fig. 1). The observed genetic similarities between the two geographically close *P. falciparum* populations can be explained by the fact that *P. falciparum* populations often follow the isolation-by-distance (IBD) model of population structure (Tanabe et al. 2010) and are also maintained as genetically sub-structured populations (Anderson et al. 2000). Whether Indian *P. falciparum* populations follow the IBD and/or genetically sub-structured models of population structure remains to be seen with a larger sample size of *P. falciparum* isolates collected from a wide geographic distribution within India.

The distant placement of the PfPH10 isolate from the other *P. falciparum* isolates in the phylogenetic tree (Fig. 2) was not surprising considering the fact that this sequence bears a large number of mutations (21 nucleotide substitutions) (Table I). This corroborates earlier observations on the high sequence divergence of PfPH10 from the Blsp1 and the 3D7 isolates (Tyagi et al. 2014). Due to the unknown origin within India of the PfPH10 clinical isolate (Sharma et al. 2001) we are not in a position to discuss the observed high mt genomic differentiation among the Indian isolates, the reference 3D7 isolate and global isolates of *P. falciparum*. The placement of all global isolates in a single cluster supports the hypothesis of a comparatively lower polymorphic nature of mt genomes in relation to other recombined genomes (e.g., nuclear genomes) and reasserts the idealistic nature of mt genomes for evolutionary inferences (Conway et al. 2000). In general, the four Indian *P. falciparum* isolates possess an appreciable level of nucleotide diversity (as measured by \( \pi \)) that corresponds to the African isolates. The observed results, although with a very limited sample size, provide sufficient background information for future studies, including the sequencing more Indian

![Fig. 2: neighbor-joining phylogenetic tree based on whole mitochondrial genome sequence alignment showing genetic interrelationships among Plasmodium falciparum isolates from different endemic localities of the globe. a: Joy et al. (2003); b: Tyagi et al. (2014); c: present paper; d: Sharma et al. (2001).](image)

*P. falciparum* isolates and performing associated comparative and evolutionary genomic studies. Such studies will help to understand the population structure and demography of Indian *P. falciparum* isolates and revisit the evolutionary history of global *P. falciparum*. Most importantly, such studies will help discover whether the antimalarial drug atovaquone could be beneficial malaria chemotherapy in India. This is because the cyt b gene present in the *P. falciparum* mitochondrion is considered to be the target of atovaquone (Vaidya et al. 1993, Biagini et al. 2006) and low or no diversity in the cyt b gene in populations would be the ideal condition for using atovaquone to treat *P. falciparum* malaria.

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